# **Example Candidate Responses**

Cambridge International Level 3 Pre-U Certificate in CHEMISTRY (9791)





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 UNIVERSITY of CAMBRIDGE
 International Examinations



# Example Candidate Responses

# Chemistry (9791)

**Cambridge International Level 3 Pre-U Certificate in Chemistry (Principal)** 

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## Cambridge International Level 3 Pre-U Certificate

# Chemistry

# 9791

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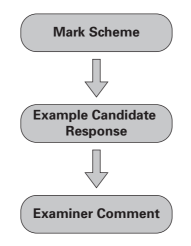
### **Introduction**

The main aim of this booklet is to exemplify standards for those teaching Cambridge Pre-U, and to show how different levels of candidates' performance relate to the subject's curriculum and assessment objectives.

Cambridge Pre-U is reported in three bands (Distinction, Merit and Pass) each divided into three grades (D1, D2, D3; M1, M2, M3; P1, P2, P3).

In this booklet a range of candidate responses has been chosen to illustrate as far as possible each band (Distinction, Merit and Pass). Each response is accompanied by a brief commentary explaining the strengths and weaknesses of the answers.

For ease of reference the following format for each paper has been adopted:



Each question is followed by an extract of the mark scheme used by Examiners. This, in turn, is followed by examples of marked candidate responses, each with an examiner comment on performance. Comments are given to indicate where marks were awarded and how additional marks could have been obtained. In this way, it is possible to understand what candidates have done to gain their marks and what they still have to do to improve their grades.

Teachers are reminded that a full syllabus and other teacher support materials are available on www.cie.org.uk. For past papers and Examiner Reports please contact CIE on international@cie.org.uk.

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## **Components at a Glance**

Component	<b>Component Title</b>	Duration	Weighting (%)	Type of Assessment
Paper 1	Part A Multiple Choice	1 hour	15	Written paper, externally set and marked
Paper 2	Part A Written	2 hours 15 minutes	35	Written paper, externally set and marked
Paper 3	Part B Written	2 hours 15 minutes	35	Written paper, externally set and marked
Paper 4	Practical	2 hours	15	Practical exam, externally set and marked

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This booklet contains a selection of example candidate responses and examiner comments for Papers 2, 3 and 4.

## Paper 2 Part A Written

#### **Question 1 Mark Scheme**

(a)	ethanol: 46 g mol <sup>-1</sup> octane: 114 g mol <sup>-1</sup> 1 mark if both numbers correct	
	no penalty for incorrect sig. figs.	[1]
(b)	density in g cm <sup>-3</sup> = 2 g mol <sup>-1</sup> / 24000 cm <sup>3</sup> mol <sup>-1</sup> = $8.3 \times 10^{-5}$ g cm <sup>-3</sup> no penalty for incorrect sig. figs.	[1]
(c)	zero (by definition) award this mark if the zero is just entered in the table	[1]
(d)	-285.8 kJ mol <sup>-1</sup> the mark is given for indicating that it is the same as the enthalpy of formatio question	on given in the [1]
(e)	$C_8H_{18}(I) + 25/2O_2(g) \rightarrow 8CO_2(g) + 9H_2O(I)$ (1) equation must be per mole of octane and correctly balanced for the first mark correct state symbols for octane, oxygen, carbon dioxide and water (1)	[2]
(f)	$\begin{split} &\Delta_c H^e \ (octane) = 8 \Delta_t H^e \ (CO_2) + 9 \Delta_t H^e \ (H_2 O) - \Delta_t H^e \ (octane) \\ &= \{(8 \times -393.5) + (9 \times -285.8) - (-250)\} \ kJ \ mol^{-1} \\ &= \{-3148.0 - 2572.2 + 250.0\} \ kJ \ mol^{-1} \\ &= -5470.2 \ kJ \ mol^{-1} \ (3) \\ &\Delta_t H^e (CO_2) \ and \ \Delta_t H^e (H_2 O) \ multiplied \ by \ 8 \ and \ 9, \ respectively \ (1) \\ &correct \ signs \ (1) \\ &2 \ marks \ given \ if \ all \ correct \ apart \ from \ an \ arithmetical \ slip \\ &2 \ marks \ given \ for \ 5470.2 \ kJ \ mol^{-1} \end{split}$	[max 3]
(a)	(i) methanol: $-726.0 \text{ kJ mol}^{-1} / 32 \text{ g mol}^{-1} = -22.69 \text{ kJ g}^{-1}$	[1]
(9)	<ul> <li>(ii) hydrogen: -285.8 kJ mol<sup>-1</sup> / 2 g mol<sup>-1</sup> = -142.9 kJ g<sup>-1</sup> accept from 2 to 5 sig. figs.</li> <li>no penalties for missing units or forgetting minus sign -1 for each wrong answer</li> </ul>	
	<ul> <li>–1 if sig. figs. outside the allowed range (only penalise once)</li> </ul>	[1]

- (h) (i) ethanol:  $-1367.3 \text{ kJ mol}^{-1} \times 0.789 \text{ g cm}^{-3} / 46 \text{ g mol}^{-1}$ =  $-23.5 \text{ kJ cm}^{-3}$ 
  - (ii) octane: -5470.2 kJ mol<sup>-1</sup> × 0.703 g cm<sup>-3</sup> / 114 g mol<sup>-1</sup>
    = -33.7 kJ cm<sup>-3</sup>
    accept only 2 or 3 sig. figs.
    no penalties for missing units or forgetting minus sign
    -1 for each error
    -1 if sig. figs. outside the allowed range (only penalise once)
    where working is correct but final answer is inexplicably wrong, allow the mark [1]
- (i) the enthalpy change of combustion value for hydrogen is for standard conditions, and so relates to gaseous hydrogen, not to liquid hydrogen
   OR no account taken of different temperatures / latent heat of vaporisation of hydrogen allow comment about how the value of the density of liquid hydrogen is unsuitable for the calculation of energy per unit volume for gaseous hydrogen allow just a reference to the different state of hydrogen

[Total: 14]

[1]

#### Example Candidate Response – Distinction (D1)

1 Table 1.1 gives some data on four fuel sources: methanol, ethanol, hydrogen and octane. Octane can serve as a rough approximation of petrol.

name	formula	molar mass /g mol <sup>-1</sup>	density /g cm <sup>-3</sup>	∆ <sub>c</sub> H <sup>⊕</sup> (298K) /kJ mol <sup>–1</sup>	∆ <sub>f</sub> H <sup>◆</sup> (298K) /kJ mol <sup>-1</sup>
methanol	CH3OH	32	0.793 <sup>a</sup>	-726.0	-239.1
ethanol	C <sub>2</sub> H <sub>5</sub> OH	46	0.789 <sup>a</sup>	-1367.3	-277.1
liquid hydrogen	H <sub>2</sub>	2	0.0711 <sup>b</sup>		
octane	C <sub>8</sub> H <sub>18</sub>	114	0.703 <sup>a</sup>	V	-250.0

#### Table 1.1

<sup>a</sup> At 298K and 1 bar pressure.

<sup>b</sup> At 20K and 1 bar pressure.

- (a) Insert the missing molar mass values in the table.
- (b) Calculate the density of gaseous hydrogen at 298 K and 1 bar pressure. Assume 1 mol of any gas occupies 24 dm<sup>3</sup> at 298 K and 1 bar pressure. Give your answer in g cm<sup>-3</sup>.

5.22 x1

-285.8

(c) What is the value of the standard enthalpy of formation of hydrogen gas, H<sub>2</sub>?

0.000

(d) Use the information in Table 1.2 to give the value of the standard enthalpy of combustion of hydrogen.

Table '	1.2
---------	-----

name	Δ <sub>f</sub> H <sup>Φ</sup> (298 K) /kJ mol <sup>−1</sup>
water	-285.8
carbon dioxide	-393.5

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[1]

g cm<sup>-3</sup> [1]

...[1]

and the second state of the second

#### Cambridge Pre-U Example Candidate Responses

(e) Write down the chemical equation that represents the standard enthalpy of combustion of octane. Include state symbols.

\$ (e) F C. H. + 12.5 02(5) 8002

(f) Use the enthalpy of formation data in Table 1.1 and Table 1.2 to calculate the standard enthalpy of combustion of octane.

 $-9 16 CQ_{(5)} + 15 H_2 O_{(5)}$ 2  $7 - 313.5 \times 16$ - 313.5+16 -393.5×16-285.8×18) [3] 5470

(g) An important property of a fuel, especially when the fuel has to be lifted (such as in aviation), is the energy released on combustion *per gram* of fuel.

Calculate the enthalpy change of combustion per gram of fuel at 1 bar pressure and 298 K for methanol and hydrogen gas.

(i) methanol  $-\frac{726 \text{ k} \text{ S}_{\text{meth}}}{232 \text{ m}^{-1}} : -22.69 \text{ k} \text{ S}_{\text{s}}^{-1}$   $-22.69 \text{ k} \text{ S}_{\text{s}}^{-1}$ (ii) hydrogen gas  $-\frac{285.8 \text{ k} \text{ S}_{\text{met}}}{2 \text{ s}_{\text{met}}} : -142.9 \text{ k} \text{ S}_{\text{s}}^{-1}$  $-143 \text{ k} \text{ S}_{\text{s}}^{-1}$  [2] (h) Another important characteristic of a fuel, especially when there is a fuel tank of limited size, is the energy released on combustion per cm<sup>3</sup> of fuel.

Calculate the enthalpy change of combustion per cm<sup>3</sup> of fuel for ethanol and octane.

-1367.3 kJml x 0.017 mlc (i) ethanol -23.45 kJ cm -23.5 kJ 0.017 ml 23.733 (ii) octane 5470 0.00617 -YOUAA 0.7035 33 ...[2] Explain why, given the data in the question, it is not strictly possible to make a fair (i) comparison of the energy released per cm<sup>3</sup> of liquid hydrogen with the other fuels. in given is at 20k rather 298K Not be consider by SU ble The Allen John .....[1] Total: 14]

#### **Examiner Comment**

The candidate was able to complete almost all of the calculations correctly but failed to read (b) with sufficient care.

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#### Example Candidate Response – Distinction

 Table 1.1 gives some data on four fuel sources: methanol, ethanol, hydrogen and octane. Octane can serve as a rough approximation of petrol.

name	formula	molar mass /g mol <sup>-1</sup>	density /g cm <sup>-3</sup>	∆ <sub>c</sub> H <sup>⊕</sup> (298K) /kJ mol <sup>-1</sup>	∆ <sub>f</sub> H <sup>+</sup> (298K) /kJ mol <sup>-1</sup>
methanol	СН3ОН	32	0.793 <sup>a</sup>	-726.0	-239.1
ethanol	C2H5OH	46	0.789 <sup>a</sup>	-1367.3	-277.1
liquid hydrogen	H <sub>2</sub>	2	0.0711		
octane	C <sub>8</sub> H <sub>18</sub>	114 V	0.703 <sup>a</sup>		-250.0

#### Table 1.1

1

<sup>a</sup> At 298 K and 1 bar pressure.

<sup>b</sup> At 20 K and 1 bar pressure.

- (a) Insert the missing molar mass values in the table.
- (b) Calculate the density of gaseous hydrogen at 298K and 1 bar pressure. Assume 1 mol of any gas occupies 24 dm<sup>3</sup> at 298K and 1 bar pressure. Give your answer in g cm<sup>-3</sup>.
  Density = Mass
  Mass of H2 m at 298K = 1×2

Density = 83.3 gcm

83.3 g cm<sup>-3</sup> [1]

=2

(c) What is the value of the standard enthalpy of formation of hydrogen gas, H<sub>2</sub>?

o kJmol"

(d) Use the information in Table 1.2 to give the value of the standard enthalpy of combustion of hydrogen.

7. 7.	1-593.5 1-155.8 name	∆ <sub>f</sub> H <sup>●</sup> (298K) /kJ mol <sup>-1</sup>	
X X	water	-285.8	
	carbon dioxide	-393.5	. /

[1]

....[1]

(e) Write down the chemical equation that represents the standard enthalpy of combustion of octane. Include state symbols.

Ce Hieres + 25/20215 & CO245+942012) ...[2]

(f) Use the enthalpy of formation data in Table 1.1 and Table 1.2 to calculate the standard enthalpy of combustion of octane.

$$C_{8}H_{18}(e) + \frac{27}{2}O_{2}(g) \xrightarrow{AHe} 8 (O_{2}(g) + 9 H_{2}O(e) 
\int -250.0 \int 0 \int -393.5 \times 8 \int -285.8 \times 9 \\
\theta C_{15} + \frac{37}{2}O_{2}(g) + 9 H_{2}cg) \\
AHe = -(-250.0) + (-393.5 \times 8) + (-285.8 \times 9) \\
= -5470.2 \ \text{kJmst}^{-1} \qquad [3]$$

(g) An important property of a fuel, especially when the fuel has to be lifted (such as in aviation), is the energy released on combustion per gram of fuel.

Calculate the enthalpy change of combustion per gran of fuel at 1 bar pressure and 298 K for methanol and hydrogen gas.

- (i) methanol  $-726.0 \text{ kJmol}^{-1}$   $= -726 \times 32$   $= -23232 \text{ kJg}^{-1}$ (ii) hydrogen gas
  - $-285.8 \text{ kJml}^{-285.8 \text{ kJml}^{-1}}$ = -285.8 × )2 = -571.6 kJg<sup>-1</sup>

N . Hr

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(h) Another important characteristic of a fuel, especially when there is a fuel tank of limited size, is the energy released on combustion per cm<sup>3</sup> of fuel.

Calculate the enthalpy change of combustion percm3 of uel for ethanol and octane. Mr x A ont. (i) ethanol Nect may st. -1367.3 KTmor = -1367.3 m x 0.789 x 46 -49624. 8 k (cm 3 = - 49624.8 kJon-3 (to 1 dp) (ii) octane -5470.2 KTmol" Verh = -5470.2 × 0.703 × 114 -438392.8 H m [2] = \_ 438392.8 kJon 3 (to lap) (i) Explain why, given the data in the question, it is not strictly possible to make a fair comparison of the energy released per cm<sup>3</sup> of liquid hydrogen with the other fuels. Liquid hydrogen's density is measured at 20 k whereas other fuels at 298K, so deasity not comparable, i.e. Ka energy[1] to not con [Votal: 14]

#### **Examiner Comment**

The candidate showed a good understanding of the principles involved in most of the calculations but in (b) failed to convert dm<sup>3</sup> into cm<sup>3</sup> and made a careless error in (g). Also a lack of appreciation of significant figures was shown in (h).

#### Question 2 Mark Scheme

(a)	178/(178 + 32) × 100% = 84.8%	[1]
(b)	correct plotting of point in van Arkel triangle (1) the point has coordinates (2.39, 2.45) half a gradation of leeway either side scale, is acceptable	e, i.e. 0.05 on the
	it is an insulator (1)	[2]
(c)	it is ionic	[1]
(d)	reaction 1: $HfO_2 + 4HCl \rightarrow HfCl_4 + 2H_2O(1)$ reaction 2: $HfCl_4 + 2Mg \rightarrow Hf + 2MgCl_2(1)$	
	ecf incorrect hafnium chloride formula in step 2 from step 1	[2]
		[Total: 6]

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84.8

2.315

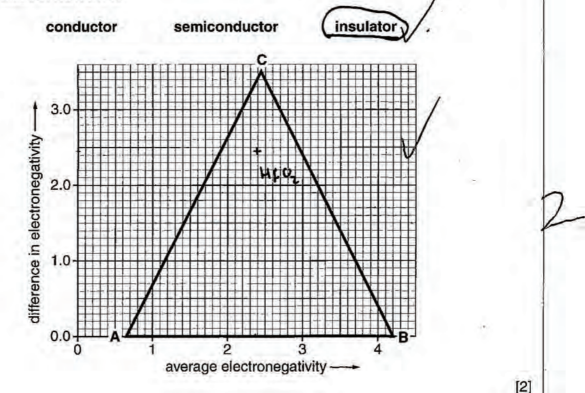
#### Example Candidate Response - Distinction

- The Intel® super-fast 45 nm Core 2 processors are based on Penryn technology. This involves 2 the use, for the first time in computer chips, of an oxide of hafnium.
  - (a) This oxide of hafnium has the formula HfO2. Calculate the percentage of hafnium by mass in this oxide.

(b) Table 2.1 provides the electronegativity data for O and Hf.

elem	ent	electronegativity	3.61-46= 2.45
0		3.61	3.6141.16 : 2.315
H	t l	1.16	

Computer chips contain electrical conductors, semiconductors and insulators. On the van Arkel triangle mark the point corresponding to the oxide of hafnium and use this point to deduce its electrical properties. Ring the correct option.



The oxide of hafnium is

(c) Use your van Arkel plot to decide whether the oxide of hafnium is best described as ionic, covalent or metallic. Ring the correct option below.

The oxide of hafnium is best described as

ionic

metallic

1.1

(d) Elemental hafnium has neutron-absorbing properties that are useful in nuclear reactors. It can be extracted from the oxide, HfO<sub>2</sub>, by the following reactions.

covalent

reaction 1 reaction with hydrochloric acid

reaction 2 reduction of a product of reaction 1 with magnesium

Write balanced equations for these reactions.

equation for reaction 1 $HfO_1 + 4HCL \rightarrow HfCL_1 + 2H_2O$	
equation for reaction 2 $M_{fCL_{4}} + 2M_{g} \rightarrow 2M_{gCL_{2}} + M_{f}$ [2]	2
[Total: 6]	(6)

#### **Examiner Comment**

A Distinction level candidate would be expected to achieve full marks on this question.

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#### **Question 3 Mark Scheme**

(a)	CsF [1]
(b)	clear that a Cs electron is in the F outer shell (1) charges shown on the ions (1) allow a Cs ion with 8 electrons shown in the outer shell no marks awarded if there is sharing of electrons [2]
(c)	largest possible difference in electronegativity between its two constituent elements owtte [1]
(d)	high boiling point / low vapour pressure / does not evaporate easily [1]
(e)	wide separation of charges / large ions AND low charges both needed for mark [1]
(f)	high surface tension (1) (relatively) high melting and/or boiling points or liquid at rtp (compared to analogous compounds) or high specific thermal capacity (1) less dense as a solid than as a liquid (owtte) or greatest density at 4° C (1) [max 2]
(g)	H bond drawn between hydrogen on one molecule and oxygen on another (1) oxygen lone pair drawn at the start of one of the H bonds and both ends of a dipole shown on each molecule (1) bond angle of 180° around an H-bonding hydrogen – shown explicitly or looks like 180° intended (1)
	intended (1) [3]
(h)	two H bonds shown between two amine groups on melamine and the carbonyl oxygens on PTCDI, and an H bond shown between the NH group on the PTCDI and the $-N=$ atom on melamine that is between the two H-bonding amine groups. (2) for both marks the H-bonds from the amine groups must be from one of the two hydrogens of each amine in particular, rather than just from $-NH_2$ otherwise just 1 mark is given ecf from (g) if the notation for an H-bond is incorrect no marks if all three H bonds aren't correctly identified small errors in copying down the structures should not be penalised as long as they don't
	affect the H-bonding interaction [2]
	[Total: 13]

[Total: 13]

#### Example Candidate Response – Distinction (D1)

- 3 The compound whose bonding most resembles pure ionic bonding is caesium fluoride.
  - (a) Write down the formula of caesium fluoride.

CsF

[ \*\* ] [ \* Cs \*]

(b) Draw a dot-cross diagram to show the bonding in caesium fluoride. Show outer electrons only.

: F :

[2] (c) Explain why caesium fluoride is the compound whose bonding most closely resembles pure ionic. The difference in electro regativity between caesium and fluorine is the frectest out of all binary compound. [1]

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.....[1]

.[1]

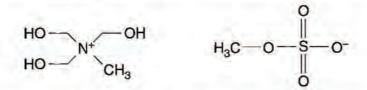
(d) Virtually all ionic compounds are solids at room temperature and pressure. However, researchers have designed ionic compounds whose ionic bonding is so weak that they are liquids under these conditions. Ionic liquids are often easy to handle as solvents as they are non-volatile; they have also recently found use in solar cells for this reason.

Explain what is meant by non-volatile.

They do not evaporate easily V

(e) In the pure ionic bonding model, the ionic bond energy is proportional to the charge on each ion and inversely proportional to the distance between the charges, which are considered to be located at the centre of ions.

The structure of an ionic substance which is a liquid at room temperature and has been used in thermometers (reported in *Green Chemistry*, 2008) is shown below.



Suggest two features of these ions that account for the compound having such a low melting point.

1. There is a large amount of steric hindrance between the charges resulting in bond strept

- 2. The charges on each ion by 14 and 1- respectively, making the ionic bonol weark [1]
- (f) Hydrogen-bonding is weaker than ionic or covalent bonding, but accounts for many important intermolecular attractions.

State two anomalous properties of water that are the result of hydrogen-bonding.

- 1. Solid water (ice) is less dense than the liquid form
- 2. Higher boiling point than all other Group 16 hydrogen compounds V [2]
- (g) Draw a second molecule of water and a hydrogen-bond between the two molecules. Indicate the bond angle around the hydrogen atom involved in the hydrogen-bond. Include all relevant lone pairs and dipoles.

[3]

(h) Hydrogen-bonding is directional (i.e. a specific link between two atoms can be drawn) and has many applications in linking together molecules in an organised way. This linking has been put to use recently by researchers designing self-assembling surface networks for applications in nanotechnology (reported in *Nature*, 2008).

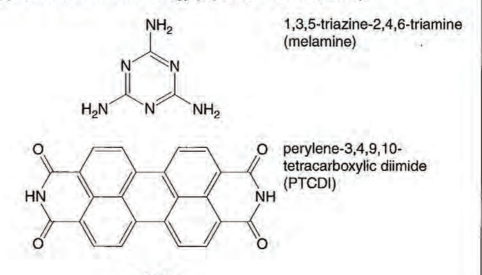
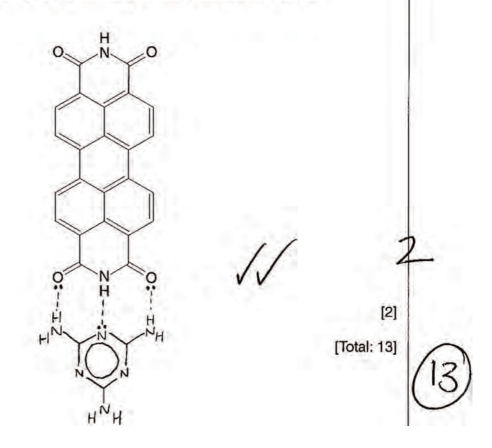


Fig. 3.1

The two molecules in Fig. 3.1 were chosen for the self-assembling network. A molecule of melamine and a molecule of PTCDI attach together strongly via three hydrogen bonds. Suggest where these **three** hydrogen-bonds form by drawing the melamine below in the correct orientation, with the hydrogen-bonds connecting the relevant atoms.



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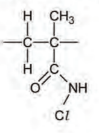
#### **Examiner Comment**

A Distinction candidate would have probably scored 10/11 marks, with marks for **(h)** being the most difficult to score.

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#### **Question 4 Mark Scheme**

(a) allow either the repeat fragment or the notation with it in brackets brackets must be used if a skeletal formula is used to represent the repeat unit only two carbons in the backbone should be shown in the fragment or between the brackets allow any unambiguous structural formula



[1]

[1]

(b) the monomer should show a C=C double bond between the two backbone carbons from the repeat unit, and an amide in place of the N-chloroamide the C=C double bond must be explicit rather than implied

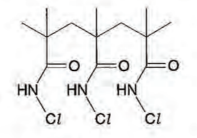


(c) (i) (+)1 [1] (ii) (+)1 ecf from (c)(i), i.e. answer to (c)(ii) should be the same as (c)(i) [1] (iii) oxidation:  $2I^- \rightarrow I_2 + 2e^-(1)$ reduction:  $C/O^- + 2H^+ + 2e^- \rightarrow C/^- + H_2O$  $HOC/ + H^* + 2e^- \rightarrow C/^- + H_2O(1)$ or [2] (iv) starch [1] (v)  $12.50 \text{ cm}^3 \times 0.100 \text{ mol } \text{dm}^{-3} = 0.00125 \text{ mol}$ no sig. figs. or unit penalties [1] (vi) 0.00125 mol / 2 = 0.000625 mol no sig. figs. or unit penalties ecf from (v), i.e. answer from (v) should be divided by 2 [1] (vii)  $0.000625 \text{ mol} \times 35.5 \text{ g mol}^{-1} \times 100 \text{ cm}^3 / 10 \text{ cm}^3 = 0.222 \text{ g} (2)$ 1 mark for multiplying answer to (vi) by 35.5 g mol<sup>-1</sup> 1 mark for scaling up by 10, even if this isn't explicitly explained. no sig. figs. or unit penalties ecf from (i) [2] [Total: 11]

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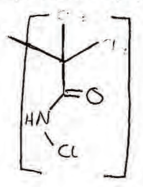
#### Example Candidate Response – Distinction

4 Chemists have recently found a way of making the strong, light-weight and thermally stable polymer, Kevlar<sup>®</sup>, to be antibacterial (reported in *Industrial & Engineering Chemistry Research*, 2008). This was achieved by coating it with another polymer, a fragment of which is shown in Fig. 4.1.





(a) Draw the repeat unit of the polymer structure shown in Fig. 4.1.



(b) The polymer shown in Fig. 4.1 is made by the following reactions:

reaction 1 reaction 2 an addition polymerisation reaction of a monomer known as MAA the substitution of a hydrogen atom in the polymer with a chlorine atom using bleach

Draw the structure of the monomer MAA.



(c) Not all the nitrogen atoms in the polymer end up bonded to a chlorine atom. The quantity of chlorine actually present in the polymer can be determined using reactions 3, 4 and 5.

A known mass of polymer (written as XCONH–Cl) is hydrolysed to convert the chlorine content of the polymer to chloric(I) acid, HOCl:

[1]

[1]

	<b>action 3</b> XCONH–C $l$ + H <sub>2</sub> O $\rightarrow$ XCONH here X = remainder of polymer)	H <sub>2</sub> + HOCl					
The	e chlorine content is then 'converted' to i	odine:					
rea	ection 4 HOC $l$ + 2KI + H <sup>+</sup> $\rightarrow$ I <sub>2</sub> + KCl	+ H <sub>2</sub> O + K <sup>+</sup>					
The	e quantity of iodine is determined using a	a titration with sodium thiosulfate:					
rea	iction 5 $I_2 + 2Na_2S_2O_3 \rightarrow 2NaI + Na_2$	S <sub>4</sub> O <sub>6</sub>					
(i)	State the oxidation number of the chlorine in HOC1.						
	+1	[1]					
(ii)	Given that <b>reaction 3</b> is a hydrolysis re in the polymer. +	action, state the oxidation number of chlorine					
(iii)	Write ionic half equations for the oxidal oxidation $\mathcal{ZI} \longrightarrow \mathcal{I}$ reduction $\mathcal{A}^+ + \mathcal{Z}e^ \mathcal{A}^+$	tion and reduction processes in reaction $f$ . $z + 2e^{-1}$					

In the analysis of a sample of polymer, 1.00 g of the polymer was hydrolysed (reaction 3). The resulting mixture was reacted with excess acidified potassium iodide (reaction 4) and then made up to 100 cm<sup>3</sup> with distilled water. 10.0 cm<sup>3</sup> of this solution reacted with exactly 12.50 cm<sup>3</sup> of sodium thiosulfate solution of concentration 0.100 mol dm-3 (reaction 5). Name a suitable indicator for this titration. (iv) Starch (v) Determine the amount (in moles) of sodium thiosulfate used in the titration., Mol = 12.50 × 0.100 1.25 × 10 ..... mol [1] (vi) Determine the amount (in moles) of iodine, I2, in the titration. = 5+ OIX251 6.25 × 10 .... mol [1] Determine the mass of chlorine present in the sample of polymer. (vii) nass = 6.75 × 109 × 35.5 0.0222g (35F Stoichiometry Jz: Cl is 1:1 so 6-25×10<sup>-9</sup> mol Cl ..[2] al: 11]

#### **Examiner Comment**

In (a) the candidate was unable to draw the repeat unit of the polymer. The reduction reaction in (c) (iii) was targeted at the most able candidates. A scaling factor of 10 was missed in (c) (vii).

#### Example Candidate Response – Merit

4 Chemists have recently found a way of making the strong, light-weight and thermally stable polymer, Kevlar<sup>®</sup>, to be antibacterial (reported in *Industrial & Engineering Chemistry Research*, 2008). This was achieved by coating it with another polymer, a fragment of which is shown in Fig. 4.1.

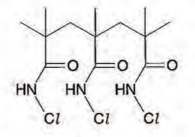
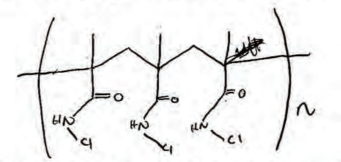


Fig. 4.1

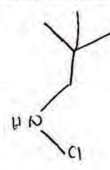
(a) Draw the repeat unit of the polymer structure shown in Fig. 4.1.



(b) The polymer shown in Fig. 4.1 is made by the following reactions:

reaction 1 reaction 2 an addition polymerisation reaction of a monomer known as MAA the substitution of a hydrogen atom in the polymer with a chlorine atom using bleach

Draw the structure of the monomer MAA.



[1]

[1]

(c) Not all the nitrogen atoms in the polymer end up bonded to a chlorine atom. The quantity of chlorine actually present in the polymer can be determined using reactions 3, 4 and 5.

A known mass of polymer (written as XCONH-Cl) is hydrolysed to convert the chlorine content of the polymer to chloric(I) acid, HOCl:

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# Cambridge Pre-U Example Candidate Responses

	<b>action 3</b> XCONH– $Cl$ + H <sub>2</sub> O $\rightarrow$ XCONH <sub>2</sub> + HOC <i>l</i> here X = remainder of polymer)
The	e chlorine content is then 'converted' to iodine:
rea	action 4 HOC $l$ + 2KI + H <sup>+</sup> $\rightarrow$ I <sub>2</sub> + KC $l$ + H <sub>2</sub> O + K <sup>+</sup> $\downarrow \downarrow \downarrow \downarrow \downarrow \downarrow \downarrow \downarrow$
	e quantity of iodine is determined using a titration with sodium thiosulfate:
rea	action 5 $I_2 + 2Na_2S_2O_3 \rightarrow 2NaI + Na_2S_4O_6$
(i)	State the oxidation number of the chlorine in HOC1.
(ii)	Given that reaction 3 is a hydrolysis reaction, state the oxidation number of chlorine in the polymer. 1
(111)	Write ionic half equations for the oxidation and reduction processes in reaction 4. oxidation $2I \xrightarrow{-} I_2 + 2e \xrightarrow{-} V$ reduction $CI^+ + 2e^{-} \xrightarrow{-} CI^- X$ [2]

In the analysis of a sample of polymer, 1.00g of the polymer was hydrolysed (reaction 3). The resulting mixture was reacted with excess acidified potassium iodide (reaction 4) and then made up to 100 cm<sup>3</sup> with distilled water. 10.0 cm<sup>3</sup> of this solution reacted with exactly 12.50 cm<sup>3</sup> of sodium thiosulfate solution of concentration 0.100 mol dm<sup>-3</sup> (reaction 5). (iv) Name a suitable indicator for this titration. .. [1] (v) Determine the amount (in moles) of sodium thiosulfate used in the titration. Mol - Volx come 1.25×10-3 mol [1] (vi) Determine the amount (in moles) of iodine, I2, in the titration. 1. 25 ×10-3/2 6.25×101 mol [1] Determine the mass of chlorine present in the sample of polymer. (vii) .....[2] 10× [Tptal: 11 **Examiner Comment** 

The candidate scored on most of the lower level questions although they failed to attempt (c) (iv) which tested basic knowledge. The overall standard of the answer was improved by scoring in (c) (ii) and for the oxidation half-equation in (c) (iii).

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Question 5 Mark Scheme

(a)	292 - 122 = 170						[1]		
(b)	9								[1]
(c)	18 ecf fr	om pa	art (b),	i.e. the	e num	ber of	eleme	ents wide = twice the number of orbitals	[1]
(d)	4p 5:	s 4d 5p	o 6s 41	f 5d					[1]
(e)	6d, 7	p, 8s a	and 5g	shoul	d be a	dded	to the	diagram as below	[1]
	1s	2s	3s	4s	5s	6s	7s	8s	
		2p	Зр	4p	5p	6p	7p		
			3d	4d	5d	6d			
				4f	5f				
					5g				
(f)	two g	y elect	rons						[1]
									[Total: 6]

### Example Candidate Response – Pass

5 The scientific community was shocked at the recent claim of the discovery of an isotope of a new element with a mass number of 292 (published in <i>arXiv</i> , 2008): this is over 50 mass units higher than uranium, the heaviest known naturally-occurring element. There is a possibility that there is an 'island of stability' beyond the known Periodic Table at some very high atomic numbers.							
1	(a)	The authors of this claim suggested that the atomic number of the element is 122. How many neutrons are there in this isotope?					
		170 [1]	1				
	(b)	If this element really exists then it will require a new block of the periodic table, corresponding to the occupancy of another type of subshell, beyond the $s$ , $p$ , $d$ and $f$ . This would be a $g$ subshell, which is predicted to be found in the 5th shell of an atom, i.e. the 5g subshell.					
		Based upon the sequence of subshells in the Periodic Table, $s$ , $p$ , $d$ , $f$ , predict how many orbitals there are in a $g$ subshell.	,				
		9 (nine) subshells [1]	1				
	(c)		I				
		<u>18[1]</u>	1				
		thich the subshells are filled by electrons (the Aufbau principle). $1s \rightarrow 2s \qquad 3s \qquad 4s \qquad 5s \qquad 6s \qquad 7s \qquad 7s \qquad 2p \qquad 3p \qquad 4p \qquad 5p \qquad 6p \qquad 7p \qquad 3d \qquad 4d \qquad 5d \qquad 6d \qquad 4d \qquad 6d \qquad 6d \qquad 4d \qquad 6d \qquad 6d \qquad 4d \qquad 6d \qquad 6$					
	(d)	List the order of filling subshells from 4p to 5d.	0				
		$\frac{4p + 5p + 6s \rightarrow 5d}{X}$					
	(e)	The subshells in the scheme above are those that are occupied by the elements up to uranium. Add to the above diagram the next four subshells that would be expected to be filled. [1]	0				
	(f)	Following the Aufbau principle, how many electrons in the 5g subshell would element 122 be expected to have?	0				
		[Total: 6]	3)				

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#### **Examiner Comment**

The candidate was able to use their knowledge of atomic structure and s, p, d, f orbitals and extend it to a new element containing a g subshell scoring the marks for (a), (b) and (e). However, the candidate was unable to apply the Aufbau principle to this new element which would have raised their level of achievement.

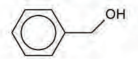
#### Question 6 Mark Scheme

decreased by 1 (1)

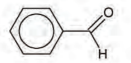
(a)	5 signals	[1]
(b)	3 isomers	[1]
(c)	any oxidising agent, e.g. manganate, dichromate, chlorate ignore omission of acidification where necessary oxidising agents that lack an oxygen atom need to be indicated as (aq) if a formula is given it must be correct to earn the mark if an oxidation number is quoted it must be correct to earn the mark	[1]
(d)	Grignard (reagent)	[1]
(e)	<ol> <li>(nucleophilic) substitution allow hydrolysis (1)</li> <li>hydrolysis (1)</li> <li>dehydration / elimination allow reduction as FGL of functional group decreased by 1 (1)</li> </ol>	carbon has

[3]

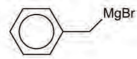
(f) A: benzyl alcohol (phenylmethanol) (1)



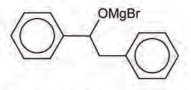
B: benzaldehyde (1)



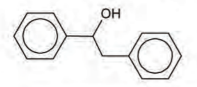
C: benzylmagnesium bromide (1)



D: PhCH(OMgBr)CH<sub>2</sub>Ph (1)



E: PhCH(OH)CH<sub>2</sub>Ph (1)



allow all structural and displayed formulae as long as structure is unambiguous penalise repeated systematic or trivial errors only once

[5]

[Total: 12]

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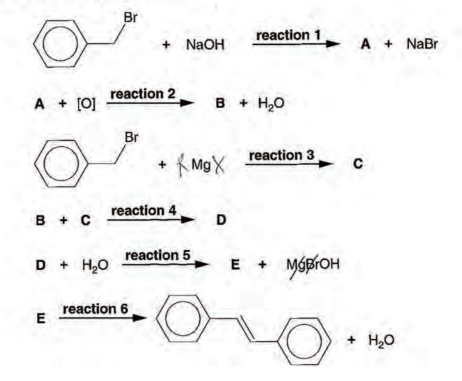
#### Example Candidate Response – Pass

- 6 A survey in 2008 of the 24 million known organic compounds identified the most common structural motifs. The 3rd most popular shape (after the hexagon and pentagon) was based on the 1,2-diphenylethane molecule.
  - (a) How many signals would there be in the carbon-13 NMR spectrum of the 1,2-diphenylethane molecule?
  - (b) If one of the hydrogen atoms on one of the phenyl groups of 1,2-diphenylethane is substituted with a chlorine atom, how many possible isomers would there be?

......

A closely related compound to 1,2-diphenylethane is 1,2-diphenylethene, commonly known as stilbene.

Stilbene can be synthesised from (bromomethyl)benzene according to the scheme below. The reactions are labelled 1 to 6 above the reaction arrows. The benzene ring does not take <sup>-</sup> part in any of these reactions.



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#### Cambridge Pre-U Example Candidate Responses

(c) Reaction 2 is a partial oxidation. In the equation [O] represents the oxygen atom provided from some suitable reagents. Suggest such a reagent. Pothisium Remangamente H/KMiloy aciditica ..[1] (d) What is the name of the type of compound produced in reaction 3? wide ...[1] ..... (e) . Classify the type of reaction in 1, 5 and 6. substitution reaction 1 ..... ...[1] reaction 5 ..... hy drolys .....[1] reaction 6 ..... Condusation X [1] (f) Draw the structures of unknowns A to E. [1] 0 в [1] C OMA [1] D [1] 04 CH E [1] al: 12]

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#### **Examiner Comment**

The candidate showed some knowledge and understanding of the chemistry of halogenoalknes and alcohols in recognising the types of reactions 1 and 5, identifying the products of reactions 1 and 2, and recalling a reagent to oxidise product A.

#### Question 7 Mark Scheme

(a)	δ+	on the I and $\delta$ - on the N	
		correct use of arrow nomenclature	[1]
(b)	(i)	$NH_3 + 3F_2 \rightarrow NF_3 + 3HF$	[1]
	(ii)	$2NF_3 \rightarrow N_2 + 3F_2$ or with stoichiometry 1:1/2:3/2	[1]
	(111)	correct dot-cross diagram, including lone pairs on the fluorines (1) shape = pyramidal (or trigonal pyramidal) (1) bond angle indicated as anything from 102 to 107° (1) (actual bond angle is 102.3 degrees: there is weak bonding pair-bor due to the electron density in the N–F bonds being shifted towards fluorine atoms)	
	(iv)	stronger N–F bonds / higher activation energy allow: since the F–F bond is weaker than the Cl–Cl bond the therma NF <sub>3</sub> will be less exothermic than for NCl <sub>3</sub>	al decomposition of [1]
(c)	(i)	$N_2O_5 + H_2O \rightarrow 2HNO_3$	[1]
	(ii)	$\begin{array}{l} 12\text{HNO}_3 + \text{P}_4\text{O}_{10} \rightarrow 6\text{N}_2\text{O}_5 + 4\text{H}_3\text{PO}_4 \ (2) \\ \text{accept } 6\text{HNO}_3 + \text{P}_2\text{O}_5 \rightarrow 3\text{N}_2\text{O}_5 + 2\text{H}_3\text{PO}_4 \\ 1 \text{ mark for correct formulae but incorrect balancing} \end{array}$	[2]
	(iii)	NO <sub>2</sub> <sup>+</sup> NO <sub>3</sub> <sup>-</sup> or NO <sub>2</sub> NO <sub>3</sub>	[1]
			[Total: 11]

#### Example Candidate Response – Distinction (D1)

- 7 Nitrogen forms a variety of oxides and halides.
  - (a) Nitrogen triiodide, NI<sub>3</sub>, is an explosive that detonates with a snap even when only touched lightly. Given that the electronegativity value for nitrogen is 3.07 and for iodine is 2.36, indicate below the dipole in an N-I bond.

δ- δ+ N–I

(b) (i) Nitrogen trifluoride, NF<sub>3</sub>, can be prepared by reacting ammonia with fluorine. In this reaction the fluorine oxidises the nitrogen in ammonia while the oxidation number of hydrogen is unchanged.

Give the equation for this reaction.

# NH3 +6F - NE +HE X [1]

(ii) Nitrogen trifluoride is used to <u>etch</u> silicon in microelectronics. It is decomposed to its elements and the fluorine is used to attack the silicon.

Give the equation for the decomposition of nitrogen trifluoride.

Nitrogen trifluoride is a molecule that has attracted controversy recently for its (iii) possible potent contribution to the greenhouse effect. Draw the dot-cross diagram of this molecule; only include outer electrons. State the shape and the bond angle.

 $2 NF_2 \rightarrow N_2 + 3F_2$ 

shape .... trigona bond angle ..... 107 [3] (iv) Whereas nitrogen trifluoride is reasonably easy to handle, nitrogen trichloride is an extremely dangerous explosive. Suggest why nitrogen trifluoride is more stable

than the other nitrogen trihalides. Because of the short N = F bond whice it makes it have a stronger bond enthal

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[1]

......[1]

- (c) N2O5 is a less well-known oxide of nitrogen.
  - (i)  $N_2O_5$  is the anhydride of nitric acid, which means that it reacts with water to produce the acid. Write an equation for  $N_2O_5$  reacting with water.

N205 + H20 -+ 2HNO2 ....[1]  $N_2O_5$  can be made by reacting nitric acid with a dehydrating agent such as (ii) phosphorus(V) oxide. Bearing in mind that phosphorus(V) oxide is the anhydride of phosphoric acid, H<sub>3</sub>PO<sub>4</sub>, write an equation for the reaction between nitric acid and phosphorus(V) oxide. 62 + 1NO2 + P2O5 -> 2H3PO4 +3N2O ...[2] In the solid state  $N_2O_5$  is an ionic compound. Given that  $N_2O_5$  is sometimes known as 'nitronium nitrate' write the ionic formula representation of  $N_2O_5$ . (iii) NO2 NO3 [1] [Total: 11]

## **Examiner Comment**

Although the overall performance on this question was excellent the candidate made a serious error in the basic chemical knowledge required for (b) (i).

#### Example Candidate Response – Merit

- 7 Nitrogen forms a variety of oxides and halides.
  - (a) Nitrogen triiodide, NI<sub>3</sub>, is an explosive that detonates with a snap even when only touched lightly. Given that the electronegativity value for nitrogen is 3.07 and for iodine is 2.36, indicate below the dipole in an N–I bond.

(b) (i) Nitrogen trifluoride, NF<sub>3</sub>, can be prepared by reacting ammonia with fluorine. In this reaction the fluorine oxidises the nitrogen in ammonia while the oxidation number of hydrogen is unchanged.

Give the equation for this reaction.

NH3 +3F2 -> NF3 + 3HF

(ii) Nitrogen trifluoride is used to etch silicon in microelectronics. It is decomposed to its elements and the fluorine is used to attack the silicon.

Give the equation for the decomposition of nitrogen trifluoride.

 $2NF_3 \longrightarrow N_2 + 3F_2$ 

(iii) Nitrogen trifluoride is a molecule that has attracted controversy recently for its possible potent contribution to the greenhouse effect. Draw the dot-cross diagram of this molecule; only include outer electrons. State the shape and the bond angle.

109.5-2.5-107

tetrahedral shar

shape . bond angle .....10

(iv) Whereas nitrogen trifluoride is reasonably easy to handle, nitrogen trichloride is an extremely dangerous explosive. Suggest why nitrogen trifluoride is more stable than the other nitrogen trihalides.

F-N bond will be very short and stron none any required to break it making [1] more stable

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[3]

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## Cambridge Pre-U Example Candidate Responses

(c)  $N_2O_5$  is a less well-known oxide of nitrogen. N2O5 is the anhydride of nitric acid, which means that it reacts with water to produce (i) the acid. Write an equation for N2O5 reacting with water. N205+H200 11172HN03 cars ..[1] N2O5 can be made by reacting nitric acid with a dehydrating agent such as (ii) phosphorus(V) exide. Bearing in mind that phosphorus(V) exide is the anhydride of phosphoric acid, H<sub>3</sub>PO<sub>4</sub>, write an equation for the reaction between nitric acid and phosphorus(V) oxide. +HNO3 -> 104 + 1NO2 .[2] In the solid state N2O5 is an ionic compound. Given that N2O5 is sometimes known (iii) as 'nitronium nitrate' write the ionic formula representation of NoO5. NON04 .....[1] NO4 SO4 [Total: 11]

## **Examiner Comment**

The candidate showed an understanding of electronegativity values and correctly predicted the dipole in an N–I bond. In (b) (i), (b) (ii) and (c) the candidate was able to write the balanced equations using the information given but was not able to write the more difficult equation in (c) (ii). Part (b) (iii) was targeted at the lower grades but the candidate only scored 2/3 marks. The candidate showed an understanding of bond lengths/strengths in a covalent compound but was unable to use the information in (c) (iii) to write an ionic formula, the more accessible and widely scored mark.

39

# Question 8 Mark Scheme

(a)	cis	(or Z)	[1]
(b)	acc	deca-6,9,12,15-tetraenoic acid cept tetrenoic instead of tetraenoic cept without the hyphens or with hyphens instead of commas.	[1]
(c)	2 <sup>5</sup> =	= 32 geometric isomers	[1]
(d)	inst	antaneous dipole – induced dipole forces	[1]
(e)		<u>3rd from last</u> carbon atom from the COOH functional group is the final one C=C double bond (owtte)	[1]
(f)	acc 1,2	CH <sub>2</sub> CHBrCH <sub>3</sub> (1) sept any unambiguous structure that is correct. -dibromopropane (1) o't penalise errors with commas, spaces or hyphens	[2]
(g)	(i) (ii)	amount of C=C bonds in 100 g = 6 × 100 g / 328 g mol <sup>-1</sup> = 1.83 mol (1) mass of iodine required = 1.83 mol × 254 g mol <sup>-1</sup> = 465 g (1) ecf for second mark if correct calculation with wrong number of C=C bonds. 1 mark lost for not quoting final answer to 3 or 4 sig. figs. no penalties for missing units oleic acid C <sub>18</sub> H <sub>34</sub> O <sub>2</sub>	[2] [1]
(h)	(i)	the I–C <i>l</i> bond has a (permanent) dipole	[1]
	(ii)	use of cervonic acid in calculation (as it reacts with the most IC <i>l</i> ) (1) vol of 25% excess IC <i>l</i> = $1.25 \times (6 \times 0.100 \text{ g} / 328 \text{ g mol}^{-1}) / 0.100 \text{ mol dm}^{-3} = 22.9 \text{ cm}^3$ (1) give this mark if a correct calculation has been performed with another fatty acid. excess IC <i>l</i> to be used = $25.0 \text{ cm}^3$ (1) ecf for rounding up the volume of IC <i>l</i> to a pipette size (10 cm <sup>3</sup> , 20 cm <sup>3</sup> or 50 cm <sup>3</sup> ) transferred using a pipette (1)	[4]

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- (iii) transfer sample directly to stoppered flask (1) 250 cm<sup>3</sup> flask (1) transfer using 1,1,1-trichloroethane washings (1) leave for 30 minutes after adding Wijs' reagent (1) measuring cylinder used for adding KI (1) measuring cylinder used for adding water (1) Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> dispensed from a burette or use of the word "titrate" or "titration" (1) starch indicator (1) blue-black to colourless (1) white tile (1) shaking of (stoppered) flask after Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> additions ("swirling" not enough) (1) working with 1,1,1-trichloroethane in a fume cupboard (1) [max 8]
- (iv) vol Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>(aq) × 0.100 mol dm<sup>-3</sup> = amount Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>(aq) (1) amount  $Na_2S_2O_3(aq) = 2 \times amount ICl (excess) (1)$ amount IC1 (reacted) = amount IC1 (initial) - amount IC1 (excess) (1) compare this amount IC1 (reacted) with n(C=C bonds) × 0.100 g / molar mass for each fatty acid (in order to identify the fatty acid, which will be the one that most closely resembles the experimental value.) (1) loss of 1 mark for confusing I2 with ICl. [4]

[Total: 27]

#### Example Candidate Response – Distinction (D1)

8 Evidence is accumulating that omega-3 oils help to protect us from schizophrenia and depression, and even improve learning and memory. Omega-3 oils are glyceryl esters of omega-3 fatty acids.

One omega-3 fatty acid is cervonic acid.

HO cervonic acid (a) Which configuration of the C=C double bonds is present in this molecule? cis ..[1] The systematic name of cervonic acid (ignoring geometric isomerism) is docosa-4,7,10,13,16,19-hexaenoic acid. "Docosa" indicates that there is a 22-carbon chain. The numbers indicate the carbon atom where the C=C double bonds start, counting from the carboxylic acid (-COOH) functional group. "Hexaen" indicates that there are six double bonds in the chain. Two other omega-3 fatty acids are stearidonic acid and timnodonic acid. stearidonic acid HO timnodonic acid "Octadeca" indicates that there is an 18-carbon chain. Write down the systematic name (b) for stearidonic acid. Ignore the geometric isomerism. butaenoic WA provenencie [1] (c) How many geometric isomers are there of timnodonic acid, including the molecule shown? 32 [1] (d) What is the name of the intermolecular force that will operate between the hydrocarbon chains? Ring the correct option. [1] permanent dipole-permanent dipole hydrogen bonding instantaneous dipole-induced dipole

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(e) Another fatty acid that is believed to have beneficial health effects is oleic acid.

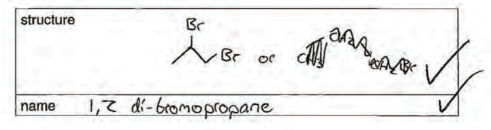
HO 
$$H_2$$
  $H_2$   $H_2$   $H_3$   $H_4$   $H_2$   $H_2$   $H_3$   
HO  $H_2$   $H_2$   $H_3$   $H_3$ 

"Omega" is the **last** letter of the Greek alphabet. By comparing oleic acid, which is **not** an omega-3 acid, with the omega-3 fatty acids in part **(a)**, suggest what "omega-3" relates to in the structure of these oils and fatty acids.

that there is a double bond 3 carbons awa Sion the end of the hydrocarbon chain ......[1]

(f) It is possible to differentiate between different types of fatty acids and oils in the laboratory by measuring the degree of unsaturation in the hydrocarbon chains. Halogens react with alkenes in an addition reaction.

Draw the structure of the product when propene reacts with Br<sub>2</sub> and name it systematically.



(g) Table 8.1 shows information about various fatty acids.

#### Table 8.1

fatty acid	molecular formula	molar mass /g mol <sup>-1</sup>	number of C=C double bonds
cervonic acid	C22H32O2	328	6
stearidonic acid	C18H28O2	, 276	4
timnodonic acid	C20H30O2	302	5
oleic acid	C18H302 1	282	1

(is not legible)

The degree of unsaturation in a fatty acid is commonly expressed by the mass of iodine that reacts with 100.0g of the acid. Use the information in Table 8.1 to calculate the mass of iodine that would react with 100.0g of cervonic acid. Use an appropriate number of significant figures in your answer.

Reacts will  $G J_z = 6 \times 0.30 \pm 9 = x \div (127 \times 2)$ Mol acid =  $\frac{100}{328} = 0.3059 \text{ mol}$ 

......[2]

464.6 g (45F

[2]

- (ii) Write the molecular formula of oleic acid in Table 8.1.
- (h) The interhalogen compound ICl also reacts with alkenes in an addition reaction. It is commonly used as Wijs' reagent, a 0.100 moldm<sup>-3</sup> solution of ICl in glacial ethanoic acid. ICl reacts faster with alkenes than the pure halogens and so can be used to determine volumetrically the unsaturation in fatty acids and oils.
  - Suggest why ICI reacts with alkenes faster than the pure halogens, Cl2, Br2 and (i) I,

[1] casily undergo dedrophilic Radition For the volumetric determination, 30 minutes is required for the reaction between the fatty acid and Wijs' reagent to go to completion. A common solvent for Wijs' reagent and the fatty acids is 1,1,1-trichloroethane. You are to design an experiment using Wijs' reagent to identify an unknown fatty acid that is one of cervonic acid, stearidonic acid or timnodonic acid. You have 0.100 g of the unknown acid dissolved in 20 cm<sup>3</sup> of 1,1,1-trichloroethane. You should aim to produce data that is as accurate as possible using conventional laboratory methods. You should use the information in Table 8.1. The first part of the experiment involves adding an accurately known excess of (ii) Wijs' reagent to the fatty acid solution. The amount of Wijs' reagent should be at

least 25% greater than the maximum amount that could be required to react with the fatty acid.

Work out a suitable volume of Wijs' reagent to add to the fatty acid solution and explain how you would add this quantity in the laboratory. Cervonic, acid has most double bends, , and Slio reade = CONC X VOIU mo Volume = 0.0183 = 18.3 cm ZSCM as way 18 most accurate measure

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[1]

After this reaction is complete,  $10 \text{ cm}^3$  of a 2.0 moldm<sup>-3</sup> aqueous solution of KI is added to the non-aqueous fatty acid solution to convert the excess IC*l* to iodine, I<sub>2</sub>.

$$ICl + KI \rightarrow l_{2} + KCl$$

This quantity of KI(aq) should be a large excess. Next, 100 cm<sup>3</sup> of deionised water is added to the mixture. The iodine is distributed between the aqueous and non-aqueous phases.

(iii) The second part of the experiment involves determining the amount of iodine generated by the excess IC*l*. This is done using a  $0.100 \text{ mol dm}^{-3}$  solution of sodium thiosulfate, Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>.

$$l_2 + 2Na_2S_2O_3 \rightarrow 2NaI + Na_2S_4O_6$$

The whole mixture is reacted with sodium thiosulfate in one batch since the iodine is distributed between the phases. In this reaction considerable shaking is needed to ensure adequate mixing of the phases.

Write out a method for both parts of the experiment that a fellow student could follow. The original fatty acid solution comes in a small sample bottle that is nearly full. Include safety considerations.

\* 025

in a MAD cm<sup>3</sup> Wardhear conical f

is a slight chance that some of the reactants there that the buna escape is securely Sule Man r wear googles dace when 10 iritants or escape .mau outernise har wash the shin CACCESS large some 8 a charoic acid 8 broell have any naked mable flames in not laborator ......[8] Explain how you would work out the identity of the fatty acid given the volume of (iv) thiosulfate obtained from the experiment. (m3) + 1000 = mol + conjection Volume insulphate. thiosulphate used using above og ". I Sind mol iomotion - we know that & this number of Because of mol and there is the same. been has TO as I, From the moles used of Wijs' solution" parti Sent ace lest over of up by thiosulphate. the. This I, used anount as with IC which in moles of Using the macled acio of acid in data the theoretical in table 8.1 calculate amount used Now compare of each acid the mol acid mol will ratio of this the acid Ka and 1:6 is the nol acid : ICI shows 6 double mal bonds are plesem acid has 6 double bonds \* then this is the one which was tested 25 [Total: 27]

\* initially

\* eserto table 8.1

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# **Examiner Comment**

This good Distinction candidate has only dropped 2 marks. Writing 'butaenoic' in part (b) was a common error even at Distinction level. Part (h) (iv) was targeted at the most able candidates and gaining 3 out of the 4 marks here was a good achievement.

#### Example Candidate Response – Distinction

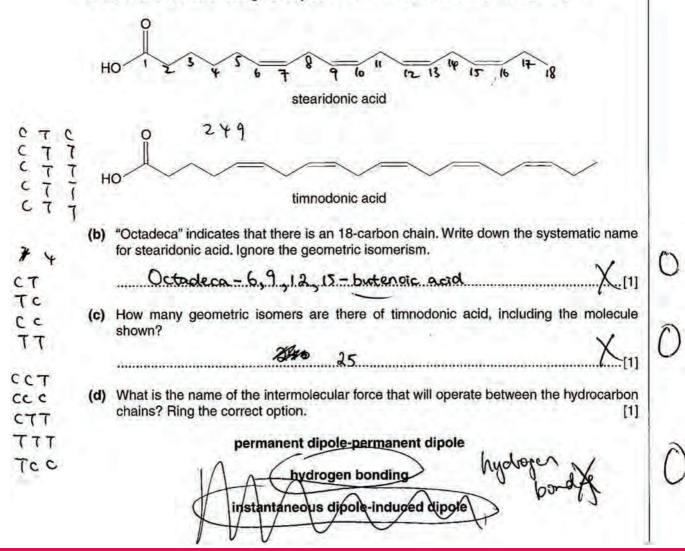
8 Evidence is accumulating that omega-3 oils help to protect us from schizophrenia and depression, and even improve learning and memory. Omega-3 oils are glyceryl esters of omega-3 fatty acids.

One omega-3 fatty acid is cervonic acid.

#### cervonic acid

(a) Which configuration of the C=C double bonds is present in this molecule?

The systematic name of cervonic acid (ignoring geometric isomerism) is docosa-4,7,10,13,16,19-hexaenoic acid. "Docosa" indicates that there is a 22-carbon chain. The numbers indicate the carbon atom where the C=C double bonds start, counting from the carboxylic acid (-COOH) functional group. "Hexaen" indicates that there are six double bonds in the chain. Two other omega-3 fatty acids are stearidonic acid and timnodonic acid.



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Y

(e) Another fatty acid that is believed to have beneficial health effects is oleic acid.

$$f_{10} + f_{11} + f_{12} + f_{12} + f_{13} + f$$

(i) The degree of unsaturation in a fatty acid is commonly expressed by the mass of iodine that reacts with 100.0g of the acid. Use the information in Table 8.1 to calculate the mass of iodine that would react with 100.0g of cervonic acid. Use an appropriate number of significant figures in your answer.

Cervonic acid + 6 Iz -> N= 0.305 ×6 =1.829 Mass = 1 (00) 1.829×127×2 (to Idp [Turn over = 0.305

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- (ii) Write the molecular formula of oleic acid in Table 8.1.
- (h) The interhalogen compound ICl also reacts with alkenes in an addition reaction. It is commonly used as Wijs' reagent, a 0.100 mol dm-3 solution of ICl in glacial ethanoic acid. ICl reacts faster with alkenes than the pure halogens and so can be used to determine volumetrically the unsaturation in fatty acids and oils.
  - (i) Suggest why ICI reacts with alkenes faster than the pure halogens, Cl<sub>2</sub>, Br<sub>2</sub> and 12.

ICL is polar, with de CL ians pure halogen has no dipole moments and are non-polor. Jo of in ICE can attack alkene Ti-bond

agier to break it, actip as exclusiophility & aget faster. [1]

For the volumetric determination, 30 minutes is required for the reaction between the fatty acid and Wijs' reagent to go to completion. A common solvent for Wijs' reagent and the fatty acids is 1,1,1-trichloroethane.

You are to design an experiment using Wijs' reagent to identify an unknown fatty acid that is one of cervonic acid, stearidonic acid or timnodonic acid. You have 0.100 g of the unknown acid dissolved in 20 cm3 of 1,1,1-trichloroethane.

You should aim to produce data that is as accurate as possible using conventional laboratory methods.

You should use the information in Table 8.1.

(ii) The first part of the experiment involves adding an accurately known excess of Wijs' reagent to the fatty acid solution. The amount of Wijs' reagent should be at least 25% greater than the maximum amount that could be required to react with the fatty acid.

Work out a suitable volume of Wijs' reagent to add to the fatty acid solution and explain how you would add this quantity in the laboratory.

Cervonic and has the most number of add vonte acto ervonic acid) 65×103 Cract Vol of ICL + 0.0365 dr 1+25 5 cm3 x ( to 3 s. f.) [4]

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[1]

After this reaction is complete, 10 cm<sup>3</sup> of a 2.0 moldm<sup>-3</sup> aqueous solution of KI is added to the non-aqueous fatty acid solution to convert the excess IC*l* to iodine, I<sub>2</sub>.

$$ICl + KI \rightarrow I_2 + KCl$$

This quantity of KI(aq) should be a large excess. Next, 100 cm<sup>3</sup> of deionised water is added to the mixture. The iodine is distributed between the aqueous and non-aqueous phases.

(iii) The second part of the experiment involves determining the amount of iodine generated by the excess IC*l*. This is done using a 0.100 mol dm<sup>-3</sup> solution of sodium thiosulfate,  $Na_2S_2O_3$ .

$$I_2 + 2Na_2S_2O_3 \rightarrow 2NaI + Na_2S_4O_6$$

The whole mixture is reacted with sodium thiosulfate in <u>one batch</u> since the iodine is distributed between the phases. In this reaction considerable shaking is needed to ensure adequate mixing of the phases.

Write out a method for both parts of the experiment that a fellow student could follow. The original fatty acid solution comes in a small sample bottle that is nearly full. Include safety considerations.

(read coeffly ad record The inital volum of Nasce 03 on piret (2) Holding I drop of Norsen at a time top into mature Stop when the most colour is pute yellow B) Usif a dropper ( graduate and t cm3) add 2 and of starch solution into motive, it should non (4) contrail ripaty intil mixture to (5) record fal volum on purette and colculate 0 - Titre (6) Repeat expressed until a constrant reading of time whin ± 0.1 cm (iv) Explain how you would work out the identity of the fatty acid given the volume of thiosulfate obtained from the experiment. - throsulfate - x cm3 forming voluce of mile = 0. molo . 1000 = 1×104 hole 1=2 n(Iz) = 5x10 x mole n (eness ICe) = 5x 105 x n (ICl wed to react with fatty .bbx1.0 let fatty and Mr be y nole offalt Cervente actol used is 12x Stearidonic acred ..... timpolor acid. l'ine in At Sacid and see which is with T with be - 72 en ratio . [Total: 2]

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# **Examiner Comment**

The candidate performed well in part (h) (iv) suggesting how a relevant but unfamiliar calculation could be carried out but was unable to do the more accessible calculation in part (h) (ii). An incorrect answer was given to part (d) which would not have been expected at this level.

# Paper 3 Part B Written

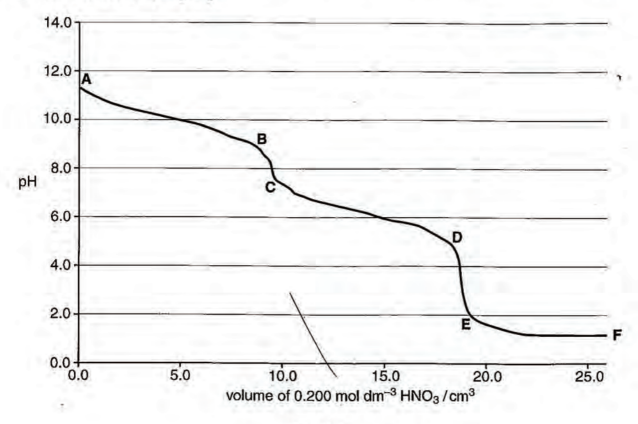
# Question 1 Mark Scheme

Question Number	Expected Answer	Max Marks	Rationale	
1 (a) (i)	(A and B) $H^* + CO_3^2 \rightarrow HCO_3^-$	4	1/2 for overall equation	
(ii)	(C and D) $HCO_3^* + H^* \rightarrow H_2O + CO_2$	1	1/2 for both non-ionic	
(b) (i)	methyl yellow	(1)	La state at a set of the set	
(iii)	pKa of indicator matches pH change at equivalence	4	allow pKa on vertical section of graph owtte	
(iii)	from Yellow to Red	1		
(c)	$\begin{array}{l} 18.8 \times 0.200 \ / \ 1000 = 3.76 \times 10^3 \ \text{mol of HNO}_3 \\ \text{so amount Na}_2\text{CO}_3 = 3.76 \times 10^3 \ / \ 2 = 1.88 \times 10^3 \ \text{mol so conc} = 1.88 \times 10^3 \times 1000 \ / \ 20 = 0.094 \ \text{mol dm}^3 \\ = 0.094 \times 106 = 9.964 \ \text{g dm}^3 \end{array}$	111	allow 9.96 do not allow 10/10.0 ecf ecf	
(d) (i)	$H_2O \rightleftharpoons H^* + OH^* / 2H_2O \rightleftharpoons H_3O^* + OH^*$	1		
(ii)	$\mathcal{K}_c = [H^+][OH]/[H_2O]$ [H <sub>2</sub> O] const negligible dissociation / equilibrium left	1	allow water is in large excess	
(111)	$ [H^*] = \sqrt{51.3 \times 10^{-14}} = 7.16 \times 10^{-7} \text{ (mol dm}^{-3}\text{)}  pH = -\log 7.16 \times 10^{-7} = 6.15 $	1 1+1	allow 6.14	
(iv)	(endothermic) [H <sup>+</sup> ] higher at higher temperature so equilibrium moves right with increased temperature / increased temperature favours endothermic change	1	allow ecf from (d)(iii)	
(e)	$\begin{array}{l} 5 \times 1  /  1000 = 5 \times 10^{-3} \mbox{ mol of H}^{*} \\ 0.1 \times 1.00 \times 10^{-4} = 1 \times 10^{-5} \mbox{ mol of HCl in soln} \\ 1 \times 10^{-5} + 5 \times 10^{-3} \mbox{ in 105 cm}^{3} \\ = 5.01 \times 10^{-3}  /  0.105 = 0.0477 \mbox{ mol dm}^{-3} \\ pH = -log \ 0.0477 = 1.32 \\ \mbox{ calculation of initial pH} = 4.77 \\ after addition of HCl; \ 0.01 + 5 \times 10^{-3} = 0.015 \mbox{ mol } \\ = 0.015 \times 1000  /  105 = 0.143 \mbox{ mol dm}^{-3} \mbox{ CH}_3 \mbox{ COOH} \\ and \ 0.01 - 5 \times 10^{-3} = 0.0476 \mbox{ mol dm}^{-3} \mbox{ CH}_3 \mbox{ COO} \\ pH = pK_a - log \ [acid]  / \ [salt] \\ = 4.77 - log \ 0.143  / \ 0.0476 = 4.17 \end{array}$	1 1 1 1 1 1 1 1 1	1.32 = 4 marks 4.17 = 4 marks	
		[25]		

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## Example Candidate Response – Distinction

1 Fig. 1.1 shows the pH changes during the addition of 0.200 mol dm<sup>-3</sup> HNO<sub>3</sub> to 20.0 cm<sup>3</sup> of a solution of sodium carbonate, Na<sub>2</sub>CO<sub>3</sub>.





24"+ 0032- -> 002 +420

- (a) Write ionic equations for the reactions which occur in the solution between point A and point B on the graph and between point C and point D on the graph.
  - (i) ionic equation for the reaction occurring between A and B

$$\frac{HMO_3}{D} \stackrel{\frown}{\longrightarrow} \frac{MO_3}{TH^2} \stackrel{\frown}{\longrightarrow}$$

1.4

(ii)

(b) Table 1.1 gives some information about seven different indicators.

indicator	pK <sub>a</sub>	acid form	base form
thymol blue	1.6	yellow	blue
methyl yellow	3.3	red	yellow
chlorophenol red	6.0	yellow	red
bromothymol blue	7.1	yellow	blue
cresol purple	8.3	yellow	purple
thymolphthalein	9.9	colourless	blue
alizarin yellow	11.0	yellow	red

#### Table 1.1

(i) From the information given in Table 1.1 choose the indicator that would be most suitable for determining the end-point occurring between points **D** and **E** on the graph.

methy! yellow /.....[1]

(ii) Explain your choice.

within pka value are the steepest pert of the enve the (neutralisation point) .....[1]

(iii) What colour change will be seen with this indicator at the end-point?

yellow red [1]

(c) Calculate the concentration, in gdm<sup>-3</sup>, of sodium carbonate present in the original solution, given that the end-point between D and E occurs after 18.80 cm<sup>3</sup> of HNO<sub>3</sub> have been added.

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(d) (i) Write an equation for the dissociation of water.

HD = HT + OM

(ii) Use the equation in (d)(i) to write an expression for the equilibrium constant,  $K_c$ , for this reaction. Use this expression to show that  $K_w = [H^+][OH^-]$ . Justify and explain your reasoning.

Me = [H+][OH] X [H+][OH] MW = [H+][OH] X [H+][OH] At water is a much larger spantity compared to Ht & OH - its providered confirm and hence committed [3]

(iii) At 373K the ionic product of water,  $K_w$ , has a value of 51.3 × 10<sup>-14</sup> mol<sup>2</sup> dm<sup>-6</sup>. Use this information to calculate the pH of water at 373K. Give your answer to 3 significant figures.

51.3×10" = [H1] (H+]= 7.162 ... ×10" + pH=-logu [H-] - 6-14494 ...

(iv) At 298 K the pH of water is 7.00. Use this information to state whether the dissociation of water is endothermic or exothermic and explain your answer.

From (111) we can tell that taking temperature decreased the ptt. Equilibrium tries to oppose the change, so if leas [11] is created, the towards reaction is [2] slowed and so it must be exothermice

- (e) Calculate the final pH, at 298 K, after a 5.00 cm<sup>3</sup> portion of 1.00 mol dm<sup>-3</sup> hydrochloric acid is added separately to
  - (i)  $100 \text{ cm}^3$  of a solution of  $1.00 \times 10^{-4} \text{ mol dm}^{-3}$  hydrochloric acid.

Here 
$$\vec{H} + \vec{H} + \vec{$$

(ii)  $100 \text{ cm}^3$  of a solution that contains  $0.100 \text{ mol dm}^{-3}$  ethanoic acid and  $0.100 \text{ mol dm}^{-3}$  sodium ethanoate. ( $K_a$  for ethanoic acid is  $1.70 \times 10^{-5} \text{ mol dm}^{-3}$  at 298 K)

$$\begin{array}{cccc} CH_{3}CODH \longrightarrow CH_{3}COT + H^{4} \\ \hline H^{4} J \overline{C} CH_{3}COT J & H_{4} \\ \hline H^{4} J \overline{C} CH_{3}COT J & H_{4} \\ \hline H^{4} J \overline{C} CH_{3}COT J & H_{4} \\ \hline H^{4} P H_{4} P H_{6} (hrothour control of the minimum of th$$

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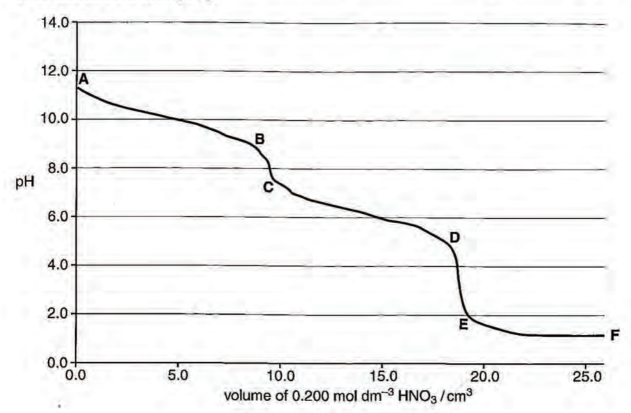
## **Examiner Comment**

The salvage mark for an overall ionic equation was earned in (a) with full marks in (b) and (c). Part (d) was generally answered well but confusion between rate and equilibrium lost marks in part (iv).

As with many candidates, part (e) (i) earned full marks but the buffer calculation in (e) (ii) proved more challenging, although 1 mark was earned here for the initial pH calculation.

## Example Candidate Response – Distinction

1 Fig. 1.1 shows the pH changes during the addition of 0.200 mol dm<sup>-3</sup> HNO<sub>3</sub> to 20.0 cm<sup>3</sup> of a solution of sodium carbonate, Na<sub>2</sub>CO<sub>3</sub>.





NO3 + Nat -> NaNoz

- (a) Write ionic equations for the reactions which occur in the solution between point A and point B on the graph and between point C and point D on the graph.
  - (i) ionic equation for the reaction occurring between **A** and **B**  $2H^{+} + CO_{3}^{2^{-}} - H_{2}O + CO_{2}$ (ii) ionic equation for the reaction occurring between **C** and **D**

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(b) Table 1.1 gives some information about seven different indicators.

indicator	pK <sub>a</sub>	acid form	base form
thymol blue	1.6	yellow	blue
methyl yellow	3.3	red	yellow
chlorophenol red	6.0	yellow	red
bromothymol blue	7.1	yellow	blue
cresol purple	8.3	yellow	purple
thymolphthalein	9.9	colourless	blue
alizarin yellow	11.0	yellow	red

## Table 1.1

(i) From the information given in Table 1.1 choose the indicator that would be most suitable for determining the end-point occurring between points D and E on the graph.

methyl yellow [1]

V

0,

(ii) Explain your choice.

(iii)

	The pka is 3.3 , and the steep relevant steep section
	of the write is between pHs 2 and 4. [1]
1	What colour change will be seen with this indicator at the end-point?
Ś	Ally yellow to red [1]

(c) Calculate the concentration, in gdm<sup>-3</sup>, of sodium carbonate present in the original solution, given that the end-point between D and E occurs after 18.80 cm<sup>3</sup> of HNO<sub>3</sub> have been added.

:. 19.9 (3)+) golm )

Ødm<sup>-3</sup> [3]

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(d) (i) Write an equation for the dissociation of water.

HO # H+ + OH-

(ii) Use the equation in (d)(i) to write an expression for the equilibrium constant,  $K_c$ , for this reaction. Use this expression to show that  $K_w = [H^+][OH^-]$ . Justify and explain your reasoning.

$$K_{c} = \frac{[Hf]_{en} [OH^{-1}_{on}]}{[H_{2}O]_{en}}$$

Res [H20] (2 is 1) since water de barely dissociates at all, Leaning Km= [H+][OH] ......[3]

(iii) At 373K the ionic product of water,  $K_w$ , has a value of 51.3  $\times$  10<sup>-14</sup> mol<sup>2</sup> dm<sup>-6</sup>. Use this information to calculate the pH of water at 373K. Give your answer to 3 significant figures.

Kw = [H+] [OH] ≈ [H+J2 : (H+)= 151.3 x 10 "moliden" = 7,162 x10-7 (4,+) pH= -logio [H+] = 1000 6.14 (3,7)

(iv) At 298 K the pH of water is 7.00. Use this information to state whether the dissociation of water is endothermic or exothermic and explain your answer.

pH lowered at higher temp, therefore forwards reaches favoured 1120 = H++0H as more are more 11 igns present. This means forwards reaction is endo thegenic. [2]

- (e) Calculate the final pH, at 298 K, after a 5.00 cm<sup>3</sup> portion of 1.00 mol dm<sup>-3</sup> hydrochloric acid is added separately to
  - (i)  $100 \text{ cm}^3$  of a solution of  $1.00 \times 10^{-4} \text{ mol dm}^{-3}$  hydrochloric acid.
    - 1 × 10<sup>-5</sup> mol present

0.005 mol HCL added

= 1,32 (30+)

- in 5.01 × 10-3 not HCL present in 0.105 dm3
  - 1. 0.04771 molden<sup>3</sup> of HU (42+) HU fully dissociates ∴ 0.04771 (43+) molden<sup>3</sup> of hydram pH= - Log 10 CH+J

(ii) 100 cm<sup>3</sup> of a solution that contains 0.100 moldm<sup>-3</sup> ethanoic acid and 0.100 mol dm<sup>-3</sup> sodium ethanoate.
 (K<sub>a</sub> for ethanoic acid is 1.70 × 10<sup>-5</sup> moldm<sup>-3</sup> at 298 K)

$$K_{n} = \frac{[H^+][CH_3CH_200]}{[CH_3CO0H]}$$

$$\approx \frac{[H+]^2}{[Cu_3 cooh]} \approx \frac{[H+]^2}{[O.1]}$$

 $1.7 \times 10^{-5} = \frac{CH+3^2}{0.1}$ 

0.005 not HCl udded

$$= 5.130 (4st) \times 10^{-4} \text{ mol} + 0.005 \text{ }$$

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= 1.31 (352) [4]

[Total: 25]

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1 .

## **Examiner Comment**

This question is from a script earning a solid Distinction overall.

As with many candidates part (a) proved tougher than expected and this became a high level discriminator. A mark was allowed here for the overall equation but two separate steps were needed.

Part (b) was correct and the only mark lost in part (c) was for the incorrect ratio. This was careless given the equation in (a) and ecf was allowed from there on.

In part (d) surprisingly few candidates recognised the reason for leaving  $[H_2O]$  out of the  $K_w$  expression and the suggestion here (that its concentration is 1) was often seen. Correct calculations in parts (iii) and (iv) gained full marks.

Most candidates were correctly able to calculate the pH in part (e) (i) but part (e) (ii) was another high level discriminator with many candidates failing to recognise the context of a buffer calculation.

The clarity of exposition in part (d) (iv) gives evidence of solving chemical problems and rationalising unfamiliar material while the clear calculation in part (e) (i) also gives evidence of good problem solving.

# Question 2 Mark Scheme

2 (a)	(i)	high temperature and ethanol as solvent	2	one mark for each solvent and third mark for any appropriate mention of temperature
	(ii)	(lower temperature) and water as solvent	1	
(b)	(i)	molecule with <u>non-superimposable mirror-image</u> form / molecule with <u>4 different groups</u> attached to the <u>same (C)</u> atom / no plane of symmetry	4	do not allow molecules allow atoms in place of groups
	(ii)	eguimolar mixture of two enantiomers / optical isomers	1	
1	(111)	stereoisomers that are not enantiomers / non- superimposable mirror images	,t	
10	(iv)	Br Br H <sub>3</sub> C CH <sub>2</sub> CH <sub>3</sub> H <sup>1000</sup> C CH <sub>2</sub> CH <sub>3</sub> H <sup>300</sup> C CH <sub>2</sub> CH <sub>3</sub>	2	
(c)	(1)	$H_{3C} \xrightarrow{1}_{H_{3C}} \xrightarrow{Br_{1}}_{C''''''H_{1}} \xrightarrow{OH}_{H_{3C}} \xrightarrow{OH}_{H_{3C}} \xrightarrow{OH}_{H_{1} \text{ for inverted}} \xrightarrow{H_{1} \text{ for inverted}}_{\text{structure}}$	4	
12.2	(ii)	S-(+)-butan-2-ol	2	R-(+)- butan-2-ol/S-(-)-butan-2-ol/S-(+)-propan-2-ol all = 1/2

(d) (i)	RX = 1 <sup>st</sup> order: Expt 1 to 2 [RX] ×3 = rate ×3 so directly	1	
(ii)	proportional	1	
(111)	OH = 1 <sup>st</sup> order: Expt 1 to 3 doubling [RX] would double rate to 8.0 × 10 <sup>-4</sup> so doubling again due to doubling [OH] so also directly proportional (2) rate = k[RX][OH] (ecf) (1) $4.0 \times 10^{-4} = k \times 0.050 \times 0.10$ (1) $k = 4.0 \times 10^{-4}/0.005 = 0.08$ (1) dm <sup>3</sup> mol <sup>-1</sup> s <sup>-1</sup> (1)	6	
(iv)	MBr or MBr	9	
		[23]	

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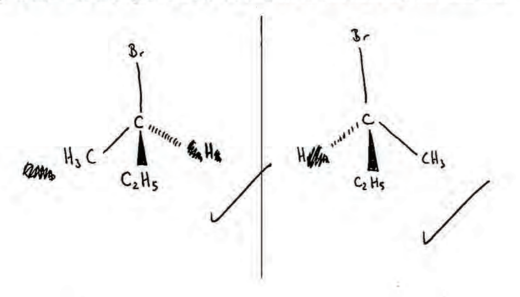
Example Candidate Response – Merit

- 2 2-bromobutane reacts with potassium hydroxide by either elimination or nucleophilic substitution depending on a combination of factors.
  - (a) State the conditions needed to bring about each of these reactions.
    - (1) elimination <u>1</u> atm, 248 k, c. H3, (11) nucleophilic substitution MASA Heat under refluer, com. H2 504 [3]
  - (b) 2-bromobutane is a *chiral* molecule and, when it is prepared by the reaction between but-1-ene and hydrogen bromide, a *racemate* is formed. The enantiomers in the *racemate* can be converted to *diastereoisomers* by covalent derivatisation with suitable *chiral* reagents. Pure samples of each of the enantiomers can then be obtained by simple separation techniques as the *diastereoisomers* have different physical and chemical properties.

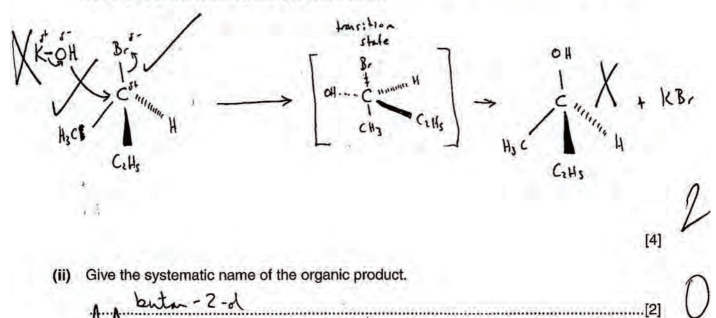
Give definitions of each of the words in italics.

(1) chiral - A central cabon atom is konded to four groups or Long. .....[1] (ii) racemate - A racemic mixture, where there are equal optical isomer present in the colution .....[1] diastereoisomers ....... (iii) containing A \* ext spatial arrangement, curring them to retate the place-planice light i opposite divertions. [1]

(iv) Draw suitable diagrams of the two different enantiomers of 2-bromobutane.



- (c) When R-(-);2-bromobutane undergoes nucleophilic substitution with potassium hydroxide under appropriate conditions the reaction proceeds predominantly by the S<sub>N</sub>2 mechanism. When the progress of the reaction is followed in a polarimeter the optical activity is seen to change gradually from -23.1° via zero to +13.5°.
  - (i) Draw a curly-arrow mechanism for the reaction that is taking place. Show the 3-D structures of the reactant and product clearly.



.....[2]

[2]

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(d) The kinetics of the reaction of a different bromoalkane (RBr) with aqueous alkali were investigated at 323 K. The results are shown in Table 2.1.

experiment	[RBr]/moldm <sup>-3</sup>	[OH <sup>-</sup> ]/moldm <sup>-3</sup>	initial rate/moldm <sup>-3</sup> s <sup>-1</sup>
1	0.05	0.10	4.0 × 10 <sup>-4</sup>
2	0.15	0,10	1.2 × 10 <sup>-3</sup>
3	0.10	0.20	1.6 × 10 <sup>-3</sup>

(i) Deduce the order of reaction with respect to RBr and with respect to the hydroxide ion, OH<sup>-</sup>.

Give reasons for each of your answers. - When the concer RB rate guadron =4 1 = 2 n (X4) TOH concident 2:=4 v =2 .....[4] (ii) Write the rate equation for the reaction. = k [RDr] [OH-] ...[1] ' (iii) Calculate the value of the rate constant, k, at 323K and give its units. Plug is values from esr. 2: 4.0×10-4 = k [0.05] [0.10] K= 16 (iv) Draw the skeletal formula of RBr, which is an isomer of 2-bromobutane.

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[1]

[Total:23]

## **Examiner Comment**

Overall this script fell in the low Merit band.

Many candidates failed to remember the crucial distinction between elimination and nucleophilic substitution of halogenoalkanes being in the choice of solvent and temperature.

The more familiar definitions were remembered in part (b) but, again as with many, the less familiar diastereoisomers caused problems and was confused with enantiomers. It was rare to see such clear 3-D diagrams in part (iv) but 2 marks were earned in this case.

Clear use of curly arrows and unambiguous representation of inversion were the keys in part (c) (i) but there was no inversion shown here and, in part (ii) the clue in the stem failed to stimulate the correct response of a full systematic name with appropriate prefixes.

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# Question 3 Mark Scheme

3 (	(a)	(i)	for CO change is from 2 moles of gas to 1 mole gas and 1 mole solid whereas for SiO change is from 2 moles of gas to 2 moles of solid owtte decrease in <u>disorder</u> / <u>randomness</u> is greater with SiO	1	
		(ii)	2 moles of solid produce 2 moles of solid owtte very little change in disorder	1	
(	(b)	(i)	$\Delta_r S = \Sigma S_{\text{products}} - \Sigma S_{\text{reactants}} = (31.1 + 55.3) - (2 \times 50) \\= -13.4 \text{ (J K}^{-1} \text{ mol}^{-1})$	2	+36.4 = 1/2 (not x2) -29.5 = 1/2 (reversed 55.3 and 55)
		(ii)	$\Delta_r G^{\bullet} = \Delta H - T\Delta S = -126.8 - (298 \times -3.4 / 1000)$ = -122.8 (kJ mol <sup>-1</sup> ) (-22800 J mol <sup>-1</sup> )	2	-118 ecf from 36.4 / -137.6 ecf from -29.5 allow -123 do not allow -122 units not required but must be consistent
(	(c)	(i)	$\begin{split} \Delta_r G^{\Phi} &= -RT ln K_p \text{ so } ln K_p = \Delta_r G^{\Phi} / -RT \\ &= -120.1 \times 10^{-3} / (-8.31 \times 298) = 48.50 \\ \text{ so } K_p &= 1.15 \times 10^{21} \end{split}$	2	3.72 × 10 <sup>21</sup> = 1 / 2 (used 122.8 from <b>3(b)(ii)</b> ) 1.05 = 1 / 2 (no × 10 <sup>3</sup> ignore units
	1	(ii)	$\Delta_r G^{\Phi} = 0 = \Delta H - T\Delta S$ so $\Delta H = T\Delta S$ and $T = \Delta H / \Delta S$ = -172500/-175.9 = 980.7 K	2	allow 981 K
		(iii)	activation energy / kinetic barrier too high	1	
				[13]	

## Example Candidate Response – Distinction

The elements of Group 14 can all form monoxides and dioxides. The stabilities of the monoxides, 3 with respect to disproportionation into the element and the dioxide, vary. The equations for the disproportionation reactions are given in Table 3.1 together with some thermodynamic data for the reactions.

disproportionation equation	∆ <sub>r</sub> S <sup>⊕</sup> (298K) / J K <sup>-1</sup> mol <sup>-1</sup>	Δ <sub>r</sub> H <sup>e</sup> (298K) / kJmol <sup>-1</sup>	Δ <sub>r</sub> G <sup>e</sup> (298K) / kJ mol <sup>-1</sup>
$2CO(g) \rightarrow C(s) + CO_2(g)$	-175.9	-172.5	-120.1
$2SiO(g) \rightarrow Si(s) + SiO_2(s)$	-362.9	-711.5	-603.4
$2\text{GeO}(s) \rightarrow \text{Ge}(s) + \text{GeO}_2(s)$		-126.8	
$2SnO(s) \rightarrow Sn(s) + SnO_2(s)$	-9.200	-9.100	-6.360
$2PbO(s) \rightarrow Pb(s) + PbO_2(s)$	-4.000	+157.2	+158.4

#### Table 3.1

(a) Explain why the entropy change for the disproportionation of

(i) SiO is so much bigger than for CO,

withow goinds . 1970 frent totalend structure, to have a Eight workent are So Sr.O changes from a gas to Silve which it is solved in they are book gases no phase charge.

(ii) PbO is so close to zero.

No phase change and the esthelpy change is the driving force so there is no need for a large entropy for the residien to be ptofensable [2] Also no change in de number of moleoj

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(b) Table 3.2 gives the standard molar entropies for germanium and its oxides.

Table 3.2

name	standard molar entropy at 298 K, S <sup>e</sup> (298 K)/J K <sup>-1</sup> mol <sup>-1</sup>		
germanium, Ge(s)	31.1		
germanium monoxide, GeO(s)	50.0		
germanium dioxide, GeO <sub>2</sub> (s)	55.3		

(i) Calculate the standard entropy change,  $\Delta_r S^{\Theta}(298 \text{ K})$ , for the disproportionation of germanium monoxide.

2Geo + Ge+ Gion 31.3+55-3 - (50.0×2) X

(ii) Calculate the standard free energy change,  $\Delta_r G^{\Theta}(298 \text{ K})$ , for the same reaction.

40/mol 

- (c) Use data from Table 3.1 to calculate
  - (i) the value of the equilibrium constant,  $K_{\rm p}$ , for the disproportionation of carbon monoxide, CO,

1.15×10" Pato 2

 (ii) the temperature above which the disproportionation of carbon monoxide ceases to be favourable.

= -192-5× LOOD - T \$ (-175-9)

(d) Explain why carbon monoxide does not spontaneously disproportionate at room temperature.

Ronatenyoutan 200 Md competion, At Penets the oxy fear and the part to produce (1) penetion to ha a very high accirculan energy [Total: 13] you have to break the triple bound GO

1

..... [2]

#### **Examiner Comment**

A Distinction level response; albeit one that started poorly. Clear references to states of matter and molar quantities were needed with the general advice being to be as specific as possible with references to given data.

A slip in part (b) (i) was the only other source of lost marks in this response with the calculations clear and a recall of the concept of the activation barrier in part (d).

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## Example Candidate Response – Distinction

3 The elements of Group 14 can all form monoxides and dioxides. The stabilities of the monoxides, with respect to disproportionation into the element and the dioxide, vary. The equations for the disproportionation reactions are given in Table 3.1 together with some thermodynamic data for the reactions.

disproportionation equation	∆, <i>S</i> <sup>⊕</sup> (298K) / J K <sup>-1</sup> mol <sup>-1</sup>	Δ <sub>r</sub> H <sup>⊕</sup> (298K) / kJmol <sup>−1</sup>	∆ <sub>r</sub> G <sup>⊕</sup> (298 K) / kJ mol <sup>-1</sup>
$2CO(g) \rightarrow C(s) + CO_2(g)$	-175.9	-172.5	-120.1
$2SiO(g) \rightarrow Si(s) + SiO_2(s)$	-362.9	-711.5	-603.4
$2\text{GeO(s)} \rightarrow \text{Ge(s)} + \text{GeO}_2(s)$		-126.8	
$2SnO(s) \rightarrow Sn(s) + SnO_2(s)$	-9.200	-9.100	-6.360
$2PbO(s) \rightarrow Pb(s) + PbO_2(s)$	-4.000	+157.2	+158.4

## Table 3.1

- (a) Explain why the entropy change for the disproportionation of
  - (i) SiO is so much bigger than for CO,

Stander Sio produces 2 solids whereas (D makes a solid orda gos. Gozes have much higher entropy so in LO the is Uses of a chonge 

(ii) PbO is so close to zero.

id to some mon solids both of which ngend from ropies, so there is not much the difference between hore low en lues as most solide line similiarly entropies [2]

(b) Table 3.2 gives the standard molar entropies for germanium and its oxides.

Table 3.2

name	standard molar entropy at 298 K, S <sup>e</sup> (298 K)/J K <sup>-1</sup> mol <sup>-1</sup>	
germanium, Ge(s)	31.1	
germanium monoxide, GeO(s)	50.0	
germanium dioxide, GeO <sub>2</sub> (s)	55.3	

 Calculate the standard entropy change, Δ<sub>r</sub>S<sup>Θ</sup>(298K), for the disproportionation of germanium monoxide.

26.00 - 6.0, +6e

(31.1+55.3)-(50.0vz) =

F nol [2]

(ii) Calculate the standard free energy change,  $\Delta_r G^{\Theta}(298 \text{ K})$ , for the same reaction.

AG = AH-TAS = -126 800, - (298x-15.6) = -122747.2

-122.7h5mol [2]

- (c) Use data from Table 3.1 to calculate
  - (i) the value of the equilibrium constant,  $K_p$ , for the disproportionation of carbon monoxide, CO, A,  $A = R T l_n k$

1-15 × 1021 [2]

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## Cambridge Pre-U Example Candidate Responses

(ii) the temperature above which the disproportionation of carbon monoxide ceases to be favourable.  $\eta = \eta = \eta$ 

AG= AH-TAS No - - 172500 - TAX-175 Solve for A6=0 AT 177500 = T+175 985.7 :

(d) Explain why carbon monoxide does not spontaneously disproportionate at room temperature.

High activition energy, which the con I be reached at room temperature [1] [Total: 13]

## **Examiner Comment**

Overall an outstanding Distinction level script, which therefore shows what a high level discriminator question **3** (a) proved to be. This candidate only earned 1/4 here as, although the end result was mentioned in part (a) (i), for each reaction there was no mention of the <u>change</u> involved.

Again, the key to success here is a specific, detailed description of the state changes that occur.

A slip in transposing data in part (c) (ii) was the only other source of a lost mark in this question but does illustrate the need for care at all levels.

#### Example Candidate Response – Merit

3 The elements of Group 14 can all form monoxides and dioxides. The stabilities of the monoxides, with respect to disproportionation into the element and the dioxide, vary. The equations for the disproportionation reactions are given in Table 3.1 together with some thermodynamic data for the reactions.

disproportionation equation	∆ <sub>r</sub> S <sup>⊕</sup> (298 K) / J K <sup>−1</sup> mol <sup>−1</sup>	Δ <sub>r</sub> H <sup>e</sup> (298K) / kJmol <sup>-1</sup>	∆ <sub>r</sub> G <sup>⊕</sup> (298K) / kJ mol <sup>−1</sup>
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$2SnO(s) \rightarrow Sn(s) + SnO_2(s)$	-9.200	-9.100	-6.360
$2PbO(s) \rightarrow Pb(s) + PbO_2(s)$	-4.000	+157.2	+158.4

## Table 3.1

(a) Explain why the entropy change for the disproportionation of

SiO is so much bigger than for CO, (i) has a hard giant lattice structure, it SIO Sime takes a lot of energy to disproportionate. .....[2] (ii) PbO is so close to zero. Since Lead has a electronegativity compared with LOW the other Group 14 elements it forms a peak intermolecular the oxygen, thereby making it ensier to 1 with the

(b) Table 3.2 gives the standard molar entropies for germanium and its oxides.

Table 3.2

name	standard molar entropy at 298 K, S <sup>e</sup> (298 K)/J K <sup>-1</sup> mol <sup>-</sup>	
germanium, Ge(s)	31.1	
germanium monoxide, GeO(s)	50.0	
germanium dioxide, GeO <sub>2</sub> (s)	55.3	

(i) Calculate the standard entropy change,  $\Delta_r S^{\Theta}(298 \text{ K})$ , for the disproportionation of germanium monoxide.

$$2 GeO_{151} - 0 Ge(5) + GeO_{151}$$

$$AS = \Delta S products - \Delta S reactants$$

$$= (31 \cdot 1 + 5S \cdot 3) - 2(50)$$

$$= -13 \cdot 6$$

-1 mol -1

(ii) Calculate the standard free energy change,  $\Delta_r G^{\Theta}(298 \text{ K})$ , for the same reaction.

$$\Delta G^{*} = \Delta H^{*} - T\Delta S^{*}$$

$$= -126.8 \text{ kJmol}^{-1} - (2.98 \text{ ksc} - 13.6 \text{ Jk}^{-1} \text{ mol}^{-1})$$

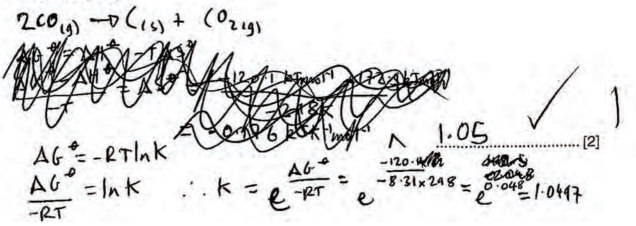
$$= -126.8 \text{ kJmol}^{-1} + 4052.8 \text{ Jmol}^{-1}$$

$$= -122747.2 \text{ kJmol}^{-1}$$

$$= -123 \text{ kJmol}^{-1}$$

(c) Use data from Table 3.1 to calculate

) the value of the equilibrium constant,  $K_p$ , for the disproportionation of carbon monoxide, CO,



the temperature above which the disproportionation of carbon monoxide ceases to be favourable. LEOD (d) Explain why carbon monoxide does not spontaneously disproportionate at room temperature. This is because the value of ArG\* is less than .....[1] 0. 1 [Total: 13] Sec. 1. - 10 Y 2 .

In part (a) there was no evidence of any recall of the key principles governing the sign and magnitude of entropy and the candidate seemed to be confused between entropy and enthalpy.

The calculations in part (b) were done well but, in part (c) (i) the candidate failed to convert  $\Delta_{r}G$  to Joules and, in (c) (ii) defining  $\Delta_{r}G$  as zero and proceeding from there was not seen.

The candidate also failed to recognise the concept of the activation energy barrier in part (d).

Consistent use of appropriate significant figures throughout part (b) provide evidence of a concern for accuracy and precision.

# Question 4 Mark Scheme

4 (a)	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	5	must see link of EFM to RFM
	bond Angle = 90"	1	ecf
(b)	(octahedral) e.g. $Co(H_2O)_8^{2^+}$ angle = 90" (tetrahedral) e.g. $CoCl_4^{2^-}$ angle = 109–110"	3	both angles needed for third mark NB CoCl₄ doesn't exist
(c)	lobes of two orbitals directed along cartesian axes lobes of three orbitals directed between cartesian axes results in split of energy levels of d orbitals promotion of electrons between orbitals <u>absorbs</u> in visible region	1 1 1 1	
1.00		[13]	

#### Example Candidate Response – Distinction

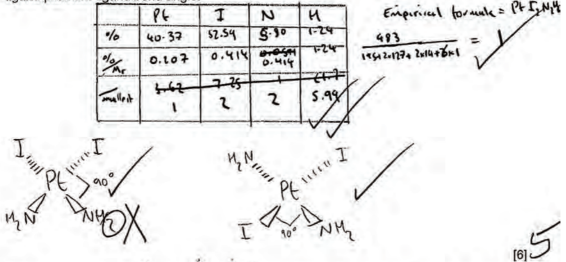
4 The complex compound diamminedichloroplatinum(II) has two isomeric forms, one of which can be prepared from potassium tetrachloroplatinate(II) as shown in the reaction sequence in Fig. 4.1.

 $\mathsf{K}_2[\mathsf{PtCl}_4] \to \mathsf{K}_2[\mathsf{PtI}_4] \to \mathbf{X} \to [\mathsf{Pt}(\mathsf{H}_2\mathsf{O})_2(\mathsf{NH}_3)_2](\mathsf{NO}_3)_2 \to \mathsf{diamminedichloroplatinum(II)}$ 

Fig. 4.1

Compound X has the composition by mass Pt 40.37%, I 52.59%, N 5.80% and H 1.24% and a relative molecular mass of 483.

(a) Calculate the molecular formula of X, draw its two possible structures and indicate the ligand-platinum-ligand bond angle.



(b) Cobalt forms both octahedral and tetrahedral complexes. Give the identities of one complex ion of cobalt with each shape and, in each case, state the ligand-cobalt-ligand bond angle.

[ (o (4)] tetrahedral octahedral [(. (ON),] X bond angle ..... 90 bond angle .... 109. [3]

(c) With reference to the 3-D orientation of d orbitals in an octahedral transition metal complex explain why such complexes are usually coloured.

ligands complexes lie at 90° to each octa hedral lines as the dry dy days dry are y dreg and day subshells be between they are slightly lower energy than coloured 2 gep is such emitted X when the electrons nove [4] Nech the shelle Total: 13]

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This was a good Distinction level script with this response exemplifying the overall standard.

In part (a) the development of the empirical formula to the molecular formula using the relative molecular mass was clearly shown and the only mark lost was for a slip in the structures – possibly the 'squashed up' molecular formula hindered back checking.

Part (b) again highlighted the need for specific recall of some example ions whilst part (c) was again only marred by an explanation couched in terms of emission.

The quality of responses to part (a) and part (c) suggest an understanding of chemical reactions, structure and the use of a model for describing colour of complexes.

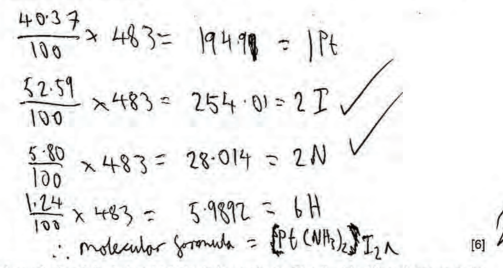
#### Example Candidate Response – Merit

4 The complex compound diamminedichloroplatinum(II) has two isomeric forms, one of which can be prepared from potassium tetrachloroplatinate(II) as shown in the reaction sequence in Fig. 4.1.

 $\mathsf{K}_{2}[\mathsf{PtC}l_{4}] \rightarrow \mathsf{K}_{2}[\mathsf{PtI}_{4}] \rightarrow \mathsf{X} \rightarrow [\mathsf{Pt}(\mathsf{H}_{2}\mathsf{O})_{2}(\mathsf{NH}_{3})_{2}](\mathsf{NO}_{3})_{2} \rightarrow \mathsf{diamminedichloroplatinum(II)}$ 

Compound X has the composition by mass Pt 40.37%, I 52.59%, N 5.80% and H 1.24% and a relative molecular mass of 483.

(a) Calculate the molecular formula of X, draw its two possible structures and indicate the ligand-platinum-ligand bond angle.



(b) Cobalt forms both octahedral and tetrahedral complexes. Give the identities of one complex ion of cobalt with each shape and, in each case, state the ligand-cobalt-ligand bond angle.

octahedral 
$$\left[C_0\left(||_{2}0\right)_{6}\right]^{3+}$$
 tetrahedral  $\left[C_0\left(L_{1+1}\right)_{7}\right]$   
bond angle 90° bond angle 109.5° [3]

(c) With reference to the 3-D orientation of d orbitals in an octahedral transition metal complex explain why such complexes are usually coloured.

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Overall this script was at the level of a high Merit and this question in itself was around the Distinction boundary.

The empirical formula was calculated correctly but the candidate failed to justify a molecular formula or to draw the isomers, which may have been due to not reading the question carefully.

An incorrect charge for  $CoCl_4^{2^{-2}}$  cost a mark in part (b) whilst a generally good account of colour in part (c) was only marred by a reference to emission of coloured light as the promoted electron 'dropped', when what is required is a reference to absorption leaving the complementary colour.

#### Example Candidate Response – Merit

4 The complex compound diamminedichloroplatinum(II) has two isomeric forms, one of which can be prepared from potassium tetrachloroplatinate(II) as shown in the reaction sequence in Fig. 4.1.

 $K_2[PtCl_4] \rightarrow K_2[PtI_4] \rightarrow X \rightarrow [Pt(H_2O)_2(NH_3)_2](NO_3)_2 \rightarrow diamminedichloroplatinum(II)$ 

#### Fig. 4.1

Compound X has the composition by mass Pt 40.37%, I 52.59%, N 5.80% and H 1.24% and a relative molecular mass of 483.

(a) Calculate the molecular formula of X, draw its two possible structures and indicate the ligand-platinum-ligand bond angle.

T P+ н 1-24 molecular formula . PEIZN2H6

(b) Cobalt forms both octahedral and tetrahedral complexes. Give the identities of one complex ion of cobalt with each shape and, in each case, state the ligand-cobalt-ligand bond angle.

Co(044)2 [Co(01+)(120)+] tetrahedral octahedral bond angle ...... 90 [3] ...... (c) With reference to the 3-D orientation of d orbitals in an octahedral transition metal complex explain why such complexes are usually coloured. (\$\frac{1}{2\frac{1}{3\pm 2}/4\pm 2}\$) explain why such complexes are usually coloured. 3 & orbitals have low every orbitals the other 2 ent along the here to there align orbitules or full orbitules ( D ) no colours will be them . If it does have severally of radiation/light can take the lower energy of to the high dow, that every is engled as light every. if comes back this leads to the complexes bety coloured . [4] gluby a colour . [Total: 13]

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[6]

Overall a Merit level script with this response on a par with the overall performance.

Again there was no progression from the empirical formula to the molecular formula shown in part (a) and unfortunately this candidate failed to recognise that  $N_2H_6$  represented two  $NH_3$  groups so was unable to make a sensible suggestion for the structure.

Part (b) illustrates the need for candidates to remember some example material for reference in questions. The concept of split energy levels was remembered in part (c) but then the origin of colour was explained in terms of emission rather than absorption.

#### Example Candidate Response – Pass

4 The complex compound diamminedichloroplatinum(II) has two isomeric forms, one of which can be prepared from potassium tetrachloroplatinate(II) as shown in the reaction sequence in Fig. 4.1.

 $\mathsf{K}_{2}[\mathsf{PtC}l_{4}] \rightarrow \mathsf{K}_{2}[\mathsf{PtI}_{4}] \rightarrow \mathsf{X} \rightarrow [\mathsf{Pt}(\mathsf{H}_{2}\mathsf{O})_{2}(\mathsf{NH}_{3})_{2}](\mathsf{NO}_{3})_{2} \rightarrow \mathsf{diamminedichloroplatinum(II)}$ 

Compound X has the composition by mass Pt 40.37%, 1 52.59%, N 5.80% and H 1.24% and a relative molecular mass of 483.

(a) Calculate the molecular formula of X, draw its two possible structures and indicate the ligand-platinum-ligand bond angle.

(b) Cobalt forms both octahedral and tetrahedral complexes. Give the identities of one complex ion of cobalt with each shape and, in each case, state the ligand-cobalt-ligand bond angle.

octahedral tetrahedral (CO (A20)6 [Cocia] bond angle .... log\* [3] With reference to the 3-D orientation of d orbitals in an octahedral transition metal complex explain why such complexes are usually coloured. being at Artain Luchelengths distance are these gaps which the light passes in whele ofthe an get through the cause all 10 ravel throug distance between orbite below) The Complex to be the some Cobour that savelona [Total: 13]

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Overall this script was just below the Merit borderline and this response exemplified the overall performance.

Specific recall of some platinum complex ion chemistry is expected (syllabus B2.4 (d)) but the candidate represented the complex as tetrahedral. The presence of only two different groups should also have served as a clue that a representation of optical isomerism would be inappropriate here. The other mark lost was for failing to show that the molecular formula was the same as the empirical formula, by reference to the relative molecular mass.

Omission of charges in part (b) was costly and part (c) required the specific recall of a principle that needs committing to memory.

# Question 5 Mark Scheme

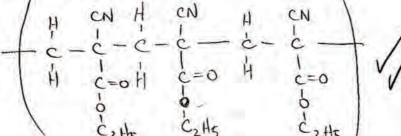
5 (	a)	C <sub>6</sub> H <sub>7</sub> NO <sub>2</sub>	1	
(	b)	<b>x</b> = 120° <b>y</b> = 104.5°	2	allow 104 - 105 for y
(	c)	1 = carboxylic Acid Level     2 = hydrocarbon Level       3 = carboxylic Acid Level	2	three = 2 two = 1
(	d) (i)	donates pair of electrons to form covalent bond	1+1	
	(11)	$\begin{array}{c ccccc} H & CO_2C_2H_5 & H & CO_2C_2H_5 \\ \hline & & & & & \\ H & & & & & \\ -C & & & & C & & \\ -C & & & & C & & \\ -C & & & & C & & \\ -C & & & & & C & \\ \hline & & & & & & \\ H & & & & & H & CN \end{array}$	2	
(	e)	$ \begin{array}{l} \mathbf{Q} = CH_2 = C(CH_2NH_2)CH_2OH \\ \mathbf{R} = CH_3CH_2OH \\ \mathbf{S} = CH_2 = C(CH_2NHOCCH_3)CH_2OCOCH_3 \\ \mathbf{T} = CH_3CH_2OCOCH_3 \\ \mathbf{U} = CH_3CH(CH_2NH_2)CO_2CH_2CH_3 \\ \hline \mathbf{Reactions of } \mathbf{Q} \mbox{ and } \mathbf{R} \mbox{ with ethanoyl chloride:} \\ \hline 2:1 \mbox{ ratio with } \mathbf{Q} \mbox{ as both } NH_2 \mbox{ and OH acylated} / \\ 1:1 \mbox{ ratio with } \mathbf{R} \mbox{ as only one OH group acylated (1)} \\ \hline Reaction \mbox{ with } NH_2 \mbox{ produces (secondary) amide (+HCl) (1)} \\ \hline Reaction \mbox{ with } NH_2 \mbox{ produce ster (+HCl) (1)} \\ \hline \frac{1}{H} \mbox{ NMR of } \mathbf{T}: \\ \mbox{ three signals indicate / due to 3 H environments (1)} \\ \mbox{ triplet } \mathbf{@} \sim 1.3 \mbox{ is protons on } CH_3 \mbox{ adj to } CH_2 \mbox{ (1)} \\ \mbox{ singlet } \mathbf{@} \sim 2.0 \mbox{ is protons on } CH_3 \mbox{ and } \mathbf{CH}_3 \mbox{ (1)} \\ \mbox{ signals near } 20 \mbox{ = Cs in the two } CH_3 \mbox{ groups (1)} \\ \mbox{ signal at } \sim 60 \mbox{ is C in } CH_2 \mbox{ (1)} \\ \mbox{ signals indicate / due to 6 H environments (1)} \\ \mbox{ disappearing signal due to } \mbox{ labile protons on } NH_2 \mbox{ (2)} \\ \end{array}$	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	do not allow 'peptide' <sup>1</sup> H NMR marks are independent of structures drawn except first mark <sup>13</sup> C NMR marks are independent of structures drawn
-	-	disappearing signal due to labile protons on NH <sub>2</sub> (2)	1+1	
			[26]	

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#### Example Candidate Response – Merit

5 Cyanoacrylate is the generic name for cyanoacrylate based fast-acting glues such as ethyl 2-cyanoacrylate. The skeletal formula of ethyl 2-cyanoacrylate is given in Fig. 5.1.

CNC(CH2)COL Fig. 5.1 (a) Give the molecular formula of ethyl 2-cyanoacrylate. CIHyNO3 (b) Give the bond angles labelled x and y. bond angle y ..... [2] (c) Write the names of the functional group levels of the carbon atoms numbered 1, 2, and 3. Carboxylic and level carbonyl level carboxylic acid 3 [2] (d) Ethyl 2-cyanoacrylate rapidly undergoes addition polymerisation in the presence of a suitable nucleophile to form long, strong chains which join the bonded surfaces together. The presence of moisture can therefore cause the glue to set, so exposure to moisture in the air can cause a tube or bottle of glue to become unusable over time. (i) Explain what is meant by the term nucleophile. lone pair dener [2] (ii) Draw the structure of part of the polymer chain that would be formed, showing three repeat units. CN CN



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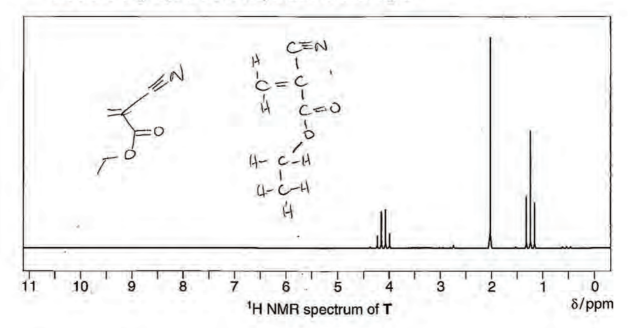
[2]

4

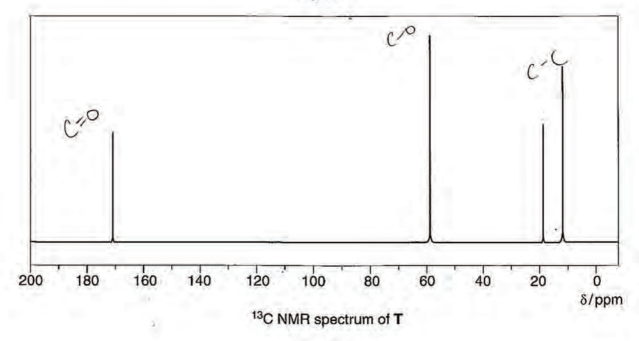
(e) The reduction of ethyl 2-cyanoacrylate with LiAIH<sub>4</sub> in ether produces two compounds, Q and R. Q reacts with ethanoyl chloride in a 1:2 molar ratio to form S. R reacts with ethanoyl chloride in a 1:1 ratio to form T. The <sup>1</sup>H NMR spectrum of T is shown in Fig. 5.2 and the <sup>13</sup>C NMR spectrum of T is shown in Fig. 5.3.

If ethyl 2-cyanoacrylate is instead reduced with hydrogen using a nickel catalyst then there is only a single product, **U**. The <sup>1</sup>H NMR spectrum of **U** has six signals, one of which disappears on shaking with D<sub>2</sub>O.

Draw the structural formula of each of the compounds **Q**, **R**, **S**, **T** and **U**. Explain the reactions and the forms of the spectra in Fig. 5.2 and Fig. 5.3. This should include the identification of the atoms or groups of atoms responsible for each signal.









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Cambridge Pre-U Example Candidate Responses

Structural formulae Q CALCERT 01 0 [5] Explanation Reactions of Q and R with ethonoyi chloride ... These are condensation Practions the pond Inere HCL forming an est elimination of Tunqual 1/eptide Q med .....[3] the signal <sup>1</sup>H NMR of T apple \$ppm Splittings hisr. is due lapelle adjacent as splithings above Carbon has no hydrogens on It. Signal with provous I ppm is dry to protons tabelled @ Tas adjacent has. ...[3] 13C NMR of T. The peak at 170 ppm is due to C=O bond and the one at around 59 ppm is due to the C-O bond The other has peaks below 20 ppm are due C-C bonds. DN ....[3] 14 NMR of U the signal that disappears is due to the an hydrogen inotope: presence Othenvise, the five other signals are due to the different, phonon. en monments, each in a differently [3] in the different) numbered arde in U. CON

Parts (a) – (d) were generally well answered as was the case for most candidates. A Distinction level candidate would have been able to identify all three functional group levels in part (c) and both bond angles in part (b).

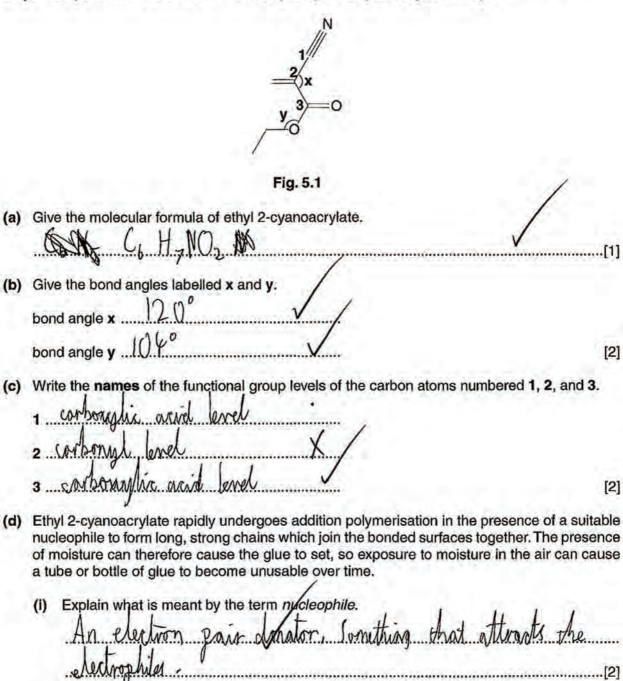
Part (e) was definitely a high level discriminator. Only 2 marks were gained for the structures and they came from ecf from incorrect original structures as the true nature of the reduction process was not recognised. Failing to distinguish between a peptide bond and an amide cost a mark early on in the description.

References to NMR spectra need to be clear and specific and the most often seen problem with references to <sup>13</sup>C spectra was, as here, with candidates who seemed to be referring to bonds as being responsible for the spectrum rather than to carbon atoms/nuclei. This could have been due to confusion with UV spectroscopy or simply poorly phrased answers.

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#### Example Candidate Response – Merit

5 Cyanoacrylate is the generic name for cyanoacrylate based fast-acting glues such as ethyl 2-cyanoacrylate. The skeletal formula of ethyl 2-cyanoacrylate is given in Fig. 5.1.



(ii) Draw the structure of part of the polymer chain that would be formed, showing three repeat units.

inAH [2]

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(e) The reduction of ethyl 2-cyanoacrylate with LiAIH<sub>4</sub> in ether produces two compounds, Q and R. Q reacts with ethanoyl chloride in a 1:2 molar ratio to form S. R reacts with ethanoyl chloride in a 1:1 ratio to form T. The <sup>1</sup>H NMR spectrum of T is shown in Fig. 5.2 and the <sup>13</sup>C NMR spectrum of T is shown in Fig. 5.3.

If ethyl 2-cyanoacrylate is instead reduced with hydrogen using a nickel catalyst then there is only a single product, **U**. The <sup>1</sup>H NMR spectrum of **U** has six signals, one of which disappears on shaking with  $D_2O$ .

Draw the structural formula of each of the compounds **Q**, **R**, **S**, **T** and **U**. Explain the reactions and the forms of the spectra in Fig. 5.2 and Fig. 5.3. This should include the identification of the atoms or groups of atoms responsible for each signal.

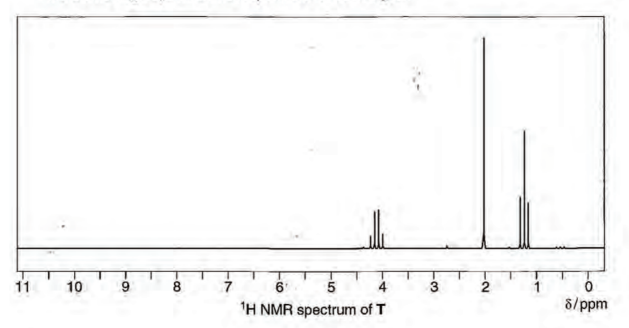


Fig. 5.2

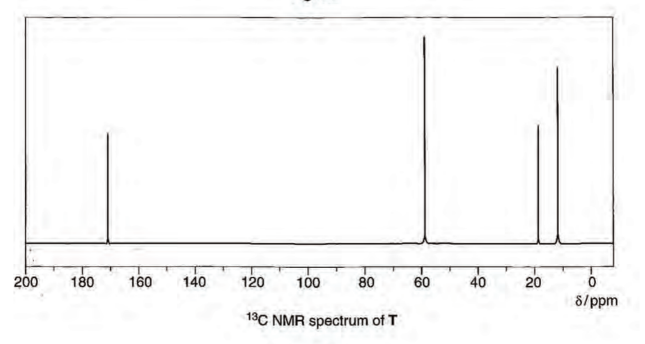


Fig. 5.3

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# Cambridge Pre-U Example Candidate Responses

Structural formulae Q Q ÓH OH 2 0 [5] Explanations (etholast) from as Conormo mathomin of purchasting hilve gre ound throw opens from the varboralism .() [3] Invironmente <sup>1</sup>H NMR Hydrogens on T 1.01 CANDON Ludwooln's on Carbon, 13C NMR of The peaks of group, the peak N h. shawig 12. gronp. 2 lid to a gro he <sup>1</sup>H NMR of **Ú** ..... freedore ... ..... ...... ٨ ...... ...... Total:26

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As with many candidates a reasonable attempt was made at parts (a) - (d) although the clue in the stem of referring to addition polymerisation was overlooked as this candidate attempted some form of condensation polymer in part (d) (ii).

Correct identification of R in part (e) should have led to a correct structure of Q but a mistake was made with regard to the chain length.

The explanations incorrectly referred to an acyl group as a carbonyl group and again lacked specificity when referring to proton NMR although the <sup>13</sup>C NMR was well explained.

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# Paper 4 Practical

## Mark Scheme

Skill	Total marks	Breakdown of marks		Qu. 1	Qu. 2	Qu. 3
Manipulation, measurement and observation	17 marks	Successful collection of data and observations	13 marks	2	8	3
		Quality of measurements or observations	2 marks	2	0	0
		Decisions relating to measurements or observations	2 marks	0	2	0
Presentation of data and observations	6 marks	Recording data and observations	2 marks	2	0	0
		Display of calculations and reasoning	2 marks	2	0	0
		Data layout	2 marks	1	0	1
Analysis, conclusions and evaluation	17 marks	Interpretation of data or observations and identifying sources of error	5 marks	5	0	0
		Drawing conclusions	10 marks	1	6	3
		Suggesting improvements	2 mark	2	0	0

- MMO = manipulation, measurement and observation collection = successful collection of data and observations quality = quality of measurements or observations decisions = decisions relating to measurements or observations
- PDO = presentation of data and observations recording = recording data and observations display = display of calculations and reasoning layout = data layout
- ACE = analysis, conclusions and evaluation interpretation = interpretation of data or observations and identifying sources of error conclusions = drawing conclusions improvements = suggesting improvements

The examiner is to check (and correct if necessary) all subtractions on the Supervisor and candidate scripts where accuracy marks are concerned.

When awarding accuracy marks in Question 1 the candidate's value for the ratio of corrected mass water: corrected mass residue should be listed on the blank page of the Supervisor script or on a sheet of paper securely attached to it.

Where Supervisor results are considered to be suspect (fewer than half the candidates score more than 1 mark), a value derived from the consistent results of the candidates may be used.

## Question 1 Mark Scheme

		Sections	Learning outcomes	Indicative material	Mark
1	(a)	PDO layout	Use the appropriate presentation medium to produce a clear presentation of the data	All balance readings clearly shown in a single table including mass of residue and mass of water lost	[1]
		PDO recording	Record raw readings of a property to the same degree of precision	All values of mass given to the same number of decimal places	[1]
		PDO recording	Use column headings that include both the quantity and the unit and that conform to accepted scientific conventions	All masses are labelled clearly with units. Must use solidus, brackets or describe units fully in words.	[1]
		MMO collection	Follow instructions given in the form of written instructions or diagrams	Final masses within 0.05 g (Award 1 mark if readings within 0.10 g)	[2]
		ACE interpretation	Calculate other quantities from data	Calculates correctly the mass of water and mass of residue (Examiner to note corrected values)	[1]
		MMO quality	Make accurate and consistent measurements and observations	Examiner to check/calculate the ratio of corrected mass water: corrected mass residue. Award 2 marks if difference is within $\pm 0.05$ of Supervisor value Award 1 mark if difference is within $\pm 0.10$ of Supervisor value.	[2]

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(b)	PDO display	Show their working in calculations, and the key steps in their reasoning	Mass of residue/120.4	[1]
	PDO display	Show their working in calculations, and the key steps in their reasoning	Mass of water/18.(0)	[1]
	ACE interpretation	Calculates other quantities from data	Calculates ratio of moles water to moles of magnesium sulfate	[1]
	ACE conclusion	Draw conclusions from interpretations of observations, data and calculated values.	States x as an integer	[1]
(c) (i)	ACE interpretation	Estimate, quantitatively the uncertainty in quantitative measurements	$\pm$ 0.005 g (or 0.0005 g) (Allow $\pm$ 0.01g (or 0.001g) as long as this value is not doubled in <b>(ii)</b> )	[1]
(c) (ii)	ACE interpretation	Express such uncertainty in a measurement as an actual or percentage error	Value from <b>(c)(i)</b> × 2 ANS / mass of water × 100	[1] [1]
(d)	ACE improvement	Suggest modifications to an experimental arrangement that will improve the accuracy of the experiment	Any sensible answer with explanation. e.g. more gentle heating or use of lid during the early stages of heating to reduce loss of solid	[1] [1]

## Example Candidate Response – Distinction

1 Epsom salts occur naturally and are a hydrated form of magnesium sulfate,  $MgSO_4.xH_2O$ . In the following experiment you will determine the value of *x*. Read all of the following instructions before you start any experimental work.

You are provided with the following:

FA 1 hydrated magnesium sulfate, MgSO<sub>4</sub>.xH<sub>2</sub>O

## Method

- 1. Weigh a clean, dry crucible.
- 2. In the crucible place the entire sample of Epsom salts, FA 1.
- 3. Reweigh the crucible.
- 4. Place the crucible in a pipe-clay triangle on top of a tripod.
- Heat the crucible gently for about 1 minute and then more strongly for a further 4 minutes.
- Allow the crucible to cool for about 1 minute and then use a pair of tongs to place the crucible on a heat proof mat.
- Leave the crucible to cool for approximately three minutes, then reweigh the crucible and its contents.
- Repeat the cycle of heating and weighing, as described in steps 4 to 7, until consecutive recorded masses do not differ by more than 0.05 g.
- (a) In a suitable table, record all masses. Calculate the mass of the residue and the mass of the water lost. Record both of these masses in the table.

	mass of crucible + residue/g	mass of residue/g	mass of
initial mass	22.09	0.00	0.00
not heated	23,60	1.2.1	0,00
heated once	22.82	0.73	0.78
keated twice	22.82	0,73	0.78
heated stimes	22.82	0,73 *	0.78"
4th heating	mm	m	m
		·	roho :: 1.07 .06 .06 .05 .00 .01 .8]

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#### Cambridge Pre-U Example Candidate Responses

(b) From the measurements you have made, determine the value of x in the formula MgSO<sub>4</sub>.xH<sub>2</sub>O.

Show your working.  $m(H_{20})_{lose} = 0.78g$ =7  $n(H_{20}) = m(H_{20}) = 0.78 = 0.0433 mol$ m (residue) final = 0.73 g n (residue) final = 0.73 g n (residue) final =  $\frac{m(residue)_{final}}{M_r(residue)_{final}} \propto = \frac{n(H_2G)}{n(residue)} = 7.1470 n_7$   $= \frac{M_r(residue)_{final}}{M_r(residue)_{final}} \propto = \frac{1}{n(residue)} = 7.1470 n_7$   $= \frac{(3.73)}{(10.73)} = \frac{1}{(10.73)} = \frac{1}{(10.73)}$ 14.5+32,1+ (15×4) (c) (i) State the uncertainty in the measurement of each mass in this experiment. uncertainty =  $\pm 0.005$  g[1] Calculate the percentage error in the mass of water that is lost. (ii) Show your working. 0.005 ×100 = 0.841 % 0.641 % [2] (d) Suggest an improvement that a student might make to the experiment and explain why this would lead to the determination of a more accurate value of x. Tepeat the experiment at least once with a new sample of Epison salts. These resertes would hope gutty confirm [2] the value of & to be 7. Ig more accurate scales were used, the masses calculated and [Total: 17] measured would be more accurate and this would lead to a higher level of accuracy when calculating ×.

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- (a) The candidate scored all 8 marks in the opening part although it was not necessary to record the mass of residue and the mass of water lost following each period of heating as was done here.
- **(b)** Again the candidate gained full marks here. Although they do not state the relative formula mass of MgSO<sub>4</sub>, they have clearly shown how the value is obtained and gain the mark.
- (c) The correct uncertainty was given in part (i) but the candidate did not double this value in part (ii) as it was not appreciated that two masses were needed to work out the mass of water lost, these being the combined mass of the crucible and FA 1, along with the combined mass of the crucible and the residue.
- (d) This answer was too vague to merit any credit. The candidate goes someway to suggesting the use of a higher precision balance but this does not merit credit.

## Example Candidate Response – Distinction

1 Epsom salts occur naturally and are a hydrated form of magnesium sulfate,  $MgSO_4.xH_2O$ . In the following experiment you will determine the value of *x*. Read all of the following instructions before you start any experimental work.

You are provided with the following:

FA 1 hydrated magnesium sulfate, MgSO<sub>4</sub>.xH<sub>2</sub>O

## Method

- . Weigh a clean, dry crucible.
- 2. In the crucible place the entire sample of Epsom salts, FA 1.
- . Reweigh the crucible.
  - A, Place the crucible in a pipe-clay triangle on top of a tripod.
- Heat the crucible gently for about 1 minute and then more strongly for a further 4 minutes.
- 6. Allow the crucible to cool for about 1 minute and then use a pair of tongs to place the crucible on a heat proof mat.
- 1. Leave the crucible to cool for approximately three minutes, then reweigh the crucible and its contents.
- /8. Repeat the cycle of heating and weighing, as described in steps 4 to 7, until consecutive recorded masses do not differ by more than 0.05 g.

#### (a) In a suitable table, record all masses.

Calculate the mass of the residue and the mass of the water lost. Record both of these masses in the table.

· · · · · · · · · · · · · · · ·	m	lass/g	
Crucible	2	2.36	
Crucible + hydrate MgSou(betoing))	2	3.88	
Hydrated Moso4		1.52	
Chicible + MgSQy (a)Fer heating)	23.10	23.10	23.09
Anhydrous MgSO4	(	0.73	
Water		0.79	*

Mass of anhydrous MgSO4 = (Crucible + MgSO4 lafter hearing]) - Crucible = 23.099 = 0.739 Man of water = (Man of hydrated MgSO4) - (Mass of anhydraus Mgso4) = 1.529

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[8]

(b) From the measurements you have made, determine the value of x in the formula  $MgSO_4.xH_2O$ .

Show your working.  $n(U_{2}O_{0}) = \frac{m(U_{2}O)}{mr(U_{2}O)} = \frac{0.799}{\sqrt{189}} = \frac{0.439}{n(M_{3}SO_{4})} = \frac{n(U_{2}O)}{n(M_{3}SO_{4})} = \frac{7.238}{mot} = \frac{10.73}{n(M_{3}SO_{4})} = \frac{10.73}{(water ct ct crystallisatus)}$   $n(M_{3}SO_{4}) = \frac{m(M_{3}SO_{4})}{mr(M_{3}SO_{4})} = \frac{0.73}{120.49} = 6.06 \times 10^{-3} \text{mot}$   $M_{3}SO_{4}.7U_{2}O (\text{ho nearest whole number})$ 

(c) (i) State the uncertainty in the measurement of each mass in this experiment.

uncertainty =  $\pm 0.0.5$  g[1]

6.33%

....[4]

(ii) Calculate the percentage error in the mass of water that is lost.

Show your working.

$$\frac{0.05}{0.79} \times 100 = 6.33\%$$

(d) Suggest an improvement that a student might make to the experiment and explain why this would lead to the determination of a more accurate value of x. Allow the Crucible to Cool in a dessicator in between heating so that moisture from the air does not add to the mass of the dehydrated Mg SO4.

[Total: 17]

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- (a) The candidate scored all 8 marks on the opening part. All the results were clearly displayed in a single table including the mass of residue and that of water lost. The data was noted to an appropriate level of precision. The headings were clear and units given. In addition, the final masses lie sufficiently close together and the results are in excellent agreement with those of the Supervisor.
- (b) The second mark for showing the mass of water lost divided by the relative formula mass of water given despite the slip of quoting the answer as 0.439 rather than 0.0439. The ratio was allowed despite the incorrect rounding of the answer which uses the two calculator values for the amount of water and residue rather than the values noted in the working.
- (c) As the candidate was using a balance that read to two decimal places, the answer should have been given as ±0.005 g. Another mark was lost by not doubling the value from part (i) in the following calculation. The candidate did not appreciate that two masses are required to work out the mass of water lost.
- (d) An excellent suggestion for an improvement and furthermore, the candidate gave a clear explanation of why the suggested improvement would lead to a more accurate value of *x*.

#### Example Candidate Response – Merit

1 Epsom salts occur naturally and are a hydrated form of magnesium sulfate,  $MgSO_4.xH_2O$ . In the following experiment you will determine the value of x. Read all of the following instructions before you start any experimental work.

You are provided with the following:

FA 1 hydrated magnesium sulfate, MgSO<sub>4</sub>.xH<sub>2</sub>O

#### Method

- 1. Weigh a clean, dry crucible.
- 2. In the crucible place the entire sample of Epsom salts, FA 1.
- 3. Reweigh the crucible.
- 4. Place the crucible in a pipe-clay triangle on top of a tripod.
- 5. Heat the crucible gently for about 1 minute and then more strongly for a further 4 minutes.
- Allow the crucible to cool for about 1 minute and then use a pair of tongs to place the crucible on a heat proof mat.
- Leave the crucible to cool for approximately three minutes, then reweigh the crucible and its contents.
- 8. Repeat the cycle of heating and weighing, as described in steps 4 to 7, until consecutive recorded masses do not differ by more than 0.05 g.
- (a) In a suitable table, record all masses. Calculate the mass of the residue and the mass of the water lost. Record both of these masses in the table.

	Time/mins	Mass 19
1st heating	0	20,121
2nd heating	9	19.349
3rd heating	18	19.339
4th hear bing	27	19.335

Masslost = 0.786 g water midue = 0.691 ratio = 1.14 5=0.06

[8]

## Cambridge Pre-U Example Candidate Responses

(b) From the measurements you have made, determine the value of x in the formula  $MgSO_4.xH_2O$ .

Show your working.  
I ritial mass of FAI = 1.4779  
Final mass of FAI = 0.6919  
Masslost = 0.7869  
Moles of H<sub>2</sub>O = 
$$\frac{0.786}{18} = 0.0436$$
  
moles of MgSO<sub>4</sub> =  $\frac{0.691}{120.4} = 5.00573$   
(c) (i) State the uncertainty in the measurement of each mass in this experiment.  
uncertainty =  $\pm 0.055$  g[1]

(ii) Calculate the percentage error in the mass of water that is lost.

Show your working. 
$$\pm 0.05g$$
  
1. Mass lost = 0.836g  
Molesof H<sub>2</sub>O =  $0.836$  = 0.0464  
18 = 0.0464  
Mulesof MySO4 =  $0.641$  = 0.00532  
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(d) Suggest an improvement that a student might make to the experiment and explain why this would lead to the determination of a more accurate value of *x*.

[Total: 17]

- (a) The candidate scored both recording marks by recording the mass to the appropriate number of decimal places and by including the correct units in the column headings. The column recording the time was superfluous. Two further marks were awarded for the candidate recording final masses after heating that fell within 0.05 g of one another. The final mark achieved for this section came from the level of agreement between the candidate's ratio of mass of water lost to mass of residue with that of the Supervisor's value. The agreement was not sufficiently close to award the second available mark. The candidate did not record all their results in a single table nor did they calculate the mass of the residue.
- (b) The candidate used their values from part (a) to correctly calculate an integer value of x. They have also clearly shown their working and so gained all 4 marks.
- (c) (i) The candidate may have interpreted this question as asking for the uncertainty in a single recording of mass or the uncertainty in the value of one of the masses. Acceptable answers for this candidate, who used a balance that read to three decimal places, would be either ±0.0005 g or ±0.001 g.
  - (ii) The candidate approached the question incorrectly by working out the maximum and minimum values of x that would arise from their quoted uncertainty in the mass of water lost.
- (d) The candidate's final comment was correct in that a more accurate value of x could be obtained by increasing the mass of FA 1. However, an incorrect answer had already been given in suggesting that accuracy could be improved by heating for shorter periods of time.

#### Example Candidate Response – Merit

1 Epsom salts occur naturally and are a <u>hydrated form of magnesium sulfate</u>,  $MgSO_4.xH_2O$ . In the following experiment you will determine the <u>value of x</u>. Read all of the following instructions before you start any experimental work.

You are provided with the following:

FA 1 hydrated magnesium sulfate, MgSO<sub>4</sub>.xH<sub>2</sub>O

## Method

- 1. Weigh a clean, dry crucible. 16.7489
- 2. In the crucible place the entire sample of Epsom salts, FA 1.
- 3. Reweigh the crucible. 18-237
- 4. Place the crucible in a pipe-clay triangle on top of a tripod.
- Heat the crucible <u>gently</u> for about 1 minute and then more strongly for a (urther 4) minutes.>
- Allow the crucible to <u>cool</u> for about 1 minute and then use a pair of tongs to place the crucible on a heat proof mat.
- Leave the crucible to cool for approximately three minutes, then reweigh the crucible and its contents.
- 8. Repeat the cycle of heating and weighing, as described in steps 4 to 7, until consecutive recorded masses do not differ by more than 0.05 g.
- (a) In a suitable table, record all masses.

Calculate the mass of the residue and the mass of the water lost. Record both of these masses in the table.

	Mass of the	Mass of the crucible	Mass	of the cri	icitle + FAI after heating ig
	crucible only	and FAI hefore heating	lst	2~2	
	16 744	18-237	17-466	17.402	
without the	Mass &	1-493	0.722	0.718	

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[8]

XIII

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(b) From the measurements you have made, determine the value of x in the formula MgSO4.xH2O. Show your working. Mass of MgSOU - XH2O = 18.237-16744 = 1.4939 RMM of My SOY=24.3+ 32-1+ (4×16)= 120-4 Mass of MgSO4 - XH20 after repeat of heating = 17-462-16.744 = 0.7180 Mass change = 1.493 - 0.718 = 0.7779 (420) mole =  $\frac{0}{RHM} = \frac{0.777}{16} = 0.043$  (-) (i) State the uncertainty in the mass uncertainty =  $\pm \Omega \Omega \Omega \Omega$  [1] (ii) Calculate the percentage error in the mass of water that is lost. Show your working. 0.0005 ×100 = 0.0645 %. 0.775 0.0645 %. 20.065 % .....[2] 0.065% (d) Suggest an improvement that a student might make to the experiment and explain why this would lead to the determination of a more accurate value of x. repeat the experiment a few times. By doing so, you will get value of x 3 times If they all round up or down to the same number than [2] 00 you have the right value for X. If they round up or down to different number, something is wrong with your experiment [Total: 17] also, I could if I heated for shorter time I would have got more data and more accurate results. Key From the table, it seems that each writer vapour already evaporated ever before repeating the procedure 4 to 7 to get 2nd data. Heating for shorter time will give me better idea of the mass of MgSQ1. So more can work out more accurate value of x.

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- (a) This candidate did not record the mass of water lost as instructed and so did not gain the first mark. This also meant that they were unable to gain the sixth marking point for correctly calculating the mass of water lost and the mass of residue. The marks for recording all mass readings to an appropriate level of precision and with the correct units were awarded, as were the marks for heating to near constant mass. This candidate also scored both marks for the quality of their data compared to that of the Supervisor.
- (b) The candidate's method was correct but at one stage the candidate rounded the number of moles of magnesium sulfate to just one significant figure which leads to an incorrect ratio using the data from part (a).
- (c) The uncertainty was correctly given in part (i) but the candidate did not appreciate that in order to work out the mass of water lost, two measurements of mass are required and so the uncertainty must be doubled before being used in the calculation in part (ii).
- (d) The suggestion to repeat the experiment was a response given by a number of candidates but did not merit any credit as it does not necessarily lead to a more accurate value of x.

# Question 2 Mark Scheme

2	(a)	MMO collection	Use their apparatus to collect an appropriate quantity of data or observations, including differences in colour, solubility or quantity of materials	Observation for FA 2: white ppt with FA 6 (award only if no reaction recorded with FA 3, FA 4 and FA 5. Ignore slight ppt if recorded with FA 5)	[1]
				ppt is soluble in excess	[1]
				Observations for FA 3: turns orange in FA 4	[1]
				yellow ppt with FA 5	[1]
				bright yellow ppt with <b>FA 6</b> (must have a comparison between the 2 yellow ppts)	[1]
				Observations for FA 4 white ppt with FA 5 white ppt with FA 6 Observation for FA 5:	[1] [1]
-	1 = 1			white ppt with FA 6	[1]
	(b)	ACE conclusion	Draw conclusion from interpretation of observations	H <sup>+</sup> : <b>FA 4</b> CrO <sub>4</sub> <sup>2-</sup> : <b>FA 3</b> SO <sub>4</sub> <sup>2-</sup> : <b>FA 4</b> Pb <sup>2+</sup> : <b>FA 6</b> Ba <sup>2+</sup> : <b>FA 5</b> OH <sup>-</sup> : <b>FA 2</b>	[1] [1] [1] [1] [1] [1]
	(c)	MMO decision	Identify where confirmatory tests are appropriate and the nature of such tests	Heat with hydroxide and aluminium foil and test gas with damp red litmus paper No pale brown gas liberated by dilute acids	[1] [1]

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## Example Candidate Response – Distinction

- FA 2, FA 3, FA 4, FA 5 and FA 6 contain a number of different ions. By making observations when each pair of solutions is mixed it is possible to determine which solution contains which ion. The following ions are present: H<sup>+</sup>, CrO<sub>4</sub><sup>2-</sup>, SO<sub>4</sub><sup>2-</sup>, Pb<sup>2+</sup>, Ba<sup>2+</sup> and OH<sup>-</sup>.
  - (a) Complete the following table by recording your observations on mixing each pair of solutions.

	FA 3	FA 4	FA 5	FA 6
FA 2	Solution stays the same yellow colour. Thue is the slight formation of an oil on shaling	No observation. Solution remains Colourless	white B pit. formed it Dispolves after a far minutes (soluble in excess)	white ppt formed which dissolve d about immediated to give a A Colourbus solution
FA 3		solution turns from the v yellow to v orange in colour	yellow ppt	Bright yellow geldionoui gept formed
FA 4			White gelaterous ppt. formed	white gel atinour plt. former
FA 5				while pot

[8]

## Cambridge Pre-U Example Candidate Responses

on	H+	CrO42-	SO42-	Pb <sup>2+</sup>	Ba <sup>2+</sup>	OH-
solution	FA4	FA 3	FA 4	FA 6	FA 5	FA\$ 2

(b) From your observations, identify which of the solutions contain the following ions.

(c) FA 6 is an aqueous solution of a nitrate. Explain how you would confirm the presence of the nitrate anion. Do not carry out any experimental work.

tule containing FAG with OH copy and Al and test the grs literated with damp red litmes gape. the litrus paper turns blue the solution ......[2] [Total: 16]

## **Examiner Comment**

- (a) Overall this is a very good answer with clear recordings of all observations and sound use of the appropriate technical terms. The only mark not to be awarded was the one for noting that the white precipitate formed by mixing FA 2 with FA 6 was soluble in an excess of FA 2. It was apparent from this candidate's answer that they added too much FA 2 in the first place so their initial white precipitate dissolved almost at once.
- (b) All the ions were correctly assigned.
- (c) As occurred on many of the scripts only the first mark could be awarded here. The second mark in this part relied on candidates realising that both nitrate and nitrite salts evolve ammonia on warming with aluminium and sodium hydroxide solution. To be certain that a nitrate was present it would be necessary to note that no pale brown gas was liberated when the sample was treated with dilute acid.

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## Example Candidate Response – Merit

- 2 FA 2, FA 3, FA 4, FA 5 and FA 6 contain a number of different ions. By making observations when each pair of solutions is mixed it is possible to determine which solution contains which ion. The following ions are present: H<sup>+</sup>, CrO<sub>4</sub><sup>2-</sup>, SO<sub>4</sub><sup>2-</sup>, Pb<sup>2+</sup>, Ba<sup>2+</sup> and OH<sup>-</sup>.
  - (a) Complete the following table by recording your observations on mixing each pair of solutions.

	FA 3	FA 4	FA 5	FA 6
FA 2	Solution in cle pale yellow bon FA3; no Change on moving	Colourless solution; no change on moung.	Pale white precipitate (From colourless) on mining i	Solution is colourless the Change on mining A
FA 3		(Part)	Mill Cream/op- white popt on ming throm Jule yellow)	pit on moren
FA 4			Opaque lite ppt on monthy (from colourless)	Opaque thite 1 pt on motion ( non is lovely
FA 5				Solid white saturate produces on mining X

[8]

(b) From your observations, identify which of the solutions contain the following ions.

SO42-Pb2+ H<sup>+</sup> CrO,2-Ba2+ ion OHsolution [6]

(c) FA 6 is an aqueous solution of a nitrate. Explain how you would confirm the presence of the nitrate anion. Do not carry out any experimental work.

Na OHIag) and Coul so Ulan. in 01 ...[2] tal: 161 th D 4

### **Examiner Comment**

- (a) The candidate observed the correct initial products when combining FA 2 with the other unknown solutions but does not go on to note that the white precipitate formed on mixing FA 2 with FA 6 is soluble in an excess of FA 2. No precipitate results from the mixing of FA 3 and FA 4 as recorded by the candidate and only shades of yellow were acceptable as descriptions for the colour of barium chromate, formed by mixing FA 3 and FA 5. Despite not being awarded the mark for the colour of barium chromate, the use of bright to describe the colour of lead chromate was sufficient to merit the mark for the comparison of the two shades of yellow. Overall the candidate uses slightly confusing terms to describe the observations. For example, it is not necessary to describe a precipitate as being opaque although this did not mean the loss of any marks. In the final part, however, with the mixing of FA 5 and FA 6, the candidate stated that a white saturate was produced which was deemed insufficient to convey the sense of a precipitate.
- (b) All ions were correctly assigned.

(c) The second mark in this part relied on candidates realising that both nitrate and nitrite salts evolve ammonia on warming with aluminium and sodium hydroxide solution. To be certain that a nitrate was present it would be necessary to note that no pale brown gas was liberated when the sample was treated with dilute acid.

### Example Candidate Response – Merit

- 2  $M_2$ ,  $M_3$ ,  $M_4$ , FA 5 and FA 6 contain a number of different ions. By making observations when each pair of solutions is mixed it is possible to determine which solution contains which ion. The following ions are present:  $M_7$ ,  $Cr_{Q_4}^{2-}$ ,  $SO_4^{2-}$ ,  $Pb^{2+}$ ,  $Ba^{2+}$  and  $OH^-$ .
  - (a) Complete the following table by recording your observations on mixing each pair of solutions.

	FA 3	FA 4 B	FA 5	FA 6
Ьа FA 2	The FA3 colur remains on addition effect shight emulsion forma Ne dissaperus;	•	torm while we percipited form while we pot change i	while is formed. while PPt is formed
FA 3		oranga coloured omulsion is formed	lime green forme a. lime green forme a. lime green pet formulio	yellow ppl formed,
FA 4			creamy emulsion formed. uhile pp-1 formed	creangy entirence pormer while percipted formed
FA 5				treany 8 annulsion formel, White percipitate formed. [8

solution FA4 FA36 FA5 FA6 FA2 FA44 X X X X [6] FA 6 is an aqueous solution of a nitrate. Explain how you would confirm the presence of the nitrate anion. Do not carry out any experimental work. NH3 IS Liberated when reached witte with oit A and you can Nitz Hurns damp red litmus blue [2] [Total: 16]	H*	CrO42-	SO42-	Pb <sup>2+</sup>	Ba <sup>2+</sup>	OH-	Use
FA 6 is an aqueous solution of a nitrate. Explain how you would confirm the presence of the nitrate anion. Do not carry out any experimental work. NHZ IS LIB erated when reacted whether with with oit - A and you can Mitz furns damp red litmus blue [2]	FA4	FAS	FAS	FA6	FAZ	FAG	
the nitrate anion. Do not carry out any experimental work. NUZ IS liberated when reached watte with oit - A and you can NItz A Jurns damp red litmus blue [2] C	1		X X	1	X×	//	[6]
the nitrate anion. Do not carry out any experimental work. NUZ IS liberated when reacted watte with oit - A and you can NItz furns damp red litmus blue. [2]							4
with oit - A and you can Nitz Jurns damp red litmus blue. [2] C						irm the prese	nce of
0	to amon b	o not bailing a					
0		가는 것을 안을 했다.		ien re	actec	1. co Arto	
0	3 15 1	1 berat	ed wh				
(9	3 15 1	1 berat	ed wh				
	3 15 1	1 berat	ed wh			21+3	[2]
	3 15 1	1 berat	ed wh			21+3	[2]

identify which of the colutions contain the following

## **Examiner Comment**

- (a) All the marks were awarded for the observations of FA 2 with each of the other unknowns, apart from the solubility of lead hydroxide in an excess of FA 2. For the observation on combining FA 3 and FA 4, although the colour was given correctly, the description of the solution as an emulsion meant that the mark was not awarded. The candidate also described the colour of barium chromate as lime green which is incorrect. The mark for noting the colour of lead chromate was awarded as the candidate has recorded it as being different to the colour of barium chromate.
- (b) The candidate incorrectly assigned the barium ion to FA 2 despite noting correctly that FA 2 did not form a precipitate with FA 3 which was correctly identified as containing the chromate ion. The candidate also incorrectly assigned the sulfate ion to FA 5 despite having correctly assigned FA 4 as containing H<sup>+</sup> which would not have given a white precipitate on mixing with sulfate. The source of these errors lies in the observation of a white precipitate with FA 2 and FA 5 which can occur if the solutions are not pure.
- (c) The candidate failed to pick up either mark here. They forgot to note that aluminium as well as sodium hydroxide solution must be added before warming if ammonia gas is to be liberated. The second and much more challenging mark was for noting the need to distinguish between nitrate and nitrite which the candidate does not do.

## Example Candidate Response – Merit

- 2 FA 2, FA 3, FA 4, FA 5 and FA 6 contain a number of different ions. By making observations when each pair of solutions is mixed it is possible to determine which solution contains which ion. The following ions are present:  $H^+$ ,  $CrO_4^{2-}$ ,  $SO_4^{2-}$ ,  $Pb^{2+}$ ,  $Ba^{2+}$  and  $OH^-$ . Examiner
  - (a) Complete the following table by recording your observations on mixing each pair of solutions.

	FĄ 3 Cr Dy 2-	FA4 ⊬ <sup>+</sup>	FA 5 but?	FA 6 (6"
FA 2	The solution remains yellow no visible change.	No visible change Solu- tion stays colourless.	The solution goes milay. Formation of a white precipitate.	Formation of a white precipitate
, FA 3		The solution changes colour to arange.		A bright yellow precipitate forms.
FA 4	*		A white precipitate forms.	A white precipitate forms.
FA 5				A UMAte milky precipitation which dissolve on when adding excess

[8]

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## Cambridge Pre-U Example Candidate Responses

(b)	From your	observations,	identify w	nich of	the solutions	contain the	following	ions.
-----	-----------	---------------	------------	---------	---------------	-------------	-----------	-------

ion	H+	CrO42-	SO42-	Pb <sup>2+</sup>	Ba <sup>2+</sup>	OH-
solution	FA4	FA3	FA 2	FAG	FA 5	FA5
	V	V	X	V	V	X

(c) FA 6 is an aqueous solution of a nitrate. Explain how you would confirm the presence of the nitrate anion. Do not carry out any experimental work.

I would pour about 2 cm<sup>3</sup> of FA6 into a boiling tube and add about 5 drops of NaOH (aq.) then put a small A foil and heat the tube with a Bursen burner. NHz should [2] come ait which can be tested with a damp red litmus paper. The gas coming off should turn the litmus paper blue. [Total: 16]

## **Examiner Comment**

- (a) Two marks were not awarded in part (a). The candidate did not test the solubility of the precipitate of lead hydroxide, formed by mixing FA 2 and FA 6, in an excess of FA 2 and so did not observe that it was soluble. On mixing FA 5 and FA 6 the candidate records a milky solution rather than a white precipitate or white solid.
- (b) The candidate assigned the hydroxide ion to be present in FA 5 despite noting correctly that FA 4 and FA 5 when mixed give a white precipitate. FA 2 was stated as containing sulfate ions. This error arose from not having noted the solubility of the lead hydroxide in an excess of hydroxide solution which would rule out the white solid as being lead sulfate.
- (c) The candidate correctly describes a positive test for nitrates or nitrites but then does not go on to say how adding dilute acid and looking for the presence of brown fumes can distinguish between them.

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# Question 3 Mark Scheme

(a)	PDO la	ayout	prese	he appropriate ntation mediur ce a clear pres data	n to	Draws up a clear table showing at least 3 observations	[1]
	MMO o	collection	an ap data d includ colou	heir apparatus propriate quar or observations ling differences r, solubility or o terials	ntity of s, s in	Observations for FA 7: colour change to green with acidified dichromate (not green ppt) AND silver mirror / black or dark brown solution or ppt with Tollens'	[1]
<b>F</b>		1 distance	1	Tallanat	1	Observations for FA 8:	[1]
FA 7 butan	al	dichror ✓	nate	Tollens' ✓		colour change to green with acidified dichromate (not green ppt but do not	
FA 8 butan	-1-ol	1		x		penalise again if mentioned in test for FA 7)	[1]
FA 9 butan	one	×		x		AND no reaction with Tollens'	
	_					Observations for FA 9: no reaction with either reagent	
(b)	ACE co	onclusion	And the second	conclusion fro retation of obs		FA 8 is butan-1-ol Must refer to positive test with dichromate and no reaction with Tollens'	[1]
						<b>FA 7</b> is batanal Sufficient to refer only to positive result with Tollens' so long as positive test was observed with dichromate and no other positive tests for Tollens' were observed	[1]
						FA 9 is butanone Sufficient to refer only to negative result with dichromate so long as no reaction with Tollens' was observed and no other negative results are recorded for dichromate. No ecf. Explanation must be based on correct observations	[1]

[Total: 7]

### Example Candidate Response – Distinction

Before starting question 3, half-fill a 250 cm<sup>3</sup> beaker with water and heat it with a Bunsen burner to between 50 °C and 70 °C. You will use this as a hot water-bath in this question. Turn off the Bunsen burner.

3 FA 7, FA 8 and FA 9 each contain a single compound which could be <u>butan-1-ol</u>, <u>butanal or</u> <u>butanone</u>. By reacting each of the compounds first with acidified potassium dichromate(VI) and then with Tollens' reagent you should identify each of the three samples.

Tollens' reagent must be prepared immediately before use.

### Method

### Test with acidified potassium dichromate(VI)

- 1. Into separate test-tubes, to a depth of approximately 1 cm, pour FA 7, FA 8 and FA 9.
- 2. Add approximately 1 cm depth of dilute sulfuric acid to each test-tube.
- 3. Add a few drops of aqueous potassium dichromate(VI).
- 4. If no initial reaction is seen, warm the test-tube in the hot water-bath.

#### Preparation of Tollens' reagent

- 5. Pour aqueous silver nitrate into a boiling tube to a depth of approximately 2 cm.
- 6. Add approximately 0.5 cm depth of aqueous sodium hydroxide.
- Add aqueous ammonia a little at a time with continuous shaking until the brown precipitate just dissolves. Be careful not to add an excess of aqueous ammonia.

### **Test with Tollens' reagent**

- 8. Into separate, clean, dry test-tubes, to a depth of approximately 1 cm, pour FA 7, FA 8 and FA 9.
- 9. Add a few drops of Tollens' reagent.
- 10. If no initial reaction is seen, warm the test-tube in the hot water-bath.

	0	oservations with	
Procedure	FA7	FA8	FAY
Test with ocidified K2Cr2O7	Mixture fums pale yellow/ green slowly Becomes pote blue offer bet - ing in water bath	Mixture starys orange contribut Bloated in water both. It then turns green/blue	Mixture Strays orange even when heated in water bath
Test with Tollers' reagent	Forms black ppt some of which coats fest trace Turns more thing on hearting.	No observable reaction, even when heated in when heated in	No observable reading even when begind In writer both .

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## Cambridge Pre-U Example Candidate Responses

(b) Identify each sample. In each case explain the observations leading to your conclusion. FA ... S... contains butan-1-ol. explanation Because it is oxidise Tokens reagent connot oxidice FA . Z ... contains butanal. it is oxidised quickel explanation Because positie & by dictromate and giresa .[1] FA ..... contains butanone Becare there is no oxide explanation ... or Tollans' reagent so it cannot be but in- 1-of or bu ...[1] [Total: 7]

- (a) The candidate set out the results in a very clear format. It was not necessary to note the colours both before and after warming each solution with acidified dichromate but so doing clearly indicates a student who is observing closely and working carefully through the series of tests. All observations were correct so the candidate gained full marks for this part.
- (b) The identities were given correctly and there was a sound explanation of how each assignment matches the observations made in the opening part.

### Example Candidate Response – Merit

Before starting question 3, half-fill a 250 cm<sup>3</sup> beaker with water and heat it with a Bunsen burner to between 50°C and 70°C. You will use this as a hot water-bath in this question. Turn off the Bunsen burner.

3 FA 7, FA 8 and FA 9 each contain a single compound which could be butan-1-ol, butanal or butanone. By reacting each of the compounds first with acidified potassium dichromate(VI) and then with Tollens' reagent you should identify each of the three samples.

Tollens' reagent must be prepared immediately before use.

### Method

### Test with acidified potassium dichromate(VI)

- 1. Into separate test-tubes, to a depth of approximately 1 cm, pour FA 7, FA 8 and FA 9.
- 2. Add approximately 1 cm depth of dilute sulfuric acid to each test-tube.
- Add a few drops of aqueous potassium dichromate(VI).
- 4. If no initial reaction is seen, warm the test-tube in the hot water-bath.

#### Preparation of Tollens' reagent

- 5. Pour aqueous silver nitrate into a boiling tube to a depth of approximately 2 cm.
- 6. Add approximately 0.5 cm depth of aqueous sodium hydroxide.
- Add aqueous ammonia a little at a time with continuous shaking until the brown precipitate just dissolves. Be careful not to add an excess of aqueous ammonia.

#### **Test with Tollens' reagent**

- Into separate, clean, dry test-tubes, to a depth of approximately 1 cm, pour FA 7, FA 8 and FA 9.
- 9. Add a few drops of Tollens' reagent.
- 10. If no initial reaction is seen, warm the test-tube in the hot water-bath.
- (a) Record all your observations in a single table.

FA	Acidified Potossium Dichromate	Biller's Reagonst	
7	two clear layers to Slightly yellow clear layers. X	gruy deposit	
8	two clear byus turns to durk. green/durkblue sols	N/A .	
9	two clear ligers to Orange . largers .	N/A .	

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(b)	Identify each sample. In each case explain the observations leading to your conclusion.	
	FA	
	explanation alcohols react to form a green/the	1
	Solt. 1	0
	FA	
	explanation because only Aldehydes reach with follows reasont to form the "silve mirror" A [1]	0
	FA	
	explanation because it did not affect either tests	
	and by process of elimination is he engage left. [1]	
		M
		9

- (a) The candidate constructed a clear table for the results and so gained the first mark. Although the observation was correct for FA 7 and Tollen's reagent, it appears that there was no reaction between FA 7 and the acidified dichromate solution. Here the candidate's background knowledge let them down as clearly none of the possible compounds could give such a result. The other observations with acidified dichromate were correct although no detail was given about whether the change happened before or after warming. The use of N/A to indicate no reaction is not recommended when describing what happens on mixing each reagent with Tollen's solution.
- (b) The assignments were correct but credit can only be given when they are supported by the correct observations. Hence, it is insufficient to say that FA 8 is an alcohol simply because it reacted with the acidified dichromate as this would also be true for the aldehyde. The assignment of FA 7 is not supported by the observation of no reaction with acidified dichromate in part (a). Consequently, only the mark for assigning FA 9 to the ketone was awarded.

### Example Candidate Response – Unclassified

Before starting question 3, half-fill a 250 cm<sup>3</sup> beaker with water and heat it with a Bunsen burner to between 50°C and 70°C. You will use this as a hot water-bath in this question. Turn off the Bunsen burner.

3 FA 7, FA 8 and FA 9 each contain a single compound which could be butan-1-ol, butanal or butanone. By reacting each of the compounds first with acidified potassium dichromate(VI) and then with Tollens' reagent you should identify each of the three samples.

Tollens' reagent must be prepared immediately before use.

### Method

### Test with acidified potassium dichromate(VI)

1. Into separate test-tubes, to a depth of approximately 1 cm, pour FA 7, FA 8 and FA 9.

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- 2. Add approximately 1 cm depth of dilute sulfuric acid to each test-tube.
- Add a few drops of aqueous potassium dichromate(VI).
- If no initial reaction is seen, warm the test-tube in the hot water-bath.

#### Preparation of Tollens' reagent

- 5. Pour aqueous silver nitrate into a boiling tube to a depth of approximately 2 cm.
- 6. Add approximately 0.5 cm depth of aqueous sodium hydroxide.
- Add aqueous ammonia a little at a time with continuous shaking until the brown precipitate just dissolves. Be careful not to add an excess of aqueous ammonia.

#### Test with Tollens' reagent

- 8. Into separate, clean, dry test-tubes, to a depth of approximately 1 cm, pour FA 7, FA 8 and FA 9.
- 9. Add a few drops of Tollens' reagent.
- 10. If no initial reaction is seen, warm the test-tube in the hot water-bath.

(a) Record all your observations in a single table. Observation Observabion with Tollens recigent Potassium dichouch () Salution Orange FA7 to a blue light Golou formed Orange solution clear so lution 8 FA remains (no change orany subside stars an orange is produced [4]

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(b) Identify each sample. In each case explain the observations leading to your conclusion. explanation ..... os it showed a por fino sue. Schufie the atallium dichomate showing a primary the t....[1] which a a primary alc contains butanal. FA ... explanation .....le cause A a 20541  $\mathcal{O}$ Tollens reagant (black manetale .....[1] FA ... Q. dontains butanone. explanation . D.c. re results for either tests [1] [Total: 7]

- (a) The first mark was awarded for presenting the results in a single table. The candidate did not gain any of the 3 remaining marks as one of the pair of observations for each unknown was incorrect. The candidate's background knowledge was clearly unsound as the observations for FA 9 could not be true for any of the possible unknowns.
- (b) No marks could be awarded here as in each case the assignment does not match the observations made in the opening section.

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