



Cambridge International Examinations

Cambridge International Advanced Subsidiary and Advanced Level

CANDIDATE NAME					
CENTRE NUMBER			CANDIDATE NUMBER		

CHEMISTRY 9701/41

Paper 4 A Level Structured Questions

October/November 2018

2 hours

Candidates answer on the Question Paper.

Additional Materials: Data Booklet

READ THESE INSTRUCTIONS FIRST

Write your Centre number, candidate number and name on all the work you hand in.

Write in dark blue or black pen.

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

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

DO NOT WRITE IN ANY BARCODES.

Answer **all** questions.

Electronic calculators may be used.

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

A Data Booklet is provided.

At the end of the examination, fasten all your work securely together.

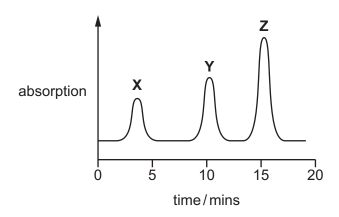
The number of marks is given in brackets [] at the end of each question or part question.



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

1 (a) An aldehyde, an alkane and a carboxylic acid, all of similar volatility, are mixed together. The mixture is then analysed in a gas chromatograph.

The gas chromatogram produced is shown.



The separation of the compounds depends on their relative solubilities in the stationary phase. The stationary phase is a liquid alcohol.

(i) Complete the table to suggest which compound in the mixture is responsible for each peak **X**, **Y** and **Z**. Explain your answer by reference to the intermolecular forces of the compounds.

peak	organic compound	explanation
X		
Υ		
z		

[2]

(ii) A student calculates the areas underneath the three peaks in the chromatogram.

peak	X	Y	Z
area/mm²	19	32	47

The area underneath each peak is proportional to the mass of the respective compound.

Calculate the percentage **by mass** in the original mixture of the compound responsible for peak **Z**.

% of mixture responsible for peak Z =	% of mixture responsible for peak Z =	[1]
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(b) (i) The mass spectrum of a halogenoalkane containing one chlorine atom **or** bromine atom will show an additional peak at M+2.

State the isotopes of chlorine and bromine responsible for M+2 peaks.

chlorine	bromine	
		[1]

(ii) The mass spectrum of bromochloromethane, CH_2BrCl , has a molecular ion peak, M, at an m/e value of 128. It also has M+2 and M+4 peaks.

Suggest the identity of the molecular ions that give rise to these peaks.

M peak				
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M+2 peak

M+4 peak

[2]

(c) Halogenoalkanes can be formed from the reaction of an alkene with a hydrogen halide.

Methylpropene reacts with hydrogen bromide to form 2-bromo-2-methylpropane.

$$H_2C = C + HBr \rightarrow H_3C - C - CH_3$$
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3

methylpropene

2-bromo-2-methylpropane

(i) Draw the mechanism of this reaction. Include all relevant curly arrows, dipoles and charges.

[3]

(ii) 1-bromo-2-methylpropane is also formed in this reaction.

Explain why 2-bromo-2-methylpropane will be the major product in this reaction.

(d) (i)	Explain what is meant by the term <i>partition coefficient</i> , $K_{\text{partition}}$.
	[2]
(ii)	The partition coefficient of organic compound H between dichloromethane and water is 4.75.
	• 2.50 g of compound H was dissolved in water and made up to 100 cm ³ in a volumetric flask.
	• 50 cm³ of this aqueous solution were shaken with 10 cm³ of dichloromethane.
	Calculate the mass of compound H that was extracted into the dichloromethane.
	mass of compound H extracted = g [2]
	[Total: 14]

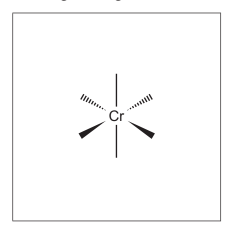
2 (a) E	Ethanedioate ions.	$C_2O_4^{2-}$, are	bidentate ligands.
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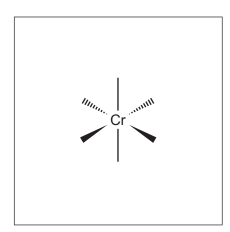
Explain what is r	meant by the	term ligand.
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(b) $Cr^{3+}(aq)$ and $C_2O_4^{\ 2-}(aq)$ ions form the complex ion $[Cr(C_2O_4)_2(H_2O)_2]^{-}$.

Draw two stereoisomers of this complex ion.

You may use \bigcap_{O} to represent $C_2O_4^{2-}$.





[2]

- (c) The solubility of calcium ethanedioate, CaC_2O_4 , is $6.65\times10^{-3}\,g\,dm^{-3}$ at 298 K.
 - (i) Write an expression for the solubility product, $K_{\rm sp}$, of CaC $_{\rm 2}$ O $_{\rm 4}$. Include its units.

$$K_{\rm sp}$$
 =

(ii) Calculate the numerical value of $K_{\rm sp}$ CaC $_{\rm 2}$ O $_{\rm 4}$ at 298 K. Give your answer in **standard form** to **two** significant figures.

$$K_{sp} CaC_2O_4 = \dots$$
 [2]

[Total: 7]

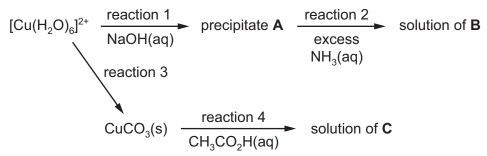
3 (a) Complete the table to show the total number of **unpaired** electrons in the 3d and 4s orbitals of each isolated gaseous atom.

	number of unpaired electrons		
	3d	4s	
Cr			
Mn			
Fe			

[2]

(b)			manganate(VII), de, potassium mang			heating	to	form
	Constru	uct an equation	on for this reaction.					
						 		[2]
(c)	Explain	the origin of	colour in transition	element cor	nplexes.			
						 		[3]

(d) The reaction scheme shows some reactions of $[Cu(H_2O)_6]^{2+}$.



	(i)	Write the formulae of	
		precipitate A,	
		complex ion B ,	
		compound C .	[3]
	(ii)	Identify a suitable reagent for reaction 3.	L°.
			[1]
(i	iii)	Write an equation for reaction 4.	
			[1]
(i	iv)	Describe two visual observations that would be made during reaction 4.	
			[1]
(e)	Pla	tin, $Pt(NH_3)_2Cl_2$, is a neutral complex of platinum(II).	
	Exp	plain why $Pt(NH_3)_2Cl_2$ has no charge.	
			[4]

(f)	(i)	Pt($(NH_a)_aCl_a$	displays	cis-trans	isomerism.
١	•	' \	•,		(11113)2002	alopiayo	ore trarre	1001110110111

Draw the structure of *trans*-platin. State its shape and the C*l*–Pt–C*l* bond angle.

	shape C <i>l</i> –Pt–C <i>l</i> bond angle	[2]
(ii)	Cis-platin is an effective anti-cancer drug.	
	Describe the action of <i>cis</i> -platin in this role.	
		[2]

(g) The use of cis-platin can cause side effects so nedaplatin has been developed.

Nedaplatin can be synthesised from cis-platin, $Pt(NH_3)_2Cl_2$, by replacing the two chloride ion ligands with a **single** bidentate ligand as shown.

Suggest the structure for nedaplatin.

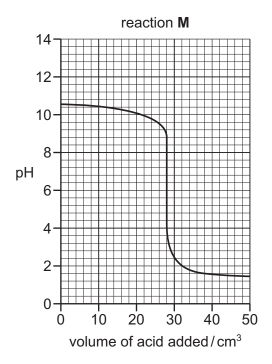
[1]

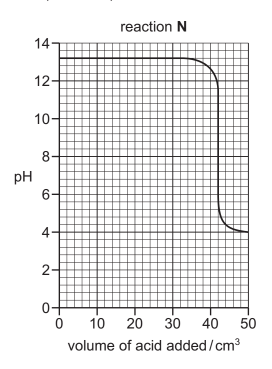
[Total: 19]

4	(a)		e enthalpy change of solut wn the group.	ion, $\Delta H_{\rm sol}^{\bullet}$, of th	e Group 2 sulfate	s becomes more	endothermic		
		Sta	te and explain the trend in	the solubility of	the Group 2 sulfa	tes down the grou	o.		
							[3]		
	(b)	(i)	Write the expression for h	$\zeta_{_{\! w}}$, the ionic pro	duct of water.				
			K _w =						
							[1]		
		(ii)	The numerical value of K_{ζ}	, increases with	increasing tempe	rature.			
				ppropriate column in each row to show the effect of increasing the pH and on the ratio $[H^{+}]$: $[OH^{-}]$.					
			effect of increasing temperature of water	decrease	stay the same	increase			
			рН						
			ratio [H⁺]∶[OH⁻]						
							[2]		
	(c)	An	aqueous solution of sodiur	n hydroxide has	s a pH of 13.25 at	298 K.			
		Cal	culate the concentration of	this sodium hy	droxide solution.				
					concentration =		moldm ⁻³ [2]		

(d)	Buf ran		ons a	are used to	regulate th	ne pH o	of a soluti	on to keep it	s pH	l value within a narro	W
				ions to de e pH of blo		hydro	gencarbo	onate ions, F	НСО	₃ -, and carbonic aci	d,
										[2]
(e)	The	$e K_a$ for et	hand	oic acid is	1.75 × 10⁻⁵ı	mol dm	⁻³ at 298	K.			
	(i)	When et pairs is t			dissolved in	ı water,	an equil	brium mixtur	e co	ntaining two acid-bas	se
					his equilibri his equilibri		the boxe	es label each	spe	ecies acidic or basic	to
С	H₃C	O ₂ H	+			\rightleftharpoons			+		
]	2]
	(ii)				prepared b 0.15 moldm	•	_		nol d	m ⁻³ ethanoic acid, a	an
		Calculat	e the	pH of the	buffer solu	ition fo	med at 2	98K. Give y	our a	answer to one decim	al

(f) Titration curves for two different acid-base reactions, **M** and **N**, are shown.





(i) Use the titration curve for reaction **M** to deduce the volume of acid added at the end-point for this titration.

volume of acid added at the end-point = cm³ [1]

(ii) The table shows some acid-base indicators.

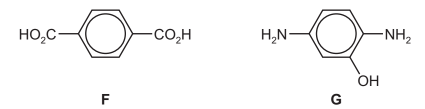
name of indicator	pH range of colour change
malachite green	0.2–1.8
bromocresol green	3.8–5.4
bromothymol blue	6.0–7.6
thymolphthalein	9.3–10.6

Name a suitable indicator for each of the acid-base titrations ${\bf M}$ and ${\bf N}$. Explain your answers.

reaction M	reaction N
explanation	
	[2]

[Total: 19]

5 (a) Polyhydroxyamide is a fire-resistant polyamide which is formed from the two monomers, **F** and **G**.



(i) Predict the number of peaks that will be seen in the carbon-13 NMR spectra of **F** and **G**.

	number of peaks
F	
G	

[2]

(ii) Draw the repeat unit of polyhydroxyamide. The amide bond should be shown displayed.

[2]

(b) When poly(ethene) is formed from ethene, many bonds are broken and formed.

Place **one tick** (\checkmark) in **each row** of the table to indicate the types of bonds broken and formed in this process.

	σ-bonds only	π -bonds only	both σ - and π -bonds
bonds broken			
bonds formed			

[2]

- (c) Addition polymers can be classified into two types.
 - homopolymer a polymer made up of the same monomer unit
 - copolymer a polymer made up of two or more different monomer units

The reaction of propene, CH₃CH=CH₂, with phenylethene, C₆H₅CH=CH₂, gives a copolymer.

Draw a length of the chain of this copolymer that contains one molecule of **each** monomer.

[2]

[Total: 11]

(d)	(i)	Polyalkenes biodegrade very slowly.	
		Explain why by referring to the structures of the polymers.	
			[1]
	(ii)	Some polymers will degrade in the environment.	
		Describe two processes by which this occurs.	
		1	
		2	
			[2]

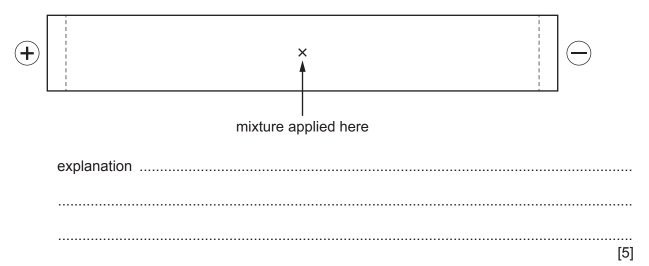
6	(a)	Ise the Data Booklet to draw the structure of the dipeptide val-lys. The peptide bond should
		e shown displayed.

[2]

(b) The isoelectric point is the pH at which an amino acid exists as a zwitterion. The isoelectric point of valine is 6.0 and of lysine is 9.8.

A mixture of the dipeptide, val-lys, and its two constituent amino acids, valine and lysine, was analysed by electrophoresis using a buffer at pH 6.0.

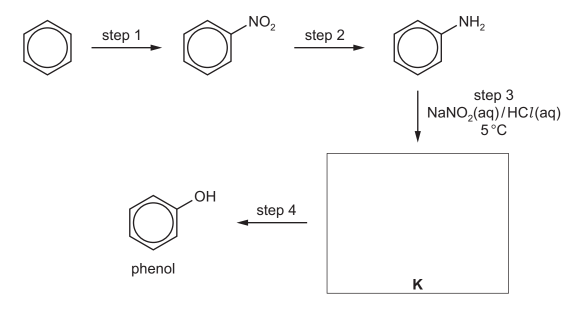
Draw and label **three** spots on the diagram of the electrophoresis paper to indicate the likely position of each of these three species after electrophoresis. Explain your answer.



[Total: 7]

7	(a)		and phenol both show a lack of reactivity towards reactants that cause the X - X bond (X = Cl or OH).	ıe
		Explain why.		
			[i	 3]
	(b)	When phenol is r C_6H_4 BrOH, are for	eacted with bromine dissolved in an inert solvent, two isomeric bromophenols ormed.	S,
		Suggest structur	es for these products. Name each compound.	
			name:	
			name:[2	2]

(c) A student suggested that phenol can be prepared from benzene by the method shown.



(i) Suggest reagents and conditions for each of the following steps.

	step 1	
	step 2	
	step 4	[3]
(ii)	Deduce the structure for K and draw its structural formula in the box.	[1]
(iii)	Name the mechanism for step 1.	

......[1]

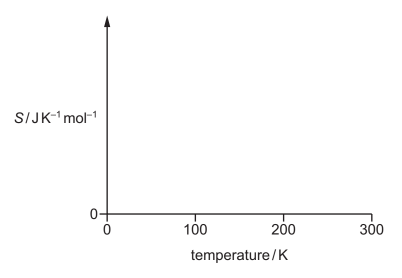
(iv) Write an equation for step 2. Use [H] for the reducing agent in this equation.

[1]

[Total: 11]

- 8 Entropy is a measure of the disorder of a system.
 - (a) Assume the entropy, S, for H₂O is zero at 0 K.

Sketch a graph on the axes to show how the entropy changes for H₂O between 0 K and 300 K.



[2]

(b) Place **one tick** (\checkmark) in **each row** of the table to show the sign of the entropy changes, $\triangle S$.

	ΔS is negative	ΔS is positive
solid dissolving in water		
water boiling to steam		

[1]

(c) The equation for a reaction that produces methanol is shown.

$$CO_2(g) + 3H_2(g) \rightarrow CH_3OH(g) + H_2O(g)$$

Use relevant bond energies from the *Data Booklet* to calculate the enthalpy change, ΔH , for this gas phase reaction.

$$\Delta H = \text{kJ mol}^{-1} [2]$$

(d) At 298 K, both products of this reaction are liquid.

$$CO_2(g) + 3H_2(g) \rightarrow CH_3OH(I) + H_2O(I)$$
 $\Delta H^{\circ} = -131 \text{ kJ mol}^{-1}$

Standard entropies are shown in the table.

substance	CO ₂ (g)	H ₂ (g)	CH ₃ OH(I)	H ₂ O(I)
S ^e /JK ⁻¹ mol ⁻¹	+214	+131	+127	+70

(i) C:	alculate the	standard	entropy	change,	ΔS ^e ,	for this	reaction
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$$\Delta S^{e} = J K^{-1} mol^{-1}$$
 [2]

(ii) Calculate the standard Gibbs free energy change, ΔG° , for this reaction at 298 K.

$$\Delta G^{\circ} = \text{kJ mol}^{-1}$$
 [2]

(iii) Predict the effect of increasing the temperature on the feasibility of this reaction.

(e) In a methanol-oxygen fuel cell, CH₃OH(I) and O₂(g) are in contact with two inert electrodes immersed in an acidic solution.

The half-equation for the reaction at the methanol electrode is shown.

$$CH_3OH + H_2O \rightleftharpoons CO_2 + 6H^+ + 6e^- \qquad E^{\circ} = -0.02V$$

(i)	Use the Data Booklet to write an equation for the overall cell reaction.			
	[1	1]		
(ii)	Use E° values to calculate the E°_{cell} for this reaction.			

 $E_{\text{cell}}^{\Theta} = \dots V [1]$

[Total: 12]

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