



Cambridge International Examinations

Cambridge International Advanced Subsidiary and Advanced Level

CANDIDATE NAME			
CENTRE NUMBER		CANDIDATE NUMBER	
CHEMISTRY			9701/43
Paper 4 A Level	Structured Questions	Oc	tober/November 2017
			2 hours
Candidates answ	er on the Question Paper.		
Additional Materia	als: 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	The compound	nitrosyl bromide,	NOBr, can be	formed by the	reaction shown.
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$$2NO + Br_2 \rightleftharpoons 2NOBr$$

(a)	Using oxidation numbers, explain why this reaction is a redox reaction.

(b) Nitrosyl bromide contains a trivalent nitrogen atom.

Draw the 'dot-and-cross' diagram for NOBr. Show outer electrons only.

[2]

(c) The rate of the reaction was measured at various concentrations of the two reactants, NO and Br_2 , and the following results were obtained.

experiment	[NO]/moldm ⁻³	[Br ₂]/moldm ⁻³	initial rate / mol dm ⁻³ s ⁻¹
1	0.03	0.02	3.4×10^{-3}
2	0.03	0.04	6.8 × 10 ⁻³
3	0.09	0.04	6.1 × 10 ⁻²
4	0.12	0.06	to be calculated

The general form of the rate equation for this reaction is as follows.

rate =
$$k[NO]^a[Br_2]^b$$

(i)	What is meant by the term <i>order of reaction</i> with respect to a particular reagent?							
		[1						

	(ii)	Use the data in the table to deduce the values of <i>a</i> and <i>b</i> in the rate equation. Show your reasoning.					
			[2]				
	(iii)	Use the data in the table to calculate the initial rate for experiment 4.					
		initial rate = $mol dm^{-3} s^{-1}$	[1]				
	(iv)	Use the results of experiment 1 to calculate the rate constant, k , for this reaction. Include the units of k .					
		rate constant, <i>k</i> = units	[2]				
	(v)	By considering the rate equation, explain why the rate decreases with decreasi temperature.	ng				
			[1]				
(d)	The	e reaction between X and Y was studied.					
		$2X + Y \rightarrow Z$					
	The	e following sequence of steps is a proposed mechanism for the reaction.					
		$\begin{array}{lll} \text{step 1} & 2X \rightarrow V \\ \text{step 2} & V + Y \rightarrow Z \end{array}$					
	The	e general form of the rate equation for this reaction is as follows.					
		$rate = k[X]^m[Y]^n$					
	Ste	p 1 is the slower step in the mechanism.					
	Dec	duce the values of m and n in the rate equation.					
	m =	= n =					
			[1]				

[Total: 12]

2 (a) The table lists values of solubility products, $K_{\rm sp}$, of some Group 2 carbonates.

	solubility product in water at 298 K, $K_{\rm sp}/{\rm mol^2dm^{-6}}$
MgCO ₃	1.0 × 10 ⁻⁵
CaCO ₃	5.0 × 10 ⁻⁹
SrCO₃	1.1 × 10 ⁻¹⁰

			SrCO ₃	1.1 × 10 ⁻¹⁰	
		the data in the tab group.	ole to describe th	ne trend in the solubility o	f the Group 2 carbonates down
					[1]
(b)	(i)	Write an equation Include state sym		uilibrium for the solubility	product for MgCO ₃ .
					[1]
	(ii)				observed when a few cm³ of ution of MgCO₃. Explain your
					[2]
(c)	Use	the data in the tal	ble to calculate t	he solubility of $MgCO_3$ in	water at 298 K, in g dm ⁻³ .
				solubility of MgCO ₃	= g dm ⁻³ [2]

(d)	(i)	Magnesium nitrate decomposes at a lower temperature than barium nitrate.
		Explain why.
		[2]
(ii)	A sample of barium nitrate was heated strongly until no further change occurred. A white solid was formed.
		Write an equation for the action of heat on barium nitrate.
		[1]
(i	ii)	When water was added to the white solid produced in (d)(ii) , an alkaline solution was produced. Adding sulfuric acid to this solution produced a white precipitate.
		Write equations to explain these observations.
		[2]
		[Total: 11]

3 (a	1)	Def	ine the te	erm <i>standard cell po</i>	tential.		
(b))	(i)		d electrode potentia	im of the experimen al of the Pb ²⁺ (aq)/l		
							[4
	(ii)	Sugges	t how the <i>E</i> for this e	electrode is -0.13 velectrode would diffed dicate this by placing	er from its <i>E</i> ° value it	
			[more negative	no change	less negative	
			Explain	your answer.			

(c) Car batteries are made up of rechargeable lead-acid cells. Each cell consists of a negative electrode made of Pb metal and a positive electrode made of PbO₂. The electrolyte is H₂SO₄(aq).

When a lead-acid cell is in use, Pb²⁺ ions are precipitated out as PbSO₄(s) at the negative electrode.

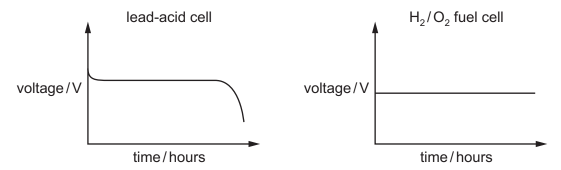
$$Pb(s) + SO_4^{2-}(aq) \rightarrow PbSO_4(s) + 2e^{-}$$

(i) Calculate the mass of Pb that is converted to PbSO₄ when a current of 0.40A is delivered by the cell for 80 minutes.

(ii) Complete the half-equation for the reaction taking place at the positive electrode.

$$PbO_{2}(s) + SO_{4}^{2-}(aq) + \dots + PbSO_{4}(s) + \dots$$
 [1]

(d) The diagrams show how the voltage across two different cells changes with time when each cell is used to provide an electric current.



Suggest a reason why

•	the voltage of the lead-acid cell changes after several hours,	
•	the voltage of the fuel cell remains constant.	
		[2]

[Total: 13]

4	(a)	Describe and explain how the density and melting point of cobalt compare to those of calcium.					
		den	sity of cobalt				
		ехр	lanation				
			ting point of cobalt				
		exp	lanation				
				 [3]			
	(b)	Tra	nsition metals can form complexes.				
			at is meant by the term transition metal complex?				
	(c)	(i)	Cobalt can form the compounds $[Co(NH_3)_5Br]SO_4$ and $[Co(NH_3)_5SO_4]Br$. These two compounds are structural isomers.				
			Define the term structural isomer.				
		(ii)	Draw a three-dimensional diagram to show the structure of the ion $[Co(NH_3)_5Br]^{2+}$. Name its shape.				
			$[Co(NH_3)_5Br]^{2+}$				
			shape	[1]			
	((iii)	State the type of bonding between the cobalt ion and NH_3 groups in the $[Co(NH_3)_5Br]^{2+}$ ions are the type of bonding between the cobalt ion and NH_3 groups in the $[Co(NH_3)_5Br]^{2+}$)n.			

((iv)	Sta	ate the oxidation number of co	oba	lt in			
		•	$[Co(NH_3)_5Br]^{2+}$		oxidation	numbe	er of Co =	
		•	$[Co(NH_3)_5 SO_4]^+$		oxidation	numbe	er of Co =	
								[1]
(d)	eac	h ot	ns of the compounds [Co(NH ther by simple chemical tests that any species bonded to					
		•	ete the table with two differ und. Give the expected obse				d to positive	ely identify each
			test		observation with [Co(NH ₃) ₅ Br]SO ₄ (a			vation with ,) ₅ SO ₄]Br(aq)
								[2]
(e)	The	two	compounds [Co(NH ₃) ₅ Br]S0	O₄ a	and [Co(NH ₃) ₅ SO ₄]E	Br are c	lifferent colo	urs.
	Ехр	lain	why the colours of the two c	om	pounds are differer	nt.		
								[2]
(f)	Son	ne t	ransition metals and their co	mno	ounds act as cataly	rsts Th	ne catalysis (can be classified
(.)			rogeneous or homogeneous		ouride det de outary	0.0. 11	io catalyolo (
	Con		ete the table by placing one ti n.	ck ((✓) in each row to in	ndicate	e the type of	catalysis in each
					heterogeneous	hom	ogeneous	
			Fe in the Haber process					
			Fe ²⁺ in the I ⁻ /S ₂ O ₈ ²⁻ reaction	n				
			NO ₂ in the oxidation of SO	2				
			V ₂ O ₅ in the Contact proces	s				
								[2]

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[Total: 14]

5 Compound P contains several functional groups.

(a)	Name the functional groups present in P .	
(b)	Compound P can be polymerised.	
	Draw a section of the polymer of P showing two repeat units. Name the type of polymerisation.	
	type of polymerisation	

[2]

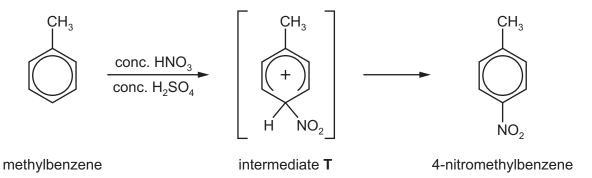
(c) Complete the following table to show the structures of the products formed and the *type of organic reaction* when **P** reacts with the four reagents.

reagent	structure(s) of product(s)	type of organic reaction
excess Br₂(aq)		
excess hot, concentrated, acidified MnO ₄ ⁻ (aq)		
excess hot HC1(aq)		
excess H ₂ /Pt catalyst		

[8]

[Total: 12]

6 (a) 4-nitromethylbenzene can be prepared via an electrophilic substitution reaction as shown.



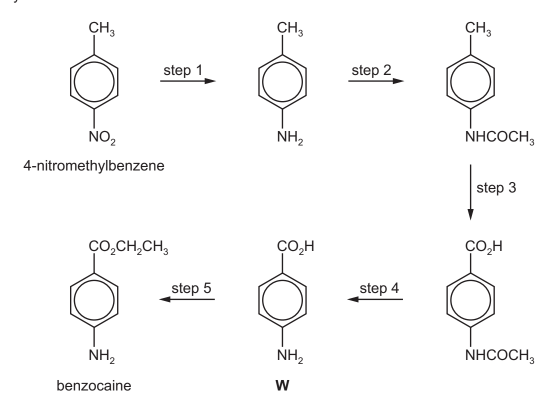
(i) This reaction also forms an isomer of 4-nitromethylbenzene as a by-product.

Draw the structure of this by-product.

(ii)	Write an equation for the reaction between $\mathrm{HNO_3}$ and $\mathrm{H_2SO_4}$ that forms the electrophile this reaction.	e fo
		[1
(iii)	Describe how the structure and bonding of the six-membered ring in intermedia differs from that in methylbenzene.	te 1
		[3

[1]

(b) Benzocaine is used as a local anaesthetic. It can be synthesised from 4-nitromethylbenzene by the route shown.



(i)	Give the systematic name	e of compound W.

	1.3
(ii)	Suggest the reagents and conditions for steps 1–5.
	step 1
	step 2
	step 3
	step 4

[6]

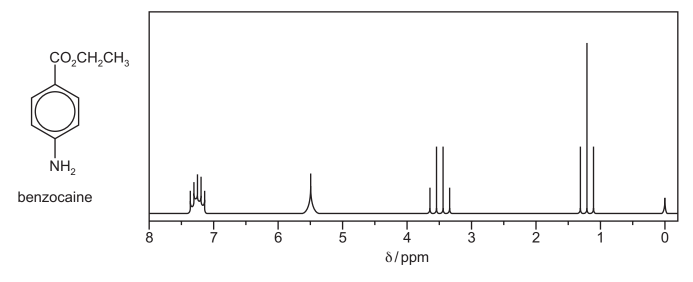
(c) Suggest how the basicity of benzocaine would compare to that of ethylamine. Explain your answer.

[2]

(d)	A sample of benzocaine,	shown below,	was	analysed	by	proton	NMR	and	carbon-13	NMR
	spectroscopy.									

(i) Predict the number of peaks that would be seen in the carbon-13 NMR spectrum.

(ii) Benzocaine was dissolved in ${\rm CDC} l_3$ and the proton NMR spectrum of this solution was recorded



Suggest why ${\rm CDC}\,l_3$ and not ${\rm CHC}\,l_3$ is used as the solvent when obtaining a proton NMR spectrum.

(iii) Use the *Data Booklet* and the spectrum in (d)(ii) to complete the table for the proton NMR spectrum of benzocaine. The actual chemical shifts, δ , for the four absorptions have been added.

δ/ppm	group responsible for the peak	number of ¹ H atoms responsible for the peak	splitting pattern
1.2			
3.5			
5.5			
7.1–7.4			multiplet

[4	1
┰	

(iv) Explain the splitting pattern for the absorption at $\delta 1.2$ ppm.

.....[1]

				15			
	(v)	The proton NMR	spectrum of be	enzocaine dissol	ved in D ₂ O was	recorded.	
		Suggest how this Explain your answ		ld differ from the	e spectrum in (c	l)(ii).	
							[1]
(e)	Ber	nzocaine can also l	oe used to syn	thesise the dyes	stuff S by the fo	llowing route.	
		CO ₂ CH ₂ CH ₃ NH ₂ benzocaine	step 1	step 2	R NaOH(aq), 〈	DH phenol	
	(!\	Commont the enter			<u> </u>		
	(i)	Suggest the reage	ents used for s	пер Т.			

(ii) Suggest structures for compounds **R** and **S** and draw them in the boxes. [2] [Total: 25]

.....[1]

7	(a)	Complete the following electronic structures.

• the iron atom, Fe 1s²2s²2p⁶

• the iron(III) ion, Fe³⁺ 1s²2s²2p⁶

[1]

(b) Solutions of iron(III) salts are acidic due to the equilibrium shown.

$$[Fe(H_2O)_6]^{3+}(aq) \iff [Fe(H_2O)_5(OH)]^{2+}(aq) + H^+(aq) \qquad K_a = 8.9 \times 10^{-4} \, \text{mol dm}^{-3}$$

Calculate the pH of a $0.25 \, \text{mol dm}^{-3} \, \text{FeC} \, l_3$ solution.

(c) The table shows numerical values of the stability constants for the following equilibrium where M can be one of the metal ions listed and L one of the ligands which replaces **one** H₂O molecule.

$$[M(H_2O)_6]^{n+}(aq) + L^{-}(aq) \rightleftharpoons [M(H_2O)_5L]^{(n-1)+}(aq) + H_2O(I)$$

metal ion, M	ligand, L	stability constant, K _{stab}
Fe ³⁺	F-	1.0 × 10 ⁶
Fe ³⁺	Cl-	2.5 × 10 ¹
Fe ³⁺	SCN-	9.0 × 10 ²
Hg ²⁺	Cl-	5.0 × 10 ⁶

(i)	What is meant by the term <i>stability constant</i> , K_{stab} ?
	[1]

- (ii) Use the data in the table to predict the formula of the complex formed in the greatest amount when
 - a solution containing equal concentrations of both F⁻ and SCN⁻ ions is added to Fe³⁺(aq),

• a solution containing equal concentrations of both Fe^{3+} and Hg^{2+} ions is added to $Cl^{-}(aq)$.

[1]

Ethanedioate ions, $^{-}O_{2}CCO_{2}^{-}$, are bidentate ligands. The abbreviation ed^{2-} can be used to represent ethanedioate ions.

(d) The complex $[Fe(ed)_2Cl_2]^{3-}$ can be formed according to the equation shown.

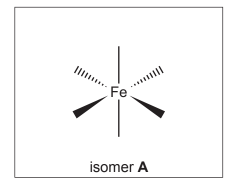
$$[Fe(H_2O)_4Cl_2]^+(aq) + 2ed^{2-}(aq) \rightleftharpoons [Fe(ed)_2Cl_2]^{3-}(aq) + 4H_2O(l)$$

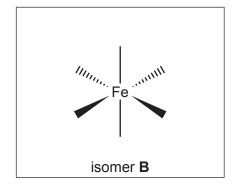
Write the expression for the equilibrium constant, K_{stab} , and state its units.

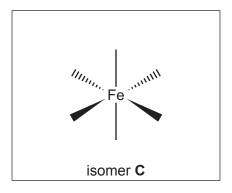
$$K_{\text{stab}} =$$

units[2]

- (e) $[Fe(ed)_2Cl_2]^{3-}$ shows geometrical and optical isomerism.
 - (i) Complete the three-dimensional diagrams to show the three stereoisomers of $[Fe(ed)_2Cl_2]^{3-}$. You may use $_{-}O^{-}$ to represent ed^{2-} .







[3]

(ii) Give the letters of two isomers of $[Fe(ed)_2Cl_2]^{3-}$ which are geometrical isomers of each other.

.....[1]

(iii) Give the letters of the two isomers of $[Fe(ed)_2Cl_2]^{3-}$ which show optical isomerism.

.....[1]

(iv) Give the letter of the isomer which has **no** dipole moment.

......[1]

[Total: 13]

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