Cambridge International Advanced Level

MARK SCHEME for the October/November 2014 series

9701 CHEMISTRY

9701/42

Paper 4 (A2 Structured Questions), maximum raw mark 100

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Question	Marking point	Marks	Marks total
1 (a) (i)	[NO] 2^{nd} order and the concentration is $\times 2$, rate $\times 4$	1	
	$[O_2]$ 1 st order and evidence of using expt 1 & 2 when the concentration is ×2, rate doubles	1	
(ii)	(0.00408×27) rate = <u>0.11</u> (mol dm ⁻³ s ⁻¹) to 2sf	1	
(iii)	(Rate =) $k [O_2][NO]^2$	1	
(iv)	k = 332(.03125) mol ⁻² dm ⁶ s ⁻¹	1 1	[6]
(b) (i)	labelled axes <i>x</i> -axis: energy (KE) and <i>y</i> -axis: molecules or particles two curves: starts origin; not touching <i>x</i> -axis again; no levelling out; curves only intersecting once curves labelled and T2 is to the right and lower max than T1	1 1 1	
(ii)	rate increases and energy of the particles increases	1	
	more particles have E_{a}	1	[5]
(c)	1 mole of F_2 and 1 mole NO reacting in the slow step	1	
	a balanced mechanism consistent with overall equation	1	
	e.g. $F_2 + NO \rightarrow NOF + F$ OR $F_2 + NO \rightarrow NOF_2$ NO + F \rightarrow NOF NOF NOF ₂ \rightarrow 2NOF		[2]
Total			[13]

2 (a)	3d 4s	1	
	(Ni) $\uparrow \downarrow$ $\uparrow \downarrow$ $\uparrow \downarrow$ \uparrow \uparrow	1	[2]
	(Ni ²⁺) $\uparrow \downarrow$ $\uparrow \downarrow$ $\uparrow \downarrow$ \uparrow		
(b) (i)	degenerate	1	
(ii)	2 upper orbitals and 3 lower orbitals	1	
(iii)	correct upper orbital diagram	1	
	correct lower orbital diagram	1	[4]
(c)	electron(s) move from lower to upper level	1	
	absorb (red/blue) light/photon	1	
	complementary colour (green) is seen OR green light is transmitted	1	[3]

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(d)	A Ni(OH B [Ni(NH) ₂ OR Ni(OH) ₂ (H ₂ O) ₄ H ₃) ₆] ²⁺ OR [Ni(NH ₃) _n (H ₂ O) _{6-n}] ²⁺ OR [Ni(NH ₃) _n (H ₂ O) _{4-n}] ²⁺			1 1	
	OR [Ni(H ₂ OR [Ni(H ₂	$\begin{array}{rcl} OH^- \to & Ni(OH)_2 \\ O_{6} \end{bmatrix}^{2+} & + & 2OH^- \to & Ni(OH)_2 & + & 6H_2O \\ {}_2O_{6} \end{bmatrix}^{2+} & + & 2NH_3 \to & Ni(OH)_2 & + & 4H_2O & + & 2NH_4^+ \\ O_{6} \end{bmatrix}^{2+} & + & 2OH^- \to & Ni(OH)_2(H_2O)_4 + & 2H_2O \end{array}$			1	
	Ni(OH) ₂ + OR Ni(H ₂ ($(6NH_3 \rightarrow [Ni(NH_3)_6]^{2+-} + 2OH^-)_6]^{2+-} + 6NH_3 \rightarrow [Ni(NH_3)_6]^{2+-} + 6H_2O$			1	[4]
Total						[13]

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3 (a) (i)	$101 = P^{35}Cl^{35}Cl$ $103 = P^{35}Cl^{37}Cl$ $105 = P^{37}Cl^{37}Cl$				1 1 1	
(ii)	9:6:1				1	[4]
(b) (i)	PCl_5 5 bonding p	pairs around P			1	
(ii)					1 1	[3]
(c) (i)	P_4O_6 structure where $O=P-O-F$	here each P has three P-O bonds and each O has two P-O bonds $P = 0$	onds e.g.		1	
(ii)	(molecule/ion/spe ion)	ecies) that donates a lone pair of electrons (to a central transi	ition metal at	om or	1	[2]
(d) (i)	$K_{sp} = [Ca^{2+}]^3 [PO_4^3]$	-] ²			1	

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(ii)	$[PO_4^{3-}] =$	$3 \times 2.50 \times 10^{-6} = 7.50 \times 10^{-6} \text{ mol dm}^{-3}$ $2 \times 2.50 \times 10^{-6} = 5.00 \times 10^{-6} \text{ mol dm}^{-3}$ $10^{-6})^{3}(5.00 \times 10^{-6})^{2}$			1	
	= 1.05(1. mol ⁵ dm ⁻¹	1) × 10 ⁻²⁶			1 1	[4]
(e) (i)	· · ·	r change) when 1 mole of an ionic compound d from its gaseous ions			1 1	
(ii)	Mg ²⁺ has OR Mg ²⁺	a smaller (ionic) radii than Ca ²⁺ is smaller than Ca ²⁺			1	[3]
Total						[16]
4 (a) (i)		$ HNO_3 \rightarrow 2HSO_4^- + NO_2^+ + H_3O^+ $ $ O_4 + HNO_3 \rightarrow HSO_4^- + NO_2^+ + H_2O $			1	

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(ii)	 int cu pr 	e of inly arrow from inside the benzene ring to NO_2^+ group termediate – penalise NO_2 connectivity or missing methyl group (ond inly arrow from C-H bond into ring oduct + H ⁺ (or as diagram –H ⁺) and 3-substituted nitromethylbenzene) $\downarrow -H^+$ $H^- NO_2$ NO_2^+	e)		3	[4]
(b) (i) (ii)	C <i>l</i> acidity of	$C_1CH_2CO_2H > CH_3CO_2H$ AND ($C_1CH_2CO_2H$) as an electronegative/ephenol > CH_3CH_2OH AND electrons on oxygen (on phenol) delocal			1 1	
		ene ring withdraws electrons from oxygen acid linked to weakening O-H bond/anion being stabilised			1	[3]

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(c)	Na	o o ONa (or ionic)	redox/reduction		
	Br ₂		(electrophilic) substitution		
	NaOH	OH and OH [1]	hydrolysis/ acid-base/		
		or ionic ONa ONa			
	1 mark for for reaction	r each correct structure on types, 2 correct = 1 mark, 3 correct = 2 r	narks	4 2	[6]

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Tot	al			13
5	(a)	$CH_{3}CH_{2}COCl > CH_{3}CH_{2}CH_{2}Cl > C_{6}H_{5}Cl$	1	
		 any two of: C-Cl bond strength is weakest in CH₃CH₂COCl ora In C₆H₅Cl (no hydrolysis) C-Cl bond is part of delocalised system OR p-orbital on Cl overlaps with π system OR electrons from Cl overlap with π system CH₃CH₂COCl carbon in C-Cl bond is more electron deficient since it is also attached to an oxygen atom ora 	1+1	[3]
	(b)	ketone, amine, carboxylic acid two correct 1 mark, all three 2	2	[2]
	(c) (i)	dipole on C-Br curly arrow breaking C-Br bond curly arrow from lone pair on N to carbon in C-Br bond H_2N H_2C H_3C CH_2 δ^{+} δ^{+}	1 1 1	
	(ii)	nucleophilic substitution	1	
	(iii)	HBr or hydrogen bromide	1	[5]

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(d)	$\mathbf{Y} = \begin{array}{c} 0 \\ H_2N \\ OH \end{array} OH$		[3]
	$W = \bigcup_{\substack{H_3N^+ \\ (Cl^-) \\ 0}} OH \qquad X = \bigcup_{\substack{H_3C^- \\ H_3C^- \\ 0}} OH OH$		
	each structure 1 mark		
(e)	$\begin{array}{ c c c c c }\hline O & O & O & O & O \\ \hline \square & -C - CH_2 - CH$	1	
	correct polyamide with two repeat units	1	[2]
Total	Total		
6 (a)	 (move in different directions) some amino acids have a different charge (move at different speeds) some amino acids have a different size/different charge (some amino acids do not move at all) some amino acids exist as a zwitterions/have no net(overall) charge/neutral/both NH₂/COOH are charged in amino acids 		[3]
(b) (i)	mobile – solvent or water stationary – alumina/silica (supported on glass/plastic/AI)		
(ii)	by adsorption	1	[3]

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(c)	any three of: (all can be awarded from a clear, labelled diagram)				
	 (base pairing) A to T OR C to G H-bonds between bases two/double stranded/chains anti-parallel strands (general structure) sugar-phosphate backbone OR BASE-SUGAR-PHOSPHATE bonded in a diagram 				
(d)	van der Waals' forces lost (in val) H-bonding gained (in ser)	1 1	[2]		
Total					
7 (a)	amide group circled OR indicated as diagram ester group circled OR indicated as diagram $H_{3}C \xrightarrow{O} + F_{3}C \xrightarrow{O} + F_{3}C$	1 1	[2]		
(b)	lower doses of the drug required OR improved activity of the drug OR reduced side effects	1	[1]		

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	(c)	decreases enzyme activity OR decreases rate at which product is formed				
		binds with the enzyme's active site OR has a complementary shape to active site OR similar shape to substrate				
		(competitive inhibition can be overcome by) increasing [substrate] OR increasing substrate concentration				
	(d)	energy source/carrier OR releases energy when hydrolysed				
To	Total					
8	(a)	M:M+1 = 100/(1.1 x n) 20.4/0.9 = 100/(1.1 x n)	1			
		x =4	1			
	(ii)	C ₄ H ₁₀ O				
	(b) (i)	(i) 2-methylpropan-1-ol OR correct structure				
		CH ₃ H ₃ C OH				
	(ii)	0.9-1.0 is (2 x)CH ₃ R/CH ₃ /RCH multiplet/1.8 is CHR/R ₃ CH	1			
		singlet/2.5 is OH 3.4 is CH_2O/CH_3O	1 1			
	(iii)	(iii) doublet 1H/one proton on adjacent carbon				

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(iv)	OH proto	or one peak disappears n is labile or exchanges for D of D ₂ O equation e.g. D ₂ O + OH \rightarrow DOH + OD as a minimum			1 1	[9]
Total						12
						100