

Cambridge International Examinations Cambridge International Advanced Subsidiary and Advanced Level

CHEMISTRY

9701/42 May/June 2016

Paper 4 A Level Structured Questions MARK SCHEME Maximum Mark: 100

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Question	Answer	Marks
1 (a) (i)	dative (covalent) or coordinate	2
	Hydrogen/H (boding)	
(ii)	octahedral	1
(iii)	$\begin{array}{rcl} Mg(NO_3)_2.6H_2O &\rightarrow & Mg(NO_3)_2 + & 6H_2O \\ Mg(NO_3)_2 &\rightarrow & MgO + & 2NO_2 + & \frac{1}{2}O_2 \end{array}$	4
	any three of (solid) dissolves/turns to liquid condensation on tube <u>white</u> solid (forms/remains) brown fumes (evolved) gas formed that relights a glowing splint	
(iv)	M_r values: Mg(NO ₃) ₂ .6H ₂ O = 256.3 MgO = 40.3 or (loss in molar mass = 256.3 - 40.3 =) 216 percentage loss = $100 \times 216/256.3 = 84.3/84.4\%$	2
(b)	(cat)-ionic radius/ion size increases (down the group) less polarisation/distortion of nitrate ion/NO ₃ ⁻	2
(c)	$2AgNO_3 \rightarrow 2Ag + 2NO_2 + O_2$	1
		[Total: 12]
2 (a) (i)	(an acid that is) partially/incompletely ionised/dissociated	1
(b) (i)	$pK_a = -\log K_a$ or $K_a = 10^{-pK_a}$	1

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Question	Answer	Marks
(ii)	ethanoic acid (1) is more acidic than propanoic acid (2) due to smaller electron-donating (R/alkyl) group/less electron-donating (R/alkyl) group(s)	3
	2-chloropropanoic acid (3) is more acidic than propanoic acid (2) due to electron-withdrawing/electronegative (Cl /chlorine) atom	
	2-chloropropanoic acid (3) is more acidic than 3^{-} -chloropropanoic acid (4) since the C1/chlorine/electronegative atom is closer to the CO ₂ ⁻ /acid	
(c) (i)	H ₂ (g) H ₂ (g) H ₁ (aq) H ⁺ (aq) M1: voltmeter/V and salt bridge labelled M2: Cu and Cu ²⁺ /CuSO ₄ (any soluble Cu(II) salt) M3: H ₂ (arrow in) and H ⁺ /HC <i>I</i> /H ₂ SO ₄ /any mineral acid M4 Pt and one solution at 1 M/1 moldm ⁻³ OR H ₂ at 1 atm	4
(ii)	$E_{cell}^{\Theta} = 0.34 \text{ (V)}$ and $(Cu^{2+})/Cu$ is the positive electrode	1
d (i)	$\begin{array}{l} K_{a}=1.23\times10^{-5}\\ [H^{+}]=\sqrt{(K_{a}.c)}=\sqrt{(1.23\times10^{-5}\times0.1)}=1.11\times10^{-3}\text{mol}\text{dm}^{-3} \end{array}$	2
	pH = 3.0 (2.96) ecf from [H ⁺]	

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Question	Answer	Marks
(ii)	$E = 0.0 + 0.059\log(1.11 \times 10^{-3}) \text{ OR} = -0.17(4)\text{V}$	2
	so new <i>E</i> _{cell} = 0.34 + 0.17 = 0.51V ecf from (d)(i)	
		[Total: 14]
3 (a) (i)	(CH ₃) ₂ CHCN	1
(ii)	reaction 1: NH_3 (in ethanol) under pressure (+ heat) or heat NH_3 in a sealed tube	3
	reaction 2: KCN/NaCN and heat/reflux (in ethanol)	
	reaction 3: H_2 + Ni <i>or</i> LiA lH_4	
(b) (i)	$CH_{3}CH_{2}NH_{2} + H_{2}O \rightarrow CH_{3}CH_{2}NH_{3}^{+} (+) OH^{-}$	1
(ii)	ethylamine is more basic than ammonia because of electron-donating (alkyl/ethyl/R) group (in ethylamine)	2
	which makes the lone pair (on N) more available for donation	
	or the <u>lone pair</u> (on N) more available for a proton/H ⁺	
(c) (i)	A solution which resists/minimises/roughly maintains changes in <u>pH</u> when (small amounts of) H ⁺ or OH ⁻ are added	1
(ii)	$CH_3NH_2 + H^+ \rightarrow CH_3NH_3^+$	2
	$CH_3NH_3Cl + OH \rightarrow CH_3NH_2 + H_2O + Cl$	
		[Total: 10]

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Question	Answer	Marks
4 (a) (i)	CI//////Pt	2
	(cis) (trans)	
(ii)	<i>cis</i> is (more) polar due to both Cl ^(δ−) on same side or <i>cis</i> is (more) polar as dipoles do not cancel/unsymmetrical or trans is non-polar as it is bond dipoles cancel	1
(iii)	 (This can only be <i>cis</i>) its mirror image is the same/superimposable or the distance between two coordinating nitrogens/oxygens is too small to bond <i>trans</i> or difficult for the NH₂ and O to change places (since 5-memebered rings can only bridge adjacent positions) 	1
(b) (i)	It's not square planar or it's tetrahedral	1
(ii)	must be 3D structure (i.e. tetrahedral-like) $R_{3}P$ Ni or $R_{3}P$ Ni Or $R_{3}P$ Ni Or $R_{3}P$ Ni Or $R_{3}P$ Or $R_{3}P$ Or $R_{3}P$ Or PR_{3} Or $PR_{$	1
		[Total: 6]

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Question	Answer	Marks
5 (a) (i)	$K_{\text{stab}} = \frac{[Cd(CH_3NH_2)_4^{2^+}]}{[Cd^{2^+}] [CH_3NH_2]^4}$	2
	units: mol ⁻⁴ dm ¹²	
(ii)	$\begin{array}{rcl} Cd^{2^{+}} + 4CH_{3}NH_{2} \rightleftharpoons [Cd(CH_{3}NH_{2})_{4}]^{2^{+}} \\ \text{at start: } 1 \times 1^{-4} & 0 \\ \text{at eqm: } 1 \times 10^{-7} & \textbf{y} & 1 \times 10^{-4} - 1 \times 10^{-7} \\ & & \textbf{or } 9.99 \times 10^{-5} \text{ or } 1.0 \times 10^{-4} \end{array}$	2
	$9.99 \times 10^{-5} / (y^4 \times 10^{-7}) = 3.6 \times 10^6$ and y = $\sqrt[4]{(9.99 \times 10^{-5})} / (1 \times 10^{-7} \times 3.6 \times 10^6) = 0.129 / 0.13$	
(b) (i)	(each complex is formed by) making (4 ×)N-Cd bonds and breaking (6 ×) O-Cd bonds or same types of/similar bonds forming/breaking or same number of bonds forming/breaking	1
(ii)	$\Delta S = (\Delta H - \Delta G) / T = (60.7 - 56.5) \times 1000 / 298 = (+)14 / (+)14.1$	1
(iii)	fewer moles (of solutes) are forming (one mole of) the complex (so less loss of disorder) or one en displaces two H ₂ O whereas one CH ₃ NH ₂ only displaces one H ₂ O	1
(iv)	The $[Cd(H_2NCH_2CH_2NH_2)_2]^{2+}$ / equilibrium 2 complex (is more stable) because: either K_{stab} is greater or ΔG^e is more negative.	1
		[Total: 8]

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Question	Answer	Marks
6 (a)	 essential mark M1 the reactants/substrate has a shape complementary/specific to <u>active site</u> – can be awarded from a labelled diagram as below or diagrams showing this specificity clearly any two of M2: reactants/substrate binds to/fits into the <u>active site</u> of the enzyme M3: (Interaction with site) causes a specific bond to be weakened, (which breaks) or lowers activation energy 	3
	M4: forms an E-S complex M5: products released from enzyme/active site	
	enzyme substrate (products)	
(b) (i)	δ 26 is CH3-CO δ 52 is CH3-O δ 169 is CH3CO δ 167 is phenyl-CO	2
	Phenyl ethanoateis Bmethyl benzoateM1 = any two correct δ linked to phenylethanoate/methyl benzoate	
	M2 = the rest correct	
(ii)	heat with H_3O^+ (to hydrolyse the ester)	3
	then add Br ₂ (aq)/bromine water	
	decolourises/gives white ppt. (with phenol from B)	
		[Total: 8]

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Question	Answer	Marks
7 (a) (i)	<i>labelled with</i> M1: <u>DC</u> power supply + and –/battery/cell/+ and – sign (on cell/electrodes) with a complete circuit	3
	M2: buffer solution/electrolyte labelled	
	M3: (amino acid) mixture/x on (filter) paper/gel/agarose	
	d.c. power supply electrolyte amiso acid mixture placed here Elter paper soaked in buffer solution	
(ii)	direction of movement related to charge (of amino acids)	2
	distance travelled depends on charge / M _r (of amino acids)	
(b) (i)	Asp + Val: pH 12 because Asp will be –CH ₂ COO ⁻ (R-group) moves further (to positive electrode than Val) or pH 12 Asp more negative so moves further (to positive electrode) or pH 12 because Asp has a charge of 2– but Val has a charge of 1– or best at pH 7 because Asp will be negatively charged (anionic) but Val neutral	1
(ii)	Lys + Ser: pH 2 because Lys will be (CH ₂) ₄ NH ₃ ⁺ (R-group) moves further (to negative electrode than Ser) or pH 2 Lys more positive so moves further (to negative electrode) or pH 2 because Lys has a charge of 2+ and Ser has a charge of 1+ or pH 7 because Lys is positively charged (cationic) but Ser neutral/zwitterionic	1

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Question	Answer	Marks
(iii)	Tyr + Phe: pH 12 because Tyr will be $C_6H_5CH_2O^-$ (R-group) moves further/more/faster (to positive electrode than Phe) or pH12 because Tyr has a charge of 2– but Phe has a charge of 1–	1
(c) (i)	$H_2N \xrightarrow{CH_2OH}_{H_2N} \xrightarrow{CH_2OH}_{CH_2} \xrightarrow{CH_2OH}_{CO_2H}$ M1: for -CONH- as shown above M2: for rest of molecule and correct connectivity of the bonds	2
(ii)	from the IR spectrum E is O-H or N-H (allow NH₂) F is C=O G is C-O 	2
		[Total: 12]
8 (a)	M1: solubility increases (down the group)	3
	M2: because lattice energy decreases faster than does ΔH_{hyd}	
	M3: ΔH_{sol} /enthalpy of solution becomes more exothermic/less endothermic	
(b) (i)	Should be the same/similar (enthalpy change), as (both acids) are fully ionised/strong acids	1

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Page 11	Mark Scheme	Syllabus	Paper
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Question	Answer	Marks
(b)	H is J is CO_2H CO_2H CN OH $J1$ $J2$	2
(c)	step 1: $(CH_3)_2CHCH_2Cl + AlCl_3 (+ heat)$ step 2: $CH_3COCl + AlCl_3 (+ heat)$ step 3: $HCN + NaCN \text{ or } HCN + base \text{ or } HCN + CN^-$ (steps 4 and 5 could be reversed on J) If J1 step 4 then step 5 J2 step 5 then step 4 step 4: $H_3O^+ + heat/aqueous HCl + heat$ step 5: conc $H_2SO_4 + heat/conc H_3PO_4 + heat$ or $Al_2O_3 + heat$ step 6: $H_2 + Ni (+ heat)$	6
(d)	step 1: electrophilic substitution <i>or</i> alkylation step 6: reduction/hydrogenation/addition	2
		[Total: 11]

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10 (a) (i)	Fe is3s ² 3p ⁶ 3d ⁶ 4s ²	1
(ii)		1
(b)	E° values: Sn ⁴⁺ /Sn ²⁺ = +0.15(V); Fe ³⁺ /Fe ²⁺ = +0.77(V) or E°_{cell} = +0.62 (V)	2
	(Sn ²⁺ will reduce Fe ³⁺) Sn ²⁺ + 2Fe ³⁺ \rightarrow 2Fe ²⁺	
(c) (i)	essential mark $K_{stab}/stability: [Fe(H_2O)_5F]^{2+} > [Fe(H_2O)_5SCN]^+$ $(> [Fe(H_2O)_6]^{2+})$	4
	observations (violet) \rightarrow deep-red (deep-red) \rightarrow colourless	
	(violet) \rightarrow colourless which stays colourless/does not change	
(ii)	ligand displacement/exchange/substitution	1
		[Total: 9]