CAMBRIDGE INTERNATIONAL EXAMINATIONS

Cambridge International Advanced Level

MARK SCHEME for the October/November 2014 series

9701 CHEMISTRY

9701/43

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

This mark scheme is published as an aid to teachers and candidates, to indicate the requirements of the examination. It shows the basis on which Examiners were instructed to award marks. It does not indicate the details of the discussions that took place at an Examiners' meeting before marking began, which would have considered the acceptability of alternative answers.

Mark schemes should be read in conjunction with the question paper and the Principal Examiner Report for Teachers.

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Question	Marking point				Marks	Marks total
1 (a) (i)		m/e	identity			
		35	³⁵ C1			
		37	³⁷ C <i>l</i>			
		70	³⁵ Cl ³⁵ Cl or ³⁵ Cl ₂			
		72	³⁷ C <i>l</i> ³⁵ C <i>l</i>			
		74	³⁷ C <i>l</i> ³⁷ C <i>l</i> or ³⁷ C <i>l</i> ₂			
	35, 37, 70, 72, 74 correct formulae at least one struct	ure as a posi	tive ion		1 1 1	
(ii)	9:6:1				1	[4]
(b) (i)	correct charges		_		1	
	correct electrons				1	
(ii)	Lattice energy = $\Delta H_f(SrC l_2) - (\Delta l_2) = +(-830) - (+164 + 548 + 106) = -2146 \text{ (kJ mol}^{-1})$			$_{\text{om}}(C\mathit{l})$ + $2\Delta H_{\text{ea}}(C\mathit{l}))$	1 1 1	[5]
(c) (i)	$SrCO_3 + 2HNO_3 \rightarrow Sr(NO_3)_2$	+ CO ₂ + H ₂ 0)		1	

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	(ii)	$Sr(NO_3)_2 \rightarrow SrO + 2NO_2 + 0.5 O_2$	1	[2]
	(d)	(down the group) nitrates become more stable / require a higher temperature to decompose as size/radius of ion increases OR charge density of ion decreases so polarisation/distortion of anion/nitrate ion/NO ₃ ⁻ /NO bond decreases	1 1 1	[3]
2	(a)	$BrO_3^- + 5Br^- + 6H^+ \rightarrow 3Br_2 + 3H_2O$ five correct species correct balancing	1	[2]
	(b) (i)	[BrO ₃ ⁻] 1 st order and the concentration is x2, rate doubles OR evidence using expt 1 & 4 eg ratios [H ⁺] 2 nd order and the concentration is x2, rate x4 OR evidence using expt 1 & 2 [Br ⁻] 1 st order and the concentration is x4, rate x4 OR evidence using expt 1 & 3 eg ratios	1 1 1	
	(ii)	(Rate =) $k [BrO_3^-][Br^-][H^+]^2$	1	
	(iii)	k = 1.32 $mol^{-3} dm^{9} s^{-1}$	1	[6]
3	(a) (i)	chromium and copper	1	
	(ii)	(all orbitals have the) same energy	1	
	(iii)	correct id of one higher energy d orbital the other higher energy d orbital	1	[4]

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(b) (i)	pale blue precipitate A solution B solution C	$Cu(OH)_2$ OR $[Cu(OH)_2(H_2O)_4]$ $[Cu(NH_3)_4(H_2O)_2]^{2+}$ OR $[Cu(NH_3)_4]^{2+}$ $[CuC_4]^{2-}$	1 1 1	
(ii)	solution B solution C	royal/deep/dark blue OR violet-blue yellow/green	1	
(iii)	redox OR oxidation of O AND reducing agent/reducing		1	[6]
(c)	3d-shell is full/3d ¹⁰ /no vac electrons cannot move bet	cant d-orbital/d-orbital s full ween orbitals OR transitions cannot occur	1 1	[2]
(d)	green/yellow orange/red AND blue/viol	et light is <u>absorbed</u>	1 1	[2]
4 (a)	(HC <i>l</i>) strong er acid/more (HC <i>l</i> has) more ions/highe	dissociated/ionised in solution er concentration of ions	1 1	[2]
(b) (i)		nges in the pH/keeps pH <i>fairly</i> constant ounts/vols of acid/H ⁺ or base/OH ⁻ are added	1 1	
(ii)	add (ethanoic acid) to NaO excess (ethanoic acid) OR mix with sodium ethan		1 1	[4]
(c)	CH ₃ CH(NH ₂)COOH + H ⁺ → CH ₃ CH(NH ₂)COOH + OH ⁻	CH ₃ CH(NH ₃ ⁺)COOH \rightarrow CH ₃ CH(NH ₂)COO ⁻ + H ₂ O	1 1	[2]

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(d) (i)	pKa 2.99 HO OH O OH O OH O OH O OH OH OH OH OH O	1	
	pKa 4.40 HO OH O OH O OH O OH O	1	
(ii)	HO COOH HOOC OH HOOC	2	[4]
5 (a)	 any five of these seven points. σ-bonds are between C-C OR C-H carbons are sp² rings of charge above and below the ring must be in diagram presence of σ-bonds electrons/bonds are delocalised planar molecule/bond angles 120° all C-C are the same length/have intermediate bond length between C-C & C=C 	5	[5]

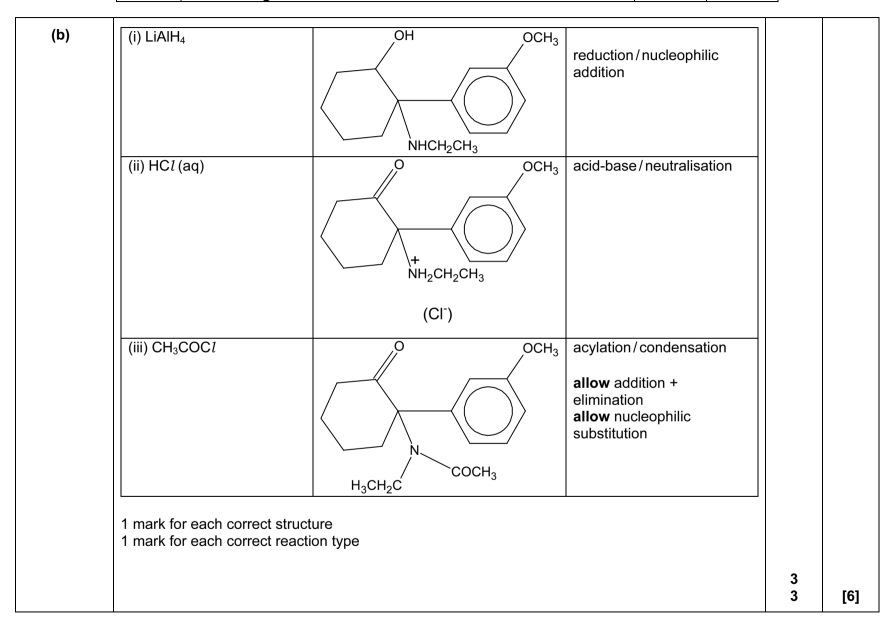
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(b)	Reagent X e.g. Br_2 , HNO_3 , Na , $NaOH$, benzenediazonium salt/ion; $RCOCl$; Fe^{3+} ; H_2+Ni substituted product for L-DOPA & vanillin (examples given are for X = Br_2 and $NaOH$)	1	
	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2	
	Reagent Y e.g. HC <i>l</i> ; Na ₂ CO ₃ , Mg, SOC <i>l</i> ₂ , PC <i>l</i> ₅ , ROH + c.H ₂ SO ₄ ; HC <i>l</i> +NaNO ₂ / HNO ₂ ; CH ₃ C <i>l</i> Correct substituted product for L-DOPA	1	
	HO NH_3^+ $COOH$		
	Reagent Z e.g. acidified Cr ₂ O ₇ ²⁻ ; 2,4-DNPH, hydrazine ; Fehling's, Tollens'; HCN; HCN + NaCN; NaBH ₄ ;	1	
	correct substituted product for vanillin	1	[7]
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6 (a) (i)	C ₁₅ H ₂₁ NO ₂	1	

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(ii)	O OCH3	1	
	* NHCH ₂ CH ₃		
(iii)	any two of ketone, amine or ether	2	[4]

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7	(a)	(ratio of) the concentrations/distribution/amount/mass of solute in two (immiscible) solvents at equilibrium OR equilibrium constant OR includes expression with <i>K</i>	1	[2]
	(b)	K_{pc} = [J in ether]/[J in H ₂ O] = $(2.14/20)/(5-2.14/75)$ = 2.81 OR 2.82	1	[2]
	(c)	1 st extraction: 2.81 =(x/10)/(5.0-x)/75 2.81(5-x) = 7.5x x= 1.36 g 2 nd extraction: 2.81 =(y/10)/(3.64-y)/75 2.81(3.64-y) = 7.5y y= 0.99 g	1	[2]
	(d) (i)	water/solvent/named solvent	1	
	(ii)	non-volatile liquid, for example mineral oil or at least a C ₁₅ hydrocarbon oil	1	
	(iii)	R _f (retardation factor) or distance travelled by solute and distance by solvent retention time	1	[4]

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	(e)	CO ₂ H ²	1	[1]
		CH ₂ OH 1		
		CO ₂ H 3		
8	(a)	C = 33 % A = T = 17 %	1	[2]
	(b) (i)	only one isomer may be active/be of therapeutic benefit	1	
	(ii)	the other (stereo) isomer may cause harm/side effects	1	[2]

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(c) (i)	structures of the following aldehydes: CHO CHO CHO		
	two correct structures = 1 mark two further correct structures – 1 mark	1 1	
(ii)	3-methylbutanal	1	
(iii)	pentanal 5 absorptions 2-methylbutanal 5 absorptions dimethylpropanal 2 absorptions	1 1 1	[6]
9 (a)	nylon, terylene – condensation; PVC – addition – all three correct	1	[1]
(b)	correct fully displayed formula of -CO-NH- unit correct polymer structure	1 1	[2]
(c)	sequence/order of amino acids (in the polypeptide chain)	1	[1]
(d)	hydrogen bond C=O and N-H in two different amino acids in the backbone diagram	1 1	[2]

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(e) (i)	disrupts hydrogen/ionic bonds as $-COOH/NH_3^+$ is deprotonated OR $-NH_3^+ + OH^- \rightarrow NH_2 + H_2O$ linked to hydrogen/ionic bond disrupted OR $-COOH + OH^- \rightarrow -COO^- + H_2O$ linked to hydrogen/ionic bond disrupted	1	
(ii)	Hg ²⁺ interferes with/breaks the disulfide bond/bridge not sulfite, sulfate, sulfur, sulfide OR -S-S- shown with Hg ²⁺ in an equation OR disrupting ionic interactions linked to carboxyl/COO– groups	1	
(iii)	(Heat to 70 °C) breaks the van der Waals' forces/hydrogen bonding	1	[3]