**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

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

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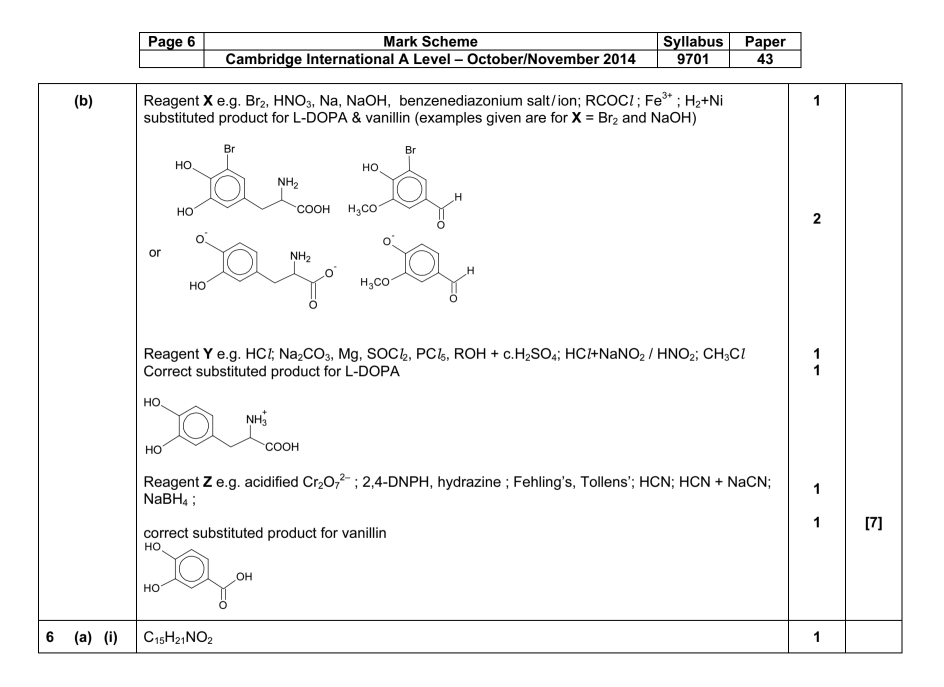
(	(ii)	$Sr(NO_3)_2 \Rightarrow SrO + 2NO_2 + 0.5 O_2$	1	[2]
(d)		(down the group) nitrates become more stable / require a higher temperature to decompose	1	
		as size/radius of <b>ion</b> increases <b>OR</b> charge density of <b>ion</b> decreases so polarisation/distortion of anion/nitrate ion/NO <sub>3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

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	u ala blua una sisitata A			
(b) (i)	pale blue precipitate <b>A</b> solution <b>B</b>	$Cu(OH)_2$ <b>OR</b> [Cu(OH)_2(H_2O)_4] [Cu(NH_3)_4(H_2O)_2]^{2+} <b>OR</b> [Cu(NH_3)_4]^{2+}		
	solution <b>C</b>	$[Cu(NH_3)_4(H_2O)_2]$ <b>OR</b> $[Cu(NH_3)_4]$ $[CuC_4]^{2-}$	1	
	Solution C			
(ii)	solution <b>B</b>	royal/deep/dark blue <b>OR</b> violet-blue	1	
	solution <b>C</b>	yellow/green	1	
(iii)	redox <b>OR</b> oxidation of	Cu <b>OR</b> reduction of Cu <sup>2+</sup>	1	
	AND reducing agent/redu			[6]
(c)	3d-shell is full/3d <sup>10</sup> /no va	cant d-orbital/d-orbital <b>s</b> full	1	
	electrons cannot move be	tween orbitals <b>OR</b> transitions cannot occur	1	[2]
(d)	green/yellow		1	
.,	orange/red AND blue/vio	let light is <u>absorbed</u>	1	[2]
4 (a)	(HC <i>l</i> ) stronger acid/more	dissociated / ionised in solution	1	
( )	(HCl has) more ions/high		1	[2]
(b) (i)	A solution that resists cha	nges in the pH/keeps pH <i>fairly</i> constant	1	
		nounts/vols of acid/H⁺ or base/OH⁻ are added	1	
(ii)	add (ethanoic acid) to Na	OH <b>OR</b> an equation	1	
( )	excess (ethanoic acid)	•	1	[4]
	<b>OR</b> mix with sodium etha	noate		
(c)	CH <sub>3</sub> CH(NH <sub>2</sub> )COOH + H <sup>+</sup> •		1	
	CH <sub>3</sub> CH(NH <sub>2</sub> )COOH + OH	$\rightarrow$ CH <sub>3</sub> CH(NH <sub>2</sub> )COO <sup>-</sup> + H <sub>2</sub> O	1	[2]

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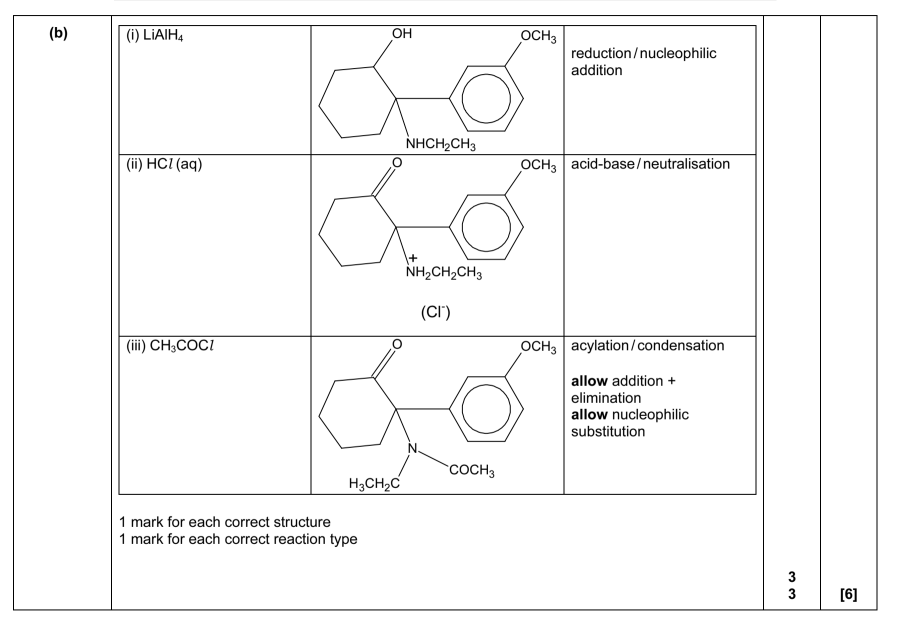
(d) (i)	pKa 2.99 HO $HO$ $OH$ $OH$ $OH$ $OH$ $OH$ $OH$	1	
	$pKa 4.40 \qquad HO \qquad \longrightarrow \qquad OH \qquad OH \qquad OH \qquad OH \qquad OH \qquad OH $	1	
(ii)	$\begin{array}{cccc} HO \\ HOOC \\ H$	2	[4]
5 (a)	<ul> <li>any five of these seven points.</li> <li>σ-bonds are between C-C OR C-H</li> <li>carbons are sp<sup>2</sup></li> <li>rings of charge above and below the ring must be in diagram</li> <li>presence of σ-bonds</li> <li>electrons/bonds are delocalised</li> <li>planar molecule/bond angles 120°</li> <li>all C-C are the same length/have intermediate bond length between C-C &amp; C=C</li> </ul>	5	[5]



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(ii)	O OCH <sub>3</sub>	1	
	* NHCH <sub>2</sub> CH <sub>3</sub>		
(iii)	any <b>two</b> of ketone, amine or ether	2	[4]

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7	(a)	(ratio of) the concentrations/distribution/amount/mass of <b>solute</b> in <b>two</b> (immiscible) <b>solvents</b> at equilibrium <b>OR</b> equilibrium constant <b>OR</b> includes expression with $K$	1 1	[2]
	(b)	$K_{pc} = [J \text{ in ether}]/[J \text{ in } H_2O]$ = (2.14/20)/(5–2.14/75) =2.81 <b>OR</b> 2.82		[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		
	(ii)	non-volatile liquid, for example mineral oil or at least a $C_{15}$ hydrocarbon oil		
	(iii)	1. $R_f$ (retardation factor) or distance travelled by solute <b>and</b> distance by solvent 2. retention time		[4]

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	(e)		CO <sub>2</sub> H <sup>2</sup>			1	[1]
			CH <sub>2</sub> OH 1				
			CO <sub>2</sub> H 3				
			CO <sub>2</sub> t				
8	(a)	C = 33 % A = T = 17 %				1 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:		
		two correct structures = 1 mark two further correct structures – 1 mark	1	
	(ii)	3-methylbutanal	1	
	(iii)	pentanal5 absorptions2-methylbutanal5 absorptionsdimethylpropanal2 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 H H H H H H H H H H	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)		ydrogen/ionic bonds as $-COOH/NH_3^+$ is deprotonated $A_2^+ + OH^- \rightarrow NH_2 + H_2O$ linked to hydrogen/ionic bond disrupted				
		$H + OH^{-}$ $\to COO^{-} + H_{0}O$ linked to by drog on / ionic bond disrupted			1	

(iii)	(Heat to 70 °C) breaks the van der Waals' forces/hydrogen bonding	1	[3]
(ii)	Hg <sup>2+</sup> interferes with/breaks the disulfide bond/bridge <b>not</b> sulfite, sulfate, sulfur, sulfide <b>OR</b> -S-S- shown with Hg <sup>2+</sup> in an equation <b>OR</b> disrupting ionic interactions linked to carboxyl/COO– groups	1	
	<b>OR</b> –COOH + OH <sup>-</sup> $\rightarrow$ –COO <sup>-</sup> + H <sub>2</sub> O linked to hydrogen/ionic bond disrupted	1	