## **CAMBRIDGE INTERNATIONAL EXAMINATIONS**

**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

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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Page 2	Mark Scheme	Syllabus	Paper
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Question	Marking point	Marks	Marks total
1 (a) (i)	[NO] $2^{nd}$ order <b>and</b> the concentration is $\times 2$ , rate $\times 4$	1	
	$[O_2]$ 1 <sup>st</sup> order <b>and</b> evidence of using expt 1 & 2 when the concentration is ×2, rate doubles	1	
(ii)	$(0.00408 \times 27)$ rate = $0.11$ (mol dm <sup>-3</sup> s <sup>-1</sup> ) to 2sf	1	
(iii)	(Rate =) $k [O_2][NO]^2$	1	
(iv)	k = 332(.03125) $mol^{-2} dm^6 s^{-1}$	1	[6]
(b) (i)	labelled axes x-axis: energy (KE) and y-axis: molecules or particles two curves: starts origin; not touching x-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 <sub>2</sub> and 1 mole NO reacting in the <b>slow</b> step	1	
	a balanced mechanism consistent with overall equation	1	
	e.g. $F_2 + NO \rightarrow NOF + F$ <b>OR</b> $F_2 + NO \rightarrow NOF_2$ $NO + F \rightarrow NOF$ $NO + NOF_2 \rightarrow 2NOF$		[2]
Total			[13]

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2 (a)	3d4s	1	
		1	[2]
	(Ni <sup>2+</sup> ) $\uparrow \downarrow \uparrow \downarrow \uparrow \uparrow$		
(b) (i)	degenerate	1	
(ii)	2 upper orbitals and 3 lower orbitals	1	
(iii)	correct lower orbital diagram  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)	$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	1	
	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	1	
	$Ni(OH)_2 + 6NH_3 \rightarrow [Ni(NH_3)_6]^{2^{+-}} + 2OH^-$ $OR \ Ni(H_2O)_6]^{2^{+-}} + 6NH_3 \rightarrow [Ni(NH_3)_6]^{2^{+}} + 6H_2O$	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)	PC <i>l</i> ₅ 5 bonding pairs around P	1	
(ii)		1 1	[3]
(c) (i)	$P_4O_6$ structure where each P has three P-O bonds and each O has two P-O bonds e.g. $O=P-O-P=O$	1	
(ii)	(molecule/ion/species) that <b>donates</b> a lone pair of electrons (to a central transition metal atom or ion)	1	[2]
(d) (i)	$K_{\rm sp} = [{\rm Ca}^{2+}]^3 [{\rm PO_4}^{3-}]^2$	1	

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(ii)	$[Ca^{2+}] = 3 \times 2.50 \times 10^{-6} = 7.50 \times 10^{-6} \text{ mol dm}^{-3}$ $[PO_4^{3-}] = 2 \times 2.50 \times 10^{-6} = 5.00 \times 10^{-6} \text{ mol dm}^{-3}$	1	
	= $(7.50 \times 10^{-6})^3 (5.00 \times 10^{-6})^2$ = $1.05(1.1) \times 10^{-26}$ $mol^5 dm^{-15}$	1	<u>[4]</u>
(e) (i)	(enthalpy change) when 1 mole of an ionic compound is formed from its gaseous ions	1	
(ii)	Mg <sup>2+</sup> has a smaller (ionic) radii than Ca <sup>2+</sup> <b>OR</b> Mg <sup>2+</sup> is smaller than Ca <sup>2+</sup>	1	[3]
Total			[16]
4 (a) (i)	$2H_2SO_4 + HNO_3 \rightarrow 2HSO_4^- + NO_2^+ + H_3O^+$ <b>OR</b> $H_2SO_4 + HNO_3 \rightarrow HSO_4^- + NO_2^+ + H_2O$	1	

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(ii)	<ul> <li>any three of <ul> <li>curly arrow from inside the benzene ring to NO<sub>2</sub><sup>+</sup> group</li> <li>intermediate – penalise NO<sub>2</sub> connectivity or missing methyl group (once)</li> <li>curly arrow from C-H bond into ring</li> <li>product + H<sup>+</sup> (or as diagram –H<sup>+</sup>)</li> </ul> </li> <li>allow 2- and 3-substituted nitromethylbenzene)</li> </ul>	3	[4]
(b) (i) (ii)	acidity of $ClCH_2CO_2H > CH_3CO_2H$ <b>AND</b> ( $ClCH_2CO_2H$ ) as an electronegative/electron withdrawing $Cl$ acidity of phenol $> CH_3CH_2OH$ <b>AND</b> electrons on oxygen (on phenol) delocalised into ring	1	
	OR benzene ring withdraws electrons from oxygen stronger acid linked to weakening O-H bond/anion being stabilised	1	[3]

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(c)	Na	O ONA (or ionic)	redox/reduction		
	Br <sub>2</sub>	Br OH OH	(electrophilic) substitution		
	NaOH	OH and OH [1]	hydrolysis/ acid-base/		
	1 mark for	r each correct structure on types, 2 correct = 1 mark, 3 correct = 2 r	marks	4 2	[6]

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Total			13
5 (a)	$CH_3CH_2COCl > CH_3CH_2Cl > C_6H_5Cl$	1	
	<ul> <li>any two of:         <ul> <li>C-Cl bond strength is weakest in CH<sub>3</sub>CH<sub>2</sub>COCl ora</li> </ul> </li> <li>In C<sub>6</sub>H<sub>5</sub>Cl (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</li> <li>CH<sub>3</sub>CH<sub>2</sub>COCl carbon in C-Cl bond is more electron deficient since it is also attached to an oxygen atom ora</li> </ul>	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 <b>lone pair</b> on N to carbon in C-Br bond  H <sub>2</sub> N  H <sub>3</sub> C  CH <sub>2</sub> Br  S+	1 1 1	
(ii)	nucleophilic substitution	1	
(iii)	HBr or hydrogen bromide	1	[5]

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(d)	$\mathbf{Y} = \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	3	[3]
(e)	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	1 1	[2]
Total			15
6 (a)	<ul> <li>(move in different directions)     some amino acids have a different charge</li> <li>(move at different speeds)     some amino acids have a different size/different charge</li> <li>(some amino acids do not move at all)     some amino acids exist as a zwitterions/have no net(overall) charge/neutral/both     NH<sub>2</sub>/COOH are charged in amino acids</li> </ul>	1 1 1	[3]
(b) (i)	mobile – solvent <b>or</b> water stationary – alumina/silica (supported on glass/plastic/Al)	1	
(ii)	by adsorption	1	[3]

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(c)	any three of: (all can be awarded from a clear, labelled diagram)		
	<ul> <li>(base pairing) A to T OR C to G</li> <li>H-bonds between bases</li> <li>two/double stranded/chains</li> <li>anti-parallel strands</li> </ul>		
	<ul> <li>(general structure) sugar-phosphate backbone OR BASE-SUGAR-PHOSPHATE bonded in a diagram</li> </ul>	3	[3]
(d)	van der Waals' forces lost (in val) H-bonding gained (in ser)	1	[2]
Total			11
7 (a)	amide group circled <b>OR</b> indicated as diagram ester group circled <b>OR</b> indicated as diagram	1	[2]
	$H_3C$ $H_3C$ $H_3C$ $CH_3$ $H_3C$ $CH_3$		
(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 <b>OR</b> decreases rate at which product is formed	1	
	binds with the enzyme's active site <b>OR</b> has a complementary shape to active site <b>OR</b> similar shape to substrate	1	
	(competitive inhibition can be overcome by) increasing [substrate] <b>OR</b> increasing substrate concentration	1	[3]
(d)	energy source/carrier OR releases energy when hydrolysed	1	[1]
Total			7
8 (a)	$M:M+1 = 100/(1.1 \times n)$ $20.4/0.9 = 100/(1.1 \times n)$	1	
	x = 4	1	
(ii)	C <sub>4</sub> H <sub>10</sub> O	1	[3]
(b) (i)	2-methylpropan-1-ol <b>OR</b> correct structure  CH <sub>3</sub> OH	1	
(ii)	0.9-1.0 is (2 x)CH <sub>3</sub> R/CH <sub>3</sub> /RCH multiplet/1.8 is CHR/R <sub>3</sub> CH singlet/2.5 is OH 3.4 is CH <sub>2</sub> O/CH <sub>3</sub> O	1 1 1 1	
(iii)	doublet 1H/one proton on adjacent carbon	1 1	

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(iv)	OH peak or one peak disappears	1	
	OH proton is labile $\textbf{\textit{or}}$ exchanges for D of D <sub>2</sub> O $\textbf{\textit{or}}$ as an equation e.g. D <sub>2</sub> O + OH $\rightarrow$ DOH + OD as a minimum	1	[9]
Total			12
			100