CAMBRIDGE INTERNATIONAL EXAMINATIONS

GCE Advanced Level

MARK SCHEME for the October/November 2013 series

9701 CHEMISTRY

9701/41

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.

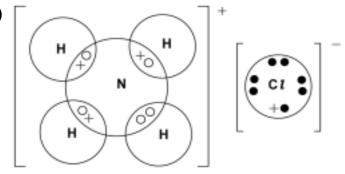
Cambridge will not enter into discussions about these mark schemes.

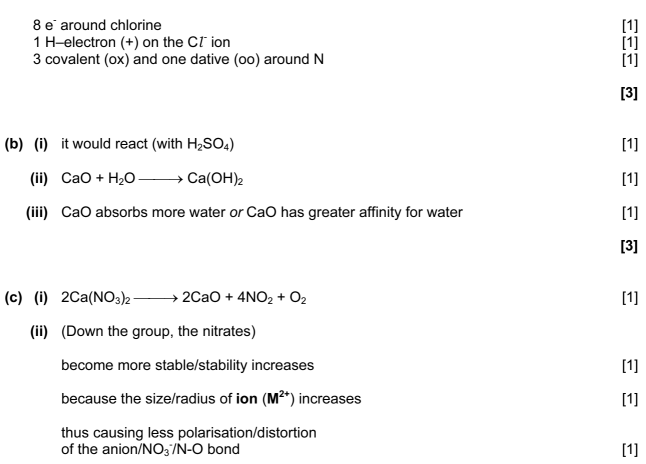
Cambridge is publishing the mark schemes for the October/November 2013 series for most IGCSE, GCE Advanced Level and Advanced Subsidiary Level components and some Ordinary Level components.



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1 (a)





[Total: 10]

[4]

Page 3	Mark Scheme	Syllabus	Paper
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- 2 (a) (i) Si-Si bonds are weaker (than C-C bonds) [1]
 - (ii) metallic (Sn) is weaker than (giant) covalent (Ge) [1]

[2]

(b) (i)
$$SiCl_4 + 2H_2O \longrightarrow SiO_2 + 4HCl$$

 $or SiCl_4 + 4H_2O \longrightarrow Si(OH)_4 + 4HCl$
 $or SiCl_4 + 3H_2O \longrightarrow H_2SiO_3 + 4HCl$
(partial hydrolysis is *not sufficient* e.g. to $SiCl_3OH + HCl$) [1]

(ii)
$$PbCl_4 \longrightarrow PbCl_2 + Cl_2$$
 [1]

(iii)
$$SnCl_2 + 2FeCl_3 \longrightarrow SnCl_4 + 2FeCl_2$$
 [1]

(iv)
$$SnO_2 + 2NaOH \longrightarrow Na_2SnO_3 + H_2O$$

 $or SnO_2 + 2NaOH + 2H_2O \longrightarrow Na_2Sn(OH)_6$
 $or ionic equation $SnO_2 + 2OH^- \longrightarrow SnO_3^{2-} + H_2O$ [1]$

[4]

[Total: 6]

	Page 4	ı	Mark Scheme	Syllabus	Paper	
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3	(a) (i)		+ HZ \longrightarrow NH ₄ ⁺ + Z ⁻ OH + HZ \longrightarrow CH ₃ OH ₂ ⁺ + Z ⁻			[1] [1]
	(ii)	_	+ B ⁻ \longrightarrow NH ₂ ⁻ + BH OH + B ⁻ \longrightarrow CH ₃ O ⁻ + BH			[1] [1]
						[4]
((b) (i)	a rea	action that can go in either direction			[1]
	(ii)		of forward = rate of backward reaction rward/back reactions occurring but concentrations of a	all species do no	t change	[1]
						[2]
((c) (i)	a so	lution that resists changes in pH			[1]
		whe	n small quantities of acid or base/alkali are added			[1]
	(ii)	in th	e equilibrium system HZ + $H_2O = Z^- + H_3O^+$			[1]
			tion of acid: reaction moves to the left [†] combines with Z ⁻ <u>and</u> forms HZ			[1]
			tion of base: the reaction moves to the right [†] combines with OH ⁻ and more Z ⁻ formed			[1]
					[5 ma	x 4]
((d) (i)	[H ⁺]	= $\sqrt{(0.5 \times 1.34 \times 10^{-5})}$ = 2.59 × 10 ⁻³ (mol dm ⁻³)			[1]
		pH =	2.59/2.6 (min 1 d.p)		ecf	[1]
	(ii)	CH ₃ ($CH_2CO_2H + NaOH \longrightarrow CH_3CH_2CO_2Na + H_2O$			[1]
	(iii)	n(ac	id) in 100 cm ³ = 0.5 × 100/1000 = 0.05 mol id) remaining = 0.05 – 0.03 = 0.02 mol I remaining] = 0.2 (mol dm ⁻³)			[1]
			vise, n(salt) = 0.03 mol + 0.3 (mol dm ⁻³)			[1]
	(iv)	pH =	= 4.87 + log(0.3/0.2) = 5.04–5.05		ecf	[1]
						[6]
(CH ₂ COC <i>l</i>			
	J is	NaC		00D= N D \		[2]
	(or	corres	sponding Br compounds for G , H and J ; CH₃CH₂COBr	, oubr ₂ , nabr)	[Total:	18]

Page 5	Mark Scheme	Syllabus	Paper
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- (a) (the energy change) when 1 mol of bonds [1] is broken in the gas phase
 - [2]
 - **(b) (i)** (C-X bond energy) decreases/becomes weaker (from F to I) [1]
 - due to bond becoming longer/not such efficient orbital overlap [1]
 - (ii) (as the bond energy of C-X decreases) the halogenalkanes become more reactive (answer must imply that it is from F to I) [1]
 - (c) The C-Cl bond is weaker than the C-F and C-H bonds or C-Cl bond (E = 340) and C-H (E = 410) [1]
 - so is (easily) broken to form Cl*/Cl radicals/Cl atoms [1] causing the breakdown of O₃ into O₂ [1]
 - (d) Cl-CH₂CH₂-CO₂H [1] HO-CH₂CH₂CH₂-C1 [1]

- (e) (i) light/UV/hv or 300°C [1]
 - (ii) (free) radical substitution [1]
 - (iii) $\Delta H = E(C-H) E(H-Cl) = 410 431 = -21 \text{ kJ mol}^{-1}$ [1]
 - (iv) $\Delta H = E(C-H) E(H-I) = 410 299 = +111 \text{ kJ mol}^{-1}$ ecf [1]
 - (v) The reaction with iodine is endothermic or ΔH is positive or requires energy [1]
 - (vi) $Cl_2 \longrightarrow 2Cl^*$ [1] [1] [1]

[8]

[3]

[3]

[3]

[Total: 19]

Page 6	Mark Scheme	Syllabus	Paper
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5 (a) (i) many monomers form a polymer

[1]

- (ii) addition [1]
- (iii) C=C/double/ π bond is broken **and** new C-C single bond<u>s</u> are formed or double bond breaks and forms single bonds with other monomers

[1] [3]

(b) propenoic acid [1]

[1]

(c) (i) CO₂Na CO₂Na

carbon chain **and** CO₂H [1]

- at least one sodium salt
- (ii) 120° to 109(.5)° [1] due to the change from a trigonal/sp² carbon to a tetrahedral/sp³ carbon [1]

[4]

(d) (i) $\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$

Any four:

hydrogen bond labelled

water H-bonded to O through H atom $\delta+/\delta$ - shown on each end of a H-bond

lone pair shown on O or C=O or H₂O on a correct H-bond

Na⁺ shown as coordinated to a water molecule

[3]

(ii) Solution became paler **and** Cu⁽²⁺⁾ swapped with Na⁽⁺⁾ or darker in colour **and** polymer absorbs water

[1]

[4]

Page 7	,	Mark Scheme	Syllabus	Paper
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(e) (i)	alke	ne(1), amide(1)		[2]
(ii)	NH_3			[1]
(iii)	H ₂ O			[1]
(iv)		(aq)/H₃O ⁺ and heat/reflux (not warm) H⁻ (aq), heat and acidify		[1]
	<i>0,</i> 0	· (aq), noat and adiany		[5]
				[Total: 17]

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Section B

6 (a) (i) **six/6** (gsv, sgv, gvs, vgs, svg, vsg)

[1]

(ii) H_3C CH_3 OH OH

(iii) valine (allow glycine) [1]

(iv) any two of: hydrogen bonds and CO_2H or OH or NH_2 or CONH or CO or NH or CO_2^- ionic bonds and NH_3^+ or $CO_2^$ van der Waals' and $-CH_3$ or -H

2 × [1]

[6]

(b) (i) same shape/structure as substrate

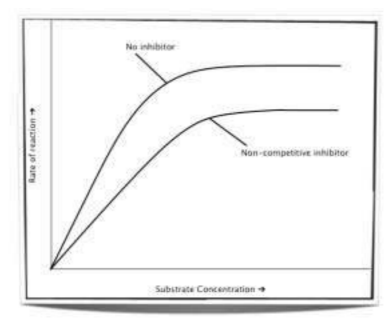
[1]

(inhibitor) competes/blocks/binds/bonds to **active site** *or* substrate cannot bind to **active site**

[1] [1]

(ii) binds with enzyme and changes shape/3D structure (of enzyme/active site)

(iii)



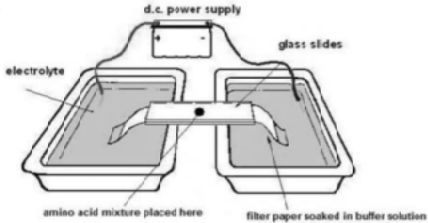
[1]

[4]

[Total: 10]

Page 9	Mark Scheme	Syllabus	Paper
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7 (a)



power supply (idea of complete circuit) electrolyte/buffer solution gel/filter paper/absorbent paper (amino acid) sample/mixture [centre of plate]

4 × [1]

[4]

(b) any two from:

size/ $M_{\rm r}$ (of the amino acid species) charge (on the amino acid species) temperature

2 × [1]

[2]

(c) Ratio of the <u>concentration</u> of a solute in each of two (immiscible) solvents or equilibrium constant representing the distribution of a solute between two solvents or $PC = [X]_a/[X]_b$ (at a constant temperature)

[1]

[1]

(d) (i) $K_{pc} = [Z \text{ in ether}]/[Z \text{ in H}_2O] - \text{allow reverse ratio}$ 40 = (x/0.05)/((4-x)/0.5)

[1]

= 3.2 g

ecf [1]

(ii) First extraction 40 = (x/0.025)/((4-x)/0.5)x = 2.67 g

ecf [1]

(iii) Second extraction: 1.33g remain in solution Second extraction 40 = (y/0.025)/((1.33-y)/0.5) y = **0.887** g

mass extracted = 2.67 + 0.89 = 3.56/3.6 g

ecf [1]

[4]

[Total: 11]

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- 8 (a) (i) (nitrates are) soluble [1]
 - (ii) $Ba^{(2+)}$ and $Pb^{(2+)}$ [1]
 - SO₄⁽²⁻⁾ [1]
 - BaCO₃/PbCO₃/CaSO₄ are insoluble [1]
 - (b) (i) fertilisers/animal manure [1]
 - (ii) washing powder/detergents/fertilisers/animal manure [1]
 - (iii) growth/production of algae/weeds/plants or eutrophication [1]
 - (c) (i) any one of:

$$2SO_2 + O_2 \longrightarrow 2SO_3$$
 and $SO_3 + H_2O \longrightarrow H_2SO_4$

or
$$SO_2 + NO_2 \longrightarrow SO_3 + NO$$
 and $SO_3 + H_2O \longrightarrow H_2SO_4$

$$or SO_2 + \frac{1}{2}O_2 + H_2O \longrightarrow H_2SO_4$$
 [1]

(ii) roasting sulfide ores/extraction of metals from sulfide ores [1]

[2]

[4]

[3]

[Total: 9]