## UNIVERSITY OF CAMBRIDGE INTERNATIONAL EXAMINATIONS

GCE Advanced Subsidiary Level and GCE Advanced Level

## MARK SCHEME for the October/November 2007 question paper

## 9701 CHEMISTRY

9701/04

Paper 4 (Theory 2), 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.

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		GCE A/AS LEVEL – October/November 2007 9701 04	4
1 (a	) (i)	$K_{a} = [H^{\dagger}][RCO_{2}^{-}]/[RCO_{2}H]$	[1]
	(ii)	$pK_a = -log_{10}K_a or -logK_a or log [H^+]^2/[RCO_2H] NOT ln;$	[1] <b>[2]</b>
(b	) (i)	acid strength <u>increases</u> from no. 1 to no. 3 <i>or</i> down the table <i>or</i> as $Cls$ increase due to the electron-withdrawing effect/electronegativity of chlorine (atoms) stabilising the anion <i>or</i> weakening the O-H bond NOT $H^+$ more available	[1] [1] [1]
	(ii)	chlorine atom is further away (from O-H) in no. 4, so has less influence	[1]
	(iii)	either: pH = $\frac{1}{2}$ (p $K_a$ – log <sub>10</sub> [acid]) or $K_a$ = $10^{-pKa}$ = $1.259 \times 10^{-3}$ = $\frac{1}{2}$ (4.9 + 2) [H <sup>+</sup> ] = $\sqrt{(K_a. c)}$ = $3.55 \times 10^{-4}$ = 3.4 (allow 3.5) pH = 3.4 ([1] for correct expression & values; [1] for correct working)	[1] ecf [1] <b>[6]</b>
(c	) (i)	catalyst	[1]
	(ii)	$CH_3CH_2CO_2H + Cl_2 \longrightarrow CH_2CHClCO_2H + HCl$	[1]
	(iii)	nucleophilic substitution NOT addition/elimination	[1]
	(iv)	$M_r(CH_3CH_2CO_2H) = 74$ $M_r(CH_2CH(NH_2)CO_2H) = 89$ $\therefore 10.0 \text{ g should give } 10 \times 89/74 = 12.03 \text{ g}$	[1]
		∴ percentage yield = 100 x 9.5/12.03 = <b>79</b> %  ([2] for correct a	ecf [1] answer) <b>[5]</b>
(d		$H_3$ -CH(CH $_3$ )-CO $_2$ correct atoms charges on H of $H_3$ N, and –COO but not –C-O-O correct charges	

Mark Scheme

**Syllabus** 

Paper

Page 2

[Total: 15]

[2]

Page 3	Mark Scheme	Syllabus	Paper
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$$\label{eq:mgC2O4(aq) + Ca(NO3)2} \begin{split} \text{Mg(NO3)}_2 + \text{CaC}_2\text{O}_4(s) & & \text{[1]} \\ \textbf{(B)} & & \text{[max 3]} \end{split}$$

(c) (i) 
$$(K_{sp} =) [Mg^{2+}][OH^{-}]^{2}$$
 [1] units are  $mol^{3}dm^{-9}$  ecf from  $K_{sp}$  [1]

(ii) 
$$(call [Mg(OH)_2(aq)] = [Mg^{2+}] = x)$$
  $\therefore K_{sp} = 2 \times 10^{-11} = 4x^3$  [1]

$$\therefore x = 1.71 \times 10^{-4} \text{ mol dm}^{-3}$$
 ecf [1]

(iii) less soluble because of the common ion effect   
or the equilibrium 
$$Mg(OH)_2(s) \Rightarrow Mg^{2+}(aq) + 2OH^{-}(aq)$$
 is moved to the left [1]

[5] [Total: 12]

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(b) 
$$K_2Cr_2O_7 + 2HCl \longrightarrow 2KCrClO_3 + H_2O$$
 [1]

(ii) 
$$E^{\circ}$$
 data and half equations:  $Cr_2O_7^{2-} + 14H^{+} + 6e^{-} \longrightarrow 2Cr^{3+} + 7H_2O$   $E^{\circ} = 1.33 \text{ V}$  [1]  $Cl_2 + 2e^{-} \longrightarrow 2 Cl^{-}$   $E^{\circ} = 1.36 \text{ V}$  [1] overall ionic equation:

(iii) (dilution will) lower E<sup>e</sup> for 
$$Cr_2O_7^{2-}/Cr^{3+}$$
 or raise E<sup>e</sup> for  $Cl_2/Cl$  }

or lower [Ct] or  $[H^{\dagger}]$  will shift equilibrium in eqn to the left hand side

 $Cr_2O_7^{2-} + 6Cl + 14H^+ \longrightarrow 2Cr^{3+} + 3Cl_2 + 7H_2O$ 

(iv) 
$$Br_2/Br^- = +1.07 \text{ V}$$
, so  $Cr(VI)$  would oxidise  $Br^-$  (easily) [1]

[Total: 9]

[1]

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4 (a)  $CCl_4$  is unreactive. (The rest react (with increasing vigour))

no d-orbitals or available/low-lying empty orbitals in carbon or unable to expand octet

e.g.  $SiCl_4 + 2H_2O \longrightarrow SiO_2 + 4HCl$ ( $or GeCl_4$  etc)  $or Si(OH)_2Cl_2$   $or Si(OH)_4$ (allow balanced equations for partial hydrolysis)

[1]

(b) (i) 
$$E(Cl-Cl) = 244 \text{ kJ mol}^{-1}$$
;  $2 E(C-Cl) = 2 \times 340 = 680 \text{ kJ mol}^{-1}$   
 $\therefore \Delta H = -436 \text{ (kJ mol}^{-1})$  [1]

(ii) 
$$\Delta H = 359 - 329 = +30 \text{ (kJ mol}^{-1})$$

(iii) since reaction (ii) is endothermic, the +4 oxidation state is less stable or the +2 oxidation state is more stable (down the group)

[1] [3] [Total: 6]

[3]

5 (a) 
$$2 \text{ MnO}_4^- + 5 \text{ H}_2\text{O}_2 + 6 \text{ H}^+ \longrightarrow 2 \text{ Mn}^{2+} + 8 \text{ H}_2\text{O} + 5 \text{ O}_2$$
 [1]

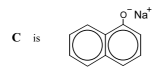
(b) 
$$E_{cell}^{\theta} = 1.52 - 0.68 = +0.84 \text{ (V)}$$
 [1]

- (c) (i) (as KMnO<sub>4</sub> is added), colour changed (from purple) to colourless *NOT* pink or effervescence/bubbles (of O<sub>2</sub>) are produced [1] at end-point, change is to (first) pink [1]
  - (ii)  $n(MnO_4^-) = 0.02 \times 15/1000 = 3 \times 10^{-4}$  [1] since  $H_2O_2 : MnO_4^- = 5:2$ ,  $\Rightarrow n(H_2O_2) = (5/2) \times 3 \times 10^{-4} = 7.5 \times 10^{-4} \text{ in } 25 \text{ cm}^3$   $\therefore [H_2O_2] = 7.5 \times 10^{-4} \times 1000/25 = 3.0 \times 10^{-2} \text{ mol dm}^{-3}$  [1] [4]

[Total: 6]

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## 6 (a) (i)



allow ONa but no covalent O-Na bond

[1]

(ii) amide, ester 2 x [1]

(iii)  $CO_2 \ or \ H_2CO_3 \ or \ Na_2CO_3$  [1]  $CH_3NH_2 \ or \ CH_3NH_3^{\dagger}CI^{-}$  [1]

(iv) 
$$H_3O^+$$
 and heat >80° or  $OH^-$ (aq) and heat >80° [1]

(ii) 
$$tin/Fe + HCl NOT LiAlH_4$$
 [1]

(iii)

(d) (i) (allow any orientation of groups)

(ii)  $[Cu(NH_3)_4]^{2+}$  or  $[Cu(NH_3)_4(H_2O)_2]^{2+}$  NOT  $[Cu(NH_3)_6]^{2+}$  [1]

(iii) ligand substitution/exchange [1]

[Total: max 15]

2 x [1] **[4]** 

[1]

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7 (a) 
$$HNO_3 + H_2SO_4$$
 [1] at  $50 - 60^{\circ}C$  (or  $\leq 60^{\circ}C$ ) not dilute or (aq) [2]

(b) 
$$2H_2SO_4 + HNO_3 \longrightarrow 2HSO_4^- + H_3O^+ + NO_2^+$$
 (allow equ. with only one  $H_2SO_4$ , giving  $H_2O$ ) [1]



reaction I:  $Cl_2 + AlCl_3/accept$  other halogen carriers NOT aq, nor u.v.

reaction II:  $KMnO_4 + H^+NOTHClnorHNO_3$ reaction III:  $KMnO_4 + H^+NOTHClnorHNO_3$ 

reaction IV:  $Cl_2 + AlCl_3/accept$  other halogen carriers NOT aq, nor u.v.

<u>both</u> I + IV [1] <u>both</u> II + III [1] **[4]** 

[Total: 7]

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8	(a)	(i)	Two	interlinked spirals or chains or strands woven round e	ach other	[1]
		(ii)	By h	ydrogen bonds between bases		[1] <b>[2]</b>
	(b)	Tra	nscrip	otion – (1)DNA/RNA/nucleic acid unravels – (2)strand is used as a template – (3)mRNA reads the sequence on this strand/		[1] [1]
				produces complementary strand		[1]
		Tra	nslati	<ul><li>– (5)tRNA translates the codon from mRNA</li></ul>	a a ta shain	[1] [1]
				<ul> <li>(6)tRNA carries amino acids to ribosome/adds</li> </ul>	a.a. to chain	[1] [max 4]
	(c)	(i) (ii)	(cou	uption of the secondary/tertiary/quaternary/3D structure ld be answered in terms of bonds e.g. hydrogen bonds covalent/peptide bonds in the (protein) chain are too s	s break)	[1] [1] <b>[2]</b>
	(d)	ATI ATI	P (+ F P is p	s provided by the breakdown/hydrolysis of adenosine t d₂O) → ADP + P <sub>i</sub> (+ energy) or in words roduced during respiration/Krebs cycle/oxidation of glu londria/ADP is recycled		P) [1]

Mark Scheme

Syllabus

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[Total: 11]

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9	(a) Suitable diagram showing origin of two energy states/or description Needs to mention applied magnetic field/electron transfer negates Indication that energy difference is in the radio frequency range				[1] [1]
			on that frequency of absorption <i>or</i> gap between the 2 e		ניו
		depend	s on the nature of nearby atoms <i>or</i> the chemical enviro	nment of the <sup>1</sup> H	[1] <b>[3]</b>
	(b)	They ar	o not damage tissues/X-rays harmful/NMR of lower ene e not obscured by bones/skeleton in be tuned to examine particular tissues/tumours/orgai		[1] [1] [1] <b>[max 2]</b>
	(c)	(i) M:	M+1 = 100/(1.1n)		
			$\frac{0.66 \times 200}{14.5 \times 1.1} = \frac{66}{15.95} = 4.14 = 4$ carbon atoms eck for 1.1 in divisor, if missing, penalise		[1]

Quartet at  $\delta$  4 suggests a –CH<sub>2</sub>- group (adjacent to a –methyl group)

Triplet at  $\delta$  1.2 suggests a methyl group (adjacent to a –CH<sub>2</sub>–)

**G** is ethyl ethanoate (or structure)/if methyl propanoate given here

**Mark Scheme** 

(ii) Singlet at  $\delta$  2 suggests methyl adjacent to C=O

cannot score first marking point

(allow -OCH<sub>2</sub>-)

Page 9

[Total: 10]

[1]

[1]

[1]

[1] **[5]** 

Syllabus

Paper

	Page 10		)	Mark Scheme	Syllabus	Paper	
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10	(a)	Iron	Iron is higher in the reactivity series than copper (owtte)/allow use of E <sup>e</sup>			[1]	
		Cu <sup>2</sup>	$Cu^{2+}(aq) + Fe(s) \rightarrow Cu(s) + Fe^{2+}(aq)$ If conversion to $Fe^{3+}$ given, $E_{cell}$ is $-0.38$				
	(b)	It do	es n	ot require investment in machinery/labour		[1]	
		It re	quire	s little energy		[1]	
			•	produces little/no pollution/noise ccept comparison with electrolytic method		[1] [max 2]	
	(c)	The	proc	ess takes a long time/requires smaller workforce		[1] <b>[1]</b>	
	(d)	(i)	0.75	% is 7.5 kg in every tonne of ore			
				ce 150,000 tonnes of ore yield $\frac{7.5 \times 150000}{1000}$ tonnes 125 tonnes Cu			
			1125	5 x 0.6 = 675 tonnes (accept 680)		[1]	
				x 0.17 = 76.5 tonnes (accept 77) 125 x 0.17 = 191.25 tonnes (accept 191) – this is an ed	of if 675 not in <b>(i)</b>	[1] <b>[2]</b>	
	(e)	bone	ds wi	m is too high in the reactivity series/very reactive/alum th oxygen which are too strong/aluminium ore doesn't			
		/Fe	unab	le to displace A <i>l</i>		[1] <b>[1]</b>	
	(f)		Control the pH ( <i>greater</i> than pH 6.0) Bioremediation/growth of special plants (to remove heavy metals)				
		Other reasonable suggestions such as displacement by a more reactive metal/ precipitation/ion exchange		[1]			
						[2] [Total: 9]	