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

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- (a) (i) $K_a = [H^{\dagger}][RCO_2^{-}]/[RCO_2H]$ [1] (ii) $pK_a = -\log_{10}K_a \text{ or } -\log K_a \text{ or } \log [H^+]^2/[RCO_2H] \text{ NOT } ln;$ [1] [2] (b) (i) acid strength increases from no. 1 to no. 3 or down the table or as Cls increase [1] due to the electron-withdrawing effect/electronegativity of chlorine (atoms) [1] stabilising the anion or weakening the O-H bond NOT H⁺ more available [1] (ii) chlorine atom is further away (from O-H) in no. 4, so has less influence [1] $K_{\rm a} = 10^{-pKa} = 1.259 \times 10^{-3}$ [H⁺] = $\sqrt{(K_{\rm a}. c)} = 3.55 \times 10^{-4}$ (iii) either: pH = $\frac{1}{2}$ (p K_a – log₁₀[acid]) or $= \frac{1}{2} (4.9 + 2)$ ecf [1] = 3.4 (allow 3.5)([1] for correct expression & values; [1] for correct working) [6] (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) $M_r(CH_3CH_2CO_2H) = 74 M_r(CH_2CH(NH_2)CO_2H) = 89$ [1] \therefore 10.0 g should give 10 x 89/74 = 12.03 g :. percentage yield = $100 \times 9.5/12.03 = 79\%$ ecf [1] ([2] for correct answer) [5]
 - (d) [†]NH₃-CH(CH₃)-CO₂⁻ Allow charges on H of H₃N, and –COO but not –C-O-O

correct atoms [1] correct charges [1]

[2]

[Total: 15]



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2	(a)	latt sol but	ubility decreases (down Group II) ice energy decreases vation/hydration energy (of cation) decreases more so than does lattice energy/is not able to overcome LE soln becomes more endothermic/positive/less exothermic	[1] [1] [1] [1] [max 4]
	(b)		Intities of A and B $I(OH)_2 + H_2C_2O_4 \longrightarrow MgC_2O_4(aq) + 2H_2O$ (A)	2 x [1] [1]
		Mg	$C_2O_4(aq) + Ca(NO_3)_2 \longrightarrow Mg(NO_3)_2 + CaC_2O_4(s)$ (B)	[1] [max 3]
	(c)	(i)	$(K_{sp} =) [Mg^{2+}][OH^{-}]^{2}$ units are $\underline{\text{mol}^{3}\text{dm}^{-9}}$	[1] ecf from K _{sp} [1]
		(ii)	(call [Mg(OH) ₂ (aq)] = [Mg ²⁺] = x) $\therefore K_{sp} = 2 \times 10^{-11} = 4x^3$	[1]
			$x = 1.71 \times 10^{-4} \text{ mol dm}^{-3}$	ecf [1]
		(iii)	less soluble <u>because</u> of the common ion effect or the equilibrium $Mg(OH)_2(s) = Mg^{2+}(aq) + 2OH^{-}(aq)$ is moved to the left	[1] [5] [Total: 12]

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3 (a) K = 22.4/39.1 = 0.573 thus ratio is: Cr = 29.8/52.0 = 0.5731 Cl = 20.3/35.5 = 0.5721 O = 27.5/16.0 = 1.7193 or KCrC1O3 (scores 2) [2] [1] [1] (b) $K_2Cr_2O_7 + 2HCl \longrightarrow 2KCrClO_3 + H_2O$ [1] [1] (c) (i) redox or oxidation [1] (ii) E^o data and half equations: $Cr_2O_7^{2-} + 14H^+ + 6e^- \longrightarrow 2Cr^{3+} + 7H_2O$ $E^{\theta} = 1.33 \text{ V}$ [1] $Cl_2 + 2e^- \longrightarrow 2 Cl^ E^{e} = 1.36 \text{ V}$ [1] overall ionic equation: $Cr_2O_7^{2-} + 6Cl + 14H^+ \longrightarrow 2Cr^{3+} + 3Cl_2 + 7H_2O$ [1] (iii) (dilution will) lower E^{θ} for $Cr_2O_7^{2-}/Cr^{3+}$ or raise E^{θ} for Cl_2/Cl [1] or lower [Ct] or [H⁺] will shift equilibrium in eqn to the left hand side ∫

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

[Total: 9]

			<u> </u>
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4 (a)	no e.g	l_4 is unreactive. (The rest react (with increasing vigour)) d-orbitals or available/low-lying empty orbitals in carbon or unable to expand octet $sic l_4 + 2H_2O \longrightarrow siO_2 + 4HCl$ $sic l_4 + 2H_2O \longrightarrow siO_2 + 2HCl$ $sic l_4 = cor si(OH)_2Cl_2$ $sic l_4 = cor si(OH)_4$	[1] [1]
		(allow balanced equations for partial hydrolysis)	[1] [3]
(b)	(i)	E(C <i>l</i> -C <i>l</i>) = 244 kJ mol ⁻¹ ; 2 E(C-C <i>l</i>) = 2 x 340 = 680 kJ mol ⁻¹ $\therefore \Delta H = -436 \text{ (kJ mol}^{-1})$	[1]
	(ii)	$\Delta H = 359 - 329 = +30 \text{ (kJ mol}^{-1}\text{)}$	[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) [Total	[1] [3] al: 6]

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]

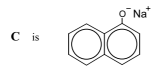
(b)
$$E_{cell}^{\theta} = 1.52 - 0.68 = +0.84 \text{ (V)}$$
 [1]

(ii)
$$n(MnO_4^-) = 0.02 \times 15/1000 = 3 \times 10^{-4}$$
 [1] since $H_2O_2 : MnO_4^- = 5:2$, $\Rightarrow n(H_2O_2) = (5/2) \times 3 \times 10^{-4} = 7.5 \times 10^{-4} \text{ in } 25 \text{ cm}^3$.: $[H_2O_2] = 7.5 \times 10^{-4} \times 1000/25 = 3.0 \times 10^{-2} \text{ mol dm}^{-3}$ [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) CO₂ or H₂CO₃ or Na₂CO₃ CH₃NH₂ or CH₃NH₃⁺C*T* [1] [1]

[1]

(iv) H_3O^+ and heat >80° or OH^- (aq) and heat >80°

[1] **[7]**

(b) (i) Br₂(aq) (or other suitable solvent)

[1]

(ii) dilute/aqueous HNO₃

[1] **[2]**

(c) (i)

[1]

(ii) tin/Fe + HCl NOT LiAlH₄

[1]

(iii)

mark each side chain separately

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

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

penalise missing H on NH₂

[1]

(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]



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7	(a)	$HNO_3 + H_2SO_4$	[1]
		at $50 - 60$ °C ($or \le 60$ °C) not dilute or (aq)	[1]
			[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: $KMnO_4 + H^+NOT HCl nor HNO_3$ reaction III: $KMnO_4 + H^+NOT HCl nor HNO_3$

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

both I + IV [1] both II + III [1] [4]

[Total: 7]

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(a)	(i) Two interlinked spirals <i>or</i> chains <i>or</i> strands woven round each other	[1]
	(ii) By hydrogen bonds between bases	[1] [2]
(b)	Transcription – (1)DNA/RNA/nucleic acid unravels – (2)strand is used as a template – (3)mRNA reads the sequence on this strand/ produces complementary strand	[1] [1] [1]
	Translation – (4)mRNA binds to the ribosome – (5)tRNA translates the codon from mRNA – (6)tRNA carries amino acids to ribosome/adds a.a. to chain	[1] [1] [1] [max 4]
(c)	(i) Disruption of the secondary/tertiary/quaternary/3D structure of the protein (could be answered in terms of bonds e.g. hydrogen bonds break)	[1]
	(ii) The covalent/peptide bonds in the (protein) chain are too strong	[1] [2]
(d)	Energy is provided by the breakdown/hydrolysis of adenosine triphosphate (ATP) ATP (+ H_2O) \rightarrow ADP + P_i (+ energy) or in words ATP is produced during respiration/Krebs cycle/oxidation of glucose, fats or prote in mitochondria/ADP is recycled	[1]

[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 Indication that frequency of absorption <i>or</i> gap between the 2 energy states	[1] [1]
		depends on the nature of nearby atoms <i>or</i> the chemical environment of the ¹ H	[1] [3]
	(b)	They do not damage tissues/X-rays harmful/NMR of lower energy They are not obscured by bones/skeleton They can be tuned to examine particular tissues/tumours/organs/protons	[1] [1] [1] [max 2]

(c) (i) M: M+1 = 100/(1.1n)

$$n = \frac{0.66 \times 200}{14.5 \times 1.1} = \frac{66}{15.95} = 4.14 = 4 \text{ carbon atoms}$$
 [1] Check for 1.1 in divisor, if missing, penalise

(ii)	Singlet at δ 2 suggests methyl adjacent to C=O	[1]
(,	Quartet at δ 4 suggests a –CH ₂ - group (adjacent to a –methyl group)	[1]
	(allow –OCH ₂ -)	r.,
	Triplet at δ 1.2 suggests a methyl group (adjacent to a –CH ₂ –)	[1]
	G is ethyl ethanoate (or structure)/if methyl propanoate given here	
	cannot score first marking point	[1]
		[5]

[Total: 10]



[Total: 9]

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10 (a) Iron is higher in the reactivity series than copper (owtte)/allow use of E^e [1] $Cu^{2+}(aq) + Fe(s) \rightarrow Cu(s) + Fe^{2+}(aq)$ [1] If conversion to Fe^{3+} given, E_{cell} is -0.38[2] [1] **(b)** It does not require investment in machinery/labour It requires little energy [1] accept it produces little/no pollution/noise [1] Do not accept comparison with electrolytic method [max 2] (c) The process takes a long time/requires smaller workforce [1] [1] (d) (i) 0.75% is 7.5 kg in every tonne of ore Hence 150,000 tonnes of ore yield $\frac{7.5 \times 150000}{1000}$ tonnes or 1,125 tonnes Cu $1125 \times 0.6 = 675 \text{ tonnes (accept } 680)$ [1] (ii) $450 \times 0.17 = 76.5 \text{ tonnes (accept 77)}$ [1] or 1125 x 0.17 = 191.25 tonnes (accept 191) - this is an ecf if 675 not in (i) [2] (e) Aluminium is too high in the reactivity series/very reactive/aluminium forms bonds with oxygen which are too strong/aluminium ore doesn't exist as sulphide /Fe unable to displace A1 [1] [1] **(f)** Control the pH (*greater* than pH 6.0) [1] Bioremediation/growth of special plants (to remove heavy metals) Other reasonable suggestions such as displacement by a more reactive metal/ precipitation/ion exchange [1]

