UNIVERSITY OF CAMBRIDGE INTERNATIONAL EXAMINATIONS

GCE Advanced Level and GCE Advanced Subsidiary Level

MARK SCHEME for the May/June 2006 question paper

9701 CHEMISTRY

9701/06

Paper 6

Maximum raw mark 40

This mark scheme is published as an aid to teachers and students, to indicate the requirements of the examination. It shows the basis on which Examiners were initially instructed to award marks. It does not indicate the details of the discussions that took place at an Examiners' meeting before marking began. Any substantial changes to the mark scheme that arose from these discussions will be recorded in the published *Report on the Examination*.

All Examiners are instructed that alternative correct answers and unexpected approaches in candidates' scripts must be given marks that fairly reflect the relevant knowledge and skills demonstrated.

Mark schemes must be read in conjunction with the question papers and the *Report on the Examination*.

The minimum marks in these components needed for various grades were previously published with these mark schemes, but are now instead included in the Report on the Examination for this session.

• CIE will not enter into discussion or correspondence in connection with these mark schemes.

CIE is publishing the mark schemes for the May/June 2006 question papers for most IGCSE and GCE Advanced Level and Advanced Subsidiary Level syllabuses and some Ordinary Level syllabuses.



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Biochemistry

1	(a)	(i)	O CH ₃ O		
			II II NH ₂ – CH ₂ – C- NH -CH – C-NH – CH ₂ – COOH	[1]	
			Displayed structure	[1]	
			One peptide linkage shown	[1]	
		(ii)	Condensation	[1]	[4]
	(b)	(i)	Weak intermolecular forces of attraction (1) Van der Waals (1)	[2]	
		(ii)	No attraction/ffinity for water	[1]	
			Non-polar structure	[1]	[4]
	(c)	(i)	Both contain the polyamide structure/-CONH-	[1]	
		(ii)	Bullet proof vests; body armour; ropes; airbags; kayaks; gloves; skis; run-flat tyres; shields for jet engines; helmets; racquets; clothing any o	ne [1]	[2]
2	(a)	Dia	gram:	[1]	
		Re	end – van der Waals forces	[1]	
		Pho	osphate end - ionic/polar	[1]	[3]
	(b)	(i)	van der Waals interactions/dipole –dipole interactions/temporary dipoles/hydrinteractions	ophobic	;
			with the hydrocarbon part of the bilayer	[1]	
		(ii)	Disrupt it/distort it/weakens the interactions between the bilayers	[1]	[3]
	(c)	K⁺ r	moves into cell, Na⁺ moves out of cell	[1]	
		Thi	s occurs by active transport	[1]	
		ATI	P/adenosine triphosphate provides the energy	[1]	
		Inte	egral proteins in the membrane are used	[1]	[4]

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Environmental Chemistry

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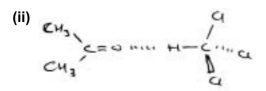
Phase Equilibria

5 (a) (i) V.P. of A = vapour pressure of A on own x mol fr. of A [1]

OR $P_A = P_A x_A$

(ii)
$$0.3 \times 48 = 14.4$$
 [1] $0.7 \times 36 = \frac{25.2}{39.6 \text{ k Pa}}$ [1]

- (iii) Raoult's law obeyed [1]
- components are similar/ideal mixture/components not interact [1] [5]
- (b) (i) Molecules attract each other OR dipoles align
 Stronger intermolecular forces than components



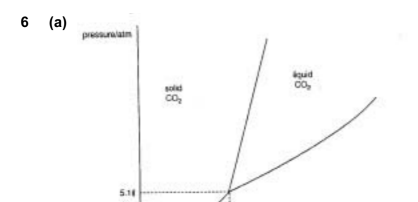
- OR Interact in 1:1 ratio [1] [2]
- (c) pure propanone [1]

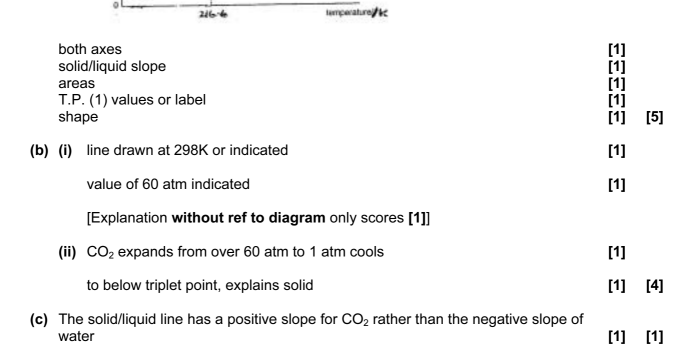
since this has lowest b.p. OR highest VP
OR is most volatile
(allow discussion of b.p./composition curves)

[1]

Then azeotrope or 0.50 composition [1] [3]

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Spectroscopy

7	(a)	Two	o absorptions	[1]	
		Asy	mmetric bend (or diagram)	[1]	
		Asy	mmetric stretch (or diagram)	[1]	[3]
	(b)	(i)	1710 cm ⁻¹ – C=O	[1]	
			2260 cm ⁻¹ – C≡N	[1]	
			3200 cm ⁻¹ – O–H	[1]	
		(ii)	NC-CH ₂ -CH ₂ -CO ₂ H	[1]	[4]
	(c)	Nm	r	[1]	
		+ in	dication of absorptions (CH ₂ \sim 1.3 δ , -O-H \sim 4.5 δ)	[2 x 1]	
		OR	Mass spectrometry	[1]	
			vo examples of likely fragmentations e.g. M-28 (loss of CN) and 7 (loss of –OH)	[2 x 1]	[3]
8	(a)	Stru	ucture II	[1]	
		Αp	eak at 3450 cm ⁻¹ is characteristic of -OH would be seen for structure II	[1]	[2]
	(b)	(i)	Triplet-quartet is characteristic of a CH ₃ next to CH ₂ group	[1]	
			Standard 1,3,3,1 and 1,2,1 diagrams	[1]	
		(ii)	Singlet (1) at δ 2.0 – 3.8 (1)	[2]	
		(iii)	Deuterium oxide will exchange protons with -OH group in structure II	[1]	
			Since deuterium does not absorb in this part of the spectrum the –OH peak would disappear	[1]	[6]
	(c)	Stru	ucture II will show (M-17) ⁺ - loss of OH		
		Stru	ucture I will show (M-31) ⁺ loss of CH ₃ O		
		Stru	ucture II will show (M-43) ⁺ loss of C ₃ H ₇	any tw o	o [2]

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Transitions Elements

(a) (i) somewhere between 4% and 20% chromium 9 [1] (ii) Cr forms its oxide/Cr₂O₃ on the steel's surface [1] which is impermeable to oxygen/hard [1] [3] **(b) (i)** Cr = 33.6/52 = 0.646O = 20.6/16 = 1.288Cl = 45.8/35.5 = 1.290[1] thus A is CrO₂Cl₂ [1] O.N. of chromium = +6[1] (ii) orange solution contains Cr₂O₇²⁻ [1] $2CrO_2Cl_2 + 3H_2O \longrightarrow Cr_2O_7^{2-} + 6H^+ + 4Cl^-$ [1] white ppte is AgCl or Ag⁺ + Cl⁻ \longrightarrow AgCl(s) [1] yellow solution contains CrO₄²⁻ [1] $Cr_2O_7^{2-} + 2OH^- \longrightarrow 2CrO_4^{2-} + H_2O$ [1] [8 max 7] [7] 10 (a) colour dues to absorption of visible light [1] d-orbitals are split into two sets at different energies [1] photon is absorbed when an electron is promoted to higher orbital [3] [1] (b) (i) [Fe(SCN]²⁺ is formed - this is red [1] F⁻ is a stronger ligand than SCN⁻ or ligand exchange occurs [1] [FeF₆]³⁻ is colourless *or* energy gap between d-orbitals is large [1] (ii) reduction occurs [1] to VO²⁺ (which is blue) [1] $SO_2 + 4H^+ + 2VO_3^- \longrightarrow SO_4^{2-} + 2VO^{2+} + 2H_2O$ [1] (further reduction to) V³⁺ (which is green) [1] $Sn^{2+} + 4H^+ + 2VO^{2+} \longrightarrow Sn^{4+} + 2V^{3+} + 2H_2O$ [1]

[8 max 7]

[7]