CAMBRIDGE INTERNATIONAL EXAMINATIONS

GCE Advanced Level

MARK SCHEME for the May/June 2014 series

9701 CHEMISTRY

9701/43

Paper 4 (Structured Questions), maximum raw mark 100

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Pag	e 2	Mark Scheme	Syllabus	Paper
		GCE A LEVEL – May/June 2014	9701	43
		Section A		
(a) ((i)	m. pt. is high(er)/large(r)/greater (for iron) density is high(er)/large(r)/greater (for iron)		
(i	ii)	(higher m. pt. due to) strong attraction between cations and electrons of more delocalised electrons	or	
		(higher density due to) greater A_{r} and smaller rad	ius	
(b) ((i)	components to be added: voltmeter <i>or</i> V salt bridge [must be labelled]		
(i	ii)	M1: A and B copper (metal) or Cu and	iron (metal) or Fe	

as 1 mol dm⁻³/1 M

Fe²⁺ or FeSO₄ etc.

or (end point is the first) permanent (pale) pink/pale purple colour

mass of Fe = $55.8 \times 1.81 \times 10^{-3} = 0.101 \text{ g} (M2 \times 55.8) \text{ ecf}$

metal atom/ion/or a lone pair donor to metal atom/ion

Cu²⁺ or CuSO₄ or CuCl₂ or Cu (NO₃)₂ etc. and

(iv) if C is Fe^{2+} ; (as [C] increases), the E of the Fe^{2+} /Fe increases/becomes more positive/

so the overall cell potential/ E_{cell} would decrease/become less positive/more

if ${\bf C}$ is Cu^{2+} ; (as $[{\bf C}]$ increases), the E of the Cu^{2+}/Cu increases/becomes more

so the overall cell potential/E_{cell} would increase/become more positive/less negative

(a) (i) A complex is a compound/molecule/species/ion formed by a central metal atom/ion surrounded by/bonded to one or more ligands/groups/molecules/anions [1]

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A ligand is a species that contains a lone pair of electrons that forms a dative bond to a

[1]

[1]

[1]

[1]

[1]

[1]

[1]

[1]

[1]

[1]

[1]

[Total: 16]

either **C** or **D**

C and D

(iii) $E_{cell}^{\theta} = 0.34 + 0.44 = 0.78 \text{ (V)}$

positive/less negative

(c) (i) (colour change is) colourless to pink/pale purple

(ii) $\{n(MnO_4^-) = 0.02 \times 18.1/1000 = 3.62 \times 10^{-4} \text{ mol}\}\$ $n(Fe^{2^+}) = 5 \times n(MnO_4^-) = 1.81 \times 10^{-3} \text{ mol}$

 $M_c = \text{mass/moles} = 0.500/1.81 \times 10^{-3} = 276.2 \text{ ecf}$

less negative

negative

or

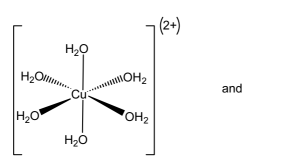
2

M2:

M3

Page 3	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2014	9701	43

(ii)



correct 3D structures: octahedral and tetrahedral [1] + [1][1]

(iii)

$$CI_{M_{1},...}$$
 CI $CI_{M_{1},...}$ $CI_{M_{1},...}$

 NH_3 -CI $\dot{N}H_3$

both structures geometric or cis-trans [1] [1]

(b) (i)
$$Cu(II)$$
 is [Ar] $3d^9$ $Cu(I)$ is [Ar] $3d^{10}$

[1] [1]

d orbitals/subshell are split (in ligand field) and (ii) Cu(II):

electron moves from lower to upper orbital or an electron is promoted/excited

in doing so it absorbs a photon/light

[2]

Cu(I): no gap in upper orbital/all orbitals are full [1]

(c) (i)
$$\Delta H^{e} = +2 \times 33.2 - 157.3 + 302.9 = (+) 212 \text{ kJ mol}^{-1} \text{ ecf}$$
 [2]

(ii)
$$\Delta H^{\text{e}} = -168.6 + 2 \times 157.3 = (+)146 \text{ kJ mol}^{-1}$$
 allow ecf from (c)(i) [1] high T/temperature since ΔH is positive/endothermic [1]

high T/temperature since ΔH is positive/endothermic

[Total: 16]

3 (a) heat in dilute HCl(aq) (or $H_2SO_4(aq)$) [1]

(b) (i) four isomers

[1]

Page 4	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2014	9701	43

(ii) must be skeletal

[1]

(iii)
$$CO_2H$$
 $+ CO_2 \text{ or } HO_2C-CO_2H$ [1]

(c) (i)
$$K_{w} = [H^{+}][OH^{-}]$$
 [1]

- (ii) In $0.15 \,\text{mol dm}^{-3} \,\text{NaOH}$, $[OH^{-}] = 0.15 \,\text{mol dm}^{-3}$ $[H^{+}] = K_{\text{w}}/[OH^{-}]$, so $[H^{+}] = 1 \times 10^{-14}/0.15 = 6.67 \times 10^{-14} \,\text{mol dm}^{-3}$ [1] $pH = -\log_{10}[H^{+}] = 13.18 \,(13.2) \,\text{ecf from } [H^{+}]$ [1]
- (iii) piperidine is a poorer proton acceptor or piperidine is partially ionised [1]
- (iv) piperidine should be a **stronger base/more basic** than ammonia because of the electron-donating (alkyl/CH₂) groups [1]

(d) (i)
$$n(HCl)$$
 at start = $0.1 \times 20/1000 = 2.0 \times 10^{-3} \text{ mol}$
 $n(HCl)$ at finish = $2 \times 10^{-3} - 1.5 \times 10^{-3} = 0.0005/5 \times 10^{-4} \text{ mol}$ [1]

(ii) this is in 30 cm³ of solution, so [HC] at finish =
$$0.5 \times 10^{-3}/0.030 = 1.67 \times 10^{-2} \text{ mol dm}^{-3}$$

pH = $-\log_{10}(1.67 \times 10^{-2}) = 1.78$ ecf from (d)(i) [1]

- (iii) pH/vol curve: start at pH 11.9 [1] vertical portion at V = 15 cm³ [1] levels off at pH 1.8
- (iv) indicator is B [1]

[Total: 16]

4 (a) three from phenol

(secondary) alcohol (primary) amine arene/aryl/benzene

3 × [1]

Page 5	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2014	9701	43

(b) (i)

step 1: HCN + NaCN or HCN + base [1]

step 2:
$$H_2$$
 + Ni or LiA lH_4 or Na + ethanol [1]

(ii) bromine decolourises or goes from orange to colourless or white ppt. formed

e.g.

HO
CHO
2 or 3 bromines in ring
Br

[1]

(c)

(i)
$$N_{AO}$$
 N_{AO} N_{AO}

(ii)
$$\stackrel{\text{HO}}{\underset{\text{HO}}{\bigvee}} \stackrel{\text{OH}}{\underset{\text{NH}_3}{\bigvee}}$$
 (or ionic) [1]

M1: amide [1]
M2: alcoholic ester [1]
M3: both phenolic esters [1]

[5] max [4]

(d) amide [1] ester

[Total: 14]

[1]

[1]

Page 6	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2014	9701	43

- **5** (a) (i) –OH *or* hydroxyl groups (allow alcohol groups) [1]
 - (ii) alkenes or C=C (double) bonds or carbon double bonds [1]
 - (iii) CH₃CH(OH) or CH₃CO- groups [1]
 - (b) V is $CH_3CH(OH)CH=CH_2$ [1]
 - **W** is CH₃CH=CHCH₂OH [1]
 - (c) compound V shows optical isomerism

(ecf for 'geometric(al)' if candidate's V is capable of cis-trans) [1]

$$H_2C$$
 CH_3 H_2C CH_3 HO CH CH CH CH CH CH CH

[Total: 8]

Page 7	Mark Scheme	Syllabus	Paper
	GCE A LEVEL - May/June 2014	9701	43

6 (a)

feature	level of bonding
formation of α-helix	secondary
formation of disulfide bonds	tertiary
formation of ionic bonds	tertiary
linking amino acids	primary

[3]

(b)

block letter	name
J	Deoxyribose
К	Cytosine
L	Phosphate
М	Thymine

4 × [1]

(c) (i) H/hydrogen (bonds between bases)

[1]

(ii) Bonds are weak **and** so require relatively little energy to break/are easily broken

[1]

(d)

	(sugar, J)	(base, M)
DNA	deoxyribose	thymine/T
RNA	ribose	uracil/U

[1]

[Total: 10]

7 (a) Expression:
$$n = \frac{100 \times 2.5}{1.1 \times 74}$$
 or equivalent [1]

n = 3.1 hence **G** has three carbon atoms

[1]

(b) (i) $(\delta 1.1)$ RCH₃ or RCH₂R or methyl or CH₃

(δ 2.2) (R)CH₂CO(R) or CH₃CO(R)

(δ 11.8) (R)COOH or (R)CONH(R)

 $3 \times [1]$

Page 8	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2014	9701	43
(ii)	The (–OH) peak at δ 11.8 (disappears)		[
	because of (O)H-D exchange <i>or</i> equation showing this (e.g. R-OH + $D_2O \rightleftharpoons R$ -OD + HOD)		['
(iii)	CH ₃ CH ₂ CO ₂ H		[
(c) (i)			
(c) (i)	H_3C O	HO HO	ОН
	or N—C H]
(ii)	If methyl ethanoate: δ 2.0–2.1 δ 3.3–4.0]
	Or if 1, 3-dioxolane: δ 3.3–4.0 δ 3.3–5.0]
	Or if 1, 2-dioxolane: δ 0.9–1.4 δ 3.3–4.0]
	Or if dihydroxycyclopropane: δ 0.9–1.4 δ 0.5–6.0		[
			[Total: 1
(a) (i)	Amide or ester or peptide		[
(ii)	Hydrolysis		[
(iii)	Drug B		[
(iv)	two ester and one amide groups circled		[
(b) (i)	At point Q because the hydrocarbon tails region is hydrophobic/non-polar/ form van der Waals only or can dissolve in the fat-soluble area		
(ii)	They all contain polar <i>or</i> hydrogen-bonding (groups)		[
	range 1×10^{-9} to 1×10^{-7} m		[

[1]

[Total: 9]

(ii) (higher frequency radiation could) cause tissue/cell damage or mutation

or harmful to cells