MARK SCHEME for the May/June 2013 series

9701 CHEMISTRY

9701/41

Paper 4 (A2 Structured Questions), maximum raw mark 100

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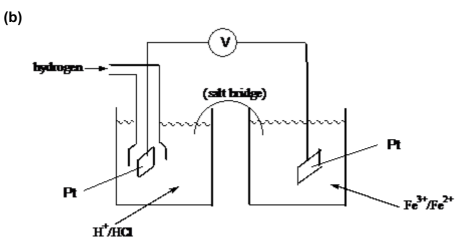
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(a) The potential of an electrode compared to that of a standard hydrogen electrode (SHE) or the EMF of a cell composed of the test electrode and the SHE
 [1]

all measurement concentrations of 1 mol dm⁻³ and 298 K/1 atm pressure [1]



 H_2 and good delivery system [1] Fe^{2+}/Fe^{3+} solution labelled [1] platinum electrodes (both) [1] salt bridge and voltmeter [1] H^+ or HCl or H₂SO₄ [1] (acid is not sufficient) [5]

(c) (i)
$$E^{\ominus} = 0.77 - 0.54 = 0.23$$
 (V) [1]

(ii) Since E^{\ominus} is positive/ $E^{\ominus} > 0$

So more products / the equilibrium will be over to the right / forward reaction is favoured ecf from (c)(i) [1]

(iii) $K_c = [Fe^{2+}]^2 [I_2] / [Fe^{3+}]^2 [I^-]^2$ [1]

units are **mol⁻¹ dm**³ ecf on expression

(iv) ([Fe²⁺] must always be twice [I₂], so) [Fe²⁺] = 0.02 (mol dm⁻³) [1]

([I⁻] must always be equal to [Fe³⁺], so) [I⁻] = 2×10^{-4} (mol dm⁻³) [1]

(v) $K_c = \{(0.02)^2 \times 0.01\} / \{(2 \times 10^{-4})^2 \times (2 \times 10^{-4})^2\}$ correct expression [1] (allow ecf from incorrect expression in (c)(iii)) (allow ecf from (c)(iv)) = $(4 \times 10^{-6}) / (1.6 \times 10^{-1.5}) = 2.5 \times 10^9 (\text{mol}^{-1} \text{ dm}^3)$ [1]

[8]

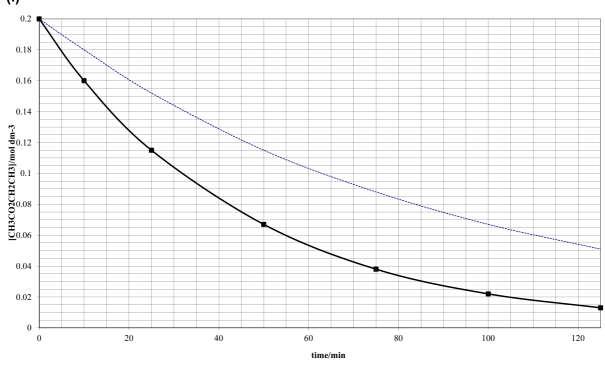
[1]

[2]

[Total: 15]

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plotting of points (-1 for any error - plotted to within 1/2 square) [1] a good best fit curve [1]

(ii) construction lines for two half-lives and $t_{\frac{1}{2}} \approx 63$ m or 32 m (±3 min) / $t_{\frac{1}{2}}$ is constant or construction lines for two tangents and mention of two values / concentration doubled, [1] rate doubled (iii) either ratio of (initial) rates (slopes) or ratio of $t_{\frac{1}{2}}$ = 2.0 [1]

so reaction is first order w.r.t. [HC1]

because HCl is a catalyst

(ii)

(iv) rate = k[CH₃CO₂CH₂CH₃][HC*l*] conditional on (a)(iii) and ecf from (a)(iii) [1]

(initial) rate = 0.2/95 or 0.2/47	
≈ 2.1×10^{-3} or 4.3×10^{-3} (mol dm ⁻³ min ⁻¹)	[1]

$$k = 2.1 \times 10^{-3} / (0.2 \times 0.1) \text{ or } 4.3 \times 10^{-3} / (0.2 \times 0.2)$$

$$\approx 0.11 \text{ (mol}^{-1} \text{ dm}^3 \text{ min}^{-1})$$
[1]

[8 max 7]

- (b) (i) because H₂O is the solvent or its concentration cannot change [1]
 - [1] [2]

[1]

[Total: 9]

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3	(a) (i)	dens	ity = mass per unit volume		[1]
	(ii)	-	s per atom <i>or A</i> _r is larger (for Fe)		
		<i>Or</i> Fe 5	5.8 and Ca 40.1		[1]
			adii/volume of atom/ion is smaller		
		or R _{Fe} =	= 0.116 nm whereas R _{Ca} = 0.197 nm		[1] [3]

(b)

reaction	acid- base	ligand exchange	precipitation	redox
$[Cu(H_2O)_6]^{2+} + 4NH_3 \rightarrow [Cu(NH_3)_4]^{2+} + 6H_2O$		\checkmark		
$[Cu(H_2O)_6]^{2+}$ + $4HCl \rightarrow [CuCl_4]^{2-}$ + $4H^+$ + $6H_2O$		\checkmark		
$2FeC\mathit{l}_2 + C\mathit{l}_2 \rightarrow 2FeC\mathit{l}_3$				\checkmark
$[Fe(H_2O)_6]^{2+} + 2OH^- \rightarrow Fe(OH)_2 + 6H_2O$	~		\checkmark	
$2Fe(OH)_2 + \frac{1}{2}O_2 + H_2O \rightarrow 2Fe(OH)_3$				\checkmark
$CrO_3 + 2HCl \rightarrow CrO_2Cl_2 + H_2O$	~	\checkmark		
$\begin{array}{c} Cr(H_2O)_3(OH)_3 + OH^- \to [Cr(H_2O)_2(OH)_4]^- + \\ H_2O \end{array}$	\checkmark	\checkmark		
$[Cr(OH)_4]^- + 1\frac{1}{2}H_2O_2 + OH^- \rightarrow CrO_4^{2-} + 4H_2O$		\checkmark		\checkmark

(Where more than one tick appears on a line in the table above – these are alternatives – but allow the mark if both are given).

[8]

(c)	n(H ₂) = 8/24 = 0.33 mol	[1]
	from equation, this is produced from 0.22 mol of A1 ecf (\times 2/3)	[1]

from equation, this is produce	d from 0.22 mol of A <i>l</i> ect (\times 2/3)	[1]

$A_r(Al) = 27$ thus mass of $Al = 27 \times 0.22 = 5.9 - 6$ g hence 5.9–6.0% ecf (× 27)	[1]
	[3]

[[]Total: 14]

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4 (a) (du	e to tl	ne) strong N≡N bond		[1] [1]
(b) (i)	e.g.	balanced equation forming a stable nitrogen oxide $N_2 + O_2 \longrightarrow 2NO$		
	or N ₂ +	$2O_2 \longrightarrow 2NO_2$		[1]
(ii)	in lig	htning		[1]
	in ar	engine/combustion of fuels (or a specific example)		[1]
(iii)	(NO ₃	, produces) acid rain <i>or</i> forms (photochemical) smog		[1] [4]
(c) (ba	se is a	a) proton acceptor		[1]
bas	sicities	: ethylamine > NH_3 > phenylamine		[1]
eth	ylamiı	ne (more basic) due to electron donating ethyl group		[1]
phe	enylar	nine (less basic) due to lone pair being delocalised int	o the ring	[1] [4]
(d) (i)	step	1: nucleophilic substitution		[1]
	step	2: hydrolysis		[1]
(ii)	step	1: KCN (in ethanol) and reflux		[1]
	step	2: H_3O^+ / aqueous acid and reflux		[1]
(iii)	T is	\sim NH_2		
	Ľ			[1]
	W is	, ⊂ CI		
	Í			
		0		[1]
				[6]
				[Total: 15]

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5 (a)

	H ₂ O	OH OH	CO ₂ H	OH
Na	H ₂	H ₂	H ₂	H ₂
KOH(aq)	х	x	x	x
Na ₂ CO ₃ (aq)	х	x	CO ₂	x

[5]

(b) (i)	(CH ₃) ₃ C–C <i>l</i> (any unambiguous structure <i>or</i> name)	[1]
(ii)	reduction or hydrogenation	[1]
(iii)	<i>either</i> CH ₃ CO ₂ H and heat with (conc) H ₂ SO ₄ or CH ₃ COC <i>l</i>	[1]
(iv)	reflux	[1]
	dilute HC1	[1] [5]

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(c) (i)

reagent and conditions	product with A	product with B
Br₂(aq)	Br C(CH ₃) ₃	no reaction
heat with HBr	no reaction	Br C(CH ₃) ₃
pass vapour over heated A <i>l</i> ₂ O ₃	no reaction	C(CH ₃) ₃
heat with acidified K ₂ Cr ₂ O ₇	no reaction	C(CH ₃) ₃

[6]

 (ii) either: Cr₂O₇²⁻/H⁺: no observation with A and goes from orange to green with B. or: Br₂(aq): white ppt. with A and no observation/ppt with B

[1] **[7]**

[Total: 17]

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6 (a)

substance	protein synthesis	formation of DNA
adenine		\checkmark
alanine	\checkmark	
aspartate	\checkmark	
phosphate		✓

[3] [3]
[1]

(b) protei	(b) protein : hydrogen bonds			
	between –NH and C=O groups on different (peptide) groups	[1]		
DNA :	hydrogen bonds	[1]		
	between bases / A & T / C & G on different chains	[1] [4]		

(c) primary: covalent bonds between (successive) amino acids

tertiary :

hydrogen bonds	between –COOH / –OH and –NH $_2$ (in side chains)
ionic bonds	between $-NH_3^+$ and $-CO_2^-$ (in side chains)
disulfide bonds	between cysteine molecules / residues / –SH groups (in side chains)
van der Waals/VDW forces	between alkyl groups / non-polar residues (in side chains)

any two rows

[2] **[3]**

[1]

[Total: 10]

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7 (a) Any four from:

- extract DNA
- use restriction enzymes (to break DNA into fragments)
- use polymerase chain reaction (to increase concentration of fragments)
- place samples on (agarose) gel
- carry out electrophoresis
- label fragments (transferred to a membrane) with radioactive isotope

[4 × 1] **[4]**

(b)

item for testing	suitable for DNA fingerprinting
human hair	\checkmark
piece of a flint tool	×
piece of Iron Age pot	×
piece of Roman leather	\checkmark

	[3] [3]
	[1]

(c)	(c) insecticides: gas-liquid or thin-layer chromatography				
	dyes	: paper <i>or</i> thin-layer chromatography	[1]		
	drugs:	gas-liquid <i>or</i> thin-layer chromatography	[1] [3]		

[Total: 10]

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	Page 10		Mark Scheme	Syllabus	Paper
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8	(a) (i)		$\begin{array}{ccc} CO_2H & CO_2H \\ I & I \\ -CH - CH_2 - CH - CH_2 - $		[1]
	(ii)	Addi	ition		[1]
	(iii)	Hydi	rogen bonding		[1] [3]
	(b) (i)	more	e / increase water absorbing properties (allow attracts	water more)	[1]
		more	e polar(ity)/more hydrophilic / has ionic side-chains (as	well as hydrophi	lic ones) [1]
	(ii)	lt sh	ould be biodegradable/decompose		[1] [3]
	(c) idea	a of ic	on exchange / replacement of Na ⁺ for Cd ²⁺ /Pb ²⁺		[1]
	(the	e meta	al ions) will be attracted to the carboxylate ions		[1] [2]
	(d) (i)	cond	densation		[1]
	(ii)		alcohol groups ighly soluble / able to form hydrogen bonds		[1] [2]
					[Total: 10]