

**MARK SCHEME for the May/June 2010 question paper
for the guidance of teachers**

9701 CHEMISTRY

9701/43

Paper 4 (A2 Structured Questions), 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, which would have considered the acceptability of alternative answers.

Mark schemes must be read in conjunction with the question papers and the report on the examination.

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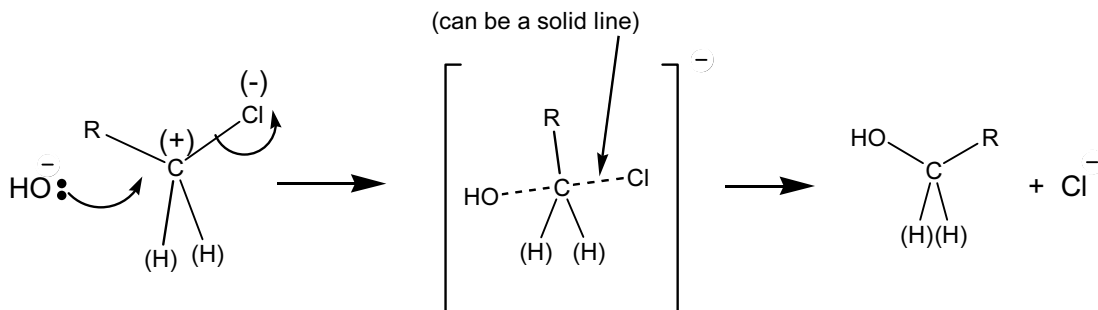
1 (a) $C_6H_5\text{-COCH}_2\text{OH}$ or $C_8H_8O_2$ and NaCl or Cl^- (1) + (1) [2]

(b) (i) the exponent / power to which a concentration is raised in the **rate equation** (or in an equation, e.g. " a " in the equ: rate = $k[A]^a$) (1)

(ii) from 1 and 2: rate increases by 50% as does $[\text{RCl}]$, so rate $\propto [\text{RCl}]^1$ (1)
from 1 and 3: rate $\propto [\text{NaOH}]^1$ (1)

(iii) (rate =) $k[\text{RCl}][\text{OH}^-]$ (1)

(iv)



marking points:

- (+) or δ^+ on C and (–) or δ^- on Cl (1)
- lone pair and charge on: OH^- (1)
- curly arrow from OH (lone pair) to $(\delta^+)C$, and either a curly arrow breaking C-Cl bond or 5-valent transition state (ignore charge) (1)
- $\text{S}_{\text{N}}1$ alternative for last mark (only award mark if candidate's rate equation shows first order reaction): curly arrow breaking C-Cl bond and carbocation intermediate. (1)

[7]

(c) (i) (add RCl / RCOCl to) (aq) Ag^+ / AgNO_3 or named indicator (e.g. MeOr) or use pH probe (1)

White ppt appears (faster with RCOCl) or turns acidic colour (e.g. red) or shows pH decrease (1)

if water is the only reagent, and no pH meter used: award only the second mark, for "steamy / white fumes"

(ii) (C=O is polarised /) carbon is more δ^+ than in R-Cl or carbon is positive or RCOCl can react via addition-elimination (mention of electronegativity on its own is not enough for the mark) (1) [3]

[Total: 12]

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2 (a) less soluble down group (1)

lattice energy and hydration energies both decrease (i.e. become less negative) (1)

but H.E. decreases more (than L.E.) or change in H.E. outweighs L.E. (1)

so ΔH_{sol} becomes more endothermic / less exothermic (1) [4]

(b) (i) for Mg: $\Delta H = 2993 - 1890 - (2 \times 550) = (+)3$ (kJ mol⁻¹) (1)

for Sr: $\Delta H = 2467 - 1414 - (2 \times 550) = -47$ (kJ mol⁻¹) (1)

(ii) Sr(OH)₂ should be **more** soluble in water, **and** ΔH is more exothermic / negative (1)

Assuming “other factors” (e.g. ΔS , or temperature etc.) are the same (1)

(iii) Sr(OH)₂ should be **less** soluble in hot water, **because** ΔH is negative / exothermic (1) [5]

(c) (i) $K_{\text{sp}} = [\text{Ca}^{2+}][\text{OH}^-]^2$ (needs the charges) units: mol³dm⁻⁹ (1) + (1)

(ii) $n(\text{H}^+) = n(\text{OH}^-) = 0.05 \times 21/1000 = 1.05 \times 10^{-3}$ mol in 25 cm³

$[\text{OH}^-] = 1.05 \times 1000/25 = 4.2 \times 10^{-2}$ (mol dm⁻³) (1)

$[\text{Ca}^{2+}] = 2.1 \times 10^{-2}$ (mol dm⁻³) (1)

$K_{\text{sp}} = 2.1 \times 10^{-2} \times (4.2 \times 10^{-2})^2 = 3.7 \times 10^{-5}$ (1)

(iii) **less** soluble in NaOH due to the common ion effect or equilibrium is shifted to the l.h.s. by high $[\text{OH}^-]$ (NOT just a mention of Le Chat^r on its own) (1) [6]

[Total: 15]

3 (a) SiF_4 is symmetrical or tetrahedral or bonds are at 109° or has no lone pair or 4 electron pairs shared equally or all Si-F dipoles cancel out, or SF_4 has a lone pair (on S). (1) [1]

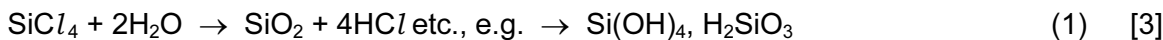
(b)

compound	molecule has an overall dipole	molecule does not have an overall dipole
BCl_3		✓
PCl_3	✓	
CCl_4		✓
SF_6		✓

mark row-by-row,

(2) [2]

(c) (i) Si and B have empty / available / low-lying orbitals or C does not have available orbitals (allow "B is electron deficient" but not mention or implication of d-orbital on B) (1)



(d) (i) $\text{Si}_3\text{Cl}_8\text{O}_2$ (this has $M_r = 84 + 280 + 32 = 396$) or $\text{Si}_4\text{Cl}_4\text{O}_9$ or $\text{Si}_8\text{Cl}_4\text{O}_2$ (1)

(ii)

mass number	structure
133	Cl_3Si
247	$\text{Cl}_3\text{Si}-\text{O}-\text{SiCl}_2$
263	$\text{Cl}_3\text{Si}-\text{O}-\text{SiCl}_2-\text{O}$

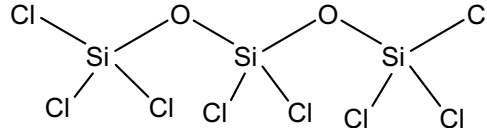
(3)

(if correct structures are not given for last 2 rows, you can award (1) mark for

two correct molecular formulae:

either $\text{Si}_2\text{Cl}_5\text{O} + \text{Si}_2\text{Cl}_5\text{O}_2$ or $\text{Si}_3\text{Cl}_8 + \text{Si}_3\text{Cl}_8\text{O}_9$ or $\text{Si}_7\text{Cl}_8 + \text{Si}_7\text{Cl}_8\text{O}_2$)

(iii)



allow ecf on the structure drawn in the third row of the table in (ii)
but any credited structure must show correct valencies for Si, Cl and O. (1) [5]

[Total: 11]

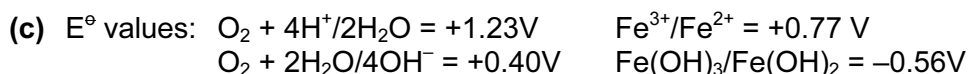
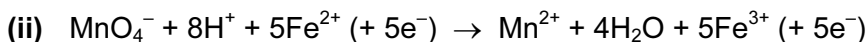
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4 (a) Cr³⁺: 1s²2s²2p⁶... 3s²3p⁶3d³
Mn²⁺: 1s²2s²2p⁶... 3s²3p⁶3d⁵
(allow (1) out of (2) for 3s²3p⁶4s²3d¹ and 3s²3p⁶4s²3d³) (1) (1) [2]

(b) (i) any three of the following points:

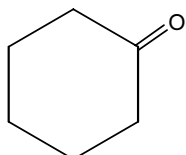
- initial (pale) green (solution)
- fades to (almost) colourless (allow yellow)
- then (permanent faint) pink
- finally (deep) purple

(3)



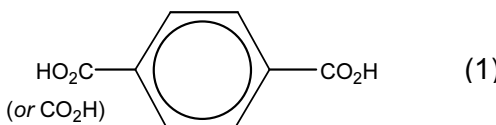
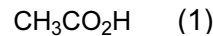
If E_{cell} is more positive it means a greater likelihood of reaction (1) [4]

(d)



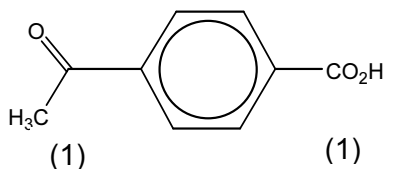
(1)

and

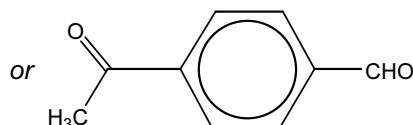


(or CO_2H)

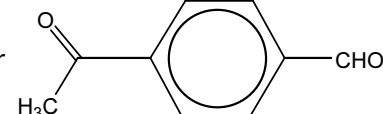
(1)



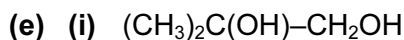
(1)



or



[5]

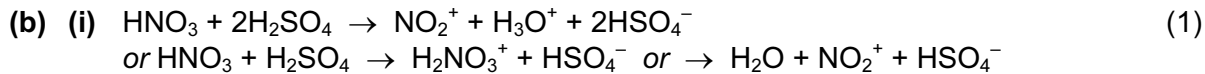


(ii) reaction I: (cold dilute) KMnO_4 ("cold" not needed, but "hot" or "warm" negates) (1)
 reaction II: $\text{Cr}_2\text{O}_7^{2-} + \text{H}^+ + \text{distil}$ (1) [3]

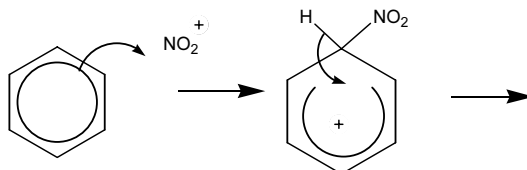
[Total: 18 max 17]

5 (a) (i) because the carbons are sp^2 / trigonal planar / bonded at 120° or are joined by π bonds / orbitals (1)

(ii) because the π electrons / double bonds are delocalised / in resonance or electrons are evenly distributed / spread out (1) [2]



(ii) electrophilic substitution mechanism: (1)



curly arrows from benzene to NO_2^+ , and showing loss of H^+ (1)
correct intermediate (with “+” in the ‘horse-shoe’) (1) [4]

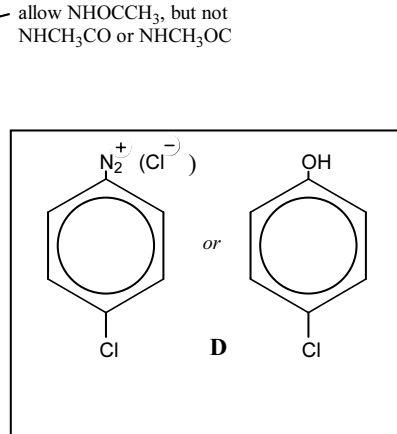
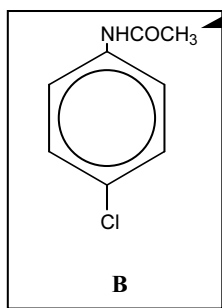
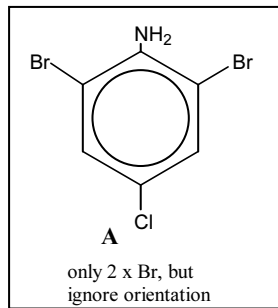
(c) $\text{Cl}_2 + \text{Al}/\text{Cl}_3 / \text{FeCl}_3 / \text{Fe} / \text{Al} / \text{I}_2$ (aq or light negates this mark) (1) [1]

(d) (i) Y is chlorobenzene (1) Z is 4-chloronitrobenzene (1) (2)

(ii) Sn / Fe + (conc) HCl (1)

HCl is conc, and second step is to add $\text{NaOH}(\text{aq})$ (1)

(iii)



no reaction
C

(4) [8]

[Total: 15]

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6 (a) (i) Primary – the **amino acid** sequence / order / chain
or diag. e.g. NH-C-CO-NH-C-CO
or amino acids bonded by covalent / amide / peptide bonds (1)

(ii) Tertiary – the coiling / folding of the protein / polypeptide chain due to interactions between side-chains on the amino acids or the structure which gives the protein its 3-D / globular shape (1) [2]

(b) (i) Diagram:
Minimum is $\text{CH}_2\text{S}-\text{SCH}_2$ (1)

(ii) Oxidation / dehydrogenation / redox (1)

(iii) Hydrogen / H bonds; ionic interactions / bonds or ion-dipole or salt bridges; van der Waals' or id-id or induced / instantaneous dipole forces (ignore hydrophobic interactions) (2) [4]

(c) (i) Hydrogen bonds (1)

(ii) Correct new strand present (see below) needed
Diagram showing C=O bonding to N-H in new strand... ✓
...and N-H bonding to C=O in new strand ✓
e.g.

New strand must contain a minimum of two amino acid residues in a single chain. Deduct a penalty of –(1) for any wrong H-bond **only** if (2) marks have already been scored. (2) [3]

(d) There are bonds or S-S bridges / linkages **between the layers / sheets** (in β -keratin) (but only van der Waals interactions between the layers in silk) (1) [1]

[Total: 10]

7 (a) The amino acid is uncharged / neutral / a zwitterion or charges balance / are equal (NOT "is non-polar")

It is equally attracted by the anode / + and the cathode / – or attracted by neither

The pH of the buffer is at the isoelectric point/IEP of the amino acid *any two* ✓✓ (2) [2]

(b) (at pH 10), $\text{H}_2\text{NCH}_2\text{CO}_2^-$ or $\text{NH}_2\text{CH}_2\text{COO}^-$ (1) [1]

(c)

amino acid	relative size	charge
A	small(est) (1)	–ve
B	large(st) (3)	–ve
C	middle (2)	+ve

(numbers are OK to show relative sizes)

Mark each row (3) [3]

(d) (i) lys – val – ser – ala – gly – ala – gly – asp (2)

(ii) gly – ala – gly (1)

(iii) aspartic acid (or lysine) (1) [4]

[Total: 10]

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8 (a) Reaction II – since electrons are used up / required / gained / received (from external circuit) (1) [1]

(b) $(Pb^{2+} + 2e^- \rightarrow Pb)$ $E^\ominus = -0.13V$
 $(PbO_2 + 4H^+ + 2e^- \rightarrow Pb^{2+} + 2H_2O)$ $E^\ominus = +1.47V$
two correct E^\ominus values (1)

Cell voltage is **1.6(0)** (V) (1) [2]

(c) (i) 3(+) (1)

(ii) They are less heavy / poisonous / toxic / polluting or are safer due to no (conc) H_2SO_4 within them (1) [2]

(d) (i) Platinum or graphite / carbon (1)

(ii) They need large quantities of **compressed** gases which take up space or the hydrogen would need to be **liquefied** or the reactant is (highly) **flammable** / **explosive** / **combustible** (1) [2]

(e) Glass: saves **energy** – the raw materials are easily accessible / cheap or making glass is energy-intensive (1)

Steel: saves **energy** – extracting iron from the ore or mining the ore is energy intensive or saves a **resource** – iron **ore** (NOT just “iron”) is becoming scarce
either one (1)

Plastics: saves a valuable / scarce **resource**: (crude) **oil** / **petroleum** (1) [3]

[Total: 10]