

UNIVERSITY OF CAMBRIDGE INTERNATIONAL EXAMINATIONS

GCE Advanced Subsidiary Level and GCE Advanced Level

MARK SCHEME for the October/November 2007 question paper

9701 CHEMISTRY

9701/02

Paper 2 (Theory 1), maximum raw mark 60

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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.

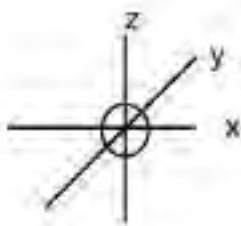
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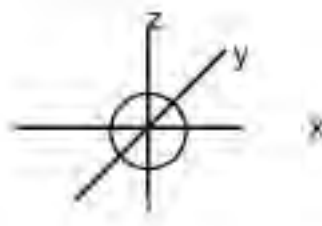
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1 (a)



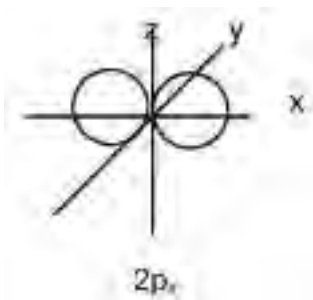
1s

spherical (1)



2s

larger spherical (1)



2p_x

double lobes along the x-axis (1)

[3]

- (b) (i) attraction between bonding electrons and nuclei (1)
attraction is electrostatic (1)

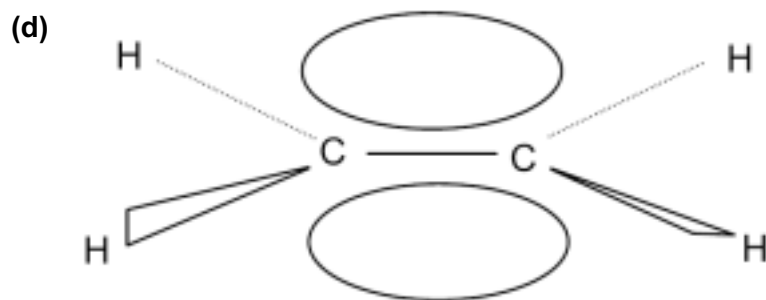
- (ii) H₂ s-s **overlap** clearly shown
must **not** be normal dot/cross diagram (1)

- HCl s-p **overlap** clearly shown
overlap must involve s and p orbitals (1) [4]

- (c) (i) bonding electrons are unequally shared **or**
the molecule has a dipole/ δ^+ and δ^- ends to molecule (1)

- (ii) the H and Cl atoms have different electronegativities
or chlorine is more electronegative than hydrogen (1) [2]

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allow two 'sausages' above **and** below the C-C axis

or two p orbitals **overlapping** sideways

to form one (localised) π bond over two carbon atoms

(1) [1]

(e) $\Delta H_f^\ominus = 2(-393.7) + 2(-285.9) - (-1411)$

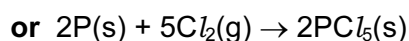
$= + 51.8 \text{ kJ mol}^{-1}$ (units given in qu.)

(3)

penalise errors: no 2 for -393.7
no 2 for -285.9
wrong sign for $-(-1411)$

[3]

[Total: 13]



equation

(1)

state symbols

(1) [2]

(b) (i) giant ionic lattice (may be in diag.)

(1)

strong ionic bonds

(1)

(ii) simple molecular **or** discrete molecules

(may be shown in a diagram)

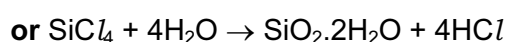
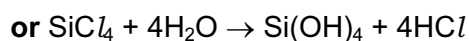
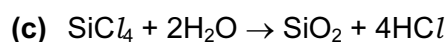
(1)

with **weak** intermolecular forces **or**

weak van der Waals' forces

between them

(1) [4]



(1) [1]

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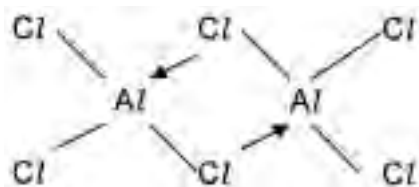
- (d) NaCl pH is 7 allow neutral (1)
- PCl_5 pH is between 1 and 4
- do **not** allow acidic (1) [2]

- (e) (i) 460 K Al_2Cl_6 (1)
- 1150 K AlCl_3 (1)

(ii) correct **dot-and-cross** diagram for AlCl_3 (1)

(iii) correct displayed structure for Al_2Cl_6 (1)

two correct co-ordinate bonds (1)



[5]

[Total: 14]

- 3 (a) P_4 (1)
- S_8 (1)
- Cl_2 (1) [3]

(b) (i) highest S_8 P_4 Cl_2 lowest (1)

allow S ... P ... Cl **or** names (1)

(ii) from S_8 to P_4 to Cl_2

there are fewer electrons in each molecule (1)

hence weaker van der Waals' forces (1) [3]

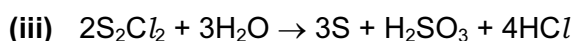
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(c) (i) $S_2Cl_2 = (2 \times 32.1) + (2 \times 35.5) = 135.2$

$$n(S_2Cl_2) = \frac{2.7}{135.2} = 0.0199 = 0.02 \quad (1)$$

$$0.02 \text{ mol } S_2Cl_2 \rightarrow \frac{0.96}{32.1} = 0.03 \text{ mol S}$$

$$1.0 \text{ mol } S_2Cl_2 \rightarrow \frac{0.03 \times 1.0}{0.02} = 1.5 \text{ mol S} \quad (1)$$



correct products (1)

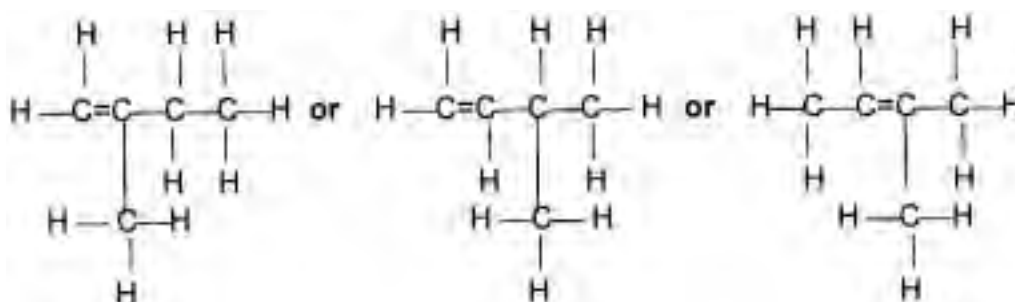
balanced equation (1) [4]

(d) oxidation product is H_2SO_3 (1)

reduction product is S (1) [2]

[Total: 12]

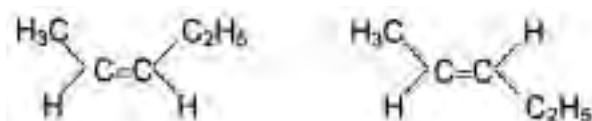
4 (a)



H atoms must be shown.

Structure must not contain any CH_3 groups (1) [1]

(b)



cis

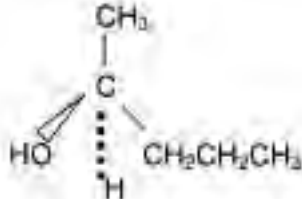
trans


(1) [2]

(c) $CH_3CH(OH)CH_2CH_2CH_3$ (1)

$CH_3CH_2CH(OH)CH_2CH_3$ (1) [2]

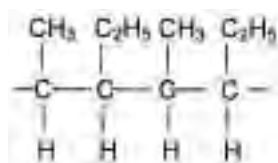
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- (d)  correct compound (1)
- correct mirror object/mirror image relationship in 3D (1) [2]

- (e)  e.g. cyclopentane structure

allow methylcyclobutane **or** dimethylcyclopropane (1) [1]

- (f) e.g.



two repeat units must be shown
relative positions of $-\text{CH}_3$ and $-\text{C}_2\text{H}_5$ may differ from those shown above (1) [1]

[Total: 9]

- 5 (a) (i) $\text{Cr}_2\text{O}_7^{2-}/\text{H}^+$ allow $\text{MnO}_4^-/\text{H}^+$ (1)
- (ii) from orange to **or** purple to colourless
green **or** green/blue (1) [2]
- (b) (i) to ensure complete oxidation of $-\text{CH}_2\text{OH}$
or to keep reactants in the reaction flask (1)
- (ii) CH_3CHO /ethanal (1) [2]
- (c) (i) CH_3I /iodomethane (1)
- (ii) nucleophilic substitution **or** hydrolysis (1) [2]

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(d) step I

red P + I₂ **or** HI(aq) **or** KBr/conc H₃PO₄ **or** PI₃ (1)

heat **but** room temperature for PI₃ (1)

step II

KCN in aqueous ethanol (1)

in aqueous ethanol, heat under reflux (1)

allow aqueous ethanol in either place

step III

aqueous mineral acid (**not** nitric acid)

or NaOH(aq) then aqueous mineral acid (1)

heat (1) [6]

[Total: 12]