# **MARKSCHEME**

**May 2006** 

**CHEMISTRY** 

**Higher Level** 

Paper 2

17 pages

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## **SECTION A**

1. (a) 
$$C_6H_{12} + 9O_2 \rightarrow 6CO_2 + 6H_2O;$$
 [1]

(b) (i) 
$$(\Delta H^{\ominus} = \sum \Delta H_{\rm f}^{\ominus}_{\rm products} - \sum \Delta H_{\rm f}^{\ominus}_{\rm reactants})$$
  
 $\Delta H^{\ominus} = (6 \times -394 + 6 \times -242) - (-43);$   
 $\Delta H_{\rm c}^{\ominus} = -3773/-3.8 \times 10^{3} \text{ (kJ mol}^{-1});$   
 $Accept \ 2, \ 3 \ or \ 4 \ sf.$   
 $Award \ [1] \ for \ +3773/+3.8 \times 10^{3} \ (kJ \ mol^{-1}).$   
 $Allow \ ECF \ from \ (a) \ only \ if \ coefficients \ used.$ 

(ii) 
$$\Delta S^{\ominus} = (S_{p}^{\ominus} - S_{r}^{\ominus}) = (6 \times 189 + 6 \times 214) - (385 + 9 \times 205);$$
  
 $\Delta S_{c}^{\ominus} = 188 \text{ (J K}^{-1} \text{ mol}^{-1});$  [2]  
Accept only 3sf.  
Award [1] for -188.  
Allow ECF from (a) only if coefficients used.

(c) 
$$(\Delta G_{c}^{\ominus} = \Delta H_{c}^{\ominus} - T\Delta S_{c}^{\ominus}) = -3800 - (298 \times 0.188);$$
  
 $= -3900 \ kJ \ mol^{-1}.$  [2]  
 $Accept - 3800 \ to -3900.$   
 $Accept \ 2, \ 3 \ or \ 4 \ sf.$   
 $Allow \ ECF \ from \ (b).$   
 $Units \ needed \ for \ second \ mark.$ 

- (d) spontaneous and  $\Delta G^{\ominus}$  negative; [1] Allow ECF from (c).
- (e)  $-1 \times \Delta H_1 / 676$ ;  $1 \times \Delta H_2 / -394$ ;  $2 \times \Delta H_3 / -484$ ;  $\Delta H_4 = -202 \text{ (kJ mol}^{-1})$ ; [4]

Accept alternative methods.

Correct answers score [4].

Award [3] for (+)202 or (+)40  $(kJ/kJ mol^{-1})$ .

2.  $A_{r}(T1) = 203 \times 0.2952 + 205 \times 0.7048 / A_{r}(T1) = 204.41;$ (a)

$$A_r(Br) = 79 \times 0.5069 + 81 \times 0.4931 / A_r(Br) = 79.99;$$

$$M_r(\text{T1Br}_3) = 204.41 + 3 \times 79.99 = 444.38 / 444.37;$$

[3]

Correct answer scores [3].

Ignore units of g or g  $mol^{-1}$ .

Apply ECF to  $M_r$  from  $A_r$  values.

(b)  $M_r$  is an <u>average</u> value (because of the isotopes);

> each HBr molecule has its own value depending on which isotopes (of H or Br) it contains/OWTTE;

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[2]

[1]

(c)  $1s^22s^22p^63s^23p^63d^{10}4s^24p^6$ ;

Do not accept noble gas shortcut. No subscripts.

- $Mg^{2+}$ ; (d) [1]
- $Al^{3+}$ ,  $O^{2-}$ , Ne,  $Na^+$ ,  $F^-$ ,  $N^{3-}$ ; (e) [2] Award [2] for any three, [1] for any two.
- **3.**  $n(Fe_2O_3) = 30 \times 10^3 \div 159.7 / n(Fe_2O_3) = 188 \text{ mol};$

 $n(C) = 5.0 \times 10^3 \div 12.01 / n(C) = 416 \text{ mol};$ 

Fe<sub>2</sub>O<sub>3</sub> is the limiting reagent or implicit in calculation;

$$n(Fe) = 2 \times n(Fe_2O_3) = 2 \times 188 = 376 \text{ mol};$$

$$m(Fe) = 376 \times 55.85 = 21 \text{ kg};$$

[5]

Accept 2sf or 3sf, otherwise use -1(SF).

Correct final answers score [5].

Allow ECF.

- 4. (a species that) gains electrons (from another species) / causes electron loss; (a) (i)
  - [1]

[2]

(ii) changes by 3;

> reduced because its oxidation number decreased  $/+6 \rightarrow +3/6+ \rightarrow 3+/$  it has gained electrons;

(i)  $C_6H_8O_6 \rightarrow C_6H_6O_6 + 2H^+ + 2e;$ (b) [1]

(ii)  $C_6H_8O_6 + 2Fe^{3+} \rightarrow C_6H_6O_6 + 2H^+ + 2Fe^{2+}$ ; [1]

### **5.** (a) same <u>general</u> formula;

successive members differ by CH<sub>2</sub>;

Do not allow elements or just "they".

similar chemical properties;

Allow same/constant.

gradual change in physical properties;

Do not allow change periodically.

same functional group;

Award [1] each for any two.

[2 max]

#### (b) add bromine (water);

alkanes – no change / stays or turns brown;

Allow red-brown or any combination of brown, orange or yellow.

alkenes – bromine (water) decolorizes;

Do not allow clear or discoloured.

or

add (acidified) KMnO<sub>4</sub>;

alkanes - no change;

alkenes – MnO<sub>4</sub> decolorizes / brown / black;

[3]

Do not accept addition of  $H_2$  or HBr.

(c) butan-1-ol: butanal;

butanoic acid;

butan-2-ol: butanone;

2 methylpropan-2-ol: no oxidation;

[4]

Also accept correct structures. Where both name and structure given structure must be correct and name largely correct.

#### **SECTION B**

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6. (a)  $K/K_c = [SO_3]^2 \div [SO_2]^2 [O_2];$  [1] Exactly as written. Accept correct  $K_p$  expression.

- (b) (i) vanadium(V) oxide / (di)vanadium pentaoxide / V<sub>2</sub>O<sub>5</sub>/Pt; [1]

  Allow just vanadium oxide but not incorrect formula.
  - (ii) catalyst does not affect the value of K<sub>c</sub>;
     forward and reverse rate increase equally/by the same factor;
     catalyst increases the rate of the reaction;
     (by providing an alternative path for the reaction with) lower activation energy; [4]
- (c) more energetic collisions / more molecules have energy greater than activation energy; more frequent collisions; [2]

  Do not accept more collisions without reference to time.
- (d) (i) shifts equilibrium position to the products/right; to the side with least gas molecules or moles / lower volume of gas; [2]
  - (ii) shifts equilibrium position to the products/right; to compensate for loss of SO<sub>3</sub> / produce more SO<sub>3</sub>; [2]
- (e) exothermic;  $K_c$  decreases with increasing temperature / back reaction favoured / heat used up / OWTTE; [2]
- (f)  $n(SO_2)_{at \text{ equilibrium}} = 1.50 0.50 = 1.00 \text{ mol};$  $n(O_2)_{at \text{ equilibrium}} = 2.00 - 0.250 = 1.75 \text{ mol};$

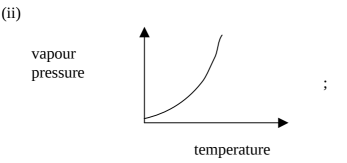
 $[SO_2] = 1.00 \div 1.50 = 0.667 \text{ mol dm}^{-3}, \ [O_2] = 1.75 \div 1.50 = 1.17 \text{ mol dm}^{-3}$  $[SO_3] = 0.500 \div 1.50 = 0.333 \text{ mol dm}^{-3};$ 

$$K_{\rm c} = (0.333)^2 \div 1.17 \times (0.667)^2;$$
  
= 0.213 dm<sup>3</sup> mol<sup>-1</sup>/0.214 dm<sup>3</sup> mol<sup>-1</sup>; [5]

If  $0.202 \text{ dm}^3 \text{ mol}^{-1}$  is given award [4], this is obtained by premature rounding. Award [5] for correct answer with units.

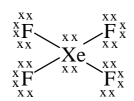
[3]

the greater the strength of the intermolecular forces the greater the enthalpy of (g) (i) vaporization/OWTTE; pentane has only van der Waals' forces between molecules; propanoic acid has H-bonding (as well as van der Waals' forces); [3]



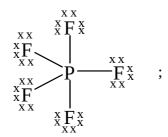
 $1^{st}$  mark: graph goes upwards with T;  $2^{nd}$  mark: curve as shown;

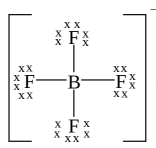
as temperature increases (more) molecules have enough energy to overcome intermolecular / attractive forces;



; lone pairs on Xe required for the mark.

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; square brackets and charge required for the mark.

Accept any combination of dots, crosses and lines. Penalise missing fluorine lone pairs once only. [3]

(b)  $XeF_4$ 

Square planar and 90°;

PF

trigonal bipyramid and 90° and 120°;

 $BF_{\scriptscriptstyle A}^{-}$ 

Tetrahedral and 109.5°/109°;

[3]

Allow <u>clear</u> suitable diagrams instead of name.

No ECF from (a).

(c) hybridization: mixing / merging of atomic orbitals;

$$N_2 - sp;$$

$$N_2H_2-sp^2$$
;

$$N_2H_4-sp^3$$
;

(d)  $\sigma$  bonds (result from the) overlapping of orbitals end to end / along inter-nuclear axis;  $\pi$  bonds (result from the) overlapping of parallel/sideways p orbitals; (single bonds)  $\sigma$  bonds only;

(double bonds) have a  $\sigma$  bond and a  $\pi$  bond;

[4]

[4]

Suitable <u>clear</u> and labelled diagrams acceptable for all marks.

(e) (i) electron removed from higher energy level / further from nucleus / greater atomic radius;

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increased repulsion by extra inner shell electrons / increased shielding effect;

(ii)  $Mg^{2+}(g) \rightarrow Mg^{3+}(g) + e;$ 

(even though) valence electrons in the same shell/main energy level / Mg<sup>2+</sup> has noble gas configuration;

Mg has greater nuclear/core charge/more protons;

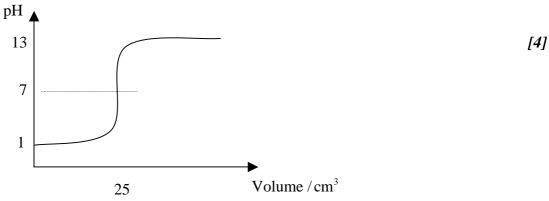
[3]

[2]

- (f) (i) Mg has twice/more delocalized electrons as Na; the ionic charge is twice as big/greater in Mg than Na; sodium ion is larger than magnesium ion; attraction of ions and electrons is less in sodium/greater in magnesium; [3 max] Correct discussion of charge density gains 2<sup>nd</sup> and 3<sup>rd</sup> mark.

  Award [1] each for any three.
  - (ii) SO<sub>2</sub> has (weak) intermolecular/van der Waals' force/dipole dipole;
     MgO has (strong) ionic bonds;
     Ionic bonding is stronger than intermolecular attraction (OWTTE);

(ii) curve should include the following: starting pH = 1; equivalence point: 25.0 cm<sup>3</sup> of NaOH; pH at equivalence point = 7; pH to finish = 12-13;



Penalise [1] if profile incorrect.

(iii) 
$$K_{\rm a} = 10^{-4.76}/1.74 \times 10^{-5}$$
; 
$$K_{\rm a} = [{\rm H}^+]^2 \div [{\rm CH_3COOH}]/1.74 \times 10^{-5} = \frac{[{\rm H}^+]^2}{0.100};$$
 
$$[{\rm H}^+] = 1.32 \times 10^{-3} \; ({\rm mol \, dm^{-3}});$$
 starting pH = 2.88; Accept 3sf. Award [4] for correct pH. Allow ECF. pH at equivalence point: 8 – 9; [5]

(b) (i) HIn is a weak acid;
 HIn 

→ H<sup>+</sup> + In<sup>-</sup> and two colours indicated;

In acid <u>equilibrium</u> moves left or vice versa; [3]

- (ii) phenolphthalein / phenol red / bromothymol blue; colour change of indicator occurs within the range of pH at equivalence point / on vertical part of graph; [2]
- (c) (i) specific examples of weak base and its salt / specific strong acid and weak base; [1]

  Name of structure acceptable.

  e.g. NH<sub>3</sub> and NH<sub>4</sub>Cl.
  - (ii) pH changes very little / most acid neutralized by base; equation from (i); [2] Any other suitable example. e.g.  $NH_3 + H^+ \rightarrow NH_4^+ / NH_4OH + H^+ \rightarrow NH_4^+ + H_2O$ .

Lewis acid electron pair acceptor;

*Brønsted-Lowry acid* Any suitable equation;

Lewis acid – BF<sub>3</sub>/AlCl<sub>3</sub>/transition metal ions that form complex ion with ligands;

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For example

BF<sub>3</sub> + NH<sub>3</sub> 
$$\rightarrow$$
 BF<sub>3</sub>NH<sub>3</sub>/Cu<sup>2+</sup> + 4NH<sub>3</sub>  $\rightarrow$  [Cu(NH<sub>3</sub>)<sub>4</sub>]<sup>2+</sup>/AlCl<sub>3</sub> + Cl<sup>-</sup>  $\rightarrow$  AlCl<sub>4</sub>; [5] Or any suitable equation.

(e) acidic;

$$[Al(H2O)6]3+ is (weak) acid due to the formation of H+ / [Al(H2O)6]3+  $\rightleftharpoons$  [Al(H<sub>2</sub>O)<sub>5</sub>(OH)]<sup>2+</sup> + H<sup>+</sup>; [2]$$

[1]

[1]

(ii)  $HOOCCHNH_2$   $CH_3$ 

Allow appropriate acyl chloride.

(iii)  $H_2N(CH_2)_6NH_2$ ;

HOOC(CH<sub>2</sub>)<sub>4</sub>COOH;

[2]

Allow correct alternative.

Accept correct names as alternatives.

If correct structure and incorrect name given, award the mark.

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Penalise COOH - C once only.

(b) (addition polymers) contain  $C = C/C \equiv C$ ;

(condensation polymers) contain two reactive/functional groups;

[2]

(c) HCOOCH<sub>3</sub>;

methyl methanoate;

[2]

Accept other correct alternative.

(d) (i) methanol / methyl alcohol;

heat and acid catalyst/H<sup>+</sup>;

$$CH_3OH + CH_3COOH \rightarrow CH_3COOCH_3 + H_2O;$$

[3]

(ii) physical properties

ethanoic acid has a higher boiling point / ester has a lower boiling point; ethanoic acid has vinegar smell, ester has sweet/fruit smell;

Must specify one smell.

ethanoic acid is more soluble in water than methyl ethanoate / methyl ethanoate is more soluble in non-polar solvents than ethanoic acid;

ethanoic acid (in water) has a pH < 7, ester (in water) has a pH = 7;

[2 max]

Award [1] each for any two.

(iii) ethanoic acid

3:1;

methyl ethanoate

1:1:

[2]

Allow 3:3.

(e) (i) 2 – chlorobutane is the optical isomer;

has a chiral carbon/asymmetric carbon atom / 4 different groups around central atom;

− 16 −

- [2]
- (ii) pass plane polarized light through (two separate) samples; each sample will rotate the polarized light in the <u>opposite</u> direction;

[2]

[2]

(iii)

Award [2] marks for 3 and [1] mark for 2 structures. Penalise missing H atoms once only.

(iv) 1-chlorobutane / 1-chloro-2-methylpropane; *Accept structures*.

[1]

[3]

### (v) mechanism

curly arrow from O of  ${}^-\text{OH}$  joined to C, and from C–Cl bond to Cl; transition state structure with partial bonds to OH and Cl and a negative charge; product: CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH / CH<sub>3</sub>CH(CH<sub>3</sub>)CH<sub>2</sub>OH;

e.g.