# MARKSCHEME 

November 2001

## CHEMISTRY

## Higher Level

## Paper 2

## SECTION A

1. 

(a) (i) $M_{\mathrm{r}}=88 / 88.17$ ..... [1]$\Delta H_{\mathrm{c}}^{\ominus}=-3325 \pm 25 \mathrm{~kJ} \mathrm{~mol}^{-1}$ (allow for ECF)(no mark if minus sign missing; no penalty if units not given)
(ii) The value should be (about) the same. ..... [1]
Same (number and type of) bonds are being broken and made. ..... [1](Do not accept "the compounds have the same relative molecular masses orsame formula".)
(b) The reaction of methanol and oxygen has a high activation energy / OWTTE. ..... [1] ..... [1]When $E \geq E_{\mathrm{a}}$ / OWTTE is provided, a (highly exothermic) reaction takes place.
(c) (i) heat evolved $=110 \mathrm{~g} \times 4.20 \mathrm{Jg}^{-1}{ }^{\circ} \mathrm{C}^{-1} \times 21.4{ }^{\circ} \mathrm{C}$ (accept (44.6-23.2)) (Award [1] for correct 4 ; [1] for 110 g , not 100 g .) ..... [2]
$=9887 \mathrm{~J}$ (accept 9890 J, allow ECF) ..... [1]
(No double jeopardy if 100 g or 10 g used; no mark without unit.)
(ii) $\quad M_{\mathrm{r}}=40.0 ; \quad \Delta H$ per mol $=-9887 \mathrm{~J} \times \frac{40.0}{10.0}=-39.5 \mathrm{~kJ} \mathrm{~mol}^{-1}($ accept -39.6$)$
(need minus sign for mark; no penalty if units not given)
(iii) Heat loss to the surrounding (thus less $\Delta H$ ) ..... [1]
Use a styrofoam/plastic cup OR insulate (and cover) beaker ..... [1]
(Accept answer that says "calculate heat gained by glass calorimeter".)
2. (a) (i) $M_{\mathrm{r}}=44.01 ; \quad \mathrm{m}_{\mathrm{C}}$ in $\mathrm{CO}_{2}$ produced $=\frac{12.01}{44.01} \times 5.470=1.493 \mathrm{~g}$

$$
\text { Percentage } \mathrm{C}=\frac{1.493}{2.036} \times 100=73.32 \%
$$

(ii) $\mathrm{m}_{\mathrm{H}}=\frac{2.02}{18.02} \times 0.697=0.0781 \mathrm{~g}$

$$
\text { Percentage } \mathrm{H}=\frac{0.0781}{2.036} \times 100=3.84 \%
$$

(b) Percentage $\mathrm{O}=100-73.32-3.84-10.75=12.09 \%$

amount, | C | H | N | O |  |
| ---: | :--- | :---: | :---: | :--- |
|  | $\frac{73.32}{12.01}$ | $\frac{3.84}{1.01}$ | $\frac{10.75}{14.01}$ | $\frac{12.09}{16.00}$ |$\quad$ (ECF if\% oxygen not worked out)

$$
=8: 5: 1: 1, \text { thus } \mathrm{C}_{8} \mathrm{H}_{5} \mathrm{NO}
$$

(c) Empirical mass $=(8 \times 12)+(5 \times 1)+14+16=131 \mathrm{~g} \mathrm{~mol}^{-1}$ (allow for $E C F$ ) [1]

This is half the $M_{\mathrm{r}}$ OR $M_{\mathrm{r}}=2 \times M_{\text {emp }}$; thus molecular formula is $\mathrm{C}_{16} \mathrm{H}_{10} \mathrm{~N}_{2} \mathrm{O}_{2}$.
3. (a) Pressure, volume and temperature of the gas. (Need all three for mark.)
$\mathrm{PV}=\mathrm{nRT}$ OR $\mathrm{n}=\frac{\mathrm{PV}}{\mathrm{RT}}[1] ; \quad \mathrm{n}=\frac{\mathrm{m}}{\mathrm{M}}$ OR $\mathrm{M}=\frac{\mathrm{m}}{\mathrm{n}}$ [1]
OR $M=\frac{\mathrm{m}}{\mathrm{V}} \frac{\mathrm{RT}}{\mathrm{P}}$ (for [2])
Where R is the Ideal Gas Constant (accept gas constant, but not just constant) ( $=8.314 \mathrm{~J} \mathrm{~mol}^{-1} \mathrm{~K}^{-1}$ )

OR calculations as appropriate using molar volume $=22.4 \mathrm{dm}^{3}$
(b) More energetic molecules escape / owtte.

Remaining molecules are less energetic (lower T)
OR Thus its average energy drops (so does its temperature)
OR Bonds are broken as molecules escape
Energy to break these bonds comes from the remaining molecules which are less energetic (thus lower T)
(c) Vapour pressures are all the same.

Vapour pressure does not depend on the:

- volume of the liquid
- volume of the container

OR As long as there is vapour-liquid equilibrium present, and the temperature remains the same, then the vapour pressure will be the same.
correct explanations in terms of dynamic equilibrium involved should be awarded marks
4. (a) Anode: $\mathrm{Cu}(\mathrm{s}) \rightarrow \mathrm{Cu}^{2+}(\mathrm{aq})+2 \mathrm{e}^{-}$(state symbols needed)

Cathode: $\mathrm{Cu}^{2+}(\mathrm{aq})+2 \mathrm{e}^{-} \rightarrow \mathrm{Cu}(\mathrm{s})$ (state symbols needed)
(b) Remains the same as $\left[\mathrm{Cu}^{2+}\right]$ does not change (reasoning needed)
(ECF if reaction in 4 (a) is incorrect)
(c) $2 \mathrm{H}_{2} \mathrm{O}(\mathrm{l}) \rightarrow \mathrm{O}_{2}(\mathrm{~g})+4 \mathrm{H}^{+}(\mathrm{aq})+4 \mathrm{e}^{-}$(states not required)
(Award [1] for balanced half equation and [1] for $\mathrm{O}_{2}$ and $\mathrm{H}^{+}$as products.)
(d) The colour intensity decreases / changes

The pH decreases / changes
(e) $\mathrm{C}=\mathrm{At}=0.180 \mathrm{~A} \times 1210 \mathrm{~s}=218 \mathrm{C}$

Number of Faradays, $F=\frac{218 \mathrm{C}}{96480 \mathrm{C}}\left(\mathrm{mole}^{-}\right)=2.26 \times 10^{-3} \mathrm{~mole}^{-}$
$2 \mathrm{~mole}^{-}$produce $1 \mathrm{~mol} \mathrm{Cu}=63.35 \mathrm{~g} \mathrm{Cu}$
thus $2.26 \times 10^{-3}$ mole $^{-}$gives $0.50 \times 63.35 \times 2.26 \times 10^{-3} \mathrm{~g}=0.0715 \mathrm{~g} \mathrm{Cu}$

## SECTION B

5. (a) (i) Average $A_{\mathrm{r}}=\frac{(32.00 \times 95.00)+(33.00 \times 0.76)+(34.00 \times 4.20)+(36.00 \times 0.020)}{100}$ [1]

$$
=32.1
$$

(ii) Number of neutrons in ${ }^{36} \mathrm{~S}=36-16=20$
(b) (i)

(Award [1] for increase from first to third I.E. and [1] for larger increase from third to fourth I.E.; for "I.E. keeps increasing" award only [1])
(ii) Boron: $1 \mathrm{~s}^{2} 2 \mathrm{~s}^{2} 2 \mathrm{p}^{1} /[\mathrm{He}] 2 \mathrm{~s}^{2} 2 \mathrm{p}^{1}$ [1]

Aluminium: $\quad 1 \mathrm{~s}^{2} 2 \mathrm{~s}^{2} 2 \mathrm{p}^{6} 3 \mathrm{~s}^{2} 3 \mathrm{p}^{1} /[\mathrm{Ne}] 3 \mathrm{~s}^{2} 3 \mathrm{p}^{1}$ [1]
Al has an $\mathrm{e}^{-}$in a higher / third energy level further away from the nucleus that is easier to remove.

Magnesium: $1 \mathrm{~s}^{2} 2 \mathrm{~s}^{2} 2 \mathrm{p}^{6} 3 \mathrm{~s}^{2} /[\mathrm{Ne}] 3 \mathrm{~s}^{2}$ [1]
$3 p^{1}$ is easier to remove than $3 s^{2}$ as it is higher in energy. [1]
(c) Spectrum showing discrete lines [1]
converging at higher energy [1]
transition of (excited) electrons from higher energy levels to lower one(s). [1]
(Only one series need be shown to score mark.)

## Question 5 continued

(d) (i) One valence electron / one electron in outer shell ..... [1]
Melting point low (compared to other metals) ..... [1]
First I.E.: low; second I.E.: very high (need both for mark) ..... [1]
For at least two full electron arrangements given. ..... [1]
(ii) Reactivity increases down the group ..... [1]
as outer electrons are further from the nucleus so easier to remove. ..... [1]
Description of reaction with water: e.g. vigorous / highly exothermic. ..... [1]
Gas produced / alkaline solution. ..... [1]
Balanced equation, e.g. $2 \mathrm{Na}(\mathrm{s})+2 \mathrm{H}_{2} \mathrm{O}(\mathrm{l}) \rightarrow 2 \mathrm{NaOH}(\mathrm{aq})+\mathrm{H}_{2}(\mathrm{~g})$ ..... [1](must be balanced; state symbols not required.)
Description of reaction with chlorine: e.g. vigorous / highly exothermic ..... [1]
white solid formed ..... [1]
Balanced equation, e.g. $2 \mathrm{Na}(\mathrm{s})+\mathrm{Cl}_{2}(\mathrm{~g}) \rightarrow 2 \mathrm{NaCl}(\mathrm{s})$ ..... [1](must be balanced, state symbols not required.)
6. (a) (i) : $\ddot{\mathrm{F}}: \mathrm{B}: \ddot{\mathrm{F}}: \quad: \ddot{\mathrm{F}}: \stackrel{\mathrm{N}}{\mathrm{N}}: \ddot{\mathrm{F}}$

(Do no penalise missing charge on ion.)
(Need lone pair on $N$ for mark; must show valence electrons on $F$ in each case. Accept dashes in place of pairs of dots, penalise only once if lone $\mathrm{e}^{-}$pairs missing on F.)
(ii) $\mathrm{BF}_{3}$ planar triangular
$\mathrm{NF}_{3}$ triangular pyramidal
$\mathrm{BF}_{4}^{-}$tetrahedral
$109 \frac{1}{2}^{\circ}$ compared to $107^{\circ} /$ bond angle in $\mathrm{BF}_{4}^{-}>$bond angle in $\mathrm{NF}_{3}$
Explanation: Lone electron pair on N produces extra repulsion that (slightly) pushes the bonding electrons closer together.
(iii) Hybridisation: mixing / merging (but not joining) of atomic orbitals (to form same number of new ones of lower energy)
$\mathrm{BF}_{3}$ sp
$\mathrm{NF}_{3} \quad \mathrm{sp}^{3}$ (hybridised orbitals)
(iv) 'Polar bond' indicates that bonding electrons are not equally distributed between the atoms in the bond
the more electronegative atom will have the greater electron density.
$B-F$ is a polar bond, because F is more electronegative than B
$\mathrm{N}-\mathrm{F}$ is a polar bond, because F is more electronegative than N
OR both bonds are polar because F is the most electronegative element.
$\mathrm{BF}_{3}$ is a non-polar molecule due to its planar triangular shape / polarity of bonds is cancelled / because of symmetrical distribution of electron cloud.
$\mathrm{NF}_{3}$ is a polar molecule due to its (trigonal) pyramidal shape and polarity of bonds does not cancel / non-symmetrical distribution of electron cloud.

## Question 6 continued

(b) (i) $\sigma$ bonds involve overlap of orbitals end-on / along the molecular axis (it has axial symmetry around axis joining the two nuclei)/ overlap of s orbitals.
$\pi$ bonds result from overlapping of parallel p orbitals / sideways overlap.
Double bond: a $\sigma$ bond and a $\pi$ bond.
Triple bond: a $\sigma$ bond plus two $\pi$ bonds.
(ii) Delocalisation: when electron pairs are not confined to two adjacent bonding atoms but extend over three or more atoms.
(iii)

(Award [1] for enthalpy diagram.)
Twice the number of bonds made and broken in 2 compared to 1 , thus twice the
energy given out.
Delocalisation present (only) in 3 .
Thus 3 is more stable / less energy is given out.
7. (a)
(i)


Heat
Acid catalyst / $\mathrm{H}^{+}$
$\mathrm{HCOOH} /$ methanoic acid
$\mathrm{CH}_{3} \mathrm{OH} /$ methanol
$\mathrm{HCOOH}+\mathrm{CH}_{3} \mathrm{OH} \rightleftharpoons \mathrm{HCOOCH}_{3}+\mathrm{H}_{2} \mathrm{O} \quad$ (accept $\rightarrow$ instead of $\left.\rightleftharpoons\right) \quad$ [1]
(ii) $\mathrm{CH}_{3} \mathrm{COOH}$ (but not $\mathrm{C}_{2} \mathrm{H}_{4} \mathrm{O}_{2}$ )

Physical:
Boiling point: acid is higher (due to greater H-bonding)/ ester is lower pH : acid $<7$; ester $=7$ (need both for mark).
OR Smell: acid: vinegar/pungent smell; ester: sweet smell.
Chemical:
acid reacts with $\mathrm{OH}^{-}$to form salt and water.
Ester reacts with $\mathrm{OH}^{-}$to form salt plus methanol / acid can be esterified; ester cannot
(b) (i) When two (small) molecules combine to form a larger one with the elimination of a smaller molecule (such as water).
The need for two functional groups on each of the two monomers.
Addition polymerisation: process in which unsaturated monomers combine to form a polymer without the elimination of any atoms/molecules.
(ii) Condensation polymer

(Award [1] for ester group and [1] for $\mathrm{CH}_{2} \mathrm{CH}_{2}$ and $\mathrm{C}_{6} \mathrm{H}_{4}$ groups.)
(c) (i)


(ii) Negatively charged or neutral species containing electron pairs / electron pair donors / Lewis bases
$\mathrm{OH}^{-}$(no mark for $\mathrm{H}_{2} \mathrm{O}$ - water does not react)
(iii) Each compound contains a C with four different groups on it / contains a chiral centre / molecules are asymmetric / contains asymmetric C / optically active starting materials.

Forming a planar carbocation.
The nucleophile can attach from either side producing an equal mixture of enantiomers / racemic mixture (thus optically inactive).
8. (a) $\frac{\left[\mathrm{H}_{2}(\mathrm{~g})\right][\mathrm{CO}(\mathrm{g})]}{\left[\mathrm{H}_{2} \mathrm{O}(\mathrm{g})\right]}$ (state symbols not needed)
moldm ${ }^{-3}$ (ECF if $K_{c}$ expression incorrect)
Endothermic.
The value of $K_{\mathrm{c}}$ increases with temperature (forward reaction is favoured).
(b) (i) The rate of the forward reaction:

Increase in temperature:
increases rate of the forward reaction
since it increases the number of collisions with $E \geq E_{\mathrm{a}}$
(thus rate of forward and reverse reaction increases).
(Award [0] for "more frequent collisions" as this is not the main reason.)
Increase in pressure:
Increases the rate of the forward reaction.
An increase in pressure increases the concentration / the same amount of molecules in a smaller volume (thus rate increases)/ more frequent collisions.

Increase in surface area of $\mathrm{C}(\mathrm{s})$ :
Increases rate of forward reaction.
The reactive surface area of $\mathrm{C}(\mathrm{s})$ increases / concentration of $\mathrm{H}_{2} \mathrm{O}(\mathrm{g})$ in contact with $\mathrm{C}(\mathrm{s})$ increases (thus rate increases)/ more frequent collisions.
(ii) $\left[\mathrm{H}_{2} \mathrm{O}\right]:\left[\mathrm{H}_{2}\right]$ ratio:

Increase in T: decreases ratio / more $\left[\mathrm{H}_{2}\right]$ compared to $\left[\mathrm{H}_{2} \mathrm{O}\right.$ ].
Favours forward / endothermic reaction by using up (some of the) heat supplied (no mark for saying "because of Le Chatelier's principle").

Increase in P:
Increases the ratio / more $\left[\mathrm{H}_{2} \mathrm{O}\right]$ compared to $\left[\mathrm{H}_{2}\right]$.
$1 \mathrm{~mol} /$ volume of gas in the reactants, $2 \mathrm{~mol} /$ volume of gases in the products / increasing pressure will move position of equilibrium to the left to relieve pressure due to more volumes of gas (in the products).
Increase in surface area of $\mathrm{C}(\mathrm{s})$ :
No effect.
Because [C(s)] is constant / position of equilibrium is unchanged.
(iii) The value of the equilibrium constant:

Increase in T: increases value of $K_{\mathrm{c}}$.
Increase in P: no effect on $K_{\mathrm{c}}$ (because $K_{\mathrm{c}}$ is only temperature dependent).
Increase in surface area: no effect ( $K_{\mathrm{c}}$ is only temperature dependent).

## Question 8 continued

(c) Correct diagram (reactants level, products level and activation energy) ..... [1]
for an endothermic change (no double jeopardy) ..... [1]
showing a lower $E_{\mathrm{a}}$ for the catalysed reaction. ..... [1]
Rate of forward reaction: increases ..... [1]
Rate of reverse reaction: increases ..... [1]
Equilibrium position: unchanged ..... [1]

