## UNIVERSITY OF CAMBRIDGE INTERNATIONAL EXAMINATIONS GCE Advanced Level

## MARK SCHEME for the May/June 2011 question paper for the guidance of teachers

## 9701 CHEMISTRY

9701/41

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

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1 (a) N≡N triple bond is (very) strongor the N₂ molecule has no polarity

[1]

(b) 
$$3Mg(s) \rightarrow 3Mg^{2+}(g)$$
  $\Delta H_1 = 3 \times 148 + 3 \times 2186 = 7002$   $N_2(g) \rightarrow 2N^{3-}(g)$   $\Delta H_2 = 994 + 2 \times 2148 = 5290$ 

LE = 
$$-\Delta H_1 - \Delta H_2 - 461$$
 =  $-12,753$  (kJ mol<sup>-1</sup>)

(-[1] for each error) [3]

(c) (i) 
$$\text{Li}_3\text{N} + 3\text{H}_2\text{O} \rightarrow \text{NH}_3 + 3\text{LiOH}$$
 (balanced equation) [1]

(ii) advantage: no high pressure/temperature/catalyst needed/standard conditions used [1]

disadvantage: Li is expensive

or Li would need to be recycled/removedor LiOH by-product is corrosive/strongly basic

**or** this would be a batch, rather than continuous process [1]

(d) (i) 
$$\text{Li}_3\text{N}: 100 \times 14/35 = 40\% \text{ N}$$
 [1] urea:  $100 \times 28/60 = 47\% \text{ N}$  [1]

(iii) 
$$NH_2CONH_2 + H_2O \rightarrow 2NH_3 + CO_2$$
  
 $or \rightarrow NH_2CO_2H + NH_3$   
 $or NH_2CONH_2 + 2H_2O \rightarrow 2NH_3 + H_2CO_3$  [1]

(iv) The LiOH would be strongly alkaline
 or would increase the pH of the soil
 or would 'burn' the crops/reduce plant growth/stunt plants
 or would contaminate the environment

[Total: 12]

[1]

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2 (a) (i) One that can go in either direction. [1]

(ii) both forward & reverse reactions are going on at the same time, but the concentrations of all species do not change (owtte) [1]

*or* rate of forward = rate of backward reaction

(b) (i) 
$$K_c = [H^{\dagger}][OH^{-}]/[H_2O]$$
 [1]

- (ii)  $K_w = [H^+][OH^-]$ [1] rearrangement of equation in (i) gives  $K_c[H_2O] = [H^+][OH^-] \& K_w = K_c[H_2O]$  (owtte) or the [H<sub>2</sub>O] is contained within K<sub>w</sub> [1]
- (iii) K<sub>w</sub> will be higher in hot water **because** reaction is endothermic [1]
- (c) (i)  $[OH^{-}] = 5 \times 10^{-2}$ ;  $[H^{+}] = (1 \times 10^{-14})/5 \times 10^{-2} = 2 \times 10^{-13}$ [1]  $pH = -log_{10}[H^{+}] = 12.7$ (correct ans = [2]) ecf [1]
  - (ii)  $[NH_4^+] = [OH^-] (= x)$ [1]  $x^2 = 1.8 \times 10^{-5} \times 0.05 \implies x (= [OH^-]) = 9.49 \times 10^{-4} (\text{mol dm}^{-3})$ (correct ans = [2])[1]
  - (iii)  $[H^{+}] = K_w/[OH^{-}] = (1 \times 10^{-14})/9.49 \times 10^{-4} = 1.05 \times 10^{-11} (\text{mol dm}^{-3})$ ecf [1]
  - (iv) pH = 11.0ecf [1]

[Total: 12 max 11]

- 3 [1] (a) (+)1; (+)2; (+)3; (+)4 O.N. corresponds to the no. of electrons in outer/valence shell/lost [1]
  - **(b)** PC*l*<sub>5</sub> fizzes or white/misty fumes or heat evolved [1]  $PCl_5 + 4H_2O \rightarrow H_3PO_4 + 5HCl$  or  $PCl_5 + 3H_2O \rightarrow HPO_3 + 5HCl$ (allow partial hydrolysis:  $PCl_5 + H_2O \rightarrow POCl_3 + 2HCl$ ) [1]
  - (c) (i) P = 30.4/31 = 0.98 Cl = 69.6/35.5 = 1.96[1] Thus E.F =  $PCl_2$ [1]

 $M_r(PCl_2) = 102$ , so  $2 \times PCl_2 = 204 \approx 200$ , so M.F. =  $P_2Cl_4$ [1]

(ii) [1] (ignore lone pairs on Cl)

(iii) O.N. = 
$$(+)2$$

(iv)  $(HO)_2P-P(OH)_2$  or H(HO)P(=O)-P(=O)(OH)Hecf from structure in (ii) [1] Allow HO-P-OH or HO-P=O Η

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**4** (a) 
$$N_2 + 2O_2 \rightarrow 2NO_2$$
 (or via NO) or  $2NO + O_2 \rightarrow 2NO_2$  [1]

(b) (i) catalytic converter **and** passing the exhaust gases over a catalyst/Pt/Rh [1]

(ii) 
$$NO_2 + 2CO \rightarrow \frac{1}{2} N_2 + 2CO_2$$
 or similar  
Allow  $2NO_2 + CH_4 \rightarrow CO_2 + N_2 + 2H_2O$  [1]

(c) No, it wouldn't be reduced. Because the reaction in (a) does not presuppose a particular fuel (owtte)
 Allow formed from N<sub>2</sub> and O<sub>2</sub> in air during combustion

(ii) 
$$NO + \frac{1}{2}O_2 \rightarrow NO_2$$
 [1]

(iii) 
$$K_p = (p_{NO}.p_{SO_3})/(p_{NO_2}.p_{SO_2})$$
 [1] units: dimensionless/none (don't accept just a blank!)

(iv) 
$$K_p = 99.8^2/0.2^2 = 2.5 \times 10^5$$
 [1]

[Total: 11]

## 5 (a)

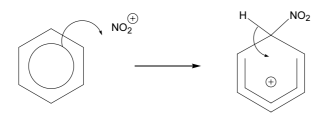
transformation	reagent + conditions
$C_2H_4 \rightarrow C_2H_5Cl$	HC <i>I</i> , no light or catalyst
$C_2H_5OH \rightarrow C_2H_5Cl$	conc $HCl + ZnCl_2$ or $SOCl_2$ or $PCl_5$ or $PCl_3$ and heat
$C_2H_6 \rightarrow C_2H_5Cl$	C 1 <sub>2</sub> + light
$C_2H_4 \rightarrow C_2H_4Cl_2$	C l <sub>2</sub> , no light or catalyst
CH₃CO₂H → CH₃COC <i>l</i>	SOC $l_2$ or PC $l_5$ or PC $l_3$ and heat
H₃C → H₃C → C1	C l <sub>2</sub> + A lC l <sub>3</sub>
— CH <sub>3</sub> — CH <sub>2</sub> C <i>l</i>	C 1 <sub>2</sub> + light or heat

[6]

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(b) (i) production of 
$$NO_2^+$$
:  $2H_2SO_4 + HNO_3 \rightarrow 2HSO_4^- + H_3O^+ + NO_2^+$  [1] (accept  $H_2SO_4 + HNO_3 \rightarrow HSO_4^- + H_2O + NO_2^+$ )



curly arrow from ring to  $NO_2^+$  and from C-H bond to ring [1] correct intermediate, including charge in the right place Note charge area must be more than half ring [1]

(ii) 
$$\mathbf{C}$$
 is  $C_6H_5CO_2H$  [1]

(iii) step 1: reagent is hot acidified or alkaline 
$$KMnO_4$$
 [1] step 2: reagent is  $Br_2 + FeBr_3/A_1C_1$  etc. ( $H_2O$  or light negates) [1]

(If  ${\bf C}$  is given as 3-bromotoluene, then allow the last [2] marks if steps 1 and 2 are reversed.)

[Total: 12]

6 (a) (i) aqueous alkaline iodine 
$$or I_2 + OH^-(aq)$$
 allow NaC1O + KI [1]

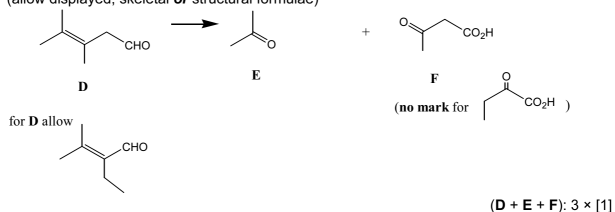
(iv)

compound	result
CH₃OH	×
CH₃CH₂OH	✓
CH₃CHO	✓
CH <sub>3</sub> CO <sub>2</sub> H	×
СНО	×
—сосн <sub>3</sub>	<b>√</b>

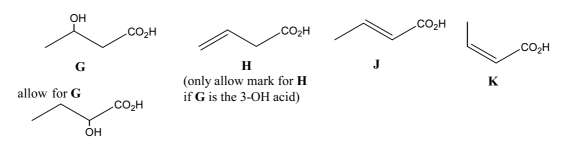
• √ • √ • √ [3]

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(b) (allow displayed, skeletal or structural formulae)



(c) (allow displayed, skeletal and structural formulae) Must be consistent with F



(N.B. letters **H**, **J**, **K** can be swapped around)

 $(G + H + J + K): 4 \times [1]$ 

geometrical or cis-trans isomerism

[1]

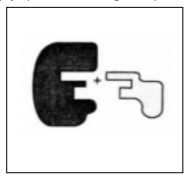
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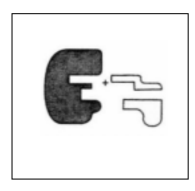
7 (a) The tertiary/3-dimensional structure/shape is held together by hydrogen/ionic/van der Waals bonds [1]

These break (relatively) easily/are weak/break at/above 45 °C

(b) (or similar diagrams)







Enzyme + substrate

Enzyme-substrate complex

Enzyme + products

3 × [1]

[1]

(c) a competitive inhibitor combines with the enzyme's active site (so preventing the substrate from binding) [1]

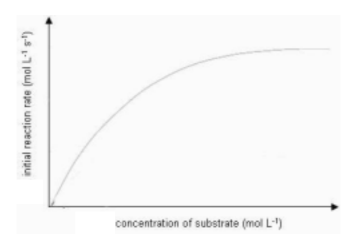
non-competitive inhibitor bonds with the enzyme away from the active site/at an allosteric site [1]

this changes the shape of the active site

[1]

**Also allow** competitive inhibition can be overcome by increasing [substrate] **or** non-competitive inhibition cannot be removed by increasing [substrate] for the 3rd mark

(d) (i)



Line must be of similar shape to original but level out below original line

[1]

(ii) Inhibitor reduces the number of enzymes with 'working' active sites (owtte)

[1]

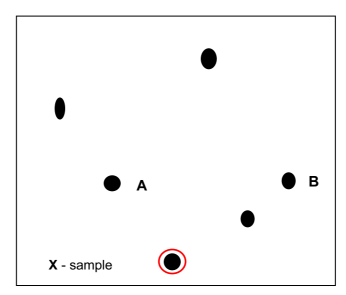
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8 (a) partition – separation due to the different solubilities of compounds in two solvents/phases

[1]

adsorption – separation due to the different attractions between the compounds and thestationary phase, relative to their solubility in the solvent[1]Note, if candidates do not refer to different solubilities and different attractionsmax 1

(b)



Ring: [1]

(c) (i) X is bromine – M and (M+2) peaks almost same height [1]

(ii) 
$$\frac{M}{M+1} = \frac{100}{1.1} \times \frac{9}{n} = \frac{100}{0.3}$$
 1.1 × n

Hence 
$$n = \frac{100 \times 0.3}{1.1 \times 9} = 3.03$$
  $p = 3$ 

(answer + working) [1]

(If the mass peak is at 122 and the compound contains Br and 3 C atoms then Q = (122 - 79 - 36)) thus Q = 7 ecf from (ii) [1]

(The compound is  $C_3H_7Br$ )

(iii) (R is at m/e 43), hence 
$$C_3H_7^+$$
 [1]

(d) Any **two** from 
$$H_2$$
,  $H_2O$ ,  $CO$ ,  $C_2H_4$ ,  $C_2H_2$ ,  $CH_4$  2 × [1]

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**9** (a) (i) One [1]

(ii) Any alkene (or allow a cyclic amide, as in caprolactam) [1]

(b) Any TWO from: addition needs unsaturated/double bonds/alkene

condensation eliminates a small molecule

condensation needs a molecule other than a hydrocarbon

empirical formula of addition polymer is the same as that of its monomer

condensation needs two different functional groups

(**NOT** – "condensation needs two different monomers") 2 × [1]

(ii)

Correct 'ester' bond [1]

'sticks' to rest of molecule [1]

Note: candidates need only show 'brackets' if more than one repeat unit shown

(iii) Polyesters [1]

(d) Monomers in *Terylene* have to alternate in order to condense out water (owtte) [1]

Alkenes can link in any order (and still form a polyalkene) (*or* diagram showing this) [1]