

**UNIVERSITY OF CAMBRIDGE INTERNATIONAL EXAMINATIONS****GCE Advanced Subsidiary Level and GCE Advanced Level****MARK SCHEME for the May/June 2008 question paper****9701 CHEMISTRY****9701/04**

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

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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.

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- 1 (a) (i) A is  $\text{Cl}_2$ /chlorine [1]
- B is  $\text{NaCl}$  or  $\text{HCl}$  or  $\text{Cl}^-$  [or words], etc. [1]
- C is salt bridge or  $\text{KCl/KNO}_3$ , etc. [1]
- D is platinum/Pt [1]
- E is  $\text{Fe}^{2+} + \text{Fe}^{3+}$  or mixture of Fe(II) + Fe(III) salts [1]
- mention of standard conditions ( $[\text{Cl}^-]$  of  $1 \text{ mol dm}^{-3}$  or  $\text{Cl}_2$  at 1 atmos  
or  $T = 25^\circ\text{C}/298 \text{ K}$ ) [1]
- (ii)  $E^\circ = E^\circ_{\text{R}} - E^\circ_{\text{L}} = 0.77 - 1.36 = (-)0.59 \text{ (V)}$  (ignore sign) [1]
- (since R.H. electrode is negative) electrons flow (from right) **to left** or to the chlorine  
electrode or anticlockwise or from (beaker) **E** to (beaker) **B** [1] [8]
- (b) (i)  $\Delta H = 3 \times (-167.2) + (-48.5) - (-399.5)$  [1]  
 $= -150.6 \text{ or } 151 \text{ (kJ mol}^{-1}\text{)}$  [1]  
 (correct ans [2])
- (ii)  $2\text{Fe}^{3+} + \text{Cu} \longrightarrow 2\text{Fe}^{2+} + \text{Cu}^{2+}$  [1]  
 (or molecular:  $2\text{FeCl}_3 + \text{Cu} \longrightarrow 2\text{FeCl}_2 + \text{CuCl}_2$ )
- $E^\circ = 0.77 - 0.34 = (+) 0.43 \text{ (V)}$  [1]  
 (no mark for  $-0.43\text{V}$ ) [4]
- [Total: 12 max 11]**
- 2 (a) (i)  $\Delta H = 4 \times 278 - 244 - 2 \times 496$  [1]  
 $= -124 \text{ (kJ mol}^{-1}\text{)}$  [1]  
 (correct ans [2])
- (ii) shape is bent/V-shaped/non-linear (or diagram) [1]  
 due to (one) lone pair and/or (1) odd/unpaired electron (or shown on diag) [1]  
 (assume electrons are on chlorine unless explicitly stated otherwise, in which case  
 award no mark)
- (iii)  $3\text{KClO}_3 + \text{H}_2\text{SO}_4 \longrightarrow \text{K}_2\text{SO}_4 + \text{KClO}_4 + \text{H}_2\text{O} + 2\text{ClO}_2$  [1] [5]
- (b) (i) coal-fired power stations; fuel in cars; car exhausts/gas emissions; other named use of a  
 fossil fuel; contact process; cement manufacture; brick manufacture; roasting of sulphide  
 ores; burning tyres (any 2) [1]  
 (NOT volcanoes etc; NOT burning of natural gas)  
 (no marks for only 1 correct source)
- (ii) causes **acid rain** [1]  
 which lower pH of lakes; leaches aluminium from soils; kills fish/plants/rainforests;  
 dissolves/corrodes/damages buildings (any 1) [1]  
 (NOT asthma etc – since this is not environmental) [3]

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- (c) (i)  $\text{CO}_2$ : simple + molecular/covalent *or* weak intermolecular forces  
 $\text{SiO}_2$ : giant/macro + molecular/covalent  
 $\text{SnO}_2$ : ionic/electrovalent (ignore “giant”) (all 3 correct) [2]  
 (2 correct = [1], 1 correct = [0])

- (ii)  $\text{SnO}_2$  is stable,  $\text{PbO}_2$  is not *or*  $\text{SnO}_2$  is the more stable [1]  
 $\text{PbO}_2 \longrightarrow \text{PbO} + \frac{1}{2} \text{O}_2$  [1]

- (iii)  $\text{H}_2\text{O} + \text{CO}_2 (\rightleftharpoons) \text{H}^+ + \text{HCO}_3^-$  [1]  
 $K_c = [\text{H}^+][\text{HCO}_3^-]/[\text{H}_2\text{O}][\text{CO}_2]$  *or*  $[\text{H}^+][\text{HCO}_3^-]/[\text{CO}_2]$  ecf [1]

- (iv)  $\text{HCO}_3^- + \text{H}^+ \longrightarrow \text{H}_2\text{CO}_3$  *or*  $\text{H}_2\text{O} + \text{CO}_2$  (or equation with  $\text{H}_3\text{O}^+$ ) [1]  
 $\text{HCO}_3^- + \text{OH}^- \longrightarrow \text{CO}_3^{2-} + \text{H}_2\text{O}$  (NB **NOT**  $\text{H}_2\text{CO}_3 + \text{OH}^- \rightarrow$ ) [1]

(words can substitute for one of the equations but not both. If two correct word descriptions are given, in the absence of at least one correct equation, award [1] mark only)

[8]

[Total: 16 max 15]

- 3 (a) tetrahedral diagram (either dashed+wedge, or similar representation) [1]  
 angles (all)  $109^\circ - 110^\circ$  [1]  
 (award [0] for part (a) if an angle of  $90^\circ$  or  $180^\circ$  is mentioned) [2]

- (b) volatility decreases *or* boiling points increase  
 (allow b.pt.  $\text{CCl}_4 > \text{SiCl}_4$  but b.pt. increases thereafter) [1]  
 due to greater van der Waals'/intermolecular forces *or* due to more electrons [1]  
 (mention of “ions” negates this mark) [2]

- (c) (i)  $\text{Pb}^{4+}/\text{Pb}^{2+}$ :  $E^\ominus = +1.69\text{V}$ ,  $\text{Sn}^{4+}/\text{Sn}^{2+}$ :  $E^\ominus = +0.15\text{V}$ , [both] [1]  
 a valid comment about relative redox power *or* stability, e.g.:  
 (hence)  $\text{Sn}^{2+}$  easily oxidised *or*  $\text{Sn}^{4+}$  is more stable than  $\text{Sn}^{2+}$  *or*  
 $\text{Pb}^{4+}$  is easily reduced *or*  $\text{Pb}^{2+}$  is more stable than  $\text{Pb}^{4+}$  *or*  
 +2 oxidation state more stable down the group [1]

- (ii)  $\text{Sn}^{2+} + \text{I}_2 \longrightarrow \text{Sn}^{4+} + 2\text{I}^-$  [1]  
 $\text{Pb}^{4+} + \text{SO}_2 + 2\text{H}_2\text{O} \longrightarrow 4\text{H}^+ + \text{SO}_4^{2-} + \text{Pb}^{2+}$  [1]  
 (N.B. no marks in (ii) for  $E^\ominus$  values) [4]

- (d) (i) for Si:  $\Delta H = 244 - 2(359) = -474$  ( $\text{kJ mol}^{-1}$ ) [1]  
 for Sn:  $\Delta H = 244 - 2(315) = -386$  ( $\text{kJ mol}^{-1}$ ) [1]  
 (allow [1] out of [2] salvage mark for 474 & 386; 962 & 874; *or* -962 & -874)

- (ii) Yes: the +4 state becomes decreasingly stable – the  $\Delta H$  is less exothermic [1]  
 (mark is for relating  $\Delta H$ s to stability: allow ecf from **d(i)** and also from **c(i)**) [3]

[Total: 11]

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- 4 (a) ester [1] [1]
- (b) reaction I: acid/ $\text{H}^+$ /HCl/ $\text{H}_2\text{SO}_4$  or alkali/ $\text{OH}^-$ /NaOH (followed by  $\text{H}^+$ ) [1]  
**heat/reflux** and aqueous (allow  $\text{H}_3\text{O}^+$  to equal  $\text{H}^+$  + aq, also assume “conc” *or* “dil” means aq (but NOT  $\text{H}_2\text{SO}_4$ ) also allow aqueous ethanol) [1]  
 (for heat: allow  $T \geq 80^\circ\text{C}$ ; **not** “warm”)
- reaction II: methanol/ $\text{CH}_3\text{OH}$  [1]  
 heat with **conc.**  $\text{H}_2\text{SO}_4/\text{H}_3\text{PO}_4$  *or* HCl(**g**) [**NOT** conc HCl] [1] [4]
- (c) (i)  $\text{BrCH}_2\text{-CHBr-CH}_2\text{Br}$  [1]
- (ii)  $\text{HO}_2\text{C-CO-CO}_2\text{H}$  [1] [2]
- (d) 890g of triglyceride produces  $3 \times 298 = 894\text{g}$  of biodiesel [1]  
 $\therefore$  500kg produces  $500 \times 894/890 = \mathbf{502\text{kg}}$  biodiesel ecf [1]  
 (correct ans [2])  
 (1004/1005kg *or* 167kg is worth [1]; 333kg is worth [0]) [2]
- (e) (i)  $\text{C}_{17}\text{H}_{35}\text{CO}_2\text{CH}_3 + 27.5 \text{ O}_2 \longrightarrow 19\text{CO}_2 + 19\text{H}_2\text{O}$  [1]  
 (*or*  $\text{C}_{19}\text{H}_{38}\text{O}_2$ )
- (ii)  $10 \times 44 \times 19/298 = \mathbf{28. (05)/28.1\text{kg}}$  ecf from equ [2]  
 (–1 for each error)  
 some ecf values:  $n = 18 \Rightarrow 26.6\text{kg}$   
 $n = 17 \Rightarrow 25.1\text{kg}$  (allow [2] for each)  
 $n = 16 \Rightarrow 23.6\text{kg}$  [3]
- (f) any one of the following.
- (saving) diminishing resources
  - economic argument (NOT just “cheaper”) – e.g. oil will become increasingly more expensive as it runs out
  - ref to  $\text{CO}_2$  cycle (e.g. no net increase in  $\text{CO}_2$ , i.e. “carbon neutral”) *or* less global warming (due to a smaller carbon “footprint”)
  - renewable/sustainable
  - the effect of biofuel cultivation on world food prices

[1] [1]

[Total: 13]

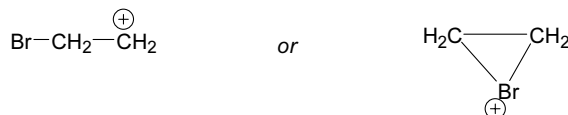
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5 (a) reaction I electrophilic addition [1]

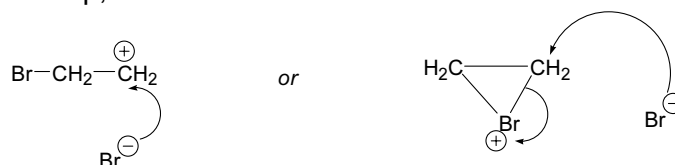
reaction II electrophilic substitution [1]

(salvage: award [1] out of [2] for “addition” + “substitution”, even if nucleophilic) [2]

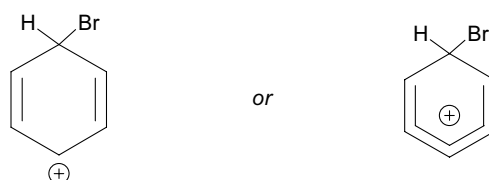
(b) reaction I: intermediate [1]



second step, attack of  $\text{Br}^-$  on bromocation. [1]



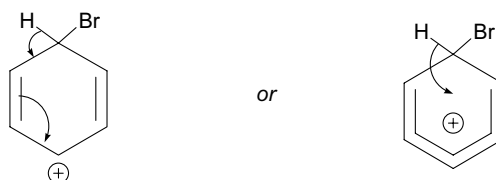
reaction II: intermediate [1]



(or with  $\oplus$  in 2-position)

(make sure  $\oplus$  is not at  $\text{sp}^3$  C-atom)

second step, loss of  $\text{H}^+$  from bromocation. [1]



[4]

(c) **Delocalised** ring of electrons (in benzene) is **stable**, (so is re-formed in second step in benzene.)

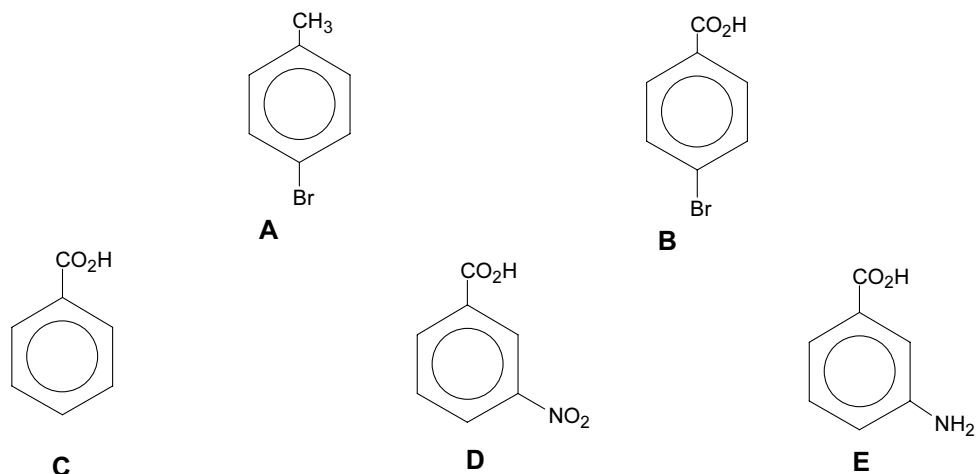
or electrons in the ethene  $\pi$  bond are localised/more available for reaction with electrophiles

[1] [1]

[Total: 7]

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6



5 x [1]

[deduct [1] mark if ring circle omitted more than once]

[allow ecf for **E** from structure of **D**]

[allow ecf for **B** from structure of **A**]

[allow -CO<sub>2</sub><sup>-</sup> for **E**]

[5]

[Total: 5]

7

| polymer | addition/condensation? | formulae of monomers                                                                                                                   |
|---------|------------------------|----------------------------------------------------------------------------------------------------------------------------------------|
| 1       | condensation           | HO <sub>2</sub> C-CO <sub>2</sub> H or ClCO-COCl<br>NH <sub>2</sub> -CH <sub>2</sub> -CH <sub>2</sub> -NH <sub>2</sub>                 |
| 2       | condensation           | HO-CH <sub>2</sub> -CH(C <sub>2</sub> H <sub>5</sub> )-CO <sub>2</sub> H<br>HO-CH <sub>2</sub> -CH(CH <sub>3</sub> )-CO <sub>2</sub> H |
| 3       | addition               | CH <sub>2</sub> =CH-CH <sub>3</sub><br>CH <sub>2</sub> =CH-CONH <sub>2</sub><br>CH <sub>2</sub> =CH-C <sub>6</sub> H <sub>5</sub>      |

↑  
[2]  
(2 correct: [1])

↑  
[6]  
(6 correct: [5])  
etc

(2 correct: [1])

(C=C bonds not needed, but penalise -[1] if C-C drawn instead of C=C)

(if more than 7 formulae drawn, then penalise -[1] for each formula in excess of 7)

[8]

[Total: 8]

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- 8 (a) primary: covalent (ignore amide, peptide etc) [1]  
 diagram showing peptide bond: (-CHR-)CONH(-CHR-) [1]
- secondary: hydrogen bonds (NOT “..between side chains”) [1]  
 diagram showing N-H...O = C [1]
- tertiary: two of the following:  
 • hydrogen bonds (diag. must show H-bonds *other* than those in  $\alpha$ -helix or  $\beta$ -pleated sheet – e.g. ser-ser)  
 • electrostatic/ionic attraction,  
 • van der Waals’/hydrophobic forces/bonds,  
 • (covalent) disulphide (links/bridges) [1] + [1]
- suitable diagram of **one** of the above [1]  
 (for disulphide: S-S **not** S=S or SH-SH) [7]
- (b) met-ala-gly-ala-gly-arg-val-lys [2]  
 any **possible** sequence with more than 8 residues, that “uses” all 6 tripeptides (overlapping or not), and that starts with *met* and ends with *lys* is worth [1] mark  
 any sequence that does **not** start with *met* or end with *lys* gets zero. [2]
- (c) CARE – this is not about DNA!  
 candidates should describe **TWO** potential effects on tertiary or quaternary structures caused by amino acid sidechains...  
 these include: disruption of H-bonding  
 disruption of disulphide bridges  
 disruption of electrostatic/ionic attraction  
 disruption of van der Waals’ forces  
 (only allow effects on the secondary structure if proline is specifically mentioned) 2 x [1]
- then award [1] mark each for **two** of the following bullet points:  
 • a description of the amino acids involved in the above, (*or* a labelled diagram) (award [1] mark for each example)  
 a description of an *effect* of interchanging amino acids, such as the..  
 • unfolding of tertiary structure/different folding/different shape (NOT denatured)  
 • inactivity of an enzyme *or* changing the active site  
 • causing of a protein to become less soluble/coagulate (e.g. sickle cells) 2 x [1]
- [4]
- [Total: 13 max 12]

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- 9 (a) (i)+(ii) any two of:  
 molecular mass/size/ $M_r$ /shape  
 (overall electrical) charge (on the species)  
 voltage/size/P.D. (of applied electric field) [1] + [1]  
 (salvage: if just “mass & charge” is mentioned, with no reference to species or molecule, award [1]) [2]
- (b) (i)  $\text{CH}_3\text{COCH}_3$  would show  
 a single peak/no splitting since all the Hs are in the same chemical environment  
 or a peak at  $\delta = 2.1$  due to  $\text{CH}_3\text{CO}$  group [1]
- $\text{CH}_3\text{CH}_2\text{CHO}$  would show 3 (sets of) peaks since there are 3 different proton environments  
 or there would be a peak at  $\delta = 9.5 - 10.0$  due to the  $-\text{CHO}$  group  
 or a peak at  $\delta = 0.9$  due to  $\text{CH}_3$   
 or a peak at  $\delta 1.3$  due to  $\text{CH}_2$  [1]
- (reasons needed for the marks. Salvage: if reasons are not given, but candidate states that propanone will have one peak and propanal three, then award [1] mark)
- (ii) different fragments:
- $\text{CH}_3\text{COCH}_3$  would form **fewer** fragments (must be stated in words)
  - $\text{CH}_3\text{COCH}_3$  would form a fragment of  $\text{CH}_3\text{CO}^+$  or at (m/e) 43
  - $\text{CH}_3\text{CH}_2\text{CHO}$  would form a fragment of  $\text{CH}_3\text{CH}_2^+$  or  $\text{CHO}^+$  at (m/e) 29
  - $\text{CH}_3\text{CH}_2\text{CHO}$  would form a fragment of  $\text{CH}_3\text{CH}_2\text{CO}^+$  or at (m/e) 57
- [charges on fragments not required for mark]
- any 3 points [3]  
[5]
- (c) (i) peaks at (m/e) 79 **and** 81 or at (m/e) 94 **and** 96 [1]
- (ii) in chlorine the M and M+2 peaks are the ratio 3:1 [1]  
 whereas in bromine they are approx. 1:1 [1] [3]

[Total: 10 max 9]

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10 (a) any **two** of the following:

- to speed delivery (of drug to target organ), i.e. faster response
- to avoid the drug being hydrolysed/reacted/decomposed (NOT digested) in the stomach
- to allow a smaller dose to be used *or* greater accuracy of dosage
- patient does not have to be conscious

2 × [1] [2]

(b) (i) spheres with a diameter of the order of nanometres/in the nanometre range/between 10 & 500 nm [1]

(ii) it is (highly) acidic *or* low pH *or* contains HCl (NOT contains enzymes) [1]

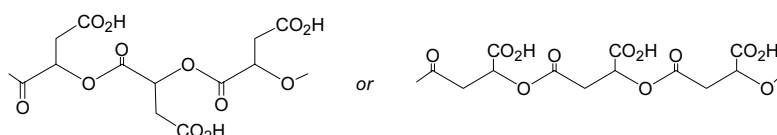
(iii) use hydrogels: of different (wall) thickness/strength (to release drug over time)  
of different chemical composition (for different breakdown times)  
incorporating pores/holes (in their walls) (any two) [1] + [1]

[4]

(c) for the **homopolymer**, **either** using the amino acid the minimum is:

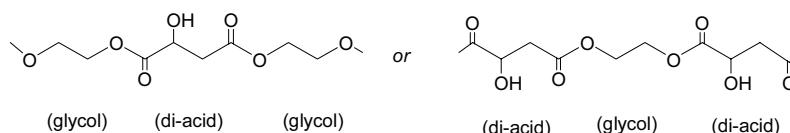


**or** using the hydroxyacid the minimum is:

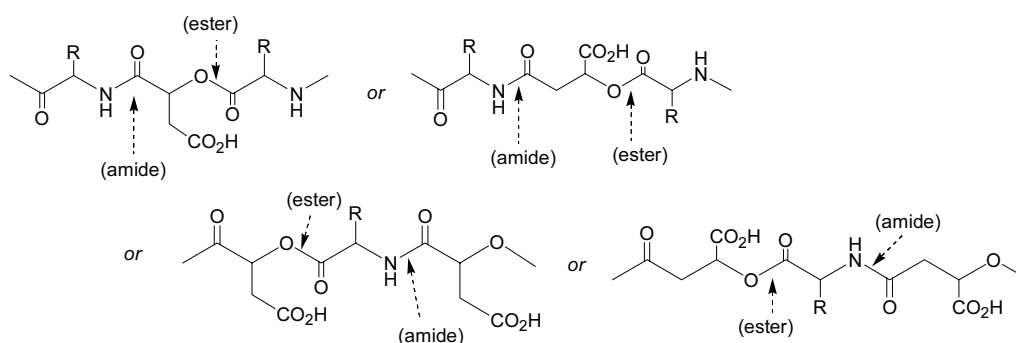


(–[1] for each error) [2]

for the **heteropolymer**, **either** using the glycol compound and the di-acid the minimum is:



**or** using the amino acid and the di-acid, the minimum is:



(A heteropolymer incorporating all three monomers can also be drawn. This should include an ester linkage between the glycol and one of the CO<sub>2</sub>H groups, and an amide linkage between the amino acid and another CO<sub>2</sub>H group. Deduct [1] mark from the whole of section (c) if complete compounds are shown rather than sections of chains. Allow 4-monomer sections instead of 3. Allow [2] marks for a polymer section even if **one** end is incomplete (e.g. is lacking an oxygen atom), but if **both** ends are incomplete deduct [1]) (–[1] for each error) [2] [4]

[Total: 10 max 9]