MARK SCHEME for the October/November 2011 question paper

for the guidance of teachers

9701 CHEMISTRY

9701/43

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

This mark scheme is published as an aid to teachers and candidates, to indicate the requirements of the examination. It shows the basis on which Examiners were instructed to award marks. It does not indicate the details of the discussions that took place at an Examiners' meeting before marking began, which would have considered the acceptability of alternative answers.

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| | Page 2 | Mark Scheme: Teachers' version | Syllabus | Paper | |
|---|--|--|----------|--------------------------|--|
| | | GCE A LEVEL – October/November 2011 | 9701 | 43 | |
| 1 | (a) Cr ³⁺ : Mn ²⁺ : | 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 3d ³ 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 3d ⁵ | | [1] [1] [2] | |

- (b) (i) Any two from
 - H^+ is on the oxidant/L.H. side of each of the $\frac{1}{2}$ -equations, or H^+ is a reactant
 - (increasing [H⁺]) will make E^e more positive
 - (increasing [H⁺]) will drive the reaction over to the R.H./reductant side or forward direction

[1] + [1]

- (ii)KMnO4:Purple/violet to colourless (allow very pale pink)[1]K2Cr2O7Orange to green[1][4]
- (c) (i) $MnO_2 + SO_2 \longrightarrow MnSO_4 (or Mn^{2+} + SO_4^{2-})$ [1]

| manganese changes/is reduced from +4 to +2 | [1] |
|--|-----|
| sulfur changes/is oxidised from +4 to +6 | [1] |

- (ii) No effect, because H⁺ does not appear in the overall equation *or* its effect on the MnO₂/Mn²⁺ change is cancelled out by its effect on the SO₂/SO₄²⁻ change [1]
 [4]
- (d) (i) $MnO_2 + 4H^+ + Sn^{2+} \longrightarrow Mn^{2+} + 2H_2O + Sn^{4+}$ [1]
 - (ii) $n(MnO_4^-) = 0.02 \times 18.1/1000 = 3.62 \times 10^{-4} \text{ mol}$ [1] $n(Sn^{2^+}) = 3.62 \times 10^{-4} \times 5/2 = 9.05 \times 10^{-4} \text{ mol}$ [1] $n(Sn^{2^+})$ that reacted with $MnO_2 = (20 - 9.05) \times 10^{-4} = 1.095 \times 10^{-3} \text{ mol}$ [1] reaction is 1:1, so this is also $n(MnO_2)$ mass of $MnO_2 = 1.095 \times 10^{-3} \times (54.9 + 16 + 16) = 0.0952 \text{ g}$ [1] \Rightarrow **95% - 96%;** 2 or more s.f. [1]

[Total: 16]

| | Page 3 | | | Mark Scheme: Teachers' version | Syllabus | Paper |
|---|--------|-------|---|--|--|---|
| | | | | GCE A LEVEL – October/November 2011 | 9701 | 43 |
| 2 | (a) | (i) | A m tł | olecule/ion/species with a lone pair (of electrons) <i>or</i> elenat bonds to a metal ion/transition element | ectron pair dono | r [1] |
| | | (ii) | by | means of a dative/coordinate (covalent) bond | | [1] [2] |
| | (b) | (i) | strai | ght line from (0, 0.01) to point at (350, 0.0028) with all | points on the lin | e [1] |
| | | (ii) | orde | er w.r.t. $Cr(CO)_6$ is 1 and order w.r.t. PR_3 is zero | | [1] |
| | | | beca or co beca of re | ause (a) $Cr(CO)_6$ graph has a constant half-life (which is construction lines on graph showing this) ause (b) PR_3 graph is a straight line (of constant slope eaction <i>or</i> no change in rate <i>or</i> shows a linear decrease | is 700 s)) <i>or</i> line shows a e | [1] a constant rate [1] |
| | | (iii) | rate | = k[Cr(CO) ₆] | | [1] |
| | | | k = (| (0.9 – 1.1) × 10 ^{−3} (s ^{−1}) (one or more s.f.) | | [1] |
| | | | eithe or | er rate ₀ = 0.01/1020 = 9.8×10^{-6} mol sec ⁻¹ when [Cr(Ce so k = $9.8 \times 10^{-6}/0.01 = 9.8 \times 10^{-4}$ t _{1/2} \approx 700 sec k = 0.693/700 = 9.9 × 10^{-4 } | O) ₆] = 0.01 mol | dm ⁻³ |
| | | (iv) | (unit | s of k are) sec⁻¹ | | [1] |
| | | (v) | N.B. <i>eithe</i> mec beca step | the chosen mechanism must be consistent with the rate r if rate = k[Cr(CO)_6] hanism B is consistent ause it's the only mechanism that does NOT involve P or only Cr(CO)_6 is involved in slow step or [PR ₃] does | ate equation in (i PR₃ in its slow/ra not affect the ra | ii). Thus: [1] te-determining te [1] |
| | | | or | | | |
| | | | if rat mec beca | e = k[Cr(CO) ₆][PR ₃], then hanism A or C or D is consistent ause both reactants are involved in slow step | | [1] [1] [9] |
| | | | | | | [Total: 11] |

| | Page 4 | Mark Scheme: Teachers' version | Syllabus | Paper | |
|---|------------------------------|--|----------|-------------------|--|
| | | GCE A LEVEL – October/November 2011 | 9701 | 43 | |
| 3 | (a) (i) E is | CH ₃ CH(NH ₂)CN | | [1] | |
| | (ii) C ₆ H | ₅CH₂CHO | | [1] [2] | |

 (b) (i) a polymer/polypeptide of amino acids, (joined by peptide bonds) (allow 'chain of amino acids' but not 'sequence': the idea of 'many' has to be conveyed)
 [1]

(ii)



peptide bond shown in full (C=O) in an ala-ala fragment in a chain [1] two repeat units [1]

Allow peptide bond shown in full (C=O) in a dipeptide ala-ala for 1 mark



[3]

(c) (i) $HCl or H_2SO_4 or NaOH or H^+ or OH^-$ reagents [1] + heat and H_2O/aq (allow H_3O^+). If T is quoted, 80 °C < T < 120 °C. NOT warm. conditions [1]

(ii)



(if a structural formula, it must have all H atoms) allow protonated or deprotonated versions [1] + [1] [max 3]

| Page 5 | Mark Scheme: Teachers' version | Syllabus | Paper |
|--------|-------------------------------------|----------|-------|
| | GCE A LEVEL – October/November 2011 | 9701 | 43 |

⁽d) (i) $NH_3^+ - CH(CH_3) - CO_2^-$

(ii)



[3] **[4]**

(e) (i) A buffer is a solution whose pH stays fairly constant or which maintains roughly the same pH or which resists/minimises changes in pH [1] when small/moderate amounts of acid/H⁺ or alkali/OH⁻ are added [1] (ii) $NH_2CH(CH_3)CO_2H + H(Cl) \longrightarrow {}^{+}NH_3CH(CH_3)CO_2H (+ Cl^{-})$ [1] (iii) blood contain HCO_3^- (or in an equation) [1] $\begin{array}{c} H^{+} + HCO_{3}^{-} \longrightarrow H_{2}CO_{3} (H_{2}O + CO_{2}) \\ OH^{-} + HCO_{3}^{-} \longrightarrow CO_{3}^{2-} + H_{2}O \end{array}$ which absorbs H^+ or equn or absorbs OHor equn [1] (iv) $[CH_3CO_2Na] = 0.05 [CH_3CO_2H] = 0.075$ [1] pH = 4.76 + log (0.05/0.075) = 4.58 or 4.6[1] [7]

[Total: 19]

[1]

| | Page 6 | Mark Scheme: Teachers' version | Syllabus | Paper |
|---|---|--|--|--|
| | | GCE A LEVEL – October/November 2011 | 9701 | 43 |
| 4 | (a) Ca(NO ₃ | $)_2 \longrightarrow CaO + 2NO_2 + \frac{1}{2}O_2$ | | [1] [1] |
| | (b) (down decomp as size/ so polai | the group) nitrates become more stable or requisose radius of (cat) ion increases or charge density of ion de risation/distortion of anion/nitrate decreases | ire a higher te ecreases | emperature to [1] [1] [1] [3] |
| | (c) (i) Li ₂ (| $CO_3 \longrightarrow Li_2O + CO_2$ | | [1] |
| | (ii) rad | ius of Li ion/Li $^{\scriptscriptstyle +}$ is less than that of Na ion/Na $^{\scriptscriptstyle +}$ (or polaris | sing power of M [*] | is greater) [1] |
| | (iii) Bro Sin car | wn/orange fumes/gas would be evolved <i>or</i> glowing splice the nitrate is likely to be thermally unstable obonate) <i>or</i> the balanced equation: $2\text{LiNO}_3 \longrightarrow \text{Li}_2\text{O} + $ | nt relights or decomposes + 2NO ₂ + ½O ₂ | [1] (just like the [1] [4] |

[Total: 8]

| | Page 7 | | Mark Scheme: Teachers' version | Syllabus | Paper |
|---|-------------------------|----------------------------------|---|----------|-------------------|
| | | GC | E A LEVEL – October/November 2011 | 9701 | 43 |
| 5 | (a) Alka or C | anes are non-p C and H have s | oolar <i>or</i> have no dipole <i>or</i> C–H bonds are s similar electronegativities | trong | [1] [1] |
| | (b) (i) | (free) radical | substitution <i>or</i> substitution by homolytic fiss | sion | [1] |
| | (ii) | initiation: | $Cl_2 \longrightarrow 2Cl^{\bullet}$ | | [1] |
| | | propagation: | $C_{l}^{l} + C_{2}H_{6} \longrightarrow C_{2}H_{5}^{\bullet} + HC_{l}^{l}$ $C_{2}H_{5}^{\bullet} + C_{l_{2}} \longrightarrow C_{2}H_{5}C_{l} + C_{l}^{\bullet}$ | | [1] |
| | | termination: | $C_2H_5^{\bullet} + Cl^{\bullet} \longrightarrow C_2H_5Cl$ or $Cl^{\bullet} + Cl^{\bullet} \longrightarrow Cl_2$ etc | | [1] |
| | | | | | all 3 names [1] |
| | (iiii) | | | | |

| structural formula of by-product | formed by |
|---|--|
| CH ₂ CI–CH ₂ CI (or isomer) | further substitution |
| CH ₃ CH ₂ CH ₂ CH ₃ | (termination of 2 ×) C ₂ H ₅ • |
| CH₃CH₂CH₂CH₂CI (or isomer) | substitution of C₄H ₁₀ by-product |

[3]

accept in the "formed by" column the formulae of radicals that will produce the compound in the "by-product" column, or the reagents, e.g. $C_4H_9^{\bullet} + Cl_2 \text{ or } C_4H_9^{\bullet} + Cl$ or $C_4H_{10} + Cl_2$ (giving $CH_3CH_2CH_2CH_2CI$).

do not allow anything more Cl-substituted than **di**chlorobutane. N.B. C_2H_5Cl is the **major** product, not a **by**-product, so do not allow C_2H_5Cl .

(iv) J/K = 2.3 : 1 or 7:3 or 21:9 [2] (reason: straightforward relative rate suggests 21:1, but there are 9 primary to 1 tertiary, so divide this ratio by 9. 21/9 = 2.33) allow [1] mark if J/K ratio is given as 21:1;

[10]



[max 5]



| | Page 8 | | | Mark Scheme: Teachers' version | | Syllabus | Paper |
|---|---|-------------|--|--|--|-----------------------------------|--------------------------|
| | | | | GCE A L | EVEL – October/November 2011 | 9701 | 43 |
| 6 | (a) | (i) (ii) | K, b | ecause it is the <i>or</i> because it c molecule: J , molecule: L , | e (only) one to contain nitrogen <i>or</i> it's a ontains CO ₂ H <i>or</i> NH groups polymer: RNA (not DNA) polymer: starch, cellulose, glycogen (not carbohydrate) | n amino acid or polysaccharide | [1] [1] [2] |
| | (b) | (i) | Cov | alent bonding | | | [1] |
| | | (ii) | Hyd | rogen bonding | | | [1] |
| (iii) Ionic/electrovalent bonding <i>or</i> disul | | | lonic | c/electrovalent | bonding <i>or</i> disulphide/–S–S– bonding | <i>or</i> van der Waals | ' forces [1] [3] |
| | (c) | (i) | Enzy | ymes | | | [1] |
| | (ii) • change in pH increase in T (NOT decrease; T > 40 °C or "too high" are OK) • addition of heavy metal ions <i>or</i> specific, e.g. Hg²⁺, Ag⁺. Pb²⁺ etc. any two bullet points | | | | | points [1] + [1] | |
| | | | char <i>or</i> m <i>or</i> m <i>or</i> he | nge in pH disru letal ions disrup letal ions disrup eating disrupts | pts ionic bonds ot ionic bonds ot –S–S– bonds hydrogen bonds | | anv one [1] |
| | Thi | s cha | anges | s: the 3D struct | ure <i>or</i> shape of the enzyme <i>or</i> the acti | ve site | [1] [max 4] |

[Total: 9]

| Page 9 | Mark Scheme: Teachers' version | Syllabus | Paper |
|--------|-------------------------------------|----------|-------|
| | GCE A LEVEL – October/November 2011 | 9701 | 43 |

7 (a)

| structural information | analytical technique | |
|--|--|-----------------|
| three-dimensional arrangement of atoms and bonds in a molecule | X-ray crystallography/diffraction | |
| chemical environment of protons in a molecule | NMR (spectroscopy) only | |
| identity of amino acids present in a polypeptide | Electrophoresis / chromatography / mass spectrometry | |
| | | [1] + [1] + [1] |

[3]

(b) (i) paper chromatography;

The components **partition** between the solvent/moving phase and the water/liquid stationary phase *or* separation relies on different solubilities (of components) in the moving solvent and the stationary water phase. [1]

| (ii) | thin-layer chromatography. | |
|------|--|-------|
| | Separation depends on the differential adsorption of the components onto the | solid |
| | particles/phase or Al_2O_3 or SiO ₂ . | [1] |

[2]

[1]

(c) (i) No. of carbon atoms present =
$$\frac{0.2 \times 100}{5.9 \times 1.1}$$
 = 3.08 hence 3 carbons [1]

- (ii) Bromine
- (iii) One bromine is present as there is only an M+2 peak / no M+4 peak *or* the M and M+2 peaks are of similar height [1]
- (iv) NMR spectrum shows a single hydrogen split by many adjacent protons and 6 protons in an identical chemical environment. This suggests...
 two –CH₃ groups and a lone proton attached to the central carbon atom [1]

Empirical formula of **N** is C_3H_7Br

Hence **N** is $(CH_3)_2CHBr$ or H CH₃—C—CH₃

Β̈́r

[1]

[1]

[6]

[Total: 11]

| | Page 10 | | Mark Scheme: Teachers' version | Syllabus | Paper |
|---|--|------------------------------|--|----------|-------------------|
| | | | GCE A LEVEL – October/November 2011 | 9701 | 43 |
| 8 | (a) (i) | Solu | ble form would be most effective | | [1] |
| | (ii) | Q , si or P | ince the 'mini-pills'/granules/powder have a larger surfa , because it has no protective casing | ace area | [1] |
| | (iii) | The | gel coat stops it being broken down while passing through the upper part of the | | |
| | | dige | stive system/stomach | | |
| | | or th | e gel coat is stable to stomach acid. | | [1] [3] |
| | | | | | |
| | (b) The drug is taken guickly/directly to the target | | | | |
| | <i>or</i> more accurate dosing can be achieved | | | | |
| | When the drug is taken by mouth it has to pass through the stomach/intestine wall to get into the bloodstream. <i>or</i> some is digested/lost to the system [1] [2] | | | | |
| | (c) (i) | cond | densation (polymerisation) | | [1] |
| | (ii) | hydr | ogen bonds <i>or</i> van der Waals' | | [1] |
| | (iii) | lt wo The | ould change the overall shape of the (drug) molecule 'fit' into the active site would be less effective | | [1] + [1] |
| | (iv) | Hydı | rolysis | | [1] [5] |
| | | | | | [Total: 10] |
| | | | | | |