MARK SCHEME for the October/November 2011 question paper

for the guidance of teachers

9701 CHEMISTRY

9701/41

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.

Mark schemes must be read in conjunction with the question papers and the report on the examination.

• Cambridge will not enter into discussions or correspondence in connection with these mark schemes.

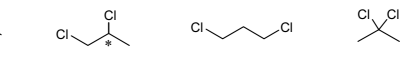
Cambridge is publishing the mark schemes for the October/November 2011 question papers for most IGCSE, GCE Advanced Level and Advanced Subsidiary Level syllabuses and some Ordinary Level syllabuses.



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|---|-----------------|------------|--|---|------------------|------------------------------|
| 1 | (a) (i) (ii) | | er burn <i>or</i> shine ligh | t/uv on mixture of $H_2 + Cl_2$ but NOT r of bromine decolourises/disappea | heat | [1] |
| | (1) | stea | my/misty/white fum ainer gets warm/ho | es produced | 115 | [2] |
| | (iii) | H-H | = 436 | C <i>l</i> -C <i>l</i> = 244 H | -Cl = 431 | |
| | | ΔH | = 436 + 244 - 2(43 | (1) = -182 kJ mol^{-1} | | [2] |
| | | H-H | = 436 | Br-Br = 193 H | -Br = 366 | |
| | | ΔH | = 436 + 193 – 2(36 | (6) = -103 kJ mol^{-1} | | [2] |
| | (iv) | H-Bi | r bond is weaker tha | an the H-C <i>l</i> bond – allow converse. | | [1] [8] |
| | (b) (i) | light | | | | [1] |
| | (ii) | | ds made = C-I & | I-I = $410 + 151 = 561$ H-I = $240 + 299 = 539$ H = $551 - 539 = +22$ kJ mol ⁻ | -1 | [2] |
| | (iii) | | overall reaction is led <i>or</i> high E _{act} | endothermic <i>or</i> no strong bond | ls/only weak bo | nds are [1] [4] |
| | (c) (i) | | olytic fission is the electron species | breaking of a bond to form (two) | radicals/neutral | species/ [1] |

- [1] (ii) •CH₂Cl [1] [3] the C-Br bond is the weakest or needs least energy to break/breaks most easily
- (d) CI

ĊI







- 4 structures: [2] 2 or 3 structures: [1]
 - [1] **[3]**

[Total: 18]

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Correct chiral atom identified

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| | | GCE A LEVE | L – October/November 2011 | 9701 | 41 |
| 2 | (a) (i) | Order w.r.t. [CH₃CHO] = Order w.r.t. [CH₃OH] = Order w.r.t. [H⁺] = 1 | | | [1] [1] [1] |
| | (ii) | rate = k[CH ₃ CHO][CH ₃ C | DH][H⁺] | | [1] |
| | (iii) | units = mol ⁻² dm ⁶ s ⁻¹ | | | [1] |
| | (iv) | rate will be 2 × 4 = 8 tim | nes as fast as reaction 1 (relative r | ate = 8) | [1] [6] |

(b)

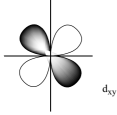
| | [CH ₃ CHO] /mol dm ⁻³ | [CH ₃ OH] /mol dm ⁻³ | [H ⁺] /mol dm ⁻³ | [acetal A] /mol dm ⁻³ | [H ₂ O] /mol dm ⁻³ |
|----------------|--|---|--|---|---|
| at start | 0.20 | 0.10 | 0.05 | 0.00 | 0.00 |
| at equilibrium | (0.20 – x) | (0.10 – 2x) | 0.05 | x | x |
| at equilibrium | 0.175 | 0.05 | 0.05 | 0.025 | 0.025 |

| (iv) | $K_c = 0.025^2 / (0.175 \times 0.05^2) = 1.4(3) \text{ (mol}^{-1} \text{ dm}^3)$ | [1] [max 9] |
|-------|---|----------------|
| (iii) | $K_c = {[acetal A][H_2O]}/{[CH_3CHO][CH_3OH]^2}$ units = mol ⁻¹ dm ³ | [1] [1] |
| (ii) | 4 values in third row | 4 x [1] |
| (i) | 3 values in second row | 3 x [1] |

[Total: 15]

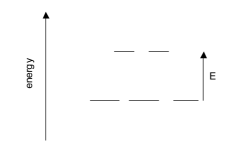
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3 (a) for example.... also allow d_{z2}



shape (4 lobes) [1] correct label e.g. d_{xy} [1] [2]

(b) (i)



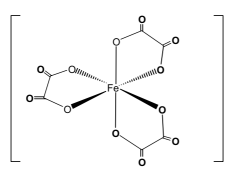
Marks are for 5 degenerate orbitals [1] and 3:2 split [1]

| (ii) | colour due to the absorption of light NOT emitted light E = hf <i>or</i> photon's energy = E in above diagram electron promoted from lower to higher orbital | [1] [1] [1] |
|------|--|--------------------------|
| | size of ΔE depends on the ligand as ΔE changes, so does f in E = hf | [1] [1] [7] |

(c) (i)
$$O.N.(carbon) = +3$$
 (4 × (-2) + 2x = -2, thus 2x = +6) [1]

3-

(iii)



[2]

(iv) $\underline{2} K_3 \operatorname{Fe}(C_2O_4)_3 \rightarrow \underline{3} K_2C_2O_4 + \underline{2} \operatorname{Fe}C_2O_4 + \underline{2} \operatorname{CO}_2$ $Or K_3 \operatorname{Fe}(C_2O_4)_3 \rightarrow \underline{3/2} K_2C_2O_4 + \operatorname{Fe}C_2O_4 + \operatorname{CO}_2$ [2]

[max 5]

[Total: 14]

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| | | | | GCE A LI | 9701 | 41 | | | |
| 4 | (a) (| | | $C_2H_5NH_2$ + HA → $C_2H_5NH_3^+$ + A ⁻ (HA can be H ₂ O, HC <i>l</i> etc.) Allow \rightleftharpoons instead of arrow | | | | | |
| | (i | ii) _ | | | | Γ | _ | | |
| | | | n | nost basic | | least basic | | | |
| | | | e | thylamine | ammonia | phenylamine | | | |
| | | - | | | | | | [1] | |
| | (ii | | ethylamine > NH ₃ due to electron-donating ethyl/alkyl group phenylamine < NH ₃ due to delocalisation of lone pair over ring | | | | | [1] [1] [4] | |
| | (b) (| (i) | C ₆ H ₅ | $_{5}\text{OH} + \text{OH}^{-} \rightarrow$ | $C_6H_5O^- + H_2O$ (or | with Na ⁺ /H₂O/A ⁻) | | [1] | |
| | (1 | • | pKa of nitrophenol is smaller/K_a is larger because it's a stronger acid/dissociates more than phenol stronger because the anionic charge is spread out moreover the NO₂ group <i>or</i> NO₂ is electron-withdrawing | | | | | [1] | |
| | (ii | ii) | рКа = 1.0 | | | | | | |
| | (i | v) | Nitro | group increas | es acidity / electro | n-withdrawing groups | increase acidit | y [1] [5] | |
| | | | | | | | | | |

(c) (i) **B** is phenyldiazonium cation, $C_6H_5-N^+\equiv N$

| (| i | i | ۱ | |
|---|---|---|---|---|
| l | l | • |) | F |

| ' - | | | | | |
|-----|----------|--|------------------------------|--|--|
| | reaction | reagent(s) | conditions | | |
| | Step 1 | NaNO ₂ + HC <i>1</i> or HNO ₂ [1] | T < 10°C [1] | | |
| | Step 2 | H₂O / aq | heat/boil/T > 10° (both) [1] | | |
| | Step 3 | HNO₃ NB HNO₃(aq) OK for both | dilute (both) [1] | | |
| | | | ٢٨) | | |

[4] **[5]**

[1]

[Total: 14]

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| | | • | • |

- 5 (a) (i) C=C double bonds / alkenes
 - (ii) -OH groups / accept alcohols or acids
 - (iii) CH₃CO– or CH₃CH(OH)– groups
 - (iv) carbonyl, >C=O, groups / accept aldehydes and ketones $4 \times [1]$

[4]



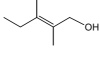


[2]

2 × [1]

(c) isomers of C





trans

| correct structure (excl. stereochemistry) | [1] |
|---|-----|
| cis and trans drawn correctly | [1] |
| type of isomerism is cis-trans or geometrical isomerism | [1] |
| | [3] |

[Total: 9]

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| 6 | (a) (i |) 2H ₂ | $NCH_2CO_2H \rightarrow H_2NCH_2CONHCH_2CO_2H + H_2O$ | | [1] |
| | (ii) |) Ske | eletal formula required | | [1] [2] |
| | (b) (i | | elix eated sheet | | [1] [1] |
| | (ii) | , For Nee with | dents should choose one of the structures belowα helix:For β pleated sheeed to show a helixNeed to show twon C=O H-Nstrands with C=O -ween turnsthem | parallel 'zig-zag' | |
| | | Wh | ichever is chosen, overall structure [1] position of H bor | nds [1] | |

[4]

(c)

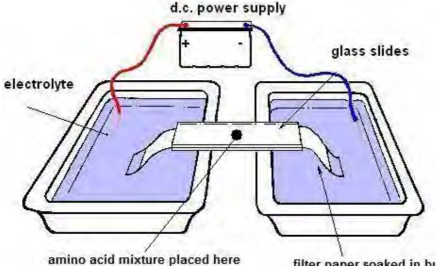
| <i>د</i> ا | | | | |
|------------|--|--|----------------------------------|--|
| | amino acid residue 1 | amino acid residue 2 | type of bonding | |
| | -HNCH(CH ₂ CH ₂ CH ₂ CH ₂ NH ₂)CO- | HNCH(CH ₂ CH ₂ CO ₂ H)CO– | lonic bonds or hydrogen bonds | |
| | -HNCH(CH ₃)CO- | –HNCH(CH ₃)CO– | van der Waals' | |
| | -HNCH(CH ₂ SH)CO- | -HNCH(CH ₂ SH)CO- | Disulfide bonds | |
| | -HNCH(CH ₂ OH)CO- | -HNCH(CH ₂ CO ₂ H)CO- | Hydrogen bonds | |

[4]

[Total: 10]

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7 (a) Sketch and label the apparatus used to carry out electrophoresis e.g



laced here filter paper soaked in buffer solution

- Marks: power supply / electrolyte + filter paper / buffer / acid mixture central 4 × [1]
 [4]
- (b) (i) pH of the buffer [1] [1] Charge on the amino acid species (ii) Size of the amino acid species / M_r [1] Voltage applied [1] Magnitude of the charge (on the amino acid species) [1] Temperature [1] (max 3) [max 3] (c) (i) They have insufficient electron density / only one electron [1] (ii) Sulfur [1]
 - because it has the greatest atomic number / number of electrons [1]
 [3]

[Total: 10]

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8 (a)

| traditional material | modern polymer used | |
|---|----------------------|--|
| Paper/cardboard/wood/leaves hessian/hemp/jute steel/aluminium | PVC in packaging | |
| Cotton/wool/linen | Terylene in fabrics | |
| Glass/china/porcelain/earthenware metal/leather | Polycarbonate bottle | |

 $3 \rightarrow 2$ marks, $2 \rightarrow 1$ mark [2]

| (b) | Rea | asons: Plastics/polymers pollute the environment for a long time do not decor biodegrade quickly They are mainly produced from oil Produce toxic gases on burning | npose/ [1] [1] [1] max two |
|-----|-------------------|--|--|
| | | ategy 1: Recycle polymer waste / use renewable resources ategy 2: Develop biodegradable polymers | [1] [1] [max 3] |
| (c) | or nyle | C mbustion would produce HC <i>l</i> / dioxins as a pollutant on/acrylic mbustion would produce HCN | [1] [1] [1] [2] |
| (d) | (i) | Polythene (or other addition polymer) | [1] |
| | (ii) | Addition polymerisation | [1] |
| | | The polymer chains don't have strong bonds between them – easy to melt Could be answered with a suitable diagram | [1] [3] |
| | | | [Total: 10] |