Cambridge International Advanced Level

MARK SCHEME for the October/November 2014 series

9701 CHEMISTRY

9701/43

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

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| Question | Marking point | | | Marks | Marks total |
|-----------|---|------------------------------------|--|--|----------------|
| 1 (a) (i) | | m/e | identity | | |
| | - | 35 | ³⁵ C1 | | |
| | - | 37 | ³⁷ C <i>l</i> | | |
| | - | 70 | ³⁵ Cl ³⁵ Cl or ³⁵ Cl ₂ | | |
| | - | 72 | ³⁷ Cl ³⁵ Cl | | |
| | - | 74 | ³⁷ Cl ³⁷ Cl or ³⁷ Cl ₂ | | |
| | 35, 37, 70, 72, 74 correct formulae at least one structu | ire as a posi | tive ion | 1 1 1 | |
| (ii) | 9:6:1 | | | 1 | [4] |
| (b) (i) | correct charges correct electrons | | _ | 1 | |
| (ii) | Lattice energy = $\Delta H_{\rm f}({\rm SrC} l_2) - (\Delta l_2) - (-164 + 548 + 106)$ = -2146 (kJ mol ⁻¹) | | | $\int_{\text{om}}(Cl) + 2\Delta H_{\text{ea}}(Cl)) \qquad 1 \\ 1 \\ 1 \end{cases}$ | [5] |
| (c) (i) | $SrCO_3 + 2HNO_3 \rightarrow Sr(NO_3)_2 -$ | + CO ₂ + H ₂ | 0 | 1 | |

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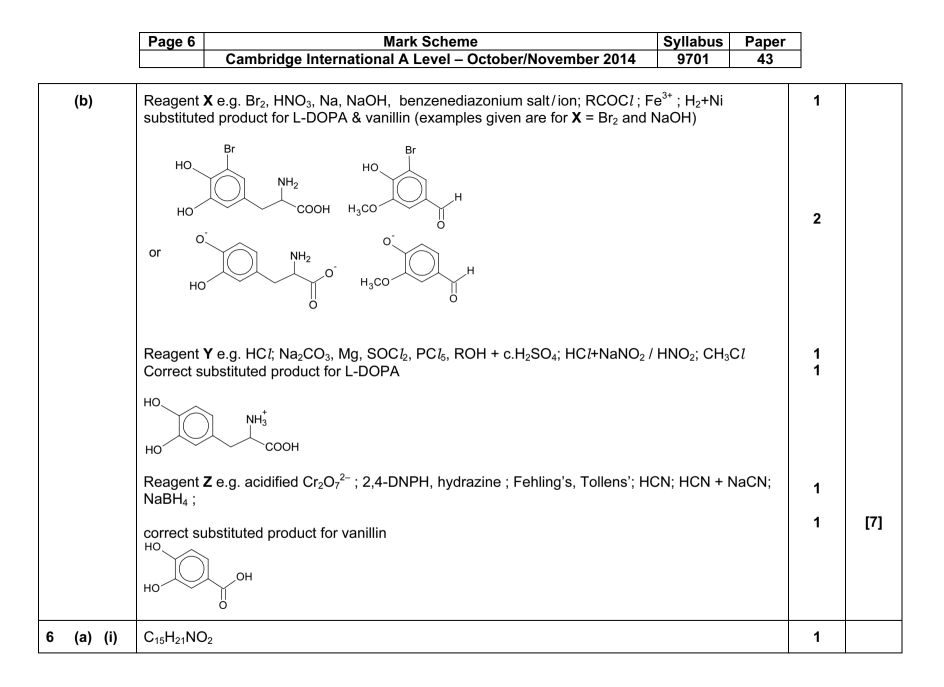
| | (ii) | $Sr(NO_3)_2 \rightarrow SrO + 2NO_2 + 0.5 O_2$ | 1 | [2] |
|------|--------|---|-------------|-----|
| (c | d) | (down the group) nitrates become more stable / require a higher temperature to decompose as size/radius of ion increases OR charge density of ion decreases | 1 1 1 | [3] |
| | | so polarisation/distortion of anion/nitrate ion/NO ₃ ⁻ /NO bond decreases | | |
| 2 (a | a) | $BrO_3^- + 5Br^- + 6H^+ \rightarrow 3Br_2 + 3H_2O$ five correct species correct balancing | 1 1 | [2] |
| (k | o) (i) | $[BrO_3^-]$ 1 st order and the concentration is x2, rate doubles OR evidence using expt 1 & 4 eg ratios $[H^+]$ 2 nd order and the concentration is x2, rate x4 OR evidence using expt 1 & 2 [Br] 1 st order and the concentration is x4, rate x4 OR evidence using expt 1 & 3 eg ratios | 1 1 1 | |
| | (ii) | (Rate =) $k [BrO_3^{-}][Br^{-}][H^{+}]^2$ | 1 | |
| | (iii) | k = 1.32 mol ⁻³ dm ⁹ s ⁻¹ | 1 1 | [6] |
| 3 (a | a) (i) | chromium and copper | 1 | |
| | (ii) | (all orbitals have the) same energy | 1 | |
| | (iii) | correct id of one higher energy d orbital the other higher energy d orbital | 1 1 | [4] |

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| (b) (i) | pale blue precipitate A solution B solution C | Cu(OH) ₂ OR [Cu(OH) ₂ (H ₂ O) ₄] [Cu(NH ₃) ₄ (H ₂ O) ₂] ²⁺ OR [Cu(NH ₃) ₄] ²⁺ [CuC l_4] ²⁻ | 1 | |
|---------|---|---|---|-----|
| (ii) | solution B solution C | royal/deep/dark blue OR violet-blue yellow/green | 1 | |
| (iii) | redox OR oxidation of AND reducing agent/redu | | 1 | [6] |
| (c) | | cant d-orbital/d-orbital s full tween orbitals OR transitions cannot occur | 1 | [2] |
| (d) | green/yellow orange/red AND blue/vio | let light is <u>absorbed</u> | 1 | [2] |
| 4 (a) | (HC <i>l</i>) strong er acid/more (HC <i>l</i> has) more ions/high | dissociated/ionised in solution er concentration of ions | 1 | [2] |
| (b) (i) | | nges in the pH/keeps pH <i>fairly</i> constant nounts/vols of acid/H⁺ or base/OH⁻ are added | 1 | |
| (ii) | add (ethanoic acid) to NaC excess (ethanoic acid) OR mix with sodium ethan | | 1 | [4] |
| (c) | $CH_{3}CH(NH_{2})COOH + H^{+} = CH_{3}CH(NH_{2})COOH + OH^{-}$ | → $CH_3CH(NH_3^+)COOH$ → $CH_3CH(NH_2)COO^- + H_2O$ | 1 | [2] |

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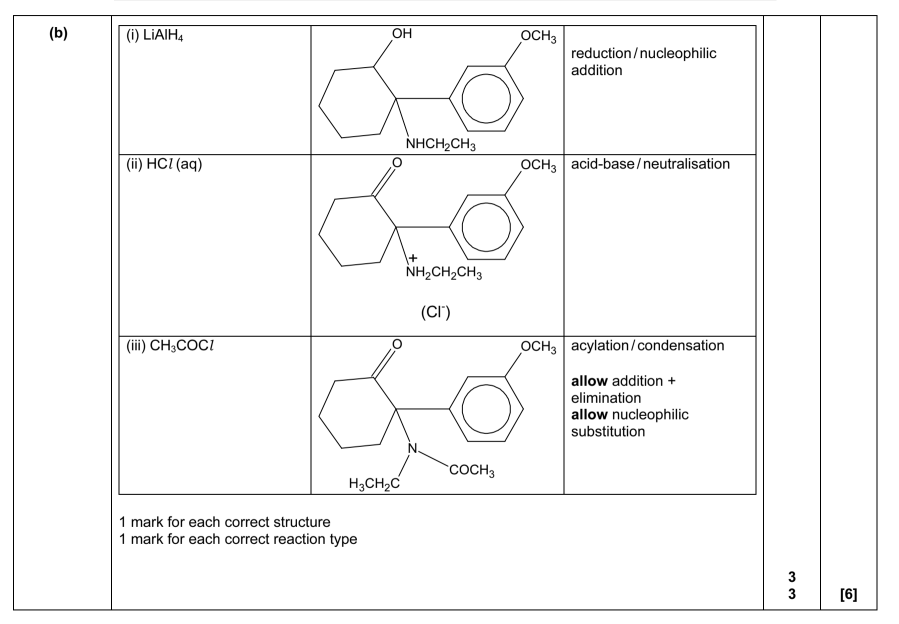
| (d) (i) | pKa 2.99 HO HO OH OH OH OH OH OH | 1 | |
|---------|---|---|-----|
| | $pKa 4.40 \qquad HO \qquad \longrightarrow \qquad OH \qquad OH \qquad OH \qquad OH \qquad OH \qquad OH $ | 1 | |
| (ii) | $\begin{array}{cccc} HO \\ HOOC \\ H$ | 2 | [4] |
| 5 (a) | any five of these seven points. σ-bonds are between C-C OR C-H carbons are sp² rings of charge above and below the ring must be in diagram presence of σ-bonds electrons/bonds are delocalised planar molecule/bond angles 120° all C-C are the same length/have intermediate bond length between C-C & C=C | 5 | [5] |



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| (ii) | O OCH ₃ | 1 | |
|-------|--|---|-----|
| | * NHCH ₂ CH ₃ | | |
| (iii) | any two of ketone, amine or ether | 2 | [4] |

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| 7 | (a) | (ratio of) the concentrations/distribution/amount/mass of solute in two (immiscible) solvents at equilibrium OR equilibrium constant OR includes expression with K | 1 1 | [2] |
|---|---------|---|--------|-----|
| | (b) | $ \begin{array}{l} \mathcal{K}_{\text{pc}} &= [J \text{ in ether}]/[J \text{ in } H_2 O] \\ &= (2.14/20)/(5-2.14/75) \\ &= 2.81 \text{ OR } 2.82 \end{array} $ | | [2] |
| | (c) | 1^{st} extraction: $2.81 = (x/10)/(5.0-x)/75$ $2.81(5-x) = 7.5x$ $x = 1.36 g$ 2^{nd} extraction: $2.81 = (y/10)/(3.64-y)/75$ $2.81(3.64-y) = 7.5y$ $y = 0.99 g$ | 1 | [2] |
| | (d) (i) | water/solvent/named solvent | | |
| | (ii) | non-volatile liquid, for example mineral oil or at least a C_{15} hydrocarbon oil | 1 | |
| | (iii) | 1. R_f (retardation factor) or distance travelled by solute and distance by solvent 2. retention time | | [4] |

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|---|---------|---|---|-------|-------|--------|-----|
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| | (e) | | CO ₂ H ² | | | 1 | [1] |
| | | | CH ₂ OH 1 | | | | |
| | | | CO ₂ H 3 CO ₂ H | | | | |
| | | | | | | | |
| 8 | (a) | C = 33 % A = T = 17 % | | | | 1 1 | [2] |
| | (b) (i) | only one isomer may be active/be of therapeutic benefit | | | | 1 | |
| | (ii) |) the other (stereo) isomer may cause harm/side effects | | | | 1 | [2] |

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| | (c) (i) | structures of the following aldehydes: | | | |
| | | $\begin{array}{c} & & \\ & & \\ & & \\ & \\ two \ correct \ structures = 1 \ mark \\ two \ further \ correct \ structures - 1 \ mark \end{array}$ | | | |
| | (ii) | 3-methylbutanal | | | |
| | (iii) | pentanal5 absorptions2-methylbutanal5 absorptionsdimethylpropanal2 absorptions | 1 1 1 | [6] | |
| 9 | (a) | nylon, terylene – condensation; PVC – addition – all three correct | 1 | [1] | |
| | (b) | correct fully displayed formula of -CO-NH- unit correct polymer structure H H H H H H H H H H | 1 1 | [2] | |
| | (c) | sequence/order of amino acids (in the polypeptide chain) | | | |
| | (d) | hydrogen bond C=O and N-H in two different amino acids in the backbone diagram | | | |

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| (e) (i) | OR –NH3 | ydrogen/ionic bonds as $-COOH/NH_3^+$ is deprotonated $A_3^+ + OH^- \rightarrow NH_2 + H_2O$ linked to hydrogen/ionic bond disrupted $H + OH^- \rightarrow -COO^- + H_2O$ linked to hydrogen/ionic bond disrupted | | | 1 | |
| (ii) | Ha ²⁺ intor | force with/broaks the disulfide hand/bridge not sulfite sulfate, sulfur | sulfido | | | |

| | Hg ²⁺ interferes with/breaks the disulfide bond/bridge not sulfite, sulfate, sulfur, sulfide OR -S-S- shown with Hg ²⁺ in an equation OR disrupting ionic interactions linked to carboxyl/COO– groups | 1 | |
|-------|--|---|-----|
| (iii) | (Heat to 70 °C) breaks the van der Waals' forces/hydrogen bonding | 1 | [3] |