## Cambridge International Examinations

Cambridge International Advanced Subsidiary and Advanced Level
AS \& A Level

## CHEMISTRY

Paper 4 A Level Structured Questions
MARK SCHEME
Maximum Mark: 100

## Published

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| Question | Answer | Marks |
| :---: | :---: | :---: |
| 1 (a) (i) | dative (covalent) or coordinate Hydrogen/H (boding) | 2 |
| (ii) | octahedral | 1 |
| (iii) | $\begin{aligned} & \mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2} \cdot 6 \mathrm{H}_{2} \mathrm{O} \rightarrow \mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2}+6 \mathrm{H}_{2} \mathrm{O} \\ & \mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2} \rightarrow \mathrm{MgO}+2 \mathrm{NO}_{2}+\frac{1}{2} \mathrm{O}_{2} \\ & \text { any three of } \\ & \text { (solid) dissolves/turns to liquid } \\ & \text { condensation on tube } \\ & \text { white solid (forms/remains) } \\ & \text { brown fumes (evolved) } \\ & \text { gas formed that relights a glowing splint } \end{aligned}$ | 4 |
| (iv) | $M_{\mathrm{r}}$ values: $\mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2} .6 \mathrm{H}_{2} \mathrm{O}=256.3 \mathrm{MgO}=40.3$ or (loss in molar mass $=256.3-40.3=$ ) 216 percentage loss $=100 \times 216 / 256.3=84.3 / 84.4 \%$ | 2 |
| (b) | (cat)-ionic radius/ion size increases (down the group) less polarisation/distortion of nitrate ion/ $\mathrm{NO}_{3}{ }^{-}$ | 2 |
| (c) | $2 \mathrm{AgNO}_{3} \rightarrow 2 \mathrm{Ag}+2 \mathrm{NO}_{2}+\mathrm{O}_{2}$ | 1 |
|  |  | [Total: 12] |
| 2 (a) (i) | (an acid that is) partially/incompletely ionised/dissociated | 1 |
| (b) (i) | $\mathrm{p} K_{\mathrm{a}}=-\log K_{\mathrm{a}}$ or $K_{\mathrm{a}}=10^{-\mathrm{p} K a}$ | 1 |


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| Question | Answer | Marks |
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| (ii) | ethanoic acid (1) is more acidic than propanoic acid (2) due to smaller electron-donating ( $\mathrm{R} /$ /alkyl) group/less electron-donating (R/alkyl) group(s) <br> 2-chloropropanoic acid (3) is more acidic than propanoic acid (2) due to electron-withdrawing/electronegative (Cl/chlorine) atom <br> 2-chloropropanoic acid (3) is more acidic than 3-chloropropanoic acid (4) since the $\mathrm{C} /$ /chlorine/electronegative atom is closer to the $\mathrm{CO}_{2}^{-} /$acid | 3 |
| (c) (i) | M1: voltmeter/V and salt bridge labelled <br> M2: Cu and $\mathrm{Cu}^{2+} / \mathrm{CuSO}_{4}$ (any soluble $\mathrm{Cu}(\mathrm{II})$ salt) <br> M3: $\mathrm{H}_{2}$ (arrow in) and $\mathrm{H}^{+} / \mathrm{HCl} / \mathrm{H}_{2} \mathrm{SO}_{4} /$ any mineral acid <br> M4 Pt and one solution at $1 \mathrm{M} / 1 \mathrm{moldm}^{-3} \mathrm{OR}_{2}$ at 1 atm | 4 |
| (ii) | $E_{\text {cell }}^{9}=0.34(\mathrm{~V})$ and $\left(\mathrm{Cu}^{2+}\right) / \mathrm{Cu}$ is the positive electrode | 1 |
| d (i) | $\begin{aligned} & K_{\mathrm{a}}=1.23 \times 10^{-5} \\ & {\left[\mathrm{H}^{+}\right]=\sqrt{ }\left(\mathrm{K}_{\mathrm{a}} \cdot \mathrm{C}\right)=\sqrt{ }\left(1.23 \times 10^{-5} \times 0.1\right)=1.11 \times 10^{-3} \mathrm{~mol} \mathrm{dm}^{-3}} \\ & \mathrm{pH}=3.0(2.96) \text { ecf from }\left[\mathrm{H}^{+}\right] \end{aligned}$ | 2 |


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| Question | Answer | Marks |
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| (ii) | $\begin{aligned} & E=0.0+0.059 \log \left(1.11 \times 10^{-3}\right) \mathrm{OR}=-0.17(4) \mathrm{V} \\ & \text { so new } E_{\text {cell }}=0.34+0.17=\mathbf{0 . 5 1 V} \\ & \text { ecf from (d)(i) } \end{aligned}$ | 2 |
|  |  | [Total: 14] |
| 3 (a) (i) | $\left(\mathrm{CH}_{3}\right)_{2} \mathrm{CHCN}$ | 1 |
| (ii) | reaction 1: $\mathrm{NH}_{3}$ (in ethanol) under pressure (+ heat) or heat $\mathrm{NH}_{3}$ in a sealed tube <br> reaction 2: $\mathrm{KCN} / \mathrm{NaCN}$ and heat/reflux (in ethanol) <br> reaction 3: $\mathrm{H}_{2}+\mathrm{Ni}$ or $\mathrm{LiAlH}_{4}$ | 3 |
| (b) (i) | $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{NH}_{2}+\mathrm{H}_{2} \mathrm{O} \rightarrow \mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{NH}_{3}^{+}(+) \mathrm{OH}^{-}$ | 1 |
| (ii) | ethylamine is more basic than ammonia... because of electron-donating (alkyl/ethyl/R) group (in ethylamine) which makes the lone pair (on N ) more available for donation or the lone pair (on N ) more available for a proton $/ \mathrm{H}^{+}$ | 2 |
| (c) (i) | A solution which resists/minimises/roughly maintains changes in pH when (small amounts of) $\mathrm{H}^{+}$or $\mathrm{OH}^{-}$are added | 1 |
| (ii) | $\begin{aligned} & \mathrm{CH}_{3} \mathrm{NH}_{2}+\mathrm{H}^{+} \rightarrow \mathrm{CH}_{3} \mathrm{NH}_{3}^{+} \\ & \mathrm{CH}_{3} \mathrm{NH}_{3} \mathrm{Cl}+\mathrm{OH} \rightarrow \mathrm{CH}_{3} \mathrm{NH}_{2}+\mathrm{H}_{2} \mathrm{O}+\mathrm{Cl} \end{aligned}$ | 2 |
|  |  | [Total: 10] |


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| Question | Answer | Marks |
| :---: | :---: | :---: |
| 4 (a) (i) |  <br> (cis) | 2 |
| (ii) | cis is (more) polar due to both $\mathrm{Cl}^{(\delta-)}$ on same side or cis is (more) polar as dipoles do not cancel/unsymmetrical or trans is non-polar as it is bond dipoles cancel | 1 |
| (iii) | (This can only be cis) its mirror image is the same/superimposable <br> or the distance between two coordinating nitrogens/oxygens is too small to bond trans or difficult for the $\mathrm{NH}_{2}$ and O to change places (since 5 -memebered rings can only bridge adjacent positions) | 1 |
| (b) (i) | It's not square planar or it's tetrahedral | 1 |
| (ii) | must be 3D structure (i.e. tetrahedral-like) <br> or <br> etc | 1 |
|  |  | [Total: 6] |


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| Question | Answer | Marks |
| :---: | :---: | :---: |
| 5 (a) (i) | $\begin{aligned} & K_{\text {stab }}=\frac{\left[\mathrm{Cd}\left(\mathrm{CH}_{3} \mathrm{NH}_{2}\right) 4^{2+}\right.}{\left[\mathrm{Cd}^{2+}\right]}\left[\mathrm{CH}_{3} \mathrm{NH}_{2}\right]^{4} \\ & \text { units: } \mathrm{mol}^{-4} \mathrm{dm}^{12} \end{aligned}$ | 2 |
| (ii) |  | 2 |
| (b) (i) | (each complex is formed by) making ( $4 \times$ )N-Cd bonds and breaking ( $6 \times$ ) O-Cd bonds or same types of/similar bonds forming/breaking or same number of bonds forming/breaking | 1 |
| (ii) | $\Delta S=(\Delta H-\Delta G) / T=(60.7-56.5) \times 1000 / 298=(+) \mathbf{1 4 / ( + ) 1 4 . 1}$ | 1 |
| (iii) | fewer moles (of solutes) are forming (one mole of) the complex (so less loss of disorder) or one en displaces two $\mathrm{H}_{2} \mathrm{O}$ whereas one $\mathrm{CH}_{3} \mathrm{NH}_{2}$ only displaces one $\mathrm{H}_{2} \mathrm{O}$ | 1 |
| (iv) | The $\left[\mathrm{Cd}\left(\mathrm{H}_{2} \mathrm{NCH}_{2} \mathrm{CH}_{2} \mathrm{NH}_{2}\right)_{2}\right]^{2+}$ /equilibrium 2 complex (is more stable) because: either $K_{\text {stab }}$ is greater or $\Delta G^{\ominus}$ is more negative. | 1 |
|  |  | [Total: 8] |


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| Question | Answer | Marks |
| :---: | :---: | :---: |
| 6 (a) | essential mark <br> M1 the reactants/substrate has a shape complementary/specific to active site - can be awarded from a labelled diagram as below or diagrams showing this specificity clearly <br> any two of <br> M2: reactants/substrate binds to/fits into the active site of the enzyme <br> M3: (Interaction with site) causes a specific bond to be weakened, (which breaks) <br> or lowers activation energy <br> M4: forms an E-S complex <br> M5: products released from enzyme/active site <br> labelled diagrams <br> (products) | 3 |
| (b) (i) | $\delta 26$ is CH3-CO $\delta 52$ is CH3-O <br> $\delta 169$ is $\mathbf{C H} 3 \mathbf{C O}$ $\delta 167$ is phenyl-CO <br> Phenyl ethanoate is $\mathbf{B} \quad$ methyl benzoate is $\mathbf{A}$ <br> M1 = any two correct $\delta$ linked to phenylethanoate/methyl benzoate <br> M2 $=$ the rest correct | 2 |
| (ii) | heat with $\mathrm{H}_{3} \mathrm{O}^{+}$(to hydrolyse the ester) <br> then add $\mathrm{Br}_{2}(\mathrm{aq}) /$ bromine water <br> decolourises/gives white ppt. (with phenol from B) | 3 |
|  |  | [Total: 8] |


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| Question | Answer | Marks |
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| $7 \quad$ (a) (i) | labelled with <br> M1: DC power supply + and -/battery / cell/+ and - sign (on cell/electrodes) with a complete circuit <br> M2: buffer solution/electrolyte labelled <br> M3: (amino acid) mixture/ $\mathbf{x}$ on (filter) paper/gel/agarose | 3 |
| (ii) | direction of movement related to charge (of amino acids) distance travelled depends on charge $/ M_{\mathrm{r}}$ (of amino acids) | 2 |
| (b) (i) | Asp + Val: <br> pH 12 because Asp will be $-\mathrm{CH}_{2} \mathrm{COO}^{-}$(R-group) moves further (to positive electrode than Val) or pH 12 Asp more negative so moves further (to positive electrode) or pH 12 because Asp has a charge of 2 - but Val has a charge of $1-$ or best at pH 7 because Asp will be negatively charged (anionic) but Val neutral | 1 |
| (ii) | Lys + Ser: <br> pH 2 because Lys will be $\left(\mathrm{CH}_{2}\right)_{4} \mathrm{NH}_{3}{ }^{+}$(R-group) moves further (to negative electrode than Ser) or pH 2 Lys more positive so moves further (to negative electrode) or pH 2 because Lys has a charge of 2+ and Ser has a charge of 1+ or pH 7 because Lys is positively charged (cationic) but Ser neutral/zwitterionic | 1 |


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| Question | Answer | Marks |
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| (iii) | Tyr + Phe: <br> pH 12 because Tyr will be $\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{CH}_{2} \mathrm{O}^{-}$(R-group) moves further/more/faster (to positive electrode than Phe) or pH 12 because Tyr has a charge of 2 - but Phe has a charge of 1 - | 1 |
| (c) (i) |  <br> M1: for - $\mathrm{CONH}-$ as shown above <br> M2: for rest of molecule and correct connectivity of the bonds | 2 |
| (ii) | from the IR spectrum <br> - E is O-H or N-H (allow $\mathrm{NH}_{2}$ ) <br> - F is $\mathrm{C}=\mathrm{O}$ <br> - $\mathbf{G}$ is $\mathrm{C}-\mathrm{O}$ | 2 |
|  |  | [Total: 12] |
| 8 (a) | M1: solubility increases (down the group) <br> M2: because lattice energy decreases faster than does $\Delta \boldsymbol{H}_{\text {hyd }}$ <br> M3: $\Delta H_{\text {sol }} /$ enthalpy of solution becomes more exothermic/less endothermic | 3 |
| (b) (i) | Should be the same/similar (enthalpy change), as (both acids) are fully ionised/strong acids | 1 |


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| Question | Answer | Marks |
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| (ii) | $\begin{aligned} & \mathrm{Ca}(\mathrm{~s})+2 \mathrm{H}^{+}(\mathrm{aq}) \longrightarrow \mathrm{Ca}^{2+}(\mathrm{aq})+\mathrm{H}_{2}(\mathrm{~g}) \\ & \boldsymbol{x}=\Delta \mathrm{H}_{\mathrm{at}}(\mathrm{Ca})+\mathrm{IE}(1)+\mathrm{IE}(2)-2 \Delta \mathrm{H}_{\mathrm{hyd}}\left(\mathrm{H}^{+}\right)+\Delta \mathrm{H}_{\text {hyd }}\left(\mathrm{Ca}^{2+}\right)-2 \mathrm{IE}(\mathrm{H})-\mathrm{E}(\mathrm{H}-\mathrm{H}) \\ & \boldsymbol{x}=178+590+1150+2(1090)-1576-2(1310)-436 \\ & \boldsymbol{x}=-534 \mathrm{~kJ} \mathrm{~mol}^{-1} \end{aligned}$ | 4 |
| (c) | $\mathrm{CH}_{3} \mathrm{CO}_{2} \mathrm{H}$ is incompletely ionised/weak acid/weaker acid enthalpy change of ionisation (of $\mathrm{CH}_{3} \mathrm{COOH}$ ) is $+2 \mathrm{~kJ} \mathrm{~mol}^{-1}$ <br> or energy needed to ionise/dissociate $\left(\mathrm{CH}_{3} \mathrm{COOH}\right)$ | 2 |
|  |  | [Total: 10] |
| $9 \quad$ (a) |  | 1 |


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| Question | Answer | Marks |
| :---: | :---: | :---: |
| (b) |  <br> H is <br> $J$ is | 2 |
| (c) | ```step 1: \(\left(\mathrm{CH}_{3}\right)_{2} \mathrm{CHCH}_{2} \mathrm{Cl}+\mathrm{AlCl}_{3}\) (+ heat) step 2: \(\mathrm{CH}_{3} \mathrm{COCl}+\mathrm{AlCl}_{3}\) (+ heat) step 3: \(\mathrm{HCN}+\mathrm{NaCN}\) or \(\mathrm{HCN}+\) base or \(\mathrm{HCN}+\mathrm{CN}^{-}\) (steps 4 and 5 could be reversed on J) If J1 step 4 then step 5 J2 step 5 then step 4 step 4: \(\mathrm{H}_{3} \mathrm{O}^{+}+\)heat/aqueous \(\mathrm{HCl}+\) heat step 5: conc \(\mathrm{H}_{2} \mathrm{SO}_{4}+\) heat/ conc \(\mathrm{H}_{3} \mathrm{PO}_{4}+\) heat or \(\mathrm{Al}_{2} \mathrm{O}_{3}+\) heat step 6: \(\mathrm{H}_{2}+\mathrm{Ni}\) (+ heat)``` | 6 |
| (d) | step 1: electrophilic substitution or alkylation <br> step 6: reduction/hydrogenation/addition | 2 |
|  |  | [Total: 11] |


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| Question | Answer | Marks |
| :---: | :---: | :---: |
| 10 (a) (i) | Fe is $\ldots .3 s^{2} 3 p^{6} 3 d^{6} 4 s^{2}$ | 1 |
| (ii) |  | 1 |
| (b) | $\begin{aligned} & E^{\ominus} \text { values: } \mathrm{Sn}^{4+} / \mathrm{Sn}^{2+}=+0.15(\mathrm{~V}) ; \mathrm{Fe}^{3+} / \mathrm{Fe}^{2+}=+0.77(\mathrm{~V}) \\ & \text { or } \\ & E_{\text {cell }}=+0.62(\mathrm{~V}) \\ & \left(\mathrm{Sn}^{2+} \text { will reduce } \mathrm{Fe}^{3+}\right) \mathrm{Sn}^{2+}+2 \mathrm{Fe}^{3+} \rightarrow 2 \mathrm{Fe}^{2+} \end{aligned}$ | 2 |
| (c) (i) | ```essential mark \(K_{\text {stab }} /\) stability: \(\left[\mathrm{Fe}\left(\mathrm{H}_{2} \mathrm{O}\right)_{5} \mathrm{~F}\right]^{2+}>\left[\mathrm{Fe}\left(\mathrm{H}_{2} \mathrm{O}\right)_{5} \mathrm{SCN}\right]^{+}\) \(\left(>\left[\mathrm{Fe}\left(\mathrm{H}_{2} \mathrm{O}\right)_{6}\right]^{2+}\right)\) observations (violet) \(\rightarrow\) deep-red (deep-red) \(\rightarrow\) colourless (violet) \(\rightarrow\) colourless which stays colourless/does not change``` | 4 |
| (ii) | ligand displacement/exchange/substitution | 1 |
|  |  | [Total: 9] |

