## Pearson Edexcel

Mark Scheme (Results)

October 2021

Pearson Edexcel International Advanced Level In Chemistry (WCH15)
Paper 01:Transition Metals and Organic Nitrogen Chemistry

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- All candidates must receive the same treatment. Examiners must mark the first candidate in exactly the same way as they mark the last.
- Mark schemes should be applied positively. Candidates must be rewarded for what they have shown they can do rather than penalised for omissions.
- Examiners should mark according to the mark scheme not according to their perception of where the grade boundaries may lie.
- There is no ceiling on achievement. All marks on the mark scheme should be used appropriately.
- All the marks on the mark scheme are designed to be awarded. Examiners should always award full marks if deserved, i.e. if the answer matches the mark scheme. Examiners should also be prepared to award zero marks if the candidate's response is not worthy of credit according to the mark scheme.
- Where some judgement is required, mark schemes will provide the principles by which marks will be awarded and exemplification may be limited.
- When examiners are in doubt regarding the application of the mark scheme to a candidate's response, the team leader must be consulted.
- Crossed out work should be marked UNLESS the candidate has replaced it with an alternative response.


## Section A (Multiple Choice)

$\left.\begin{array}{|l|l|c|}\hline \begin{array}{l}\text { Question } \\ \text { number }\end{array} & \text { Answer } & \text { Mark } \\ \hline \mathbf{1 ( a )} & \text { The only correct answer is D (Pt, Pt) } & \mathbf{1} \\ & \boldsymbol{A} \quad \text { is incorrect because both electrodes should be made of platinum } \\ & \boldsymbol{B} \text { is incorrect because both electrodes should be made of platinum } \\ \text { C is incorrect because both electrodes should be made of platinum }\end{array}\right]$

| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 ( b )}$ | The only correct answer is $\mathbf{C}\left(1.00 \mathrm{~mol} \mathrm{dm}^{-3} \mathrm{HCl}(\mathrm{aq})\right)$ | $\mathbf{1}$ |
|  | A is incorrect because $\mathrm{H}_{3} \mathrm{PO}_{4}$ is not completely ionised |  |
|  | B is incorrect because $\mathrm{H}_{2} \mathrm{SO}_{4}$ is not completely ionised |  |
| D is incorrect because $\mathrm{CH}_{3} \mathrm{COOH}$ is not completely ionised |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 ( c )}$ | The only correct answer is B (17.91 g) | $\mathbf{1}$ |
|  | A is incorrect because there should be only one mol of chromium ions per mol of dichromate ions |  |
|  | C is incorrect because there should be only one mol of chromium ions per mol of dichromate ions |  |
| D is incorrect because there should be only one mol of chromium ions per mol of dichromate ions |  |  |$\quad$.


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 ( d )}$ | The only correct answer is A (H2SO4) | $\mathbf{1}$ |
|  | $\boldsymbol{B} \quad$ is incorrect because chloride ions could be oxidised |  |
| C is incorrect because bromide ions would be oxidised |  |  |
| $\boldsymbol{D} \quad$ is incorrect because this would introduce additional chromium species into the mixture |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{2}$ | The only correct answer is C $\left(\mathrm{Pt}^{2}\left\|\mathrm{Fe}^{2+}, \mathrm{Fe}^{3+} \\|\left[\mathrm{MnO}_{4}^{-}+8 \mathrm{H}^{+}\right],\left[\mathrm{Mn}^{2+}+4 \mathrm{H}_{2} \mathrm{O}\right]\right\| \mathrm{Pt}\right)$ | $\mathbf{1}$ |
|  | $\boldsymbol{A} \quad$ is incorrect because both electrodes should be made of platinum <br> $\boldsymbol{B} \quad$ is incorrect because both electrodes should be made of platinum and the $\mathrm{MnO}_{4} / / \mathrm{Mn}^{2+}$ half-cell does not show <br> reduction <br> $\boldsymbol{D} \quad$ is incorrect because the $\mathrm{MnO}_{4} / / \mathrm{Mn}^{2+}$ half-cell does not show reduction |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{3}$ | The only correct answer is $\mathbf{D}\left(2 \mathrm{Ag}^{2+} \rightarrow \mathrm{Ag}^{+}+\mathrm{Ag}^{3+}\right)$ | $\mathbf{1}$ |
|  | $\boldsymbol{A} \quad$ is incorrect because the disproportionation is not thermodynamically feasible |  |
|  | $\boldsymbol{B} \quad$ is incorrect because the disproportionation is not thermodynamically feasible |  |
| $\boldsymbol{C} \quad$ is incorrect because the disproportionation is not thermodynamically feasible |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{4}$ | The only correct answer is A (the cathode has a more positive potential than the anode) | $\mathbf{1}$ |
|  | $\boldsymbol{B} \quad$ is incorrect because oxidation always occurs at the anode |  |
| $\boldsymbol{C} \quad$ is incorrect because oxygen is reduced at the positive electrode |  |  |
| $\boldsymbol{D} \quad$ is incorrect because the overall reaction is the same under both acidic and alkaline conditions |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{5}$ | The only correct answer is B (carbon monoxide forms stronger dative covalent bonds with haemoglobin than does <br> oxygen) | $\mathbf{1}$ |
|  | A $\quad$ is incorrect because carbon monoxide can be displaced from carboxyhaemoglobin <br> C is incorrect because the formation of carboxyhaemoglobin does not lead to an increase in the entropy of the system <br> $\boldsymbol{D} \quad$ is incorrect because the difference in bond type does not fully explain the difference in dative covalent bond strength |  |

$\left.\begin{array}{|l|l|c|}\hline \begin{array}{l}\text { Question } \\ \text { number }\end{array} & \text { Answer } & \text { Mark } \\ \hline \mathbf{6} & \text { The only correct answer is } \mathbf{D} \text { (pink solution } \rightarrow \text { blue precipitate } \rightarrow \text { yellow-brown solution) } & \mathbf{1} \\ & \boldsymbol{A} \quad \text { is incorrect because } \mathrm{CoCl}_{2}(a q) \text { is a pink solution } \\ \boldsymbol{B} \text { is incorrect because the blue precipitate dissolves in excess aqueous ammonia to form a yellow-brown solution } \\ \boldsymbol{C} \quad \text { is incorrect because } \mathrm{CoCl}_{2}(a q) \text { is a pink solution }\end{array}\right]$

| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{7}$ | The only correct answer is B $\left([\mathrm{Ni}(\mathrm{EDTA})]^{2-}\right)$ | $\mathbf{1}$ |
|  | $\boldsymbol{A} \quad$ is incorrect because chloride ions act as monodentate ligands |  |
| $\boldsymbol{C} \quad$ is incorrect because ethanedioate ions act as bidentate ligands |  |  |
| D is incorrect because 1,2-diaminoethane molecules act as bidentate ligands |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{8}$ | The only correct answer is D (none of the products are harmful to the environment) | $\mathbf{1}$ |
|  | $\boldsymbol{A} \quad$ is incorrect because the reactions occurring in catalytic converters involve heterogeneous catalysis |  |
| $\boldsymbol{B} \quad$ is incorrect because carbon monoxide is adsorbed onto the surface of the catalyst |  |  |
| $\boldsymbol{C} \quad$ is incorrect because nitrogen is desorbed from the surface of the catalyst |  |  |$\quad$.


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{9}$ | The only correct answer is $\mathbf{C}\left(\mathrm{Mn}^{2+}\right)$ | $\mathbf{1}$ |
|  | A is incorrect because $\mathrm{MnO}_{4}^{-}$ions are neither a product nor a catalyst in this reaction |  |
| $\mathbf{B} \quad$ is incorrect because $\mathrm{H}^{+}$ions are neither a product nor a catalyst in this reaction |  |  |
| $\mathbf{D} \quad$ is incorrect because $\mathrm{CO}_{2}$ is not a catalyst in this reaction |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 0}$ | The only correct answer is $\mathbf{A}\left(\mathrm{both} \mathrm{Fe}^{2+}(\mathrm{aq})\right.$ and $\mathrm{Fe}^{3+}(\mathrm{aq})$ catalyse the reaction $)$ | $\mathbf{1}$ |
|  | $\mathbf{B} \quad$ is incorrect because both $\mathrm{Fe}^{2+}(a q)$ and $\mathrm{Fe}^{3+}(a q)$ catalyse the reaction |  |
|  | $\boldsymbol{C} \quad$ is incorrect because both $\mathrm{Fe}^{2+}(a q)$ and $\mathrm{Fe}^{3+}(a q)$ catalyse the reaction |  |
| $\mathbf{D} \quad$ is incorrect because both $\mathrm{Fe}^{2+}(a q)$ and $\mathrm{Fe}^{3+}(a q)$ catalyse the reaction |  |  |


| Question | Answer | Mark |
| :--- | :--- | :--- |
| number |  |  |
| $\mathbf{1 1}$ |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 2}$ | The only correct answer is C (330.7) | $\mathbf{1}$ |
|  | A is incorrect because this is the molar mass of bromobenzene |  |
| $\mathbf{B} \quad$ is incorrect because this is the molar mass of the monosubstituted product |  |  |
| $\mathbf{D} \quad$ is incorrect because this is the molar mass of the fully substituted product |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 3}$ | The only correct answer is $\mathbf{C}\left(\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{NH}_{2}>\mathrm{NH}_{3}>\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{NH}_{2}\right)$ | $\mathbf{1}$ |
|  | $\boldsymbol{A} \quad$ is incorrect because $\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{NH}_{2}$ is the weakest base in the sequence |  |
| $\boldsymbol{B} \quad$ is incorrect because $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{NH}_{2}$ is a stronger base than $\mathrm{NH}_{3}$ |  |  |
| $\boldsymbol{D} \quad$ is incorrect because this shows the order of increasing basicity |  |  |$\quad$.


| Question <br> number | Answer | Mark |  |
| :--- | :--- | :---: | :---: |
| $\mathbf{1 4}$ |  |  | $\mathbf{1}$ |
|  | The only correct answer is B ( $\mathrm{H}_{2} \mathrm{~N}$ |  |  |
|  | A incorrect because this amine could be prepared by the reduction of butanenitrile <br> C $\quad$ is incorrect because this amine could be prepared by the reduction of 2-methylpropanenitrile <br> $\boldsymbol{D} \quad$ is incorrect because this amine could be prepared by the reduction of 2,2-dimethylpropanenitrile |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 5}$ | The only correct answer is B (4) | $\mathbf{1}$ |
|  | A is incorrect because the repeat unit of the polymer is formed from four different amino acids |  |
|  | $\boldsymbol{C} \quad$ is incorrect because the repeat unit of the polymer is formed from four different amino acids |  |
| D is incorrect because the repeat unit of the polymer is formed from four different amino acids |  |  |$\quad$.


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 6}$ | The only correct answer is D (carbon dioxide giving carboxylic acids ) | $\mathbf{1}$ |
|  | A is incorrect because Grignard reagents react with water giving alkanes |  |
|  | $\mathbf{B} \quad$ is incorrect because Grignard reagents react with methanal giving primary alcohols |  |
| C is incorrect because Grignard reagents react with ketones giving tertiary alcohols only |  |  |


| Question <br> number | Answer | Mark |
| :--- | :--- | :---: |
| $\mathbf{1 7}$ | The only correct answer is C (will be lower than the true value) | $\mathbf{1}$ |
|  | A is incorrect because using a sample that is impure would cause the value to be lower |  |
| $\boldsymbol{B} \quad$ is incorrect because using a sample that is impure would cause the value to be lower |  |  |
| $\boldsymbol{D} \quad$ is incorrect because using a sample that is impure would cause the value to be lower |  |  |

## Section B

| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 18(a) | - any indication that $\mathbf{A}$ contains $\mathrm{FeCl}_{2} /$ iron(II) chloride <br> - working to show that $\mathbf{A}$ is a tetrahydrate | Example of calculation: <br> Ignore (A contains) $\mathrm{Fe}^{2+}$ <br> Ignore $\left[\mathrm{FeCl}_{4}\right]^{2-}$ $\begin{aligned} \text { mass of water } & =198.8-(55.8+2 \times 35.5) \\ & =72.0(\mathrm{~g}) \\ \text { moles of water } & =72.0 \div 18.0=4 \end{aligned}$ <br> A is $\mathrm{FeCl}_{2} .4 \mathrm{H}_{2} \mathrm{O} /$ iron(II) chloride tetrahydrate <br> Allow $\mathrm{FeCl}_{2}\left(\mathrm{H}_{2} \mathrm{O}\right)_{4}$ | 2 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| $\mathbf{1 8 ( b )}$ | $\bullet\left[\mathrm{Fe}\left(\mathrm{H}_{2} \mathrm{O}\right)_{6}\right]^{2+}$ | Allow $\left[\mathrm{Fe}(\mathrm{OH})\left(\mathrm{H}_{2} \mathrm{O}\right)_{5}\right]^{+} /\left[\mathrm{Fe}(\mathrm{Cl})\left(\mathrm{H}_{2} \mathrm{O}\right)_{5}\right]^{+}$ <br> Ignore omission of square brackets <br> Ignore name even if incorrect | $\mathbf{1}$ |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 18(c) | - A diagram showing the octahedral shape | Example of diagram: | 1 |
|  |  |  |  |
|  |  | Accept arrows for dative covalent bonds |  |
|  |  | Allow CN for $\mathrm{C} \equiv \mathrm{N}$ <br> Do not award KCN/HCN for $\mathrm{C}=\mathrm{N}$ <br> Do not award M for Fe |  |
|  |  | Ignore connectivity of CN ligands Ignore lone pairs Ignore omission of square brackets Ignore all charges |  |
|  |  | Do not award diagrams with no 3D shape |  |


| Question Number | Answer |  | Additional guidance | Mark |
| :---: | :---: | :---: | :---: | :---: |
| 18(d) | - moles of $\mathrm{K}, \mathrm{Fe}$ | (1) | Example of calculation: | 3 |
|  |  |  | $\mathrm{mol} \mathrm{K}=35.6 \div 39.1=0.91049$ |  |
|  |  |  | Allow 0.91282 from $A_{r}$ value of 39 |  |
|  |  |  | $\mathrm{mol} \mathrm{Fe}=17.0 \div 55.8=0.30466$ |  |
|  |  |  | Allow 0.30357 from $A_{\mathrm{r}}$ value of 56 |  |
|  |  |  | Ignore SF |  |
|  | - moles of C and N <br> - calculation of K:Fe:C:N mole ratio and empirical formula | (1) | $\mathrm{mol} \mathrm{C}=21.9 \div 12.0=1.8250$ |  |
|  |  |  | $\mathrm{mol} \mathrm{N}=25.5 \div 14.0=1.8214$ |  |
|  |  |  | Ignore SF except 1 SF |  |
|  |  | (1) | $\begin{array}{c:c:c} \mathrm{K} & : \mathrm{Fe}: & \mathrm{C} \end{array}: \underset{\mathrm{N}}{\mathrm{Fe}}$ |  |
|  |  |  | $\begin{array}{cccc} 3 & : & 1 & 6 \\ \text { empirical formula is } & 6 \\ \mathrm{~K}_{3} \mathrm{FeC}_{6} \mathrm{~N}_{6} \end{array}$ |  |
|  |  |  | Allow $\mathrm{K}_{3} \mathrm{Fe}(\mathrm{CN})_{6}$ |  |
|  |  |  | Allow elements in any order |  |
|  |  |  | TE on moles of $\mathrm{K}, \mathrm{Fe}, \mathrm{C}$ and N provided empirical formula is closest whole number ratio |  |
|  |  |  | Correct answer with no working scores (3) |  |



| Question Number | Answer |  | Additional guidance |  |  |  | Mark |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 18(f) | A completed table showing: <br> - correct identification for reaction 2 <br> - correct identification for reaction 3 |  | Example of completed table: |  |  |  | 2 |
|  |  | (1) |  | Neutralisation | Ligand exchange | Redox |  |
|  |  |  | Reaction 2 |  | $\checkmark$ |  |  |
|  |  |  | Reaction 3 |  |  | $\checkmark$ |  |
|  |  |  |  | rm of positiv |  | crosses |  |
|  |  |  | Ignore any | frm of negati | dentification |  |  |
|  |  |  | Do not aw | d more than | $x$ ticked in each |  |  |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(a) | - suitable test: bromine water/ $\mathrm{Br}_{2}(\mathrm{aq})$ <br> - result of test: decolourises (from orange) with Dewar structure (and no change with benzene) | Allow bromine / $\left.\mathrm{Br}_{2}(\mathrm{l})\right) / \mathrm{Br}_{2}$ in organic solvent Do not award $\mathrm{Br} / \mathrm{Br}^{-}$ <br> Accept potassium manganate((VII))/KMnO4 and acidified/ named acid/ $\mathrm{H}^{+}$ <br> Do not award hydrogenation <br> Do not award combustion <br> Allow does not decolourise with benzene Allow brown/orange/yellow for colour of bromine water Allow red/brown/orange for colour of bromine Allow pink/purple for colour of potassium manganate((VII)) <br> Ignore colour fades for decolourises Ignore reference to addition/substitution <br> Do not award any reference to decolourisation/reaction with benzene | 2 |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(b) | - similarity: (both compounds have) one (NMR) peak <br> - difference: expected chemical shift values | Ignore any reference to IR and/or ${ }^{13} \mathrm{C}$ NMR | 2 |
|  |  | Allow (both compounds have) one proton environment |  |
|  |  | Ignore just same number of peaks |  |
|  |  | Ignore references to relative peak area/integration/splitting |  |
|  |  | Do not award any other number of peaks |  |
|  |  | chemical shift for benzene within range of 6.4 to 8.4 ppm (actual value is 7.3 ppm ) |  |
|  |  | and chemical shift for Ladenburg structure within range of |  |
|  |  | 0 to 2.3 ppm (actual value is 2.3 ppm ) |  |
|  |  | Allow any range or value within the above ranges |  |
|  |  | Ignore just benzene would have a higher chemical shift than Ladenburg structure or reverse argument |  |
|  |  | Ladenburg structure or reverse argument |  |
|  |  | Do not award additional incorrect chemical shifts |  |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(c) | An explanation that makes reference to the following points: <br> - showed that all $\mathrm{C}-\mathrm{C}$ bonds are the same length in benzene <br> - in Kekulé structure the $\mathrm{C}=\mathrm{C}$ bonds would be shorter than the $\mathrm{C}-\mathrm{C}$ bonds (or reverse argument) (1) | Ignore any reference to: <br> $\mathrm{C}-\mathrm{H}$ bonds bond strength/bond angle <br> delocalised electrons <br> Dewar/Ladenburg structures <br> Allow showed benzene is a regular hexagon <br> Allow showed benzene contains only one type of carbon-carbon bond <br> Allow benzene bond lengths are in between $\mathrm{C}=\mathrm{C}$ and $\mathrm{C}-\mathrm{C}$ <br> Ignore just benzene has no $\mathrm{C}=\mathrm{C}$ bonds <br> Do not award benzene bond lengths are longer than $\mathrm{C}-\mathrm{C} /$ shorter than $\mathrm{C}=\mathrm{C}$ <br> Allow Kekulé structure would have shown two different lengths/types of carbon-carbon bond <br> Allow Kekulé structure would have alternating carbon-carbon bond lengths <br> Ignore just Kekulé has $\mathrm{C}=\mathrm{C}$ bonds <br> Do not award $\mathrm{C}-\mathrm{C}$ bonds would be shorter than the $\mathrm{C}=\mathrm{C}$ bonds <br> If no other mark awarded, just bond lengths equal in benzene but different in Kekulé scores (1) | 2 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(d)(i) | A diagram showing: <br> - correct relative stabilities <br> - two or three numerical differences in enthalpy with appropriate arrows | Example of diagram: <br> Allow names for structures <br> If three values and arrows are given they must all be correct to score M2 <br> Allow slight imprecision in start and end of arrows in M2 <br> Ignore any x-axis label <br> Do not award double headed arrows in M2 <br> Do not award incorrect sign in M2 | 2 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(d)(ii) | An answer that makes reference to the following: <br> - pi bonds are weaker/more reactive/require less energy to break (than sigma bonds) <br> or <br> fewer bonds must break to convert the Dewar structure to benzene | Ignore just Dewar structure has pi/double bonds/is unsaturated Ignore just Dewar structure has weaker bonds <br> Do not award C=C/double bonds weaker/require less energy to break (than C-C/single bonds) <br> Accept reverse argument <br> Allow any specified numbers to indicate fewer bonds must break <br> Ignore fewer new bonds must form <br> Ignore Dewar structure is more similar to benzene <br> Ignore carbon atoms already in a ring/hexagon <br> Ignore any reference to intermolecular forces <br> Do not award (higher) ring strain in Ladenburg <br> Do not award smaller difference in enthalpy | 1 |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(e) | An explanation that makes reference to the following points: <br> E-hexa-1,4-diene <br> - twice the hydrogenation enthalpy (of hex-3-ene) as two (isolated) $\mathrm{C}=\mathrm{C}$ bonds <br> E-hexa-1,3-diene <br> - less exothermic/more stable (by $22 \mathrm{~kJ} \mathrm{~mol}^{-1}$ than E-hexa-1,4-diene and as some delocalisation of pi-bond(s) | Allow double bond for $\mathrm{C}=\mathrm{C}$ throughout <br> Accept $-118 \times 2$ (=-236) as two C=C bonds <br> Allow twice the hydrogenation enthalpy as no delocalisation of pi-bond(s) <br> Accept less negative <br> Allow more positive <br> Allow some delocalisation of double bond(s) <br> Allow double bonds/p-orbitals are conjugated <br> Allow double bonds/p-orbitals are close enough to overlap <br> Ignore just $\mathrm{C}=\mathrm{C}$ are close <br> Ignore just delocalisation of electrons <br> Ignore electron density more spread out <br> Ignore resonance stabilised | 2 |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(f)(i) | - skeletal formula of any one isomer <br> - skeletal formulae of second and third isomers | Example of correct skeletal formulae: <br> (1,2-isomer) <br> (1,3-isomer) <br> (1,4-isomer) <br> Allow Kekulé benzene ring <br> Allow structural/displayed $\mathrm{CH}_{3}$ and $\mathrm{CH}_{3} \mathrm{CO}$ groups <br> If no other mark awarded, 1,2-, 1,3- and 1,4-isomers with incorrect side chain/cyclohexane ring scores (1) | 2 |


| Question <br> Number | Answer |  | Additional guidance | Mark |
| :--- | :--- | :---: | :--- | :---: |
| 19(f)(ii) | $\bullet \quad$ (identification of $\mathbf{X}$ as) 1,4-isomer | (1) | Allow any form of identification, including (f)(i) annotation <br> Allow just '1,4' or 'para' | $\mathbf{2}$ |
|  | $\bullet$ (7 peaks consistent with) 7 carbon environments | (1) | M2 dependent on a structure containing 7 carbon <br> environments |  |
|  |  |  | Accept 1,2-isomer and/or 1,3-isomer have 9 carbon <br> environments/would have 9 peaks |  |
| Allow (4 arene peaks consistent with) 4 arene carbon <br> environments <br> Allow 1,2-isomer and/or 1,3-isomer have 6 arene carbon <br> environments/would have 6 arene peaks |  |  |  |  |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 19(f)(iii) | A mechanism including: <br> - curly arrow from on or within circle <br> to $\mathrm{C}^{+}$of $\mathrm{CH}_{3} \mathrm{CO}^{+}$ <br> - structure of intermediate ion <br> - curly arrow from $\mathrm{C}-\mathrm{H}$ bond to within ring <br> and <br> correct product <br> - balanced equation for regeneration of catalyst (1) | Example of mechanism: <br> Allow Kekulé benzene ring <br> 'Horseshoe' facing tetrahedral carbon and covering at least three carbons with some part of positive sign within 'horseshoe' <br> Allow methyl at 1,2- or 1,3-positions Ignore missing methyl substituent <br> Do not award dotted/dashed C-H/C-C bonds unless 3D structure <br> Allow 1,2- or 1,3-product from corresponding intermediate Do not award missing methyl substituent $\begin{equation*} \mathrm{AlCl}_{4}^{-}+\mathrm{H}^{+} \rightarrow \mathrm{AlCl}_{3}+\mathrm{HCl} \tag{1} \end{equation*}$ | 4 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 20(a) | An answer that makes reference to one of the <br> following points: | Ignore references to ionisation energy <br> Ignore partially full d orbital(s)/d subshell <br> Ignore more than one stable ion <br> Ignore references to heterogeneous catalysis/adsorption <br> Ignore references to alternative reaction pathways/activation energy | 1 |
|  | - variable oxidation state/oxidation number <br> or <br> (easily) oxidised and reduced (back to <br> original oxidation state) <br> or <br> (easily) donate and accept electrons <br> (from other molecules/species) | Allow can change oxidation state/oxidation number <br> Allow have different oxidation state(s)/oxidation number(s) <br> Ignore variable valency | Allow just lose and gain electrons (easily) |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :--- | :---: |
| 20(b)(i) | An answer that makes reference to the following points: | Accept coordinate for dative throughout | 2 |
|  | - monodentate: forms a single/one dative (covalent) bond (1) | Accept donates a single/one lone pair <br> Allow occupies a single/one coordination site |  |
|  | ligand: (a species with a) lone pair (of electrons) that can form <br> a dative (covalent) bond to a (central transition) metal (ion) (1) |  |  |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 20(b)(ii) | A completed diagram showing: <br> - three adjacent THF/Cl ligands | Expected diagram: <br> Ignore lone pairs <br> Ignore 1- charge on Cl ligands | 1 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 20(c)(i) | $\bullet$ yellow to (permanent pale) green | Ignore qualifiers, eg pale <br> Ignore precipitate | $\mathbf{1}$ |


| Question Number | Answer |  | Additional guidance | Mark |
| :---: | :---: | :---: | :---: | :---: |
| 20(c)(ii) | A calculation including: <br> - moles of $\mathrm{Ti}^{3+}$ in titre |  | Example of calculation: | 5 |
|  |  |  | moles of $\mathrm{Ti}^{3+}=0.085 \times \frac{20.70}{1000}=0.0017595 / 1.7595 \times 10^{-3}$ |  |
|  | - moles of $\mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2} \cdot 6 \mathrm{H}_{2} \mathrm{O}$ in $100 \mathrm{~cm}^{3}$ | (1) | moles of $\mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2} .6 \mathrm{H}_{2} \mathrm{O}=\frac{0.75}{256.3}=0.0029263 / 2.9263 \times 10^{-3}$ |  |
|  | - moles of $\mathrm{NO}_{3}{ }^{-}$in $10.00 \mathrm{~cm}^{3}$ | (1) | moles of $\mathrm{NO}_{3}{ }^{-}=\frac{0.0029263}{10} \times 2=0.00058525 / 5.8525 \times 10^{-4}$ <br> TE on moles $\mathrm{Mg}\left(\mathrm{NO}_{3}\right)_{2} .6 \mathrm{H}_{2} \mathrm{O}$ |  |
|  |  |  | Ignore SF except 1 SF in M1, M2 and M3 <br> Ignore truncation of moles in M1, M2 and M3, eg 0.0005852 |  |
|  | - $\mathrm{Ti}^{3+}$ : $\mathrm{NO}_{3}{ }^{-}$mol ratio | (1) | $\begin{aligned} \mathrm{Ti}^{3+}: \mathrm{NO}_{3}{ }^{-} \text {mol ratio } & =0.0017595: 0.00058525 \\ & =3 \quad 3: 1 \\ \mathrm{TE} \text { on moles } \mathrm{Ti}^{3+} \text { and } & \text { moles } \mathrm{NO}_{3}{ }^{-} \end{aligned}$ |  |
|  | - final oxidation state of nitrogen | (1) | final oxidation state of nitrogen $=(+) 2$ |  |
|  |  |  | TE on mol ratio provided final oxidation state of nitrogen is between -3 and +4 , eg $\begin{aligned} \mathrm{Ti}^{3+}: \mathrm{NO}_{3}{ }^{-} \text {mol ratio } & =0.0017595: 0.00029263 \\ & =6: \end{aligned}$ <br> final oxidation state of nitrogen $=-1$ |  |
|  |  |  | Do not award incorrect oxidation state of N in $\mathrm{NO}_{3}{ }^{-}$ <br> Correct answer with no working scores (1) |  |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :--- | :---: |
| 20(c)(iii) | An equation including: | Example of equation: <br> $3 \mathrm{Ti}^{3+}+\mathrm{H}_{2} \mathrm{O}+\mathrm{NO}_{3}-\rightarrow 3 \mathrm{TiO}^{2+}+2 \mathrm{H}^{+}+\mathrm{NO}$ | $\mathbf{2}$ |
|  | • selection of correct nitrogen half-equation | (1) | TE on (c)(ii) provided +3 or +4 oxidation state |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 20(c)(iv) | An answer that makes reference to the following point: | Example of calculation: |  |
|  | $\bullet$ (calculation of) $E^{\ominus}$ cell value | $\left(E_{\text {cell }}^{\ominus}=0.96-0.10=\right)(+) 0.86(\mathrm{~V})$ <br> TE on ionic equation from (c)(iii): <br> $(+) 0.7(0)(\mathrm{V})$ for $\mathrm{Ti}^{3+}+\mathrm{NO}_{3}-\rightarrow \mathrm{TiO}^{2+}+\mathrm{NO}_{2}$ <br> $(+) 0.84(\mathrm{~V})$ for $2 \mathrm{Ti}^{3+}+\mathrm{H}_{2} \mathrm{O}+\mathrm{NO}_{3}-2 \mathrm{TiO}^{2+}+\mathrm{H}^{+}+\mathrm{HNO}_{2}$ | $\mathbf{1}$ |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 20(c)(v) | An answer that makes reference to the following point: |  | $\mathbf{1}$ |
|  | $\bullet$ (heat is to) speed up/increase rate of reaction | Allow to ensure fast oxidation of Ti ${ }^{3+}$ <br> Allow to provide activation energy/ $E_{\mathrm{a}}$ <br> Allow (reaction has a) high activation energy/Ea <br> Ignore just to provide (more) energy <br> Ignore to increase collision frequency <br> Ignore to ensure complete reaction <br> Ignore any reference to thermodynamic feasibility |  |


| Question Number | Answer |  |
| :---: | :---: | :---: |
| 20(c)(vi) | This question assesses a student's ability to show a coherent and logically structured answer with linkages and fully-sustained reasoning. <br> Marks are awarded for indicative content and for how the answer is structured and shows lines of reasoning. <br> The following table shows how the marks should be awarded for indicative content. |  |
|  | Number of indicative marking points seen in answer | Number of marks awarded for indicative marking points |
|  | 6 | 4 |
|  | 5-4 | 3 |
|  | 3-2 | 2 |
|  | 1 | 1 |
|  | 0 | 0 |

The following table shows how the marks should be awarded for structure and lines of reasoning.

|  | Number of marks awarded <br> for structure and sustained <br> lines of reasoning |
| :--- | :---: |
| Answer shows a coherent and <br> logical structure with linkages and <br> fully sustained lines of reasoning <br> demonstrated throughout. | 2 |
| Answer is partially structured <br> with some linkages and lines of <br> reasoning. | 1 |
| Answer has no linkages between <br> points and is unstructured. | 0 |

The mark for indicative content should be added to the mark for lines of reasoning. For example, an answer with five indicative marking points that is partially structured with some linkages and lines of reasoning scores 4 marks (3 marks for indicative content and 1 mark for partial structure and some linkages and lines of reasoning).

If there are no linkages between points, the same five indicative marking points would yield an overall score of 3 marks ( 3 marks for indicative content and no marks for linkages).

If there is any incorrect chemistry, deduct mark(s) from the reasoning. If no reasoning mark(s) awarded, do not deduct mark(s).

Comment: Look for the indicative marking points first, then consider the mark for the structure of the answer and sustained line of reasoning.

(Total for Question $20=20$ marks)

## Section C

| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 21(a) | A completed mechanism showing: | Example of completed mechanism: | $\mathbf{1}$ |
|  | $\bullet$ curly half-arrows to show homolytic fission of O-H bond | Accept curly half-arrows originating from <br> opposite sides of the O-H bond <br> Left-hand curly half-arrow must terminate <br> between T• and H |  |
| Right hand curly half-arrow must terminate on <br> or near to O atom of $\mathrm{H}-\mathrm{O}$ |  |  |  |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 21(b) | A completed mechanism showing: | Example of completed mechanism: | 2 |


| Question <br> Number | Answer | Additional guidance |  |  | Mark |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 21(c) | A completed table showing: <br> - two or three correct answers <br> - four correct answers | Example of completed table: |  |  | 2 |
|  |  |  | [Au(Curc)2] ${ }^{+}$ | $\left[\mathrm{Al}(\mathrm{Curc})\left(\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{OH}\right)_{2}\left(\mathrm{NO}_{3}\right)_{2}\right]$ |  |
|  |  | Coordination number | 4 | 6 |  |
|  |  | $\mathrm{O}-\mathrm{M}-\mathrm{O}$ bond angle | $90^{\circ}$ | $\underset{\text { Ignore } 180^{\circ}}{\underline{90^{\circ}}}$ |  |
|  |  | Shape | square planar | octahedral |  |
|  |  | Charge on metal ion | $\underline{+3}$ | +3 |  |


| Question Number | Answer |  | Additional guidance | Mark |
| :---: | :---: | :---: | :---: | :---: |
| 21(d)(i) | - $\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ and $\mathrm{H}_{2} \mathrm{SO}_{4}$ | (1) | If name and formula given, both must be correct to score M1 | 2 |
|  |  |  | Accept names (eg sodium dichromate((VI)) and sulfuric acid) |  |
|  |  |  | Allow $\mathrm{Cr}_{2} \mathrm{O}_{7}{ }^{2-}$ and $\mathrm{H}^{+} /$acidified dichromate |  |
|  |  |  | Ignore concentration of acid |  |
|  |  |  | Do not award $\mathrm{KMnO}_{4}$ for $\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ |  |
|  |  |  | Do not award HCl for $\mathrm{H}_{2} \mathrm{SO}_{4}$ |  |
|  | - heat/reflux | (1) | M2 dependent on some mention of dichromate (or manganate) oxidising agent Ignore distillation |  |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 21(d)(ii) | - correct structure for 2-aminobenzoic acid | Example of correct structure: <br> Accept hydrochloride salt ( $-\mathrm{NH}_{3} \mathrm{Cl}$ ) <br> Allow protonated amine group ( $-\mathrm{NH}_{3}{ }^{+}$) <br> Allow any correct combination of skeletal, structural or displayed formulae <br> Allow Kekulé benzene <br> Ignore connectivity <br> Ignore name, even if incorrect | 1 |
| Question Number | Answer | Additional guidance | Mark |
| 21(d)(iii) | - $\mathrm{NaNO}_{2} /$ sodium nitrite/sodium nitrate(III) and $\mathrm{HCl} /$ hydrochloric acid | Allow $\mathrm{HNO}_{2}$ /nitrous acid <br> Allow $\mathrm{H}^{+}$and $\mathrm{NO}_{2}{ }^{-}$ <br> Ignore conditions, including concentration of HCl Ignore $\mathrm{H}_{2} \mathrm{O}$ <br> Do not award $\mathrm{NaNO}_{3} /$ sodium nitrate | 1 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :--- | :--- | :--- | :---: |
| 21(d)(iv) | • correct structure for $N, N$-dimethylphenylamine | Example of correct structure: | $\mathbf{1}$ |
|  |  |  | Allow any correct combination of skeletal, structural or <br> displayed formulae <br> Allow Kekulé benzene |
| Ignore quaternary salt (-N(CH3)3 $\left.{ }^{+}\right)$ |  |  |  |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 21(d)(v) | An explanation including: <br> - effect of temperature higher than $5^{\circ} \mathrm{C}$ <br> - effect of temperature lower than $5^{\circ} \mathrm{C}$ | (diazonium/it) decomposes / reacts with water / forms a phenol / undergoes nucleophilic substitution (above $5^{\circ} \mathrm{C}$ ) <br> Ignore byproducts form / side reactions occur / yield too low (above $5^{\circ} \mathrm{C}$ ) <br> (rate of reaction) too slow (below $5^{\circ} \mathrm{C}$ ) <br> Allow just slows down (below $5^{\circ} \mathrm{C}$ ) <br> Ignore insufficient energy for reaction to occur (below $5^{\circ} \mathrm{C}$ ) Ignore any reference to activation energy/collision frequency Ignore freezes (at $0^{\circ} \mathrm{C}$ and below) | 2 |


| Question Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 21(e)(i) | A completed equation showing: <br> - correct balancing of propanone and sodium ethanoate <br> - correct balancing of sodium hydroxide and water | Example of equation: <br> M2 dependent on M1 | 2 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 21(e)(ii) |  | Correct answer with no working scores (3) | 3 |
|  |  | Example of calculation: |  |
|  | M1: molar masses |  |  |
|  | - $\quad$ (2-nitrobenzaldehyde) and | $\begin{aligned} M(2 \text {-nitrobenzaldehyde }) & =7 \times 12.0+5 \times 1.0+1 \times 14.0+3 \times 16.0 \\ & =151\left(.0 \mathrm{~g} \mathrm{~mol}^{-1}\right) \end{aligned}$ |  |
|  | $M$ (indigotin) | $\begin{align*} M(\text { indigotin }) & =16 \times 12.0+10 \times 1.0+2 \times 14.0+2 \times 16.0  \tag{1}\\ & =262\left(.0 \mathrm{~g} \mathrm{~mol}^{-1}\right) \end{align*}$ |  |
|  | Then, for M2 and M3, either: | Allow truncation of mass/moles throughout, eg 0.03816 for 0.038168 Ignore SF except 1 SF in M2 and M3 |  |
|  | Method 1 (M2 and M3) <br> - moles indigotin in 10.0 g | moles indigotin in $10.0 \mathrm{~g}=\frac{10.0}{262}=0.038168$ |  |
|  |  | TE on $M$ (indigotin) |  |
|  | indigotin:2-nitrobenzaldehyde mol ratio | moles 2-nitrobenzaldehyde $=2 \times 0.038168$ ( $=0.076336$ ) TE on moles indigotin |  |
|  | - moles 2-nitrobenzaldehyde required | $\begin{aligned} \text { moles 2-nitrobenzaldehyde required } & =\frac{100}{85} \times 0.076336 \\ & =0.089807 \end{aligned}$ |  |
|  | and <br> mass 2-nitrobenzaldehyde required | $\begin{align*} \text { mass 2-nitrobenzaldehyde required } & =0.089807 \times 151  \tag{1}\\ & =13.561 \\ & =14(\mathrm{~g}) \end{align*}$ |  |
|  |  | TE on moles 2-nitrobenzaldehyde TE on M(2-nitrobenzaldehyde) |  |



| Question Number | Answer |  | Additional guidance | Mark |
| :---: | :---: | :---: | :---: | :---: |
| 21(f)(i) | A drawing showing: <br> - hydrolysed ester linkage <br> and <br> correct carbon frame and amine group <br> - deprotonated carboxylic acid and phenol groups | (1) <br> (1) | Example of drawing: <br> Allow -ONa <br> Do not award -O-Na <br> Allow $E$ isomer <br> Allow Kekulé benzene <br> Allow any correct combination of skeletal, structural or displayed formulae | 2 |


| Question <br> Number | Answer | Additional guidance | Mark |
| :---: | :---: | :---: | :---: |
| 21(f)(ii) | A drawing showing: <br> - correct condensation product | Example of drawing: <br> Allow Kekulé benzene <br> Allow any correct combination of skeletal, structural or displayed formulae <br> Allow diacylated product, ie | 1 |

Total for Section C = 20 marks Total for Paper = 90 marks

