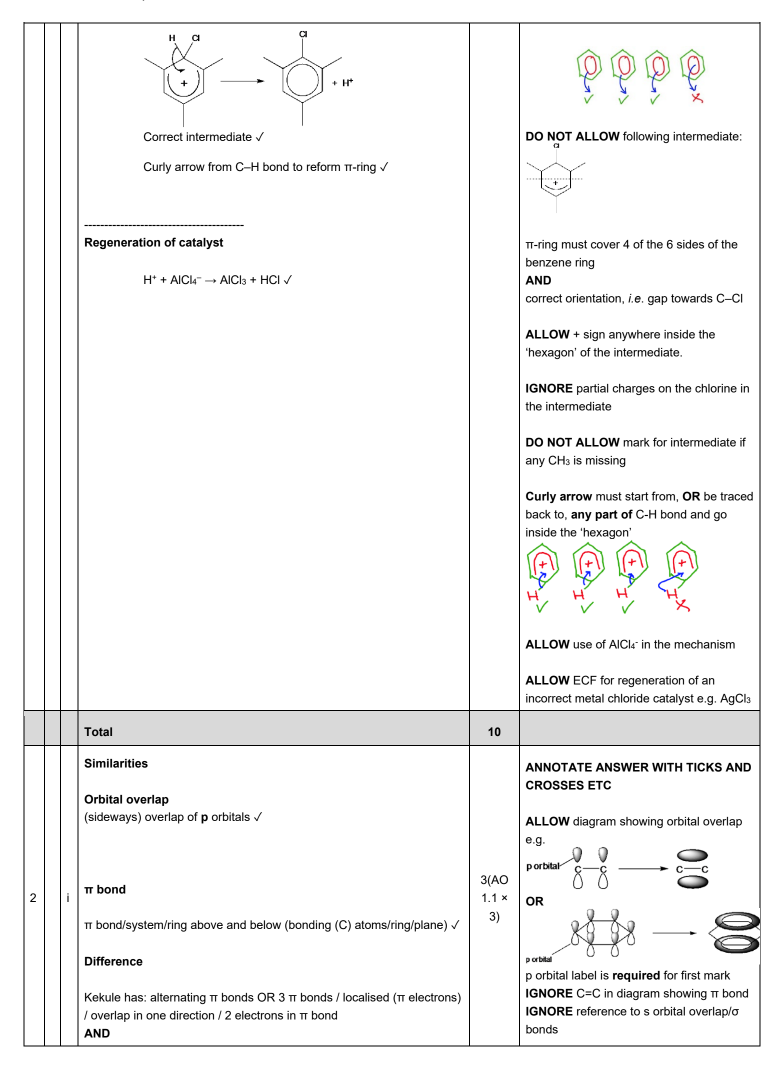
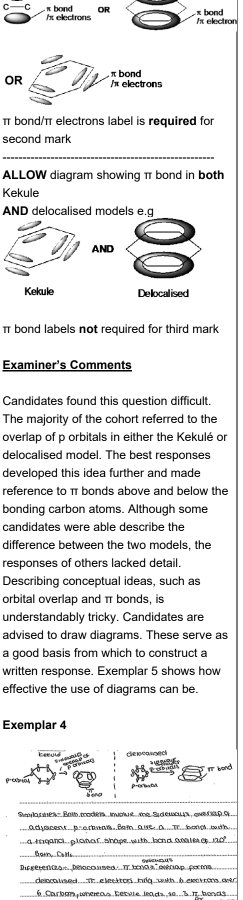
# **Mark scheme – Aromatic Compounds**

| Questi<br>on | Answer/Indicative content  | Marks  | Guidance   |
|--------------|--|--|--|
| 1 i          | Cl     Cl       Organic product with B     Organic product with C  | 2<br>(AO2.5<br>×2)   |  |
| ii           | Reactivity of B         in B electrons are localised         OR         in B π-bond is localised √         Reactivity of C         in C electrons are delocalised         OR         In C electrons are delocalised         OR         In C π-system / ring is delocalised         In B, electron density is higher         AND         B is more susceptible to electrophilic attack         OR         B attracts/accepts the electrophile/Cl₂ more         OR         B polarises the electrophile/Cl₂ more √         ORA | 3<br>(AO1.1<br>×3)   | ALLOW labelled diagram to show delocalised system         IGNORE charge density         IGNORE electronegativity         IGNORE B is more reactive/reacts more readily (no reference to electrophile)         IGNORE references to electron density spread around the π-ring         ALLOW chlorine         Examiner's Comments         Candidates answered this question well.         Many were able to correctly use the terms delocalised and localised in their responses and were able to provide comparisons for both electron density and attack of an electrophile. |
| ii<br>i      | Generation of electrophile<br>$A C _3 + C _2 \rightarrow A C _{4^-} + C ^+ \checkmark$ Attack of Cl <sup>+</sup> $\qquad \qquad $   | 5<br>(AO1.2<br>)<br>(AO1.2<br>)<br>(AO2.5<br>)<br>(AO1.2<br>)<br>(AO1.2<br>) | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES<br>ALLOW FeCl <sub>3</sub> + Cl <sub>2</sub> → FeCl <sub>4</sub> <sup></sup> + Cl <sup>+</sup><br>ALLOW use of Fe<br>NOTE: curly arrows can be straight,<br>snake-like, etc.<br>   |

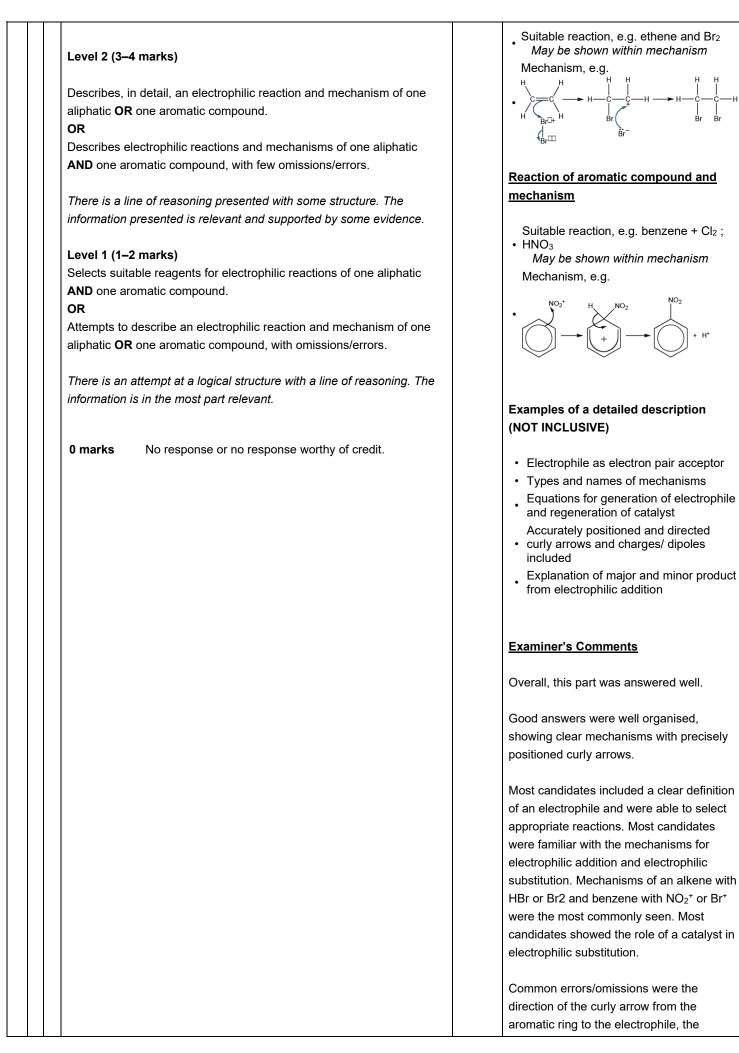


Delocalised has:  $\pi$  ring (system) / all p orbitals overlap OR ( $\pi$ electrons) spread around ring / overlap in both directions / 6 electrons **ALLOW** from labelled diagram showing  $\pi$ bond e.g. in  $\pi$  bond /



Delocaused structure ques more stabulity

|   |   |    |   |                 | This response starts with a diagram for<br>both the Kekulé and delocalised models of<br>benzene. Each diagram is clear and<br>shows the p orbitals and $\pi$ bonds<br>separately. This is an excellent response<br>for future candidates to follow.<br>The candidate's written response is<br>effectively a description of the diagrams<br>they have drawn and describes the<br>similarities and differences between these<br>two models clearly. This response scored<br>full marks and demonstrates the best style<br>of response seen in this part.<br><b>ALLOW</b> (C–C) bond enthalpy is between<br>single (C–C) and double bond (C=C)<br><b>OR</b> all (C–C) bond enthalpies are the<br>same  |
|---|---|----|---|-----------------|---|
|   | i | II | <ul> <li>Any 2 pieces of evidence from (√ √)</li> <li>Bond length (C-C) bond length is between single (C-C) and double bond (C=C) OR all (C-C) bond lengths are the same</li> <li><i>ΔH</i> hydrogenation <i>ΔH</i> hydrogenation less (exothermic) than expected</li> <li>Resistance to reaction</li> <li>Benzene is less reactive than alkenes</li> <li>OR bornination of benzene requires a catalyst/halogen carrier OR benzene does not react with/decolourise bromine (at room temperature)</li> <li>OR benzene reacts by substitution</li> <li>OR benzene does not (readily) react by addition</li> </ul> | 2(AO<br>1.1 ×2) | IGNORE enthalpy of hydration<br>Benzene is unreactive is <b>not</b> sufficient<br><i>(no comparison to alkene)</i><br>For halogen carrier,<br>ALLOW name or formula of suitable<br>catalyst<br>e.g. Fe, AICl <sub>3</sub> , FeBr <sub>3</sub><br>Examiner's Comments<br>Candidates were well prepared for this<br>question and the majority of the cohort<br>scored two marks. The most common<br>piece of evidence given was the lack of<br>reactivity of benzene with bromine, with<br>candidates citing the need for a catalyst for<br>the reaction to occur. Responses also<br>included reference to carbon-carbon bond<br>lengths as well as the enthalpy of<br>hydrogenation. A small but significant<br>proportion of the cohort referred to<br>hydration rather than hydrogenation. |
|   |   |    | Total   | 5               |   |
| 3 |   |    | Refer to marking instructions on page 5 of mark scheme for guidance<br>on marking this question.<br>Level 3 (5–6 marks)<br>Describes, in detail, electrophilic reactions and mechanisms of one<br>aliphatic AND one aromatic compound.<br>There is a well-developed line of reasoning which is clear and logically<br>structured. The information presented is relevant and substantiated.  | 6               | Indicative scientific points may include:<br><u>Explanation of role of electrophiles in</u><br><u>organic chemistry</u><br><u>Reaction of aliphatic compound and</u><br><u>mechanism</u>  |



position of the curly arrow when H<sup>+</sup> is lost from an aromatic intermediate, and not showing the lone pair when Br<sub>-</sub> attacks a carbocation.

Some answers lacked detail and gave only two mechanisms with minimal supporting words.

Lower ability candidates described nucleophilic addition or substitution as one of their mechanisms or had curly arrows going in the wrong direction.

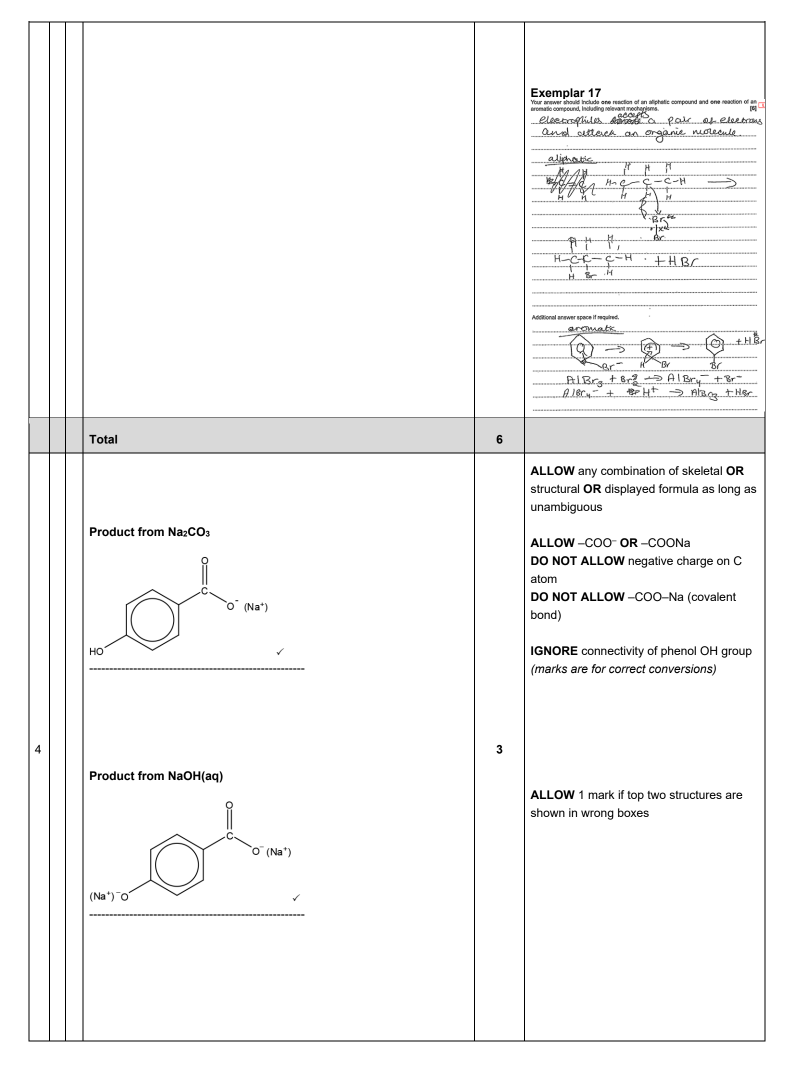
A few candidates answered in prose without including equations or diagrams. Candidates are advised that mechanisms must always be communicated in the usual diagrammatic way.

Exemplar 16 is a very clear and concise response showing all the key features of electrophilic addition and electrophilic substitution, including the role of the FeCl<sub>3</sub> catalyst. Curly arrows are precisely positioned, with correct use of lone pairs and charges. The candidate has demonstrated excellent knowledge and understanding.

The response in Exemplar 17 is clearly at a different level. The candidate has chosen an alkane rather than an alkene and has used curly arrows and charges incorrectly. This candidate appears to have been poorly prepared.

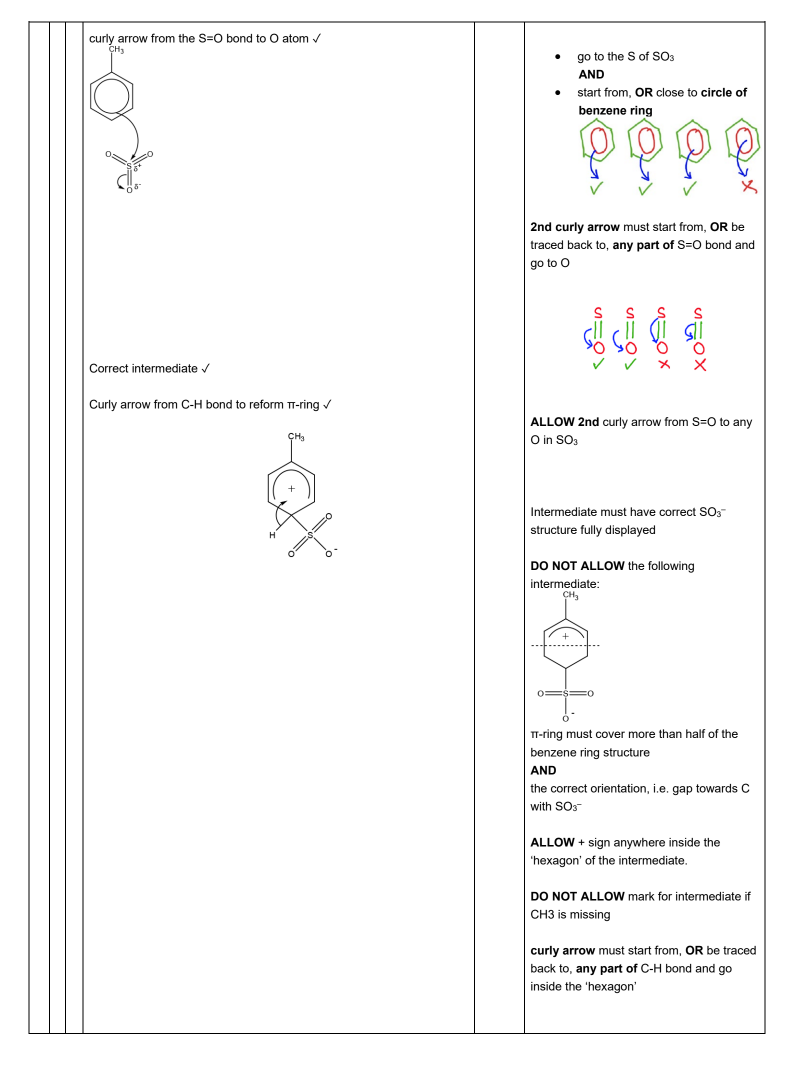
#### Exemplar 16

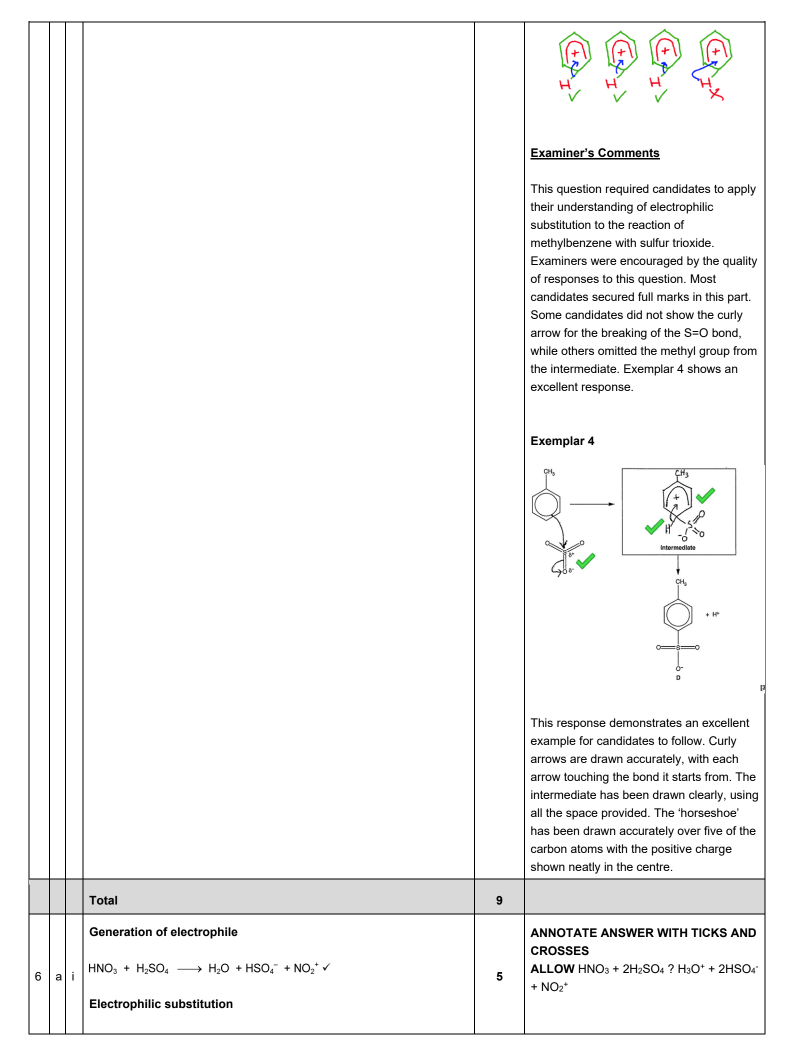
[6] luding rele Electrophiles are electron pair acceptors; and can accept or pair of electrons how 23 Aliphatic: Electrophilic addition: Ņ H-H J Co Br ٠, C2M4+ MBr , ~ C2M5Br 7. 12A ..... Aromatic: electrophilic: substitution Feel3 + Cl2 -> Feel4 + Cl C'cit Additional answer space if required.  $\rightarrow$ Fell, -> HCL + Fell, Fetta + HCC

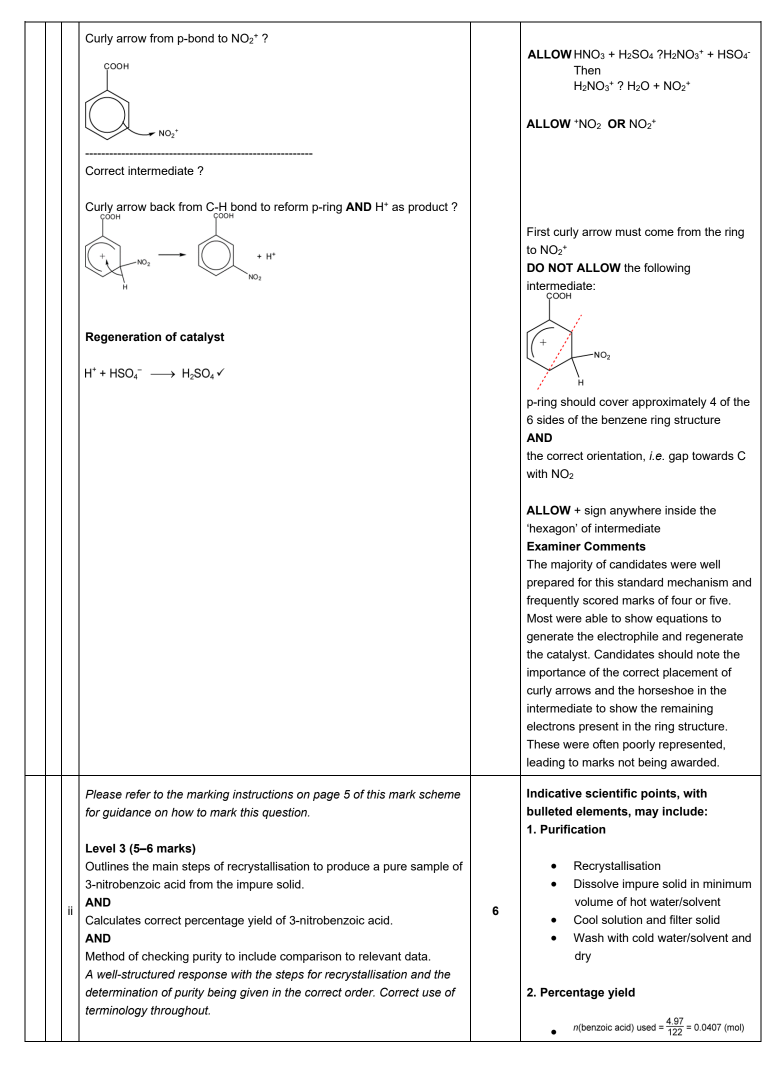


|   |   |   | Product from Br <sub>2</sub>   |   | ALLOW substitution of any H from benzene ring  |
|---|---|---|--|---|--|
|   |   |   | e.g. HO  |   | ALLOW multiple substitution, <i>i.e.</i> di-, tri-<br>and tetrabromo products.<br>IGNORE connectivity of phenol OH group<br>(marks are for correct conversions)  |
|   |   |   |  |   | Examiner's Comments  |
|   |   |   |  |   | This question assessed different reactions<br>of compound <b>H</b> , 4-hydroxybenzoic acid,<br>and discriminated well. Two of the<br>reactions focused on acid-base chemistry,<br>using the reagents Na <sub>2</sub> CO <sub>3</sub> and NaOH.<br>Many candidates recognised that the<br>carboxylic acid group would react in both<br>cases but only some managed to identify<br>when the phenol group was involved<br>correctly. A number of responses<br>suggested that a phenoxide ion was<br>formed with sodium carbonate but not with<br>sodium hydroxide.<br>The third reaction was substitution with<br>bromine. This reaction appeared more<br>familiar to all candidates with the majority<br>scoring this mark. A small proportion of<br>candidates substituted the phenol OH<br>group or carboxylic acid group. |
|   |   |   | Total  | 3 |  |
| 5 | а | i | Number of peaks2 marks2-nitrophenol AND 3-nitrophenol have six peaks/environments/types of<br>carbon √4-nitrophenol has four peaks/environments/types of carbon √4-nitrophenol has four peaks/environments/types of carbon √1 mark | 3 | IGNORE any numbers shown on<br>structures<br>ALLOW 1 mark only IF a response<br>identifies that all the compounds have 6<br>peaks/environments/types of C<br>OR all the compounds have 4<br>peaks/environments/types of carbon<br>IGNORE chemical shifts   |
|   |   |   | <ul> <li>4-nitrophenol can be distinguished</li> <li><b>OR</b></li> <li>2-nitrophenol and 3-nitrophenol cannot be distinguished √</li> </ul>   |   | <b>DO NOT ALLOW ECF</b> from an incorrect<br>number of peaks/environments/types of<br>carbon<br><u>Examiner's Comments</u>   |

|   |    |   |   | This question required candidates to apply<br>their knowledge of <sup>13</sup> C NMR spectroscopy<br>to deduce the number of different carbon<br>environments in each of the nitrophenols<br>shown. The best responses were succinct,<br>stating that it is possible to distinguish 4-<br>ntirophenol from the other two after<br>deducing the correct number of carbon<br>environments for each compound. Lower<br>ability candidates' responses did not<br>identify the symmetry in 4-nitrophenol and<br>suggested all three compounds would<br>produce six peaks, therefore making it<br>impossible to distinguish between them. |
|---|----|---|---|---|
|   | ii | (In phenol) a (Ione) pair of electrons on O is(partially)<br>delocalised/donated into the π-system / ring √<br>Electron density increases/is higher (than benzene) √<br>ORA<br>(phenol) is more susceptible to electrophilic attack<br>OR<br>(phenol) attracts/accepts electrophile/HNO <sub>3</sub> more<br>OR | 3 | ALLOW the electron pair in the p-orbitals<br>of the O atom becomes part of the π-<br>system / ring<br>ALLOW diagram to show movement of<br>lone pair into ring<br>ALLOW lone pair of electrons on O is<br>(partially) drawn/attracted/pulled/ into π-<br>system / ring<br>IGNORE activating<br>IGNORE charge density<br>IGNORE charge density<br>IGNORE phenol reacts more readily ( <i>no</i><br><i>reference to electrophile</i> )<br>ALLOW NO <sub>2</sub> <sup>+</sup> for electrophile   |
|   |    | (phenol) polarises electrophile/HNO₃<br>more √<br>ORA   |   | Examiner's Comments<br>The relative reactivity of phenol is well<br>known by candidates at this level and the<br>majority scored two or three marks.<br>Candidates who did not score full marks<br>gave imprecise or vague explanations.<br>This included reference to 'higher charge<br>density' or 'higher electronegativity' of the<br>phenol ring, rather than 'higher electron<br>density'.  |
|   |    |   |   | ANNOTATE WITH TICKS AND<br>CROSSES  |
| b |    |   | 3 | <b>NOTE:</b> curly arrows can be straight,<br>snake-like, etc. but <b>NOT</b> double headed or<br>half headed arrows  |
|   |    | Curly arrow from π-bond to S in SO <sub>3</sub><br>AND  |   | 1st curly arrow must  |







| Level 2 (3–4 marks)<br>Attempts all three scientific points but explanations may be incomplete.<br>OR  | • $n(3\text{-nitrobenzoic acid}) \text{ made } = \frac{4.85}{167} = 0.0290 \text{ (mol)}$<br>• percentage yield $= \frac{0.0290}{0.0407} \times 100 = 71.3 \text{ (%)}$   |
|--|---|
| Explains two scientific points thoroughly with very few omissions.<br>The description of checking for purity or recrystallisation is clear and<br>any calculations structured. Key terminology used appropriately.   | <b>ALLOW</b> 71 to calculator value of 71.29001554 correctly rounded.   |
| Level 1 (1–2 marks)<br>A simple explanation based on at least two of the main scientific points.<br>OR<br>Explains one scientific point thoroughly with few omissions.<br>There is an attempt at a logical structure. The description of the<br>practical techniques provides some detail but may not be in the correct<br>order.  | <b>CHECK</b> for extent of errors by <b>ECF</b><br>Alternative correct calculation may<br>calculate theoretical mass of 3-<br>nitrobenzoic acid that can be produced as<br>$0.0407 \times 167 = 6.80$ (g) followed by:<br>percentage yield $=\frac{4.85}{6.80} \times 100 = 71.3$ (%) |
| <ul> <li>Purification step is unclear with few scientific terms and little detail, e.g. just 'recrystallise'.</li> <li>Calculation is difficult to follow, may just include a calculation of moles of reactants and/or products.</li> <li>Purity check specifies a method but this is unclear with little detail, e.g. take melting point.</li> <li>0 marks</li> <li>No response or no response worthy of credit.</li> </ul> | <ul> <li>Calculation must attempt to calculate n(benzoic acid) in mol.</li> <li>3. Checking purity <ul> <li>Obtain melting point</li> <li>Compare to known values</li> <li>Pure sample will have a (sharp) melting point very close to data book value</li> </ul> </li> </ul>         |
|  | <ul> <li>ALLOW alternative approach based on spectroscopy or TLC</li> <li>Spectroscopy</li> <li>Run an NMR/IR spectrum</li> <li>Compare to (spectral) database</li> <li>Spectrum of pure sample will contain same peaks and not others</li> </ul>                                     |
|  | <ul> <li>TLC</li> <li>Run a TLC</li> <li>Compare (<i>R</i>f value) to known data</li> <li>Pure sample will have a very similar Rf</li> </ul>  |
|  | <b>Examiner Comments</b><br>This question tested some of the practical<br>techniques covered as part of the practical<br>endorsement as well as requiring<br>candidates to calculate a percentage yield<br>for the reaction. This proved to be quite a                                |

|   |   |   |   | dissolving in a hot solvent and then being<br>allowed to cool before carrying out<br>filtration. High quality answers often went<br>above and beyond the requirements of the<br>marking scheme with some candidates<br>discussing the importance of dissolving in<br>the minimum amount of hot solvent to<br>obtain a saturated solution, the need to<br>wash and dry the crystals and provided<br>detail of the apparatus and or method<br>required.<br>Most candidates discussed that purity<br>could be determined by taking the melting<br>point of the product and comparing this to<br>a value obtained from data book. The most<br>comprehensive answers gave an indicated<br>of the apparatus required to carry out the<br>melting point determination and discussed<br>how the melting point becomes higher and<br>sharper as impurities are removed.<br>Common errors included comments about<br>carrying out a boiling point determination.<br>When carrying out a percentage yield<br>calculation, it is important to round<br>answers only at the last stage of the<br>calculation. Early rounding frequently led<br>candidates to obtain answers, which did<br>not gain credit. Weaker Candidates<br>divided the mass of 3-nitrobenzoic acid by<br>the mass of benzoic acid and obtained an<br>answer of 97.6%.<br>Answer = 71.3% |
|---|---|---|---|---|
| b | i | Phenol is the most easily nitrated/most reactive<br>AND<br>Benzoic acid is the least easily nitrated/least reactive √ | 1 | e.g. nitration becomes more difficult from<br>phenol (to benzene) to benzoic acid<br><b>OR</b><br>nitration becomes easier from right to left<br>in the table<br><b>Examiner Comments</b><br>The vast majority of candidates gave a<br>good answer to indicate that it was easier<br>to carry out the nitration of phenol than  |

|    |  |   | benzene and that benzene would undergo<br>nitration easier than benzoic acid.   |
|----|--|---|---|
|    |  |   | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES   |
|    |  |   | <b>ALLOW</b> the electron pair in the p orbitals<br>of the O atom becomes part of the p-<br>system / ring   |
|    |  |   | ALLOW diagram to show movement of<br>lone pair into ring<br>ALLOW lone pair of electrons on O is<br>(partially) drawn/attracted/pulled into<br>p-system / ring  |
|    | Reactivity of phenol   |   | IGNORE activating and deactivating.   |
|    | a (lone) pair of electrons on O is (partially)<br><b>delocalised/donated</b> into the<br>p-system/ring   |   | <b>ALLOW</b> the following alternatives for susceptibility to attack:   |
|    |  |   | <ul> <li>phenol attracts electrophiles /<br/>NO<sub>2</sub><sup>+</sup> more</li> <li>phenol polarises electrophiles /<br/>NO<sub>2</sub><sup>+</sup> more</li> <li>benzoic acid attracts electrophile<br/>/ NO<sub>2</sub><sup>+</sup> less</li> </ul> |
| ii | Reactivity of benzoic acid The –COOH group on benzoic acid is an electron withdrawing group $\checkmark$ | 3 | <ul> <li>benzoic acid polarises<br/>electrophiles / NO<sub>2</sub><sup>+</sup> less</li> </ul>  |
|    | Links electron density in p-bond to reactivity   |   | Examiner Comments The most able candidates scored well or   |
|    | In phenol electron density is higher   |   | what proved a difficult question for many   |
|    | AND<br>The ring is more susceptible to attack  |   | Although the vast majority of candidates<br>knew about the reasons behind phenol's<br>increased reactivity many were unable to  |
|    | OR   |   | express themselves clearly to gain credit<br>Often answers lacked the specific detail   |
|    | In benzoic acid electron density is lower  |   | about the lone pair on the oxygen atom i  |
|    | AND<br>The ring is less susceptible to attack √  |   | the -OH group being delocalised into the<br>ring. Weaker answers discussed electron<br>being supplied to the ring, the -OH group<br>providing the electrons to the ring or just<br>that -OH is an activator. Good answers                               |
|    |  |   | expressed the increase in electron densi<br>and the subsequent increase in phenol's   |
|    |  |   | susceptibility to electrophilic attack. The most able candidates were able to   |
|    |  |   | interpret the information given in the  |
|    |  |   | question to establish that the -COOH  |
|    |  |   | group must be electron withdrawing  |
|    |  |   | leading to decreased electron density of  |
|    |  |   | the ring structure and reduced ability to   |
|    |  |   |   |

| c i | Bromination: $Br_2$ AND A/ $Br_3$ /Fe $Br_3$ /Fe $\checkmark$<br>Intermediate<br>$\downarrow \qquad \qquad$   | 3 | ALLOW any combination of skeletal OR<br>structural OR displayed formula as long as<br>unambiguous<br>ALLOW any suitable halogen carrier<br>catalyst<br>ALLOW Kekulé structure<br>IGNORE names (question asks for<br>formulae)<br>IGNORE reaction conditions even if<br>incorrect<br>IGNORE 'dilute' for HC/<br>IGNORE 'dilute' for HC/<br>IGNORE H <sub>2</sub><br>IGNORE MaOH if seen as a reagent to<br>convert nitro group into amine<br>e.g 'Sn/(concentrated) HC/ then NaOH'<br>scores the mark<br>Examiner Comments<br>Candidates were able, in the main, to<br>provide the reagents for bromination and<br>reduction. The structure of the<br>intermediate compound in the preparation<br>of 3-bromophenylamine proved to be<br>straightforward, however common errors<br>involved the omission of the halogen<br>carrier catalyst for bromination or stating<br>names rather than formulae as indicated in<br>the question. |
|-----|--|---|---|
|     | NH <sub>2</sub> is 2,4 directing $\checkmark$<br>Products (1 mark for each):<br>$\downarrow \downarrow \downarrow \downarrow$ Br<br>$\downarrow \downarrow \downarrow$ $\downarrow \downarrow$ $\downarrow \downarrow$ $\downarrow \downarrow$ $\downarrow \downarrow$ $\downarrow$ $\downarrow$ | 3 | IGNORE references to electron<br>donating/withdrawing groups<br>ALLOW –NH <sub>2</sub> activates the ring causing<br>the new group to join at positions 2 and 4.<br>ALLOW ortho and para directing for 2,4<br>directing<br>IGNORE 6-directing<br>ALLOW Kekulé structure<br>IGNORE names<br>Examiner Comments<br>The most able candidates completed this<br>question with a clear statement that the<br>–NH <sub>2</sub> group was 2,4 directing and<br>provided two clearly drawn structures of 2-<br>bromophenylamine and 4-<br>bromophenylamine. The most common<br>errors observed included drawing two<br>structures that were identical and<br>explaining the two structures in terms of<br>electron donation from the –NH <sub>2</sub>  |

|   |   |    |   |    | without any indication of positioning.<br>Candidates using the terms ortho and para<br>directing were awarded full marks for their<br>answers.   |
|---|---|----|---|----|--|
|   |   |    | Total   | 21 |  |
| 7 |   | i  | $\begin{array}{c} CI & O \\   \\ CI - C \\   \\ CI \\ H_{\checkmark} \end{array}$   | 1  | <b>ALLOW</b> any combination of skeletal <b>OR</b> structural <b>OR</b> displayed formula as long as unambiguous   |
|   |   | ii | Any <b>organic</b> reaction in which sulfuric acid is a catalyst e.g:<br>Elimination of (H <sub>2</sub> O from) alcohols<br>Nitration of benzene<br>Esterification<br>Hydrolysis of esters/amides | 1  | The answer needs to refer to the reaction:<br>i.e. 'Elimination', 'hydrolysis' are<br>insufficient but 'Esterification' describes<br>the reaction<br><b>DO NOT ALLOW</b> oxidation for alcohols/<br>aldehydes  |
|   |   |    | Total   | 2  |  |
| 8 | a |    | One mark for each correct structure/reagent/condition as shown below  | 6  | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES<br>ALLOW any combination of skeletal OR<br>structural OR displayed formula as long as<br>unambiguous<br>IGNORE names of organic compounds<br>( <i>question asks for structures</i> )<br>ALLOW aluminium(III) chloride OR<br>aluminium trichloride<br>ALLOW FeC/ <sub>3</sub> OR Fe as halogen carrier in<br>first step.<br>ALLOW sodium borohydride OR sodium<br>tetrahydridoborate<br>IGNORE [H] for reducing agent in second<br>step<br>ALLOW H <sup>+</sup> / H <sub>2</sub> SO <sub>4</sub> / H <sub>3</sub> PO <sub>4</sub> / named<br>mineral acid for reagent in third step |
|   | b |    | Use as an organic feedstock $\checkmark$  | 1  | ALLOW the production of plastics or monomers   |

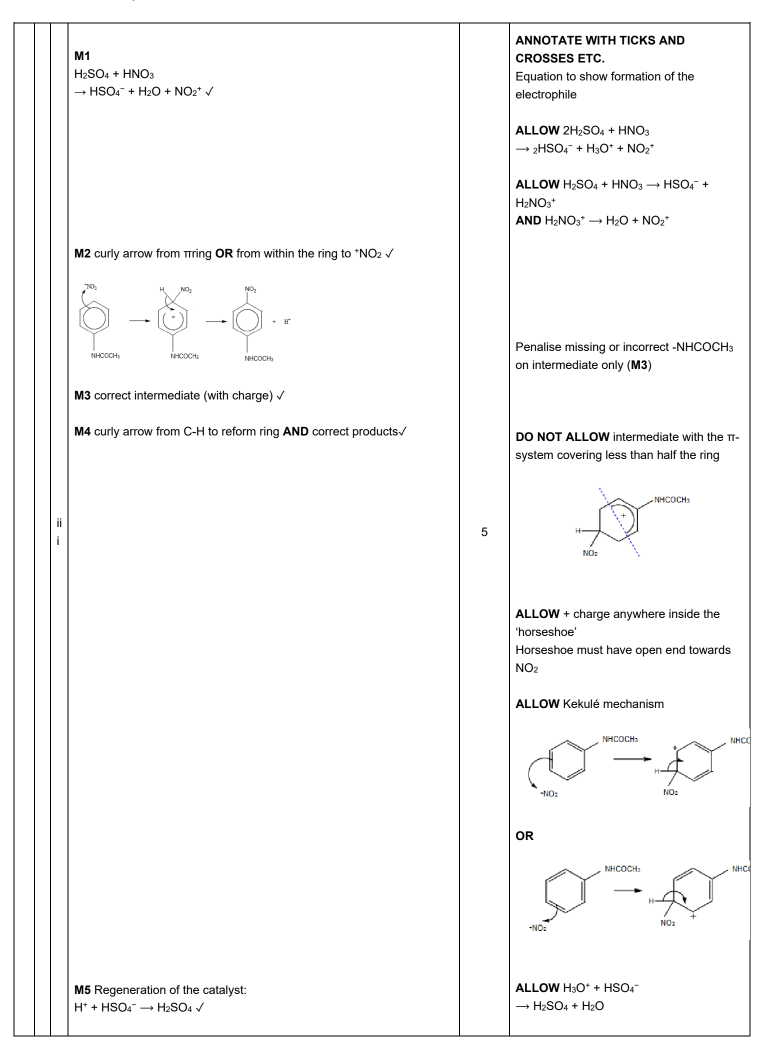
# 6.1.1 Aromatic Compounds

|     |         | OR  |   |  |
|-----|---------|---|---|--|
|     |         | UR  |   | or new polymers  |
|     |         | Combustion for energy production $\checkmark$   |   | Combustion alone is <b>not</b> sufficient  |
|     |         | Total   | 7 |  |
| 9 a | i       | Total<br>Dipole shown on C=O bond, $C^{\delta+}$ and $O^{\delta-}$ , AND curly arrow from the<br>C=O bond to the $O^{\delta-}$ atom<br>AND<br>Curly arrow from $\pi$ -bond to C in CO <sub>2</sub> $\checkmark$<br>$\int_{0}^{0} \int_{0}^{0} \int$ | 3 | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES<br>DO NOT ALLOW the following<br>intermediate:<br>$\int_{i} \int_{i} \int_$ |
|     |         |   |   | 'horseshoe' in the correct orientation, <i>ie</i><br>gap towards C with COO <sup>−</sup><br><b>ALLOW</b> + sign anywhere inside the<br>'hexagon' of intermediate   |
|     | ii      | Neutralisation ✓<br>(In Stage 1) phenol loses H <sup>+</sup><br>AND   | 2 | ALLOW acid-base ALLOW both Stage 1 AND Stage 3   |
|     |         | (In Stage 3) carboxylate ion gains $H^+ \checkmark$   |   | involve proton transfer  |
|     | ii<br>i | FIRST CHECK THE ANSWER ON THE ANSWER LINE<br>IF answer = 7.31 (g) award 3 marks   | 3 | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES  |
|     |         |   |   | ALLOW ECF at each stage  |

## 6.1.1 Aromatic Compounds

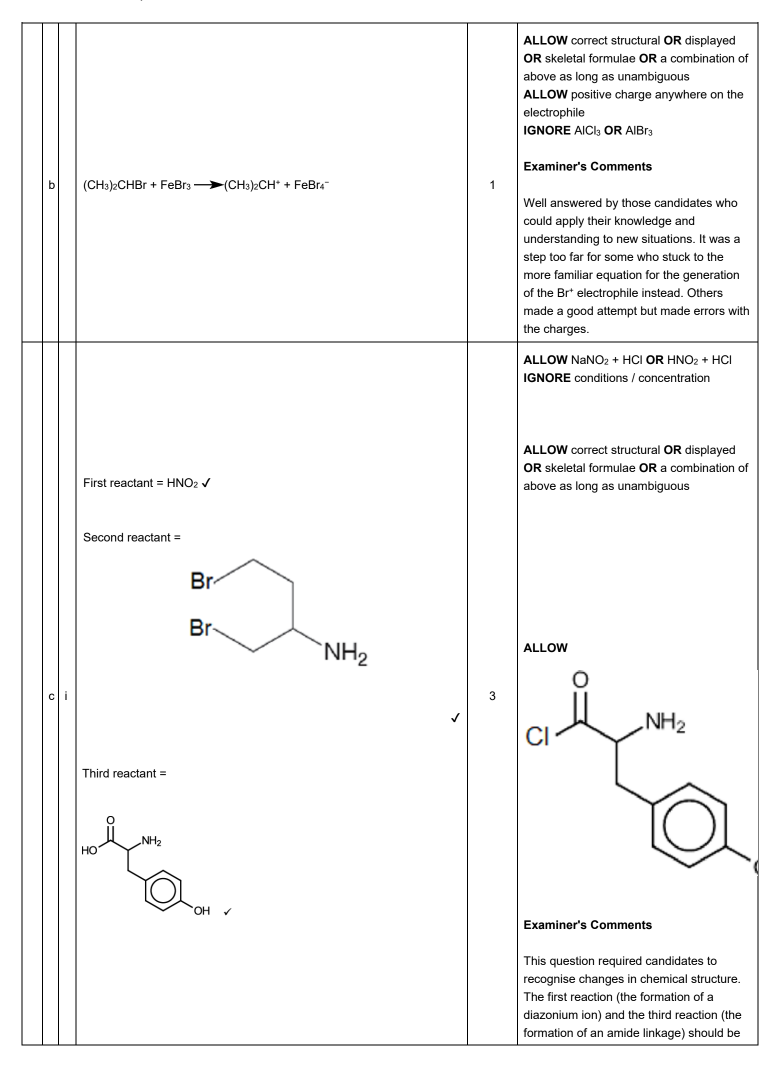
|        |   |   | actual<br>$n(\text{salicylic acid}) \text{ produced} = \frac{4.83}{138} = 0.035(0) \text{ (mol) } \checkmark$<br>theoretical<br>$n(\text{phenol}) = n(\text{salicylic acid}) = 0.035(0) \times \frac{100}{45.0} = 0.0778 \text{ (mol) } \checkmark$<br>Mass of phenol = 0.0778 × 94.0 = 7.31 (g) $\checkmark$ |    | ALLOW 3 SF up to calculator value<br>correctly rounded for intermediate values<br>100  ALLOW expected mass compound E<br>$= 4.83 \times 45.0 = 10.733 \text{ (g)}$<br>ALLOW Mass phenol reacted = $0.035 \times 94.0 = 3.29 \text{ (g)}$ |
|--------|---|---|---|----|--|
|        |   |   |   |    | ALLOW Mass of phenol used = $3.29 \times \frac{100}{45.0} = 7.31$ (g)<br>Note:<br>1.48 g would get 2 marks<br>(use of 45.0/100 instead of 100/45.0)<br>7.30 g would get 2 marks<br>(use of 0.0777 for moles phenol)                      |
|        | b |   | Skeletal formula of aspirin<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>Skeletal formula of ethanoic acid<br>$\downarrow$  | 2  | IF skeletal formulae are not used ALLOW<br>one mark if both the structures of aspirin<br>AND ethanoic acid are correct   |
| l      |   |   | Тotal   | 10 |  |
| 1<br>0 |   | i | Ethanoic anhydride<br>O $OH_{3}C O CH_{3}Other organic compound$  | 2  | ALLOW skeletal formula   |
|        |   |   |   |    | ALLOW correct structural OR displayed  |

|  |   |   | OR skeletal formulae OR a combination of<br>above as long as unambiguous<br>IGNORE names<br>Examiner's Comments<br>The structure of ethanoic anhydride was<br>well known and most candidates gave the<br>correct structure for the other organic<br>product of the reaction.   |
|--|---|---|--|
|  | FIRST CHECK THE ANSWER ON THE ANSWER LINE<br>IF answer = 2.66 (g) award 3 marks<br>IF answer = 4.36 (g) award 2 marks (% yield not used)<br>IF answer = 7.14 (g) award 2 marks (% yield used incorrectly)<br>n(phenylamine) (= $3.00/93.0$ ) = $0.0323$ mol $\checkmark$<br>n(compound A) = ( $0.0323 \times 0.61$ ) = $0.0197$ mol $\checkmark$<br>Mr (compound A) = $135$<br>AND<br>Mass of compound A = ( $135$ )( $0.0197$ ) = $2.66$ g $\checkmark$<br>OR<br>n(phenylamine) (= $3.00/93.0$ ) = $0.0323$ mol $\checkmark$<br>Mr (compound A) = $135$<br>AND<br>Theoretical mass of compound A = ( $0.0323 \times 135$ ) = $4.36 \checkmark$<br>Actual mass of compound A = ( $4.36 \times 0.61$ ) = $2.66$ g $\checkmark$ | 3 | ANNOTATE WITH TICKS AND<br>CROSSES ETC.<br>ALLOW 3 SF: 0.0323 up to calculator<br>value of 0.032258064 correctly rounded<br>ALLOW 3 SF up to calculator value<br>Penalise rounding to 2 SF once<br>ALLOW ECF on incorrectly rounded<br>values<br>Final answer must be expressed to 3<br>significant figures<br>ALLOW ecf from incorrect Mr<br>ALLOW ecf from incorrect Mr<br>IF answer = 2.65 (g) award 2 marks unless<br>this alternative method is used (3 marks)<br>93 g gives 135 g<br>3.00 g gives 135/93 x 3.00 = 4.35 g<br>4.35 x 0.61 = 2.65 g<br>Examiner's Comments<br>Once again it was clear that candidates<br>had been well prepared for this type of<br>percentage yield question. It was well<br>answered with almost all candidates<br>scoring some marks here and a high<br>proportion gaining all three marks for the<br>calculation. |



|     |   |   |  |    | Examiner's Comments   |
|-----|---|---|--|----|---|
|     |   |   |  |    | Although some lost marks through minor<br>errors on the details, most candidates<br>scored well on this question  |
|     |   |   | Total  | 10 |   |
| 1   |   |   | $ \begin{array}{c}                                     $   | 1  | <ul> <li>ALLOW correct structural OR displayed<br/>OR skeletal formulae OR a combination of<br/>above as long as unambiguous</li> <li>ALLOW disubstitution at any positions on<br/>benzene ring</li> <li>Examiner's Comments</li> <li>Generally well answered. Some<br/>candidates reacted compound B with only<br/>one mole of bromine or three moles of<br/>bromine and this underlines the<br/>importance of paying careful attention to<br/>information given in the stem of the<br/>question. Hydrogen, not HBr, was<br/>occasionally formed as the other product.</li> </ul>  |
|     |   |   | Total  | 1  |   |
| 1 2 | а | i | M1<br>p-orbitals overlap (to form pi / π-bonds) ✓<br>M2<br>π-bond(s) are delocalised in structure B ✓<br>M3<br>π-bonds are localised / between two carbons in structure A<br>✓<br>M4<br>M4<br>M4<br>Diagrams show correct position of delocalised and localised π-bonds<br>/ π-electrons | 4  | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES ETC         IGNORE p-orbitals overlap to form sigma<br>bonds         ALLOW electrons are delocalised in<br>structure B         IGNORE B has delocalised structure or<br>ring (must be electrons or π-bonds)         ALLOW π-electrons / p-orbital overlap<br>localised / between two carbons in<br>structure A         ALLOW p-orbitals overlap with one other<br>carbon         IGNORE electrons are localised OR<br>structure A has localised structure (must<br>be π-bonds / π-electrons / p-orbital<br>overlap)         ALLOW labelled diagram showing overlap<br>of p-orbitals between two carbon atoms<br>DO NOT ALLOW C=C in this diagram         Diagram for structure A must show the full<br>ring for M4 IGNORE C=C in M4 diagram |

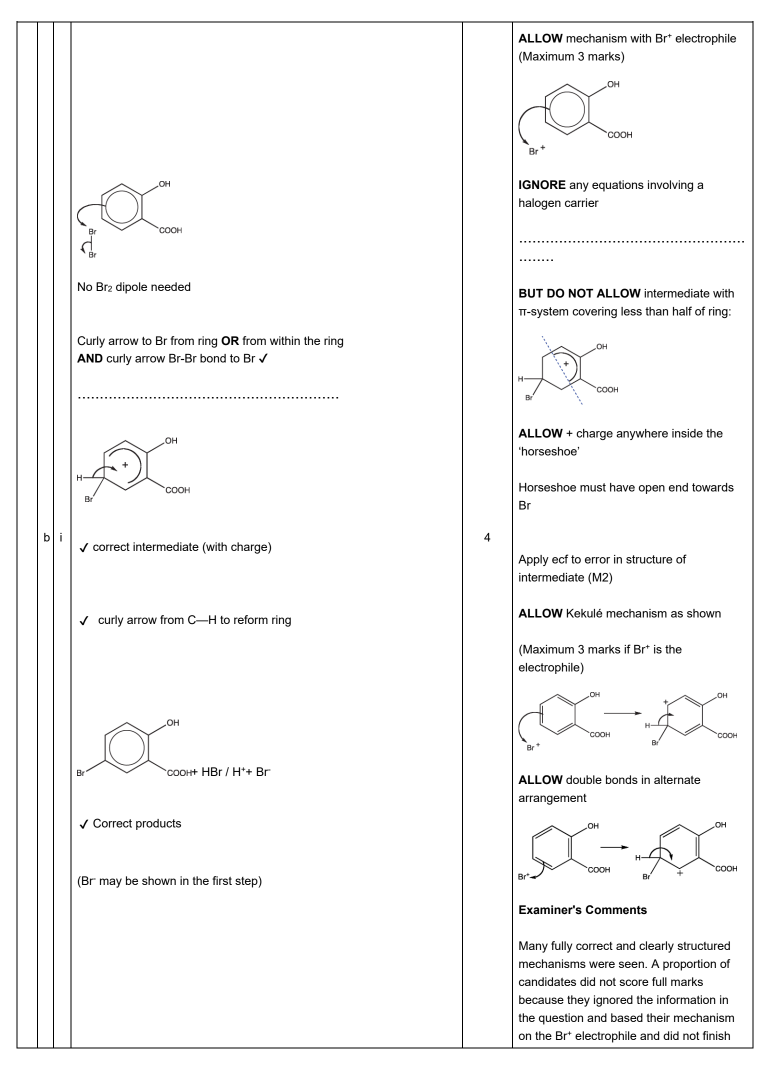
| Image: Structure B / delocalized spelled correctly and used in correct context       IGNORE charge density         Image: Structure B / delocalized spelled correctly and used in correct context       Examiner's Comments         Image: Structure B / delocalized spelled correctly and used in correct context       Examiner's Comments         Image: Structure B / delocalized spelled correctly and used in correct context       Examiner's Comments         Image: Structure B / delocalized sectors in structure B and p-otbal overlap were the most accessible marking points.       ALLOW structure B in less reactive         Image: Structure B / delocalised structure is (more) stable ✓       ALLOW structure B is less reactive for the reaction for benzene is less (negative) than 3 × (-)119         Image: Structure B is a better because (enthality change of hydrogenation for benzene is less (exothermic)) than (-) 357 (kJ mol <sup>-1</sup> ) ✓       ALLOW structure B is more stable by 149 kJ mol <sup>-1</sup> (2 marks) DO NOT ALLOW more / less energy needed for the reaction Answer must feet to data given in the question at must be a comparison (DNORE 360 kJ mol <sup>-1</sup> ) No marks can be awarded if structure A is salected         Image: Structure B is a better because (exothermic) than (-) 357 (kJ mol <sup>-1</sup> ) ✓       Ximple the data given in the question to compare the enthalpy change of hydrogenation for benzene is (exothermic) than (-) 357 (kJ mol <sup>-1</sup> ) ✓         Image: Structure B is a better because (exothermic) than (-) 357 (kJ mol <sup>-1</sup> ) ✓       Ximple the data given in the question to compare the enthalpy change is introduce A is salected         Image: Structure B is a better because (exothermic) than (-) 357 (  | <b>OR</b> correct position of p-orbital overlap  ✓  |   |  |
|---|---|---|--|
| Image: structure B / delocalised spelled correctly and used in correct context       DO NOT ALLOW electronegativity         Structures do not need to be labelled A and B if the description matches the structure       Examine's Comments         Many excellent answers with clear diagrams and explanations were seen. Delocalised electrons in structure B and portal overlap were the most accessible marking points.       ALLOW structure B is low in energy tences with clear diagrams and explanations were seen. Delocalised electrones in structure B and portal overlap were the most accessible marking points.         Image: structure B / delocalised structure is (more) stable ✓       ALLOW structure B is low in energy tences is less (exothermic) than (-) 357 (kl mor')         Image: structure B is a batter because (enthalpy change of hydrogenation for benzene is) less (exothermic) than (-) 357 (kl mor'). ✓       ALLOW enthalpy change is less than 3x enthalpy change is less than 3x enthalpy change for cyclohexene         2       Examiner's Comments       Any used the data given in the question the reaction for benzene is) less (exothermic) than (-) 357 (kl mor'). ✓         Image: structure B is a batter because (enthalpy change of hydrogenation for benzene is) less (exothermic) than (-) 357 (kl mor'). ✓       Yes and the data given in the question to comparison is low of structure A is selected         Image: structure B is a batter because (enthalpy change is heat structure B is batter the case. Some candidates contuker B to treatively the candidates conture B but relatively the candi  |   |   | IGNORE charge density  |
| Image: select context       Prequires delocalised / delocalized spelled correctly and used in correct context       Structures do not need to be labelled A and B if the description matches the structure         Examiner's Comments       Many excellent answers with clear diagrams and explanations were seen. Delocalised electrons in structure B and the data given in the analy change / hydrogenation       Delocalised electrons in structure B and portional overlap were the mast accessible marking points.         Image: Structure B is low in energy (IMOME structure B is low in an energy (IMOME structure B is low in energy (IMOME structure B is more stable by 149 kJ mol <sup>-1</sup> (2 marks)         Image: structure B is a better because (enthistly change of hydrogenation for benzene is) less (exothermic) than (~) 357 (kJ mol <sup>-1</sup> ) ✓       2         Image: structure B is a better because (enthistly change of hydrogenation for benzene is) less (exothermic) than (~) 357 (kJ mol <sup>-1</sup> ) ✓       2         Image: structure B is better because (enthistly change of hydrogenation for benzene is) less (exothermic) than (~) 357 (kJ mol <sup>-1</sup> ) ✓       2         Image: structure B is better because (is more stable by otherame to be mark for stating that structure B is better representation of benzene because (is more stable. Some candidates scored the other mark for stating that structure B is better increased is low more table. Some candidates confused is contused is comareased is more exothermic that structure A  | <i>₽</i> QWC  |   |  |
| correct context       Structures do not need to be labelled A and B if the description matches the structure         and B if the description matches the structure       Examiner's Comments         Many excellent answers with clear diagrams and explanations were seen. Delocalised electrons in structure B and p-orbital coverap were the most accessible marking points.       ALLOW structure B is low in energy (GNORE structure B is low in energy (Cover) (STALLOW entrue) (STALLOW enterue) (STALLOW entrue) (STALLOW entrue) (STALLOW entru   |   |   | DO NOT ALLOW electronegativity   |
| Image: Structure B / delocalised structure is (more) stable ✓       ALLOW structure B is a better because (enthalpy change of hydrogenation for benzene to data given in the question and must be a comparison 1QMORE 360 kL mol <sup>-1</sup> No marks can be awarded if structure A is selected         Image: Imag |   |   | and B if the description matches the   |
| ii       diagrams and explanations were seen.<br>Delocalised electrons in structure B and p-<br>orbital overlap were the most accessible<br>marking points.         ALLOW structure B is low in energy<br>IGNORE structure B is loss in energy<br>IGNORE structure B is less reactive         ALLOW enthalpy change / hydrogenation<br>for benzene is less (negative) than 3 ×<br>(-)119         IGNORE more positive than (-)357 kJ<br>mol <sup>-1</sup> ALLOW enthalpy change is less than 3x<br>enthalpy change for cyclohexene<br>ALLOW structure B is more stable by 149<br>kJ mol <sup>-1</sup> (2 marks)<br>DO NOT ALLOW more / less energy<br>needed for the reaction<br>Answer must feer to data given in the<br>question and must be a comparison<br>IGNORE 350 kJ mol <sup>-1</sup><br>No marks can be awarded if structure A is<br>selected         Iii       structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)<br>than (-) 357 (kJ mol <sup>-1</sup> ) ✓         Iii       structure B is a better because<br>(enthalpy change of hydrogenation for benzene is is) less (exothermic)<br>than (-) 357 (kJ mol <sup>-1</sup> ) ✓  |   |   | Examiner's Comments  |
| Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)<br>than (-) 357 (kJ mol <sup>-1</sup> ) √       Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)         Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)       Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)         Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)       Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)         Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)       Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)         Image: Structure B is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic)       Image: Structure B bit relatively<br>for candidates scored the other mark for<br>stating that structure B bit relatively<br>for stating that structure B bit relatively<br>for stating that structure B bit relatively<br>for stating that structure B is better<br>representation of benzene because it is<br>more stable. Some candidates confused<br>cyclohexene with structure A and<br>incorrectly concluded that the<br>hydrogenation of benzene is more<br>exothermic than structure A. Some<br>incorrectly referred to enthalpy of hydrogenation  |   |   | diagrams and explanations were seen.<br>Delocalised electrons in structure <b>B</b> and p-<br>orbital overlap were the most accessible   |
| ii       structure B / delocalised structure is (more) stable ✓       aLLOW enthalpy change is less than 3x enthalpy change is less than 3x enthalpy change for cyclohexene         iii       structure B / delocalised structure is (more) stable ✓       DO NOT ALLOW more / less energy needed for the reaction         Answer must refer to data given in the question and must be a comparison IGNORE 360 kJ mol <sup>-1</sup> No marks can be awarded if structure A is selected         2       2         Iii       structure B is a better because (enthalpy change of hydrogenation for benzene is) less (exothermic) than (-) 357 (kJ mol <sup>-1</sup> ) ✓         1ii       structure B is a better because in the question and must be a comparison IGNORE 360 kJ mol <sup>-1</sup> 1ii       structure B is a better because         (enthalpy change of hydrogenation for benzene is) less (exothermic) than (-) 357 (kJ mol <sup>-1</sup> ) ✓         2       Examiner's Comments         Many used the data given in the question to compare the enthalpy of hydrogenation of structure B but relatively few candidates scored the other mark for stating that structure B but relatively few candidates scored the other mark for stating that structure A and incorrectly concluded that the hydrogenation of benzene is nore exothermic than structure A. Some incorrectly referred to enthalpy of hydration  |   |   |  |
| reaction. Others referred to data they had<br>memorised rather than using the<br>information in the question as instructed.   | <b>structure B</b> is a better because<br>(enthalpy change of hydrogenation for benzene is) less (exothermic) | 2 | ALLOW enthalpy change / hydrogenation<br>for benzene is less (negative) than 3 ×<br>(-)119<br>IGNORE more positive than (-)357 kJ<br>mol <sup>-1</sup><br>ALLOW enthalpy change is less than 3x<br>enthalpy change for cyclohexene<br>ALLOW structure <b>B</b> is more stable by 149<br>kJ mol <sup>-1</sup> (2 marks)<br>DO NOT ALLOW more / less energy<br>needed for the reaction<br>Answer must refer to data given in the<br>question and must be a comparison<br>IGNORE 360 kJ mol <sup>-1</sup><br>No marks can be awarded if structure <b>A</b> is<br>selected<br>Examiner's Comments<br>Many used the data given in the question<br>to compare the enthalpy of hydrogenation<br>of structure <b>A</b> and structure <b>B</b> but relatively<br>few candidates scored the other mark for<br>stating that structure <b>B</b> is better<br>representation of benzene because it is<br>more stable. Some candidates confused<br>cyclohexene with structure <b>A</b> and<br>incorrectly concluded that the<br>hydrogenation of benzene is more<br>exothermic than structure <b>A</b> . Some<br>incorrectly referred to enthalpy of hydration<br>or stated that energy is required for the<br>reaction. Others referred to data they had<br>memorised rather than using the |



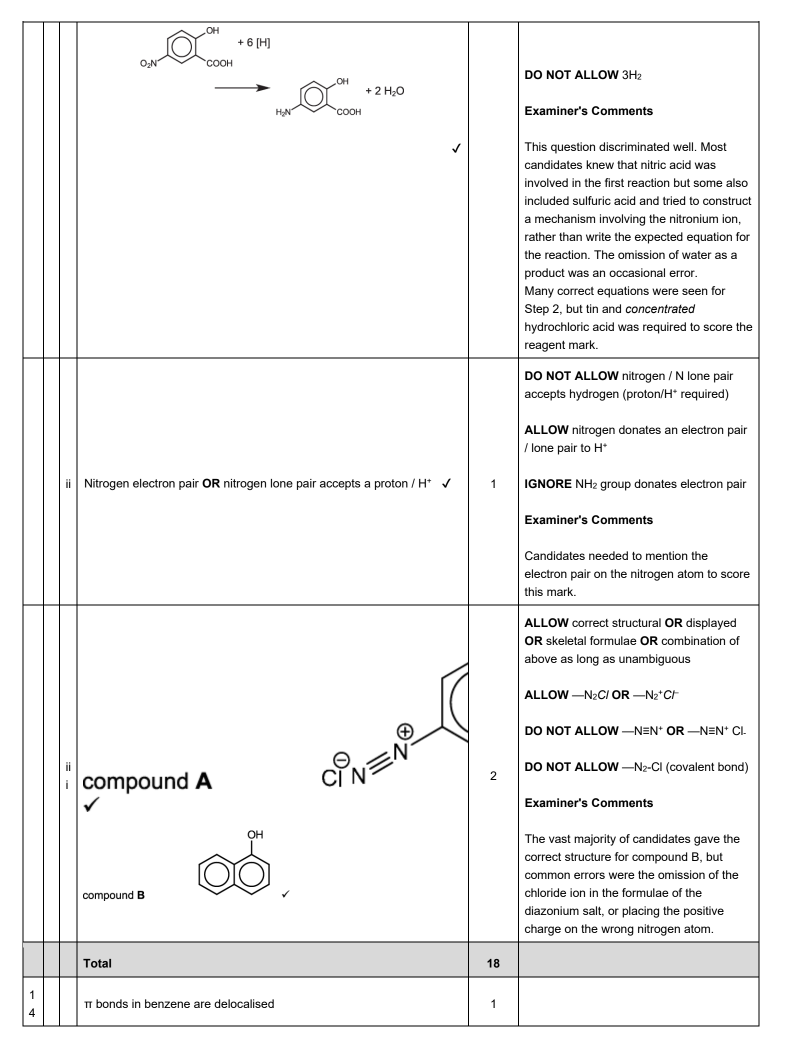
|  |         |   |   | familiar to candidates. The reagents for the<br>first reaction were very well known and<br>although the conditions for the reaction<br>were often quoted they were not required<br>for the mark. The correct structure of the<br>second reactant was rarely seen with the<br>most common incorrect responses being<br>based on cyclic structures. Most<br>candidates were able to deduce the<br>structure of the third reactant.  |
|--|---------|---|---|---|
|  | ii      | <b>FIRST CHECK THE ANSWER ON THE ANSWER LINE</b><br><b>IF</b> answer = 1.35 (g) award 3 marks<br><b>IF</b> answer = 0.54 (g) award 2 marks (no scale-up)<br><b>IF</b> answer = 0.216 (g) award 2 marks (incorrect scale-up)<br>$n(\text{compound D}) = 1.73/346 = 0.00500 \text{ mol } \checkmark$<br>$n(1,3-\text{diaminobenzene}) \text{ required } = 100/40 \times 0.005$<br>$= 0.0125 \text{ mol } \checkmark$<br>Molar mass of 1,3-diaminobenzene = 108 (g mol <sup>-1</sup> )<br><b>AND</b><br>Mass of 1,3-diaminobenzene = (108)(0.0125) = 1.35 g $\checkmark$ | 3 | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES ETC<br>If there is an alternative answer, check<br>to see if there is any ECF credit<br>possible<br>ALLOW ECF from incorrect amount,<br>scale-up or molar mass<br>Alternative 1<br>n(compound D) = 1.73/346 = 0.00500 mol<br>Molar mass of 1,3-diaminobenzene = 108<br>(g mol <sup>-1</sup> )<br>AND<br>Mass of 1,3-diaminobenzene =<br>(0.00500)(108) = 0.540 g<br>Mass of 1,3-diaminobenzene required =<br>(0.540)(100/40) = 1.35 g<br>Alternative 2<br>346 g gives 108 g<br>1.73 g gives 108/364 × 1.73 = 0.54 g<br>$0.54/40 \times 100 = 1.35 g$<br>Examiner's Comments<br>Candidates had been well prepared for<br>this type of percentage yield question and<br>it was very well answered with almost all<br>candidates scoring marks here and a high<br>proportion gaining all three marks for the<br>calculation. |
|  |         | (compound D has) <b>two</b> chiral centres <b>√</b>   |   | ALLOW (Compound D) has two<br>asymmetric carbons OR has two<br>stereocentres  |
|  | ii<br>i | Four optical isomers exist ✔  | 3 | <b>ALLOW</b> four enantiomers <b>OR</b> two pairs of enantiomers  |
|  |         | (Synthesis could) use enzymes <b>OR</b> bacteria<br><b>OR</b> use (chemical) chiral synthesis <b>OR</b> chiral catalysts  |   | INDEPENDENT MARK<br>ALLOW biological catalysts<br>ALLOW chiral transition metal complex /   |

|     |     | OR use natural chiral molecules OR single isomers (as starting materials) ✓               |    | catalyst<br>OR stereoselective transition metal<br>complex / catalyst<br>ALLOW 'chiral pool' / chiral auxiliary<br>Examiner's Comments<br>Two chiral centres and four optical isomers<br>was required in the mark scheme and less<br>specific answers did not score the first two<br>marks. Two pairs of enantiomers was an<br>accurate description worthy of the mark<br>and a reference to there being four<br>enantiomers was also given credit. The<br>majority of candidates scored only the third<br>mark with their suggestion of how to<br>improve the synthesis. Some candidates<br>missed the point here and instead<br>described techniques required to separate<br>the optical isomers. |
|-----|-----|---|----|---|
|     |     | Total   | 16 |   |
| 1 3 | a i | $O_{COONa}^{O_{Na}}$  | 1  | ALLOW correct structural OR displayed<br>OR skeletal formulae OR combination of<br>above as long as unambiguous<br>DO NOT ALLOW —O—Na OR -COO-Na<br>(covalent bond)<br>ALLOW —O <sup>-</sup><br>ALLOW —ONA ALLOW —COONA OR<br>ALLOW delocalised carboxylate<br>   |
|     |     |   |    | reaction with excess sodium hydroxide.<br>Many answers included the product<br>formed by the reaction of just one of the<br>functional groups. Most commonly the<br>phenol group was left unreacted. The mark<br>scheme permitted the omission of the<br>cation from the formula of the compound<br>but this omission was rarely seen.  |
|     | ii  | (Bromine) would be decolourised / turn (from orange / red / yellow / brown) to colourless | 1  | IGNORE goes clear   |

|  |         | OR white precipitate / solid / emulsion (formed) ✓  |   | DO NOT ALLOW other colours for<br>bromine<br>IGNORE cream precipitate<br>DO NOT ALLOW salicylic acid turns<br>colourless / decolourised<br>IGNORE temperature / fumes<br>Examiner's Comments<br>The observation for the reaction of a<br>phenol with bromine was very well known   |
|--|---------|---|---|--|
|  |         |   |   | and many candidates offered two correct<br>observations when only one was required<br>to score the mark.   |
|  | ii<br>i | $\bigcup_{COOH} + Br_2 \rightarrow \bigcup_{Br} + HBr$  | 1 | ALLOW correct structural OR displayed<br>OR skeletal formulae OR combination of<br>above as long as unambiguous<br>MUST be all correct to score mark<br>ALLOW molecular formulae, i.e.<br>$C_7H_6O_3 + Br_2 \rightarrow C_7H_5O_3Br^+ HBr$<br>Examiner's Comments<br>A very well answered question. Most<br>candidates copied the structural formulae<br>given in the question. Some made errors<br>when they unnecessarily converted the<br>structures into molecular formulae. HBr   |
|  | i<br>v  | (CH <sub>3</sub> ) <sub>2</sub> CHOH / CH <sub>3</sub> CH(OH)CH <sub>3</sub> / propan(–)2(–)ol<br>AND acid / H <sup>+</sup> / H <sub>2</sub> SO <sub>4</sub> (catalyst) ✓ | 1 | <ul> <li>was occasionally missing as a product.</li> <li>ALLOW correct structural OR displayed<br/>OR skeletal formulae OR combination of<br/>above as long as unambiguous</li> <li>ALLOW 2-propanol</li> <li>DO NOT ACCEPT incorrect name or<br/>incorrect formula of alcohol</li> <li>IGNORE reflux / concentrated (acid)</li> <li>Examiner's Comments</li> <li>Many candidates correctly gave the<br/>formula for propan-2-ol and included an<br/>acid catalyst. Common non-scoring<br/>answers omitted the acid or the alcohol or<br/>gave an incorrect name for the alcohol.</li> </ul> |

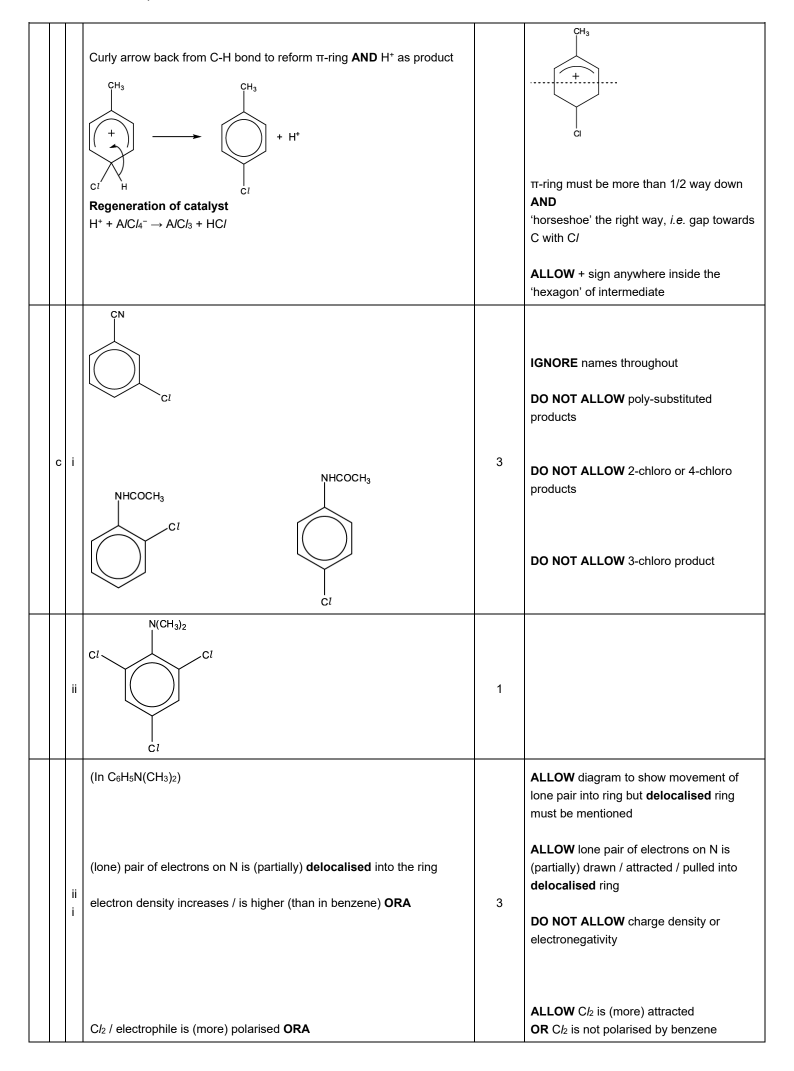


|     |    |  |   | with HBr as a product. Relatively few candidates lost marks for incorrectly positioned curly arrows.   |
|-----|----|--|---|--|
|     |    |  |   | ALLOW diagram to show movement of<br>lone pair into ring but delocalised ring must<br>be mentioned   |
|     |    | (In salicylic acid)  |   | ALLOW lone pair / pair of electrons on<br>O(H) / phenol is (partially) drawn /<br>attracted / pulled into <b>delocalised</b> ring  |
|     |    |  |   | IGNORE 'activates the ring'  |
|     |    | lone pair / pair of electrons on O(H) / phenol is $\backsim$ (partially) <b>delocalised</b> into the ring $\checkmark$ |   | ALLOW more electron rich   |
|     |    | electron density increases / is high <b>ORA √</b>  |   | <b>DO NOT ALLOW</b> charge density or electronegativity  |
| ii  | ii |  | 3 | ALLOW (salicylic acid) attracts<br>electrophiles more/more susceptible to<br>electrophilic attack  |
|     |    | Br₂ / electrophile is (more) polarised <b>ORA √</b>  |   | ALLOW Br <sub>2</sub> is (more) attracted OR Br <sub>2</sub> is<br>not polarised by benzene<br>OR induces dipoles (in bromine /<br>electrophile)   |
|     |    |  |   | Delocalise(d) needed to score the first marking point  |
|     |    | <b>QWC</b> : delocalised / delocalized / delocalise <i>etc</i> .   |   | Examiner's Comments  |
|     |    | must be spelled correctly in the correct context at least once   |   | This question was very well answered with<br>the majority of candidates scoring at least<br>two marks. The most common errors were<br>the omitting the words delocalised or lone<br>pair or failure to use the word delocalised<br>in the correct context. |
|     |    |  |   | ALLOW reagent mark if HNO₃ in equation   |
|     |    | Step 1<br>Add HNO <sub>3</sub> $\checkmark$  |   | IGNORE H <sub>2</sub> SO <sub>4</sub> (NOTE: H <sub>2</sub> SO <sub>4</sub> not required with phenols)   |
| c i | i  | $OH + HNO_3 \longrightarrow O_2N + OH + HNO_3 + O_2N + OH + O$                           | 4 | <b>IGNORE</b> concentrations of acids / temperature  |
|     |    | √<br>Step 2  |   | ALLOW correct structural OR displayed<br>OR skeletal formulae OR combination of<br>above as long as unambiguous  |
|     |    | Tin AND concentrated HC/ ✓   |   |  |
|     |    |  |   | Equations <b>MUST</b> be completely correct for <b>one</b> mark each   |



### 6.1.1 Aromatic Compounds

|        |   | Total  | 1 |  |
|--------|---|--|---|--|
| 1<br>5 |   | Electrophilic substitution means benzene ring<br>Electrophilic addition means alkene / C=C<br>Isomer of C <sub>9</sub> H <sub>8</sub> O <sub>2</sub> containing<br>C=C, benzene ring <b>AND</b> COOH<br>Correct isomer:<br>$\int_{0}^{00H} \int_{0}^{0} $ | 5 | Concluded using data provided and conclusions from 1st two marks.<br>ALLOW 1 mark for:<br>COOH<br>OR<br>(does not gain final justification mark) |
|        |   | Total  | 5 |  |
| 1<br>6 | а | <ul> <li>Experimental evidence – ANY TWO from</li> <li>carbon–carbon bond lengths are the same in benzene</li> <li>Enthalpy change of hydrogenation is less (exothermic) for benzene (than for Kekulé model)</li> <li>Discussion of named reaction to highlight greater stability, e.g. chlorination of benzene requires a catalyst whereas no catalyst is needed for alkenes</li> <li>Bonding in modern model</li> <li>p-orbitals overlap to form π bonds</li> <li>(π–)electrons are delocalised</li> </ul>   | 4 | ALLOW both marks for correctly labelled diagrams showing overlap of p-orbitals to form delocalised π–electrons                                   |
|        | b | Generation of electrophile<br>$A/Cl_3 + Cl_2 \rightarrow A/Cl_4^- + Cl^+$<br>Electrophilic substitution<br>Curly arrow from $\pi$ -bond to Cl <sup>+</sup><br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$<br>$\downarrow$   | 5 | ANNOTATE ANSWER WITH TICKS AND<br>CROSSES<br>DO NOT ALLOW the following<br>intermediate:   |



|        |    |  |    | <b>OR</b> induces dipoles (in chlorine / electrophile)   |
|--------|----|--|----|--|
|        |    | Total  | 16 |  |
| 1<br>7 | i  | HO OH HO OH  | 1  | ALLOW correct structural OR displayed<br>OR skeletal formulae OR a combination of<br>above as long as unambiguous<br>ALLOW disubstituted compound with <i>tert</i> -<br>butyl groups adjacent  |
|        | ii | (The student's friend is correct because)<br>the lone pair of electrons on the oxygen atom(s)<br>is donated to / partially delocalised into the π system<br>making quinol more susceptible to electrophilic attack | 3  | ALLOW "the oxygen p-orbital overlaps<br>with"<br>ALLOW diagrammatic answer for 1st and<br>2nd marks:<br>1st mark: π system OR 6 × p orbitals<br>shown<br>2nd mark: O lone pair OR O p-orbital AND<br>interaction<br>ALLOW undergoes electrophilic<br>substitution more easily<br>if 1st and 2nd marks achieved through<br>diagram, conclusion <b>must</b> refer to diagram<br>for 3rd mark |
|        |    | Total  | 4  |  |