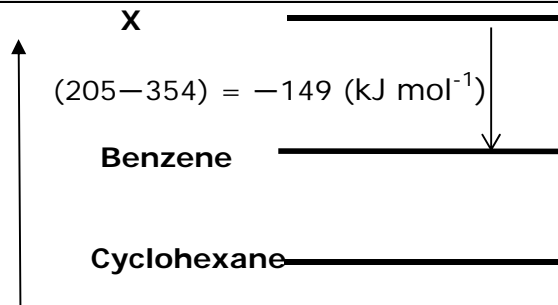
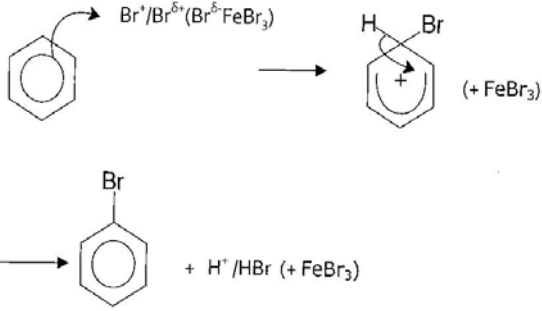
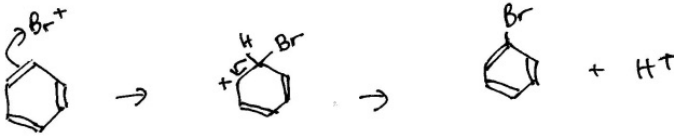


Question Number	Acceptable Answers	Reject	Mark
<b>1(a)</b>	All carbon to carbon bonds same length/ longer C-C and shorter C=C not present  IGNORE Just "benzene has a delocalised ring" Benzene does not have C=C double bonds Any references to shape/ bond angles		1

Question Number	Acceptable Answers	Reject	Mark
<b>1(b)(i)</b>	$(3 \times -118) = -354 \text{ (kJ mol}^{-1}\text{)}$		1

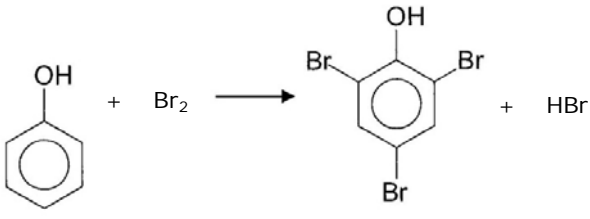
Question Number	Acceptable Answers	Reject	Mark
<b>1(b)(ii)</b>	 <p><b>First mark</b> Relative levels with names or formulae (1)</p> <p><b>Second mark</b> Value <math>-149 \text{ (kJ mol}^{-1}\text{)}</math> + arrow in correct direction ALLOW double-headed arrow (1)</p> <p>TE from value in (b)(ii) IGNORE <math>3\text{H}_2</math> if shown / cyclohexene / other arrows/values</p>	Diagram inverted scores 0          +149	2

Question Number	Acceptable Answers	Reject	Mark
<b>1(b)* (iii)</b>	The p/pi-/π/6 electrons (of carbon) (1) are delocalised in benzene (but not in X) (1)		2

Question Number	Acceptable Answers	Reject	Mark
1(c)	<p><b>First mark:</b>  <math>\text{FeBr}_3 + \text{Br}_2 \rightarrow \text{FeBr}_4^- + \text{Br}^+</math>  OR  <math>\text{Br}-\text{Br} + \text{FeBr}_3 \rightarrow \text{Br}^{\delta+} \dots \text{Br}^{\delta-}\text{FeBr}_3</math> (1)  Ignore state symbols even if wrong</p> <p><b>Second, third and fourth marks:</b>  <b>Either</b></p>  <p>Arrow from benzene ring electrons (from <b>inside</b> the hexagon) to <b>Br<sup>+</sup> / Br<sup>δ+</sup></b> (..... Br<sup>δ-</sup>FeBr<sub>3</sub>) (1)</p> <p>Correctly drawn intermediate with delocalisation covering at least three carbon atoms, but not the carbon atom bonded to the bromine, with the positive charge shown inside the horseshoe</p> <p>The bonds to H and Br may be dotted (1)</p> <p>Arrow from / close to C-H <b>bond</b> to inside the hexagon <b>and</b> H<sup>+</sup> / HBr as product (1)</p> <p><b>OR</b></p>  <p>Use of Kekulé structure for benzene and intermediate with arrow from C=C double bond to <b>Br<sup>+</sup> / Br<sup>δ+</sup></b> (..... Br<sup>δ-</sup>FeBr<sub>3</sub>) (1)</p> <p>Correctly drawn intermediate with + charge on the C atom next to the C bonded to H and Br</p>	Gap in wrong place	4

	The bonds to H and Br may be dotted (1)		
	Arrow from / close to C-H <b>bond</b> to bond beside + charged C <b>and</b> H <sup>+</sup> / HBr as product (1)		
	Each marking point is independent		

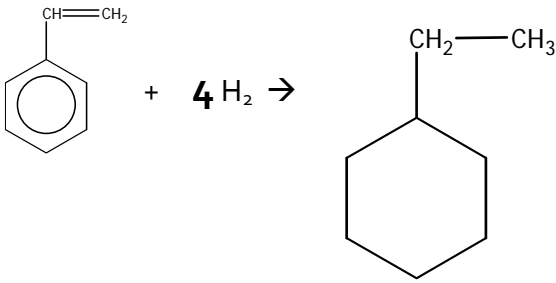
Question Number	Acceptable Answers	Reject	Mark
<b>1(d)(i)</b>	Bromine goes colourless OR It/the mixture goes from brown to colourless  ALLOW Red-brown/ Orange/ yellow/ combinations of these colours  Bromine is decolorised (1)  White precipitate/solid forms / Steamy fumes (1)  IGNORE Antiseptic smell Gets hot	Goes clear   Red to colourless  Bromine is discoloured  Effervescence	2

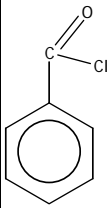
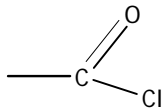
Question Number	Acceptable Answers	Reject	Mark
<b>1(d)(ii)</b>	 <p>Organic product with structure shown (1)            Rest of equation correct            ALLOW            C<sub>6</sub>H<sub>5</sub>OH or Kekule for phenol (1)             C<sub>6</sub>H<sub>5</sub>OH + 3Br<sub>2</sub> → C<sub>6</sub>H<sub>2</sub>Br<sub>3</sub>OH + 3HBr            Scores MP2 only            Substitution of 1Br or 2Br in any position in balanced equation scores MP2 only.</p>		2

Question Number	Acceptable Answers	Reject	Mark
* 1 (d) (iii)	<p><b>Lone pair of electrons on oxygen</b> (may be shown on a diagram)  <b>and</b>            EITHER            overlaps with pi cloud            OR            Feeds into / donates into / interacts with benzene ring            (1)</p> <p>Activating benzene ring / increasing electron density of ring / making attack by <b>electrophiles</b> easier (1)</p> <p>COMMENT            'Lone pair of electrons on oxygen increases electron density of ring' scores (2)</p> <p>ALLOW            benzene becomes a better nucleophile for MP2</p>	<p>OH group overlaps</p> <p>Just 'making it more reactive'.</p>	2

Question Number	Acceptable Answers	Reject	Mark
<b>2(a)(i)</b>	Addition / reduction / free-radical addition  IGNORE references to 'hydrogenation'	'redox' 'electrophilic addition' 'nucleophilic addition'	<b>1</b>

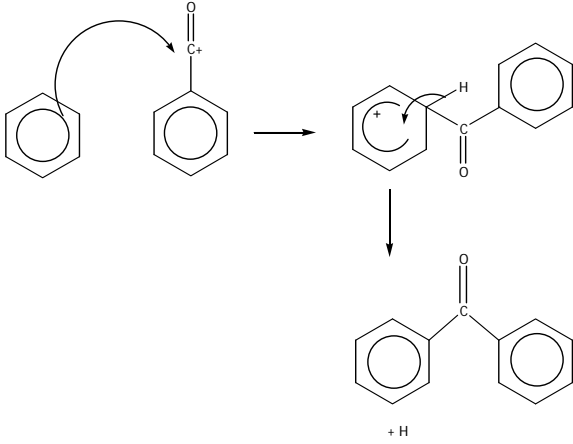
Question Number	Acceptable Answers	Reject	Mark
<b>2(a)Iii)</b>	<p><b>mark:</b> <b>Delocalization</b> (of <math>\pi/p</math> electrons in benzene ring) <b>(1)</b></p> <p>IGNORE reference to 'resonance'</p> <p><b>Second mark:</b> Results in more <b>energy</b> needed to <b>break</b> the <b>bonds</b> in benzene (compared with three separate <math>\pi</math> bonds) <b>(1)</b></p> <p>ALLOW confers <b>stability</b> on the molecule / makes benzene <b>more stable</b> (than expected)</p> <p>IGNORE Reference to carbon-carbon bond lengths Values of any enthalpy changes</p> <p>Mark the two points independently</p>		<b>2</b>

Question Number	Acceptable Answers	Reject	Mark
<b>2(a)(iii)</b>	 <p style="text-align: center;">( <math>\Delta H =</math> ) <b>— 328</b> (kJ mol<sup>-1</sup>)</p> <p><b>First mark:</b> For "4"</p> <p><b>Second mark:</b> Product as above / correct skeletal formula of product</p> <p>ALLOW Side chain written as —C<sub>2</sub>H<sub>5</sub></p> <p><b>Third mark:</b> —328 (kJ mol<sup>-1</sup>)</p> <p><b>NOTE</b></p> <p><b>One</b> H<sub>2</sub> added showing a CQ correct product with only side chain reduced and cq <math>\Delta H = -120</math> (kJ mol<sup>-1</sup>) scores <b>(2)</b></p> <p><b>Three</b> H<sub>2</sub> added showing a CQ correct product with only the benzene ring reduced and cq <math>\Delta H = -208</math> (kJ mol<sup>-1</sup>) scores <b>(2)</b></p> <p><b>Five</b> H<sub>2</sub> added with fully correct product drawn and <math>\Delta H = -448</math> (kJ mol<sup>-1</sup>) scores <b>(2)</b></p> <p><b>Three and a half</b> H<sub>2</sub> added showing a fully correct product and <math>\Delta H = -268/-293(.3)</math> (kJ mol<sup>-1</sup>) scores <b>(2)</b></p> <p><b>NOTE</b> Mark scoring points independently</p>		<b>3</b>

Question Number	Acceptable Answers	Reject	Mark
<b>2(b)(i)</b>	 <p>Mark awarded for displaying</p> 		<b>1</b>

Question Number	Acceptable Answers	Reject	Mark
<b>2(b)(ii)</b>	<p>Electrophilic substitution</p> <p><b>BOTH</b> words needed</p> <p>IGNORE references to 'acylation' and /or 'Friedel-Crafts'</p>		<b>1</b>

Question Number	Acceptable Answers	Reject	Mark
<b>2(b)(iii)</b>	<p>Friedel <b>and</b> Crafts</p> <p><b>BOTH</b> names are needed for this mark</p>		<b>1</b>

Question Number	Acceptable Answers	Reject	Mark
2(b)(iv)	<p><b>First mark:</b>  <math display="block">\text{C}_6\text{H}_5\text{COCl} + \text{AlCl}_3 \rightarrow \text{C}_6\text{H}_5\text{CO}^+ + \text{AlCl}_4^- \quad (1)</math></p> <p>+ can be anywhere on the C<sub>6</sub>H<sub>5</sub>CO in the equation for the first mark</p>  <p style="text-align: center;">(AlCl<sub>4</sub><sup>-</sup> + H<sup>+</sup> → HCl + AlCl<sub>3</sub>)</p> <p><b>NOTE:</b>  If ethanoyl chloride or any other acid chloride or the generic RCOCl is used <b>instead</b> of benzoyl chloride, no first mark can be awarded but the 2nd, 3rd and 4th marks can be awarded consequentially</p> <p><b>Second mark:</b> First curly arrow, as shown, to start from inside the hexagon to the correct C<sup>+</sup> carbon (i.e. not to the benzene ring)  Note the + must be on the C of the C=O/CO for this mark <span style="float: right;">(1)</span></p> <p><b>Third mark:</b> Intermediate correctly drawn <span style="float: right;">(1)</span></p> <p><b>NOTE</b>  + ca be shown anywhere in the ring or at the C atom where electrophile is bonded.  The 'horseshoe' in the intermediate to cover at least three carbon atoms</p> <p><b>Fourth mark:</b> Second curly arrow as shown from C—H bond to reform the ring, not from the H atom in this bond <span style="float: right;">(1)</span></p> <p><b>NOTE</b>  Products do not have to be shown nor the equation for regeneration of the catalyst given</p>		4



Question Number	Acceptable Answers	Reject	Mark
<b>2(b)(v)</b>	Absorbs / reflects / blocks / protects from / shields against / <b>uv</b> (light/ radiation) IGNORE 'non-toxic' / references to IR	<b>adsorbs</b> uv light	<b>1</b>

Question Number	Acceptable Answers	Reject	Mark										
<b>2(c)(i)</b>	<p>Any <b>TWO</b> of the following</p> <p>(1) for identifying the bond by formula as shown and (1) for wavenumber in each matching pair</p> <p><b>UNITS</b> are not required</p> <table border="1"> <thead> <tr> <th>Bond</th> <th>Wavenu range/wavenumber (cm<sup>-1</sup>)</th> </tr> </thead> <tbody> <tr> <td>C=C</td> <td>1600 / 1580 / 1500 / 1450 <b>All four values needed</b></td> </tr> <tr> <td>C=O</td> <td>1700 – 1680</td> </tr> <tr> <td>C-</td> <td>3030</td> </tr> <tr> <td>C-</td> <td>750 / 700 <b>Both values needed</b></td> </tr> </tbody> </table> <p><b>NOTE</b> ALLOW Correct wavenumber range, or any number within the correct range, <b>for C=O</b></p> <p>Mark identification of the bond and the wavenumber independently (eg a correct bond with a wrong wavenumber, or vice-versa, scores one of the two marks in each case)</p> <p>IGNORE nmr values / chemical shifts</p>	Bond	Wavenu range/wavenumber (cm <sup>-1</sup> )	C=C	1600 / 1580 / 1500 / 1450 <b>All four values needed</b>	C=O	1700 – 1680	C-	3030	C-	750 / 700 <b>Both values needed</b>		<b>4</b>
Bond	Wavenu range/wavenumber (cm <sup>-1</sup> )												
C=C	1600 / 1580 / 1500 / 1450 <b>All four values needed</b>												
C=O	1700 – 1680												
C-	3030												
C-	750 / 700 <b>Both values needed</b>												

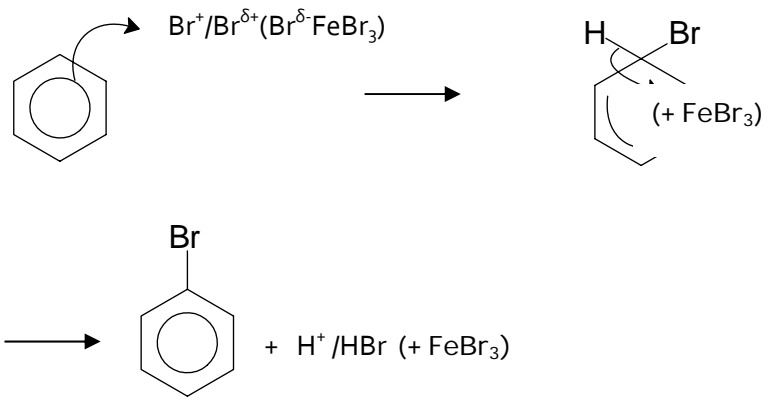
Question Number	Acceptable Answers	Reject	Mark
2(c)(ii)	<div style="text-align: center;"> </div> <p><b>First mark</b></p> <p><b>EITHER</b> Identifies correctly the <b>three</b> different proton environments</p> <p>ALLOW If the three different proton environments are only shown on one of the benzene rings</p> <p><b>NOTE</b> On right-hand ring, clockwise from C=O, positions 2, 3 and 4 And /or 2,4 and 5 are shown as different environments and /or On left-hand ring, anti-clockwise from C=O, positions 2, 3 and 4 And /or 2,4 and 5 are shown as different environments</p> <p><b>OR</b></p> <p>Identifies proton Z correctly on both benzene rings <b>(1)</b></p> <p><b>Second mark</b> Fully correct labelling both rings using the letters <b>X</b>, <b>Y</b> and <b>Z</b></p> <p><b>NOTE</b> <b>X</b> and <b>Y</b> labels are interchangeable, <b>Z</b> is not <b>(1)</b></p>		<b>2</b>

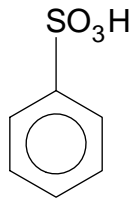
Question Number	Acceptable Answers	Reject	Mark
<b>3(a)(i)</b>	(3 x -120) = -360 (kJ mol <sup>-1</sup> )  IGNORE ΔH, and case of letters in units e.g allow KJ	No sign or + sign in answer, ie 360/+360  Any other wrong units  ΔE	<b>1</b>

Question Number	Acceptable Answers	Reject	Mark
<b>*3(a)(ii)</b>	<ul style="list-style-type: none"> <li>( Bonding in) benzene/it is more <b>stable</b> (than Kekule) by <b>152</b> kJ mol<sup>-1</sup> (consequential on (a)(i)) <b>(1)</b> IGNORE sign</li> <li><b>π /p/double bond electrons</b> are delocalized (around the ring)  OR six <b>p electrons</b> shared between six (ring) carbon atoms  OR delocalized because of overlap of <b>p orbitals</b>  OR resonance hybrid of C=C's and C-C's <b>(1)</b></li> <li><b>Substitution</b> reactions (rather than addition) <b>(1)</b></li> </ul> <p>NOTE: <b>Nucleophilic</b> substitution negates the substitution mark because it is wrong additional information</p> <ul style="list-style-type: none"> <li>Maintains/regains delocalized system OR maintains/regains stability OR maintains/regains stabilization energy <b>(1)</b></li> </ul>	Attack by electrophiles with no mention of substitution	<b>4</b>

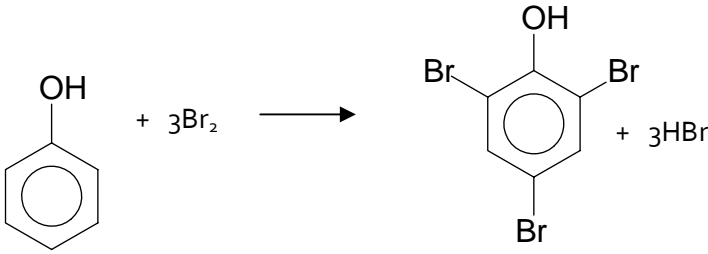
Question Number	Acceptable Answers	Reject	Mark
<b>3(b)(i)</b>	<p>Concentrated nitric acid/HNO<sub>3</sub> (1)</p> <p>Concentrated sulfuric acid/H<sub>2</sub>SO<sub>4</sub> (1)</p> <p>Allow conc or c. in place of 'concentrated'</p> <p>ALLOW Concentrated nitric acid and sulfuric acid</p> <p>OR</p> <p>Concentrated HNO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub> (2)</p> <p>Second mark depends on nitric acid</p> <p>Max. (1) if no mention of concentrated</p> <p>Nitric acid and concentrated sulfuric acid scores (1)</p> <p>NOTE: conc. HNO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub>(aq) scores (1) but conc. HNO<sub>3</sub> and conc H<sub>2</sub>SO<sub>4</sub>(aq) scores (2)</p>	Concentrated hydrochloric acid	<b>2</b>

Question Number	Acceptable Answers	Reject	Mark
<b>3(b)(ii)</b>	<p>Electrophile/electrophilic</p> <p>ALLOW Electrophyl(e)</p>	<p>Acid</p> <p>Base</p> <p>Oxidizing agent</p> <p>Reducing agent</p>	<b>1</b>

Question Number	Acceptable Answers	Reject	Mark
<b>3(b)(iii)</b>	<p> <math>\text{Br}_2 + \text{FeBr}_3 \rightarrow \text{FeBr}_4^- + \text{Br}^+</math>  OR  <math>\text{Br}-\text{Br} + \text{FeBr}_3 \rightarrow \text{Br}^{\delta+} \dots \text{Br}^{\delta-} \text{FeBr}_3</math> (1)  IGNORE state symbols even if wrong </p>  <p> Arrow from benzene ring electrons (from <b>inside</b> the hexagon) to <b>Br<sup>+</sup>/Br<sup>δ+</sup></b> (<math>\dots \text{Br}^{\delta-} \text{FeBr}_3</math>) (1) </p> <p> Correctly drawn intermediate with delocalization covering at least three carbon atoms, but not the carbon atom bonded to the bromine with the positive charge shown inside the hexagon </p> <p> The bonds to H and Br may be dotted (1) </p> <p> Arrow from or close to <b>bond</b> to H to centre of ring <b>and</b> H<sup>+</sup>/HBr as a product (1) </p> <p> ALLOW  Kekulé structure for benzene and intermediate </p> <p> <b>Each marking point is independent</b> </p>	lack of charges	<b>4</b>

Question Number	Acceptable Answers	Reject	Mark
<b>3(b)(iv)</b>	 <p>OR C<sub>6</sub>H<sub>5</sub>SO<sub>3</sub>H</p> <p>accept: displayed -SO<sub>3</sub>H</p> <p>-SO<sub>3</sub><sup>-</sup>H<sup>+</sup></p> <p>-SO<sub>2</sub>OH</p> <p>If two formulae are given both must be correct <b>(1)</b></p> <p>Penalise if bond <b>clearly</b> goes to O or H rather than S</p> <p><b>Benzenesulfonic acid</b> <b>(1)</b></p> <p>ALLOW <b>phenyl sulfonic acid</b></p>	Benzenesulfuric acid/benzosulfonic acid/benzylsulfonic acid	<b>2</b>

Question Number	Acceptable Answers	Reject	Mark
<b>3(c)(i)</b>	<p><b>Non-bonding/lone pair</b> electrons from <b>oxygen...</b> <b>(1)</b></p> <p>...are delocalized/incorporated/donated into the ring (electron system) (Could be shown in diagram)</p> <p>OR</p> <p>increases electron density on the ring <b>(1)</b></p> <p>makes it (the ring) more susceptible to electrophilic attack/makes it (the ring) a better nucleophile <b>(1)</b></p>	<p>...from methyl/methoxy</p> <p>Makes it more electronegative</p>	<b>3</b>

Question Number	Acceptable Answers	Reject	Mark
<b>3(c)(ii)</b>	<div style="text-align: center;">  </div> <p style="text-align: center;"> <span style="margin-right: 100px;"><b>(1)</b> organic formula</span> <span><b>(1)</b> balancing</span> </p> <p>ALLOW</p> <ul style="list-style-type: none"> <li>• Condensed structural formulae, for example  <math display="block">\text{C}_6\text{H}_5\text{OH} + 3\text{Br}_2 \rightarrow \text{C}_6\text{H}_2\text{Br}_3\text{OH} + 3\text{HBr}</math> <div style="text-align: center;"> <span style="margin-right: 100px;"><b>(1)</b></span> <span><b>(1)</b> balancing</span> </div> </li> <li>• multiples</li> <li>• substitution to any positions</li> </ul> <p>IGNORE: H<sub>2</sub>O Position of bond to OH</p> <p>NOTE: Correct balanced equations giving mono and disubstitution phenols score <b>1 mark</b></p>		<b>2</b>

Question Number	Acceptable Answers	Reject	Mark
<b>3(d)</b>	<p>(Chloromethyl)benzene/chloromethylbenzene/ chlorophenylmethane/ benzyl chloride OR dichloromethane <b>(1)</b></p> <p>ALLOW phenylchloromethane</p> <p>Aluminium chloride <b>(1)</b></p> <p>ACCEPT formulae eg <math>C_7H_7Cl</math>, <math>C_6H_5CH_2Cl</math>, <math>CH_2Cl_2</math>, <math>AlCl_3</math></p> <p>ACCEPT other halogen carriers eg <math>FeCl_3</math>/iron(<b>III</b>) chloride/<math>ZnCl_2</math></p> <p>ACCEPT bromine in place of chlorine for either/both marks</p> <p>Correct formula and wrong name or correct name and wrong formula or any other wrong additional information loses mark</p>	<p><math>CH_2Cl</math></p>	<b>2</b>