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

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

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|---|---------------------------------|------|
| 1(b)(ii) | X (205-354) = -149 (kJ mol ⁻¹) Benzene | Diagram inverted scores 0 | 2 |
| | Cyclohexane | | |
| | First mark Relative levels with names or formulae (1) | | |
| | Second mark Value -149 (kJ mol ⁻¹) + arrow in correct direction ALLOW double-headed arrow (1) | +149 | |
| | TE from value in (b)(ii) IGNORE 3H ₂ if shown / cyclohexene / other arrows/values | | |

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

| Question Number | Acceptable Answers | Reject | Mar k |
|--------------------|---|--------------------|----------|
| 1(c) | First mark: FeBr ₃ + Br ₂ → FeBr ₄ ⁻ + Br ⁺ OR Br-Br + FeBr ₃ → Br ^{δ+} Br ^{δ-} FeBr ₃ (1) Ignore state symbols even if wrong Second, third and fourth marks: Either Br'/Br ^{δ-} (Br ^{δ-} FeBr ₃) + H'/HBr (+FeBr ₃) Arrow from benzene ring electrons (from inside the hexagon) to Br ⁺ / Br ^{δ+} (Br ^{δ-} FeBr ₃) (1) 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 The bonds to H and Br may be dotted (1) Arrow from / close to C-H bond to inside the hexagon and H ⁺ / HBr as product (1) | Gap in wrong place | 4 |
| | Use of Kekulé structure for benzene and intermediate with arrow from C=C double bond to Br ⁺ / Br ^{δ+} (Br ^{δ-} FeBr ₃) (1) Correctly drawn intermediate with + charge on the C atom next to the C bonded to H and Br | ₹ | |

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

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

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|---|--------|------|
| 1(d)(ii) | Organic product with structure shown (1) Rest of equation correct ALLOW C_6H_5OH or Kekule for phenol (1) $C_6H_5OH + 3Br_2 \rightarrow C_6H_2Br_3OH + 3HBr$ Scores MP2 only Substitution of 1Br or 2Br in any position in balanced equation scores MP2 only. | | 2 |

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

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

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

| Question | Acceptable Answers | Reject | Mark |
|-------------------|---|--------|------|
| Number | | | |
| 2 (a)(iii) | CH=CH ₂ + 4 H ₂ → CH ₂ CH ₂ CH ₂ CH ₃ | | 3 |
| | $(\Delta H =)$ - 328 (kJ mol ⁻¹) | | |
| | First mark: For "4" | | |
| | Second mark: Product as above / correct skeletal formula of product | | |
| | ALLOW Side chain written as $-C_2H_5$ | | |
| | Third mark : —328 (kJ mol ⁻¹) | | |
| | NOTE | | |
| | One H ₂ added showing a CQ correct product with only side chain reduced and cq $\Delta H = -120$ (kJ mol ⁻¹) scores (2) | | |
| | Three H ₂ added showing a CQ correct product with only the benzene ring reduced and cq $\Delta H = -208$ (kJ mol ⁻¹) scores (2) | | |
| | Five H ₂ added with fully correct product drawn and $\Delta H = -448$ (kJ mol ⁻¹) scores (2) | | |
| | Three and a half H_2 added showing a fully correct product and $\Delta H = -268/-293(.3) (kJ \text{ mol}^{-1})$ scores (2) | | |
| | NOTE Mark scoring points independently | | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|-----------------------------|--------|------|
| 2 (b)(i) | Mark awarded for displaying | | 1 |

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

| Question | Acceptable Answers | Reject | Mark |
|-------------------|---------------------------------------|--------|------|
| Number | | | |
| 2 (b)(iii) | Friedel and Crafts | | 1 |
| | | | |
| | BOTH names are needed for this | | |
| | mark | | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|---|--------|------|
| 2 (b)(iv) | First mark: $C_6H_5COCI + AICI_3 \rightarrow C_6H_5CO^+ + AICI_4^-$ (1) | | 4 |
| | + can be anywhere on the C_6H_5CO in the equation for the first mark | | |
| | | | |
| | + H | | |
| | $(AICI_4^- + H^+ \rightarrow HCI + AICI_3)$ | | |
| | NOTE: If ethanoyl chloride or any other acid chloride or the generic RCOCI is used instead of benzoyl chloride, no first mark can be awarded but the 2nd, 3rd and 4th marks can be awarded consequentially | | |
| | Second mark: First curly arrow, as shown, to start from inside the hexagon to the correct C+ carbon (i.e. not to the benzene ring) Note the + must be on the C of the C=O/CO for this mark | | |
| | (1) Third mark: Intermediate correctly drawn | | |
| | NOTE (1) | | |
| | + 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 | | |
| | Fourth mark: Second curly arrow as shown from C— H bond to reform the ring, not from the H atom in this bond (1) | | |
| | NOTE Products do not have to be shown nor the equation for regeneration of the catalyst given | | |

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

| Question Number | Acceptable Answers | | Reject | Mark |
|--------------------|---------------------------|---------------------------|--------|------|
| 2 (c)(i) | Any TWO of the foll | owing | | 4 |
| | | | | |
| | | ne bond by formula as | | |
| | shown and (1) for w | avenumber in each | | |
| | matching pair | | | |
| | LINUTO | | | |
| | UNITS are not requ | irea | | |
| | Bond | Wavenu | | |
| | | range/wavenumber | | |
| | | (cm ⁻¹) | | |
| | C=C | 1600 / 1580 / 1500 / | | |
| | | 1450 | | |
| | | All four values | | |
| | | needed | | |
| | C=0 | 1700 – 1680 | | |
| | C- | 3030 | | |
| | C- | 750 / 700 | | |
| | | Both values needed | | |
| | NOTE | | | |
| | ALLOW | | | |
| | Correct wavenumbe | r range, or any number | | |
| | within the correct ra | inge, for C=O | | |
| | Mark identification of | of the bond and the | | |
| | wavenumber indepe | endently | | |
| | (eg a correct bond v | vith a wrong wavenumber, | | |
| | or vice-versa, score | s one of the two marks in | | |
| | each case) | | | |
| | IGNORE | | | |
| | nmr values / chemic | cal shifts | | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|--|--------|------|
| 2 (c)(ii) | Y X X Y Z Y Y First mark | | 2 |
| | EITHER Identifies correctly the three different proton environments ALLOW If the three different proton environments are only shown on one of the benzene rings | | |
| | NOTE 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 | | |
| | OR | | |
| | Identifies proton Z correctly on both benzene rings (1) | | |
| | Second mark Fully correct labelling both rings using the letters X, Y and Z | | |
| | NOTE X and Y labels are interchangeable, Z is not (1) | | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|--|---|------|
| 3(a)(i) | $(3 \times -120) = -360 \text{ (kJ mol}^{-1})$ | No sign or + sign in answer, ie 360/+360 Any other wrong units | 1 |
| | IGNORE ΔH, and case of letters in units e.g allow Kj | ΔΕ | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|--|----------------------------------|------|
| *3(a)(ii) | (Bonding in) benzene/it is more stable (than Kekule) by 152 kJ mol ⁻¹ (consequential on (a)(i)) (1) IGNORE sign | | 4 |
| | π /p/double bond electrons are delocalized (around the ring) | | |
| | OR six p electrons shared between six (ring) carbon atoms | | |
| | OR delocalized because of overlap of p orbitals | | |
| | OR resonance hybrid of C=C's and C-C's (1) | Attack by | |
| | Substitution reactions (rather than addition) (1) | electrophiles with no mention of | |
| | NOTE: Nucleophilic substitution negates the substitution mark because it is wrong additional information | substitution | |
| | Maintains/regains delocalized system OR maintains/regains stability OR maintains/regains stabilization energy | | |

| Question Number | Acceptable Answers | | Reject | Mark |
|--------------------|--|-------|---------------------------|------|
| 3 (b)(i) | Concentrated nitric acid/HNO ₃ | (1) | | 2 |
| | Concentrated sulfuric acid/H ₂ SO ₄ | (1) | Concentrated hydrochloric | |
| | Allow conc or c. in place of 'concentra | ited' | acid | |
| | ALLOW Concentrated nitric acid and sulfuric acid | | | |
| | OR | | | |
| | Concentrated HNO ₃ and H ₂ SO ₄ | (2) | | |
| | Second mark depends on nitric acid | | | |
| | Max. (1) if no mention of concentrate | d | | |
| | Nitric acid and concentrated sulfuric a scores (1) | icid | | |
| | NOTE: conc. HNO_3 and H_2SO_4 (aq) scores (1) conc. HNO_3 and conc H_2SO_4 (aq) score (2) | | | |

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

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|--|-----------------|------|
| 3 (b)(iii) | $Br_2 + FeBr_3 \rightarrow FeBr_4^- + Br^+$ OR | lack of charges | 4 |
| | Br-Br + FeBr ₃ \rightarrow Br $^{\delta +}$ Br $^{\delta -}$ FeBr ₃ (1) IGNORE state symbols even if wrong | | |
| | $Br^{+}/Br^{\delta+}(Br^{\delta-}FeBr_{3})$ H $(+ FeBr_{3})$ | | |
| | + H ⁺ /HBr (+ FeBr ₃) | | |
| | Arrow from benzene ring electrons (from inside the hexagon) to $\mathbf{Br}^+/\mathbf{Br}^{\delta^+}$ (\mathbf{Br}^{δ^-} FeBr ₃) (1) | | |
| | 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 | | |
| | The bonds to H and Br may be dotted (1) | | |
| | Arrow from or close to bond to H to centre of ring and H ⁺ /HBr as a product (1) | | |
| | ALLOW Kekulé structure for benzene and intermediate | | |
| | Each marking point is independent | | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|---|--|------|
| 3(b)(iv) | SO_3H OR $C_6H_5SO_3H$ accept: displayed - SO_3H - $SO_3^-H^+$ | | 2 |
| | -SO ₂ OH If two formulae are given both must be correct (1) Penalise if bond clearly goes to O or H rather than S | | |
| | Benzenesulfonic acid (1) ALLOW phenyl sulfonic acid | Benzenesulfuric acid/benzosulfonic acid/benzylsufonic acid | |

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

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|---|--------------|------|
| 3(c)(ii) | OH + 3Br ₂ Br Br + 3HBr | | 2 |
| | (1) (1) organic balancing formula |) | |
| | ALLOW | | |
| | Condensed structural formulae, for example C₆H₅OH + 3Br₂ → C₆H₂Br₃OH + 3HBr (1) (1) balancing | | |
| | • multiples | | |
| | substitution to any positions | | |
| | IGNORE: H ₂ O Position of bond to OH | | |
| | NOTE: Correct balanced equations giving mono and disubstitution phenols score 1 mark | | |

| Question Number | Acceptable Answers | Reject | Mark |
|--------------------|---|--------|------|
| 3(d) | (Chloromethyl)benzene/chloromethylbenzene/chlorophenylmethane/ benzyl chloride OR dichloromethane (1) ALLOW phenylchloromethane | | 2 |
| | Aluminium chloride (1) ACCEPT formulae eg C_7H_7CI , $C_6H_5CH_2CI$, CH_2CI_2 , | CH₂CI | |
| | AICI ₃ ACCEPT other halogen carriers eg FeCl ₃ /iron(III) chloride/ZnCl ₂ | 011201 | |
| | ACCEPT bromine in place of chlorine for either/both marks | | |
| | Correct formula and wrong name or correct name and wrong formula or any other wrong additional information loses mark | | |