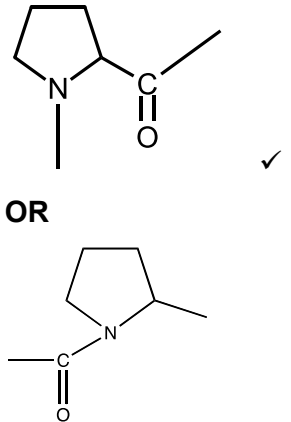
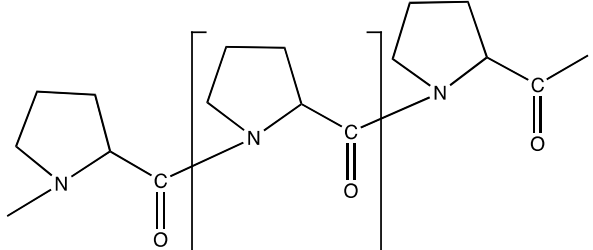
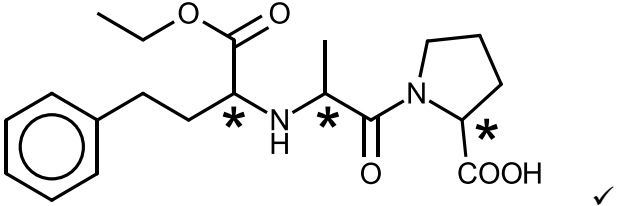
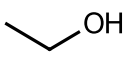
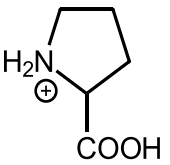
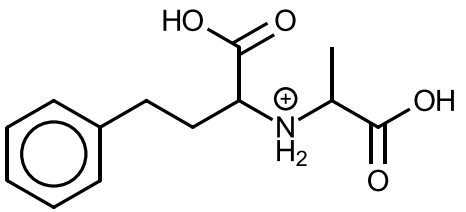
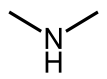
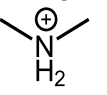
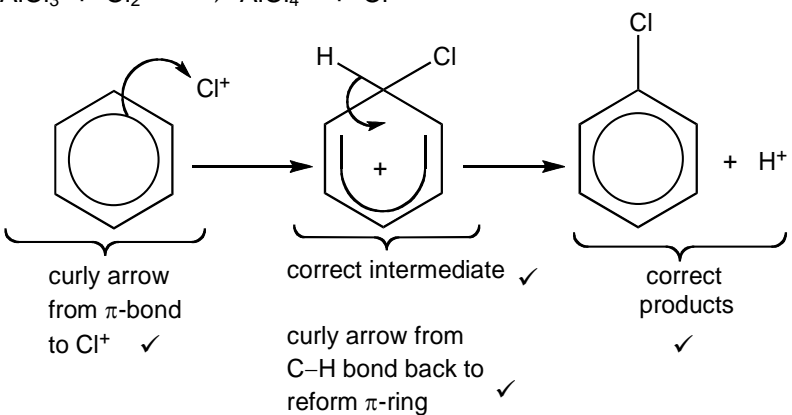
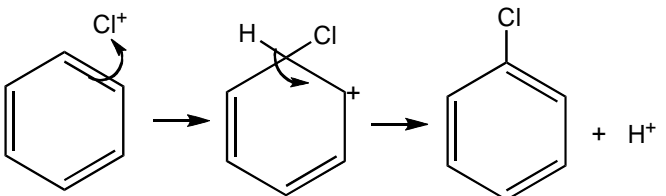
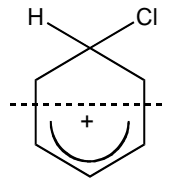
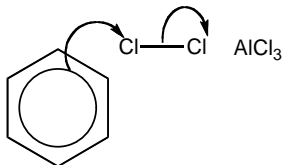


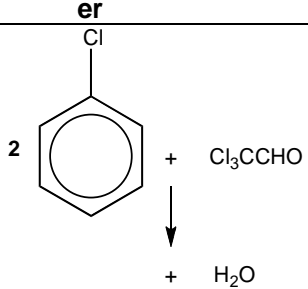
Question			Answer	Mark	Guidance
1	(a)	(i)	$\begin{array}{ccccccc} & \text{H} & \text{O} & & \text{CH}_2\text{OH} & & \\ & & & & & & \\ \text{H}_2\text{N} & -\text{C} & -\text{C} & -\text{N} & -\text{C} & -\text{COOH} & \\ & & & & & & \\ & \text{CH}_3 & & \text{H} & \text{H} & & \end{array}$ $\begin{array}{ccccccc} & \text{H} & \text{O} & & \text{CH}_3 & & \\ & & & & & & \\ \text{H}_2\text{N} & -\text{C} & -\text{C} & -\text{N} & -\text{C} & -\text{COOH} & \\ & & & & & & \\ & \text{HOH}_2\text{C} & & \text{H} & \text{H} & & \end{array}$ <div style="text-align: right;">✓</div> <div style="text-align: right;">✓</div>	2	<p>ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous</p> <p>DO NOT ALLOW peptide chains</p>
	(a)	(ii)	<p>alanine at pH 6.0</p> $\begin{array}{ccc} & \text{H} & \text{O} \\ & & \\ \text{H}_3\text{N}^{\oplus} & -\text{C} & -\text{C} & -\text{O}^{\ominus} \\ & & & \\ & \text{CH}_3 & & \end{array}$ <div style="text-align: right;">✓</div> <p>serine at pH 10.0</p> $\begin{array}{ccc} & \text{H} & \text{O} \\ & & \\ \text{H}_2\text{N} & -\text{C} & -\text{C} & -\text{O}^{\ominus} \\ & & & \\ & \text{CH}_2\text{OH} & & \end{array}$ <div style="text-align: right;">✓</div>	2	<p>ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous</p> <p>ALLOW + charge on N or H: <i>i.e.</i> $^+\text{NH}_3$ or NH_3^+</p> <p>DO NOT ALLOW ‘-’ charge on C <i>i.e.</i> ^-COO</p> <p>DO NOT ALLOW if structure is incomplete</p>

Question		Answer	Mark	Guidance
(a)	(iii)	 <p>OR</p>	1	<p>ALLOW correct structural OR displayed OR skeletal formulae</p> <p>OR combination of above as long as unambiguous</p> <p>IGNORE bond angles</p> <p>DO NOT ALLOW more than one repeat unit</p> <p>ALLOW end bonds shown as - - - -</p> <p>DO NOT ALLOW if structure has no end bonds</p> <p>IGNORE brackets unless they are used to pick out the repeat unit from a polymer chain</p> <p>IGNORE n</p> 

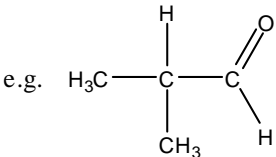
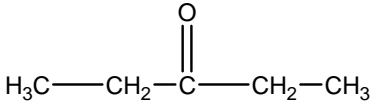
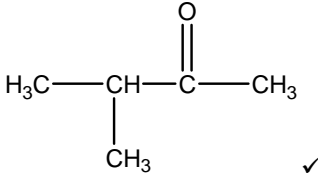
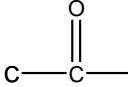
Question		Answer	Mark	Guidance									
(b)		<p style="text-align: center;">¹H NMR spectrum for serine</p> <table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th style="text-align: center;">chemical shift, δ /ppm</th> <th style="text-align: center;">relative peak area</th> <th style="text-align: center;">splitting pattern</th> </tr> </thead> <tbody> <tr> <td style="text-align: center;">2.0 to 3.0</td> <td style="text-align: center;">1</td> <td style="text-align: center;">triplet</td> </tr> <tr> <td style="text-align: center;">3.3 to 4.2</td> <td style="text-align: center;">2</td> <td style="text-align: center;">doublet</td> </tr> </tbody> </table> <p>One mark for each correct row ✓✓</p>	chemical shift, δ /ppm	relative peak area	splitting pattern	2.0 to 3.0	1	triplet	3.3 to 4.2	2	doublet	2	<p>ALLOW δ values ± 0.2 ppm, as a range or a value within the range</p> <p>ALLOW a response that implies a splitting into three for a triplet/into two for a doublet</p>
chemical shift, δ /ppm	relative peak area	splitting pattern											
2.0 to 3.0	1	triplet											
3.3 to 4.2	2	doublet											
(c)	(i)	 <p style="text-align: right;">✓</p>	1	ALL correct for one mark									
(c)	(ii)	<p><i>any two from:</i></p> <ul style="list-style-type: none"> no/fewer side effects increases the (pharmacological) activity/effectiveness Reduces/stops the need for/cost/difficulty in separating stereoisomers/optical isomers <p style="text-align: right;">✓✓</p>	2	<p>IGNORE toxic/harmful</p> <p>IGNORE a response that implies a reduced dose</p> <p>IGNORE "it takes (less) time to separate"</p>									

Question		Answer	Mark	Guidance
(c)	(iii)	   <p>✓ one mark for ethanol</p> <p>✓ one mark for proline with NH OR NH₂⁺</p> <p>✓ one mark for remaining fragment</p> <p>with  or </p> <p>✓ Fourth mark for structure of both ions shown correctly with NH₂⁺</p>	4	<p>ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous</p> <p>ALLOW + charge on H of NH₂ groups, <i>i.e.</i> NH₂⁺</p> <p>IGNORE negative (counter) ions</p>
(c)	(iv)	<p>idea of separating (the components/compounds)</p> <p>AND idea of (identifying compounds by) comparison with a (spectral) database ✓</p>	1	<p>ALLOW (identifies compounds) using fragmentation (patterns)/fragment ions (but IGNORE molecular ions)</p> <p>IGNORE retention times</p>
Total			15	

Question	Answer	Mark	Guidance
2 (a)	<p>$\text{AlCl}_3 + \text{Cl}_2 \longrightarrow \text{AlCl}_4^- + \text{Cl}^+ \checkmark$</p>  <p>curly arrow from π-bond to Cl^+ \checkmark</p> <p>correct intermediate \checkmark</p> <p>curly arrow from C-H bond back to reform π-ring \checkmark</p> <p>correct products \checkmark</p> <p>$\text{H}^+ + \text{AlCl}_4^- \longrightarrow \text{AlCl}_3 + \text{HCl} \checkmark$</p> <p>Note: 1st curly arrow should start within the ring or on the ring</p> <hr/> <p>Note: ALLOW mechanism using Kekulé structures:</p> 	6	<p>Guidance</p> <p>ANNOTATIONS MUST BE USED</p> <p>DO NOT ALLOW the following intermediate:</p>  <p>π-ring must be more than 1/2 way up AND 'horseshoe' the right way up, <i>ie</i> gap towards C with Cl</p> <p>ALLOW + sign anywhere inside the 'hexagon' of intermediate</p> <p>ALLOW 1st curly arrow starting within the hexagon</p> <p>ALLOW mechanism with $\text{Cl}-\text{Cl} \cdots \text{AlCl}_3$ for 1st 2 marks, <i>ie</i></p>  <p>Second curly arrow to either $-\text{Cl}$ or AlCl_3</p> <p>Note: If Br^+ is used, DO NOT ALLOW 1st mechanism mark but all other marks available by ECF</p>

Question		er	Mark	Guidance
(b)	(i)	 <p>2 <chem>c1ccc(Cl)cc1</chem> + <chem>ClC(Cl)(Cl)C=O</chem></p> <p>↓</p> <p>+ <chem>H2O</chem></p> <p>1st mark: reactants, correctly balanced, ✓ ie <chem>2 C6H5Cl + Cl3CCHO</chem></p> <p>2nd mark: product, (correctly balanced) ✓ ie <chem>H2O</chem></p>	2	<p>Each mark is independent of the other</p> <p>ALLOW <chem>C6H5Cl</chem> for chlorobenzene</p> <p>ALLOW any unambiguous structure for <chem>Cl3CCHO</chem>, e.g. <chem>CCl3CHO</chem></p> <p>BUT DO NOT ALLOW <chem>CCl3COH</chem></p> <p>Standalone mark</p> <p>Standalone mark</p>
	(ii)	6 ✓	1	
(c)		<p>substitution/nitration/<chem>NO2</chem> at different positions (on the ring)</p> <p>OR</p> <p>forms different isomers</p> <p>OR</p> <p>multiple substitution/nitration ✓</p>	1	<p>ALLOW examples, e.g. 1-chloro-2-nitrobenzene and 1-chloro-2-nitrobenzene</p> <p>ALLOW 'it' for nitro group</p> <p>ALLOW examples, e.g. 1-chloro-2,3-dinitrobenzene</p> <p>IGNORE nitrate/<chem>NO3</chem></p>
(d)		<p>In phenol,</p> <p>(lone) pair of electrons on O is (partially) delocalised into the ring ✓</p> <p>QWC: delocalised/delocalized/delocalise, etc must be spelt correctly in the correct context for benzene OR phenol at least once</p> <p>electron density increases/is high ✓ ORA</p> <p><chem>Cl2</chem>/electrophile is (more) polarised ✓ ORA</p>	3	<p>ANNOTATIONS MUST BE USED</p> <p>ALLOW diagram to show movement of lone pair into ring but delocalised ring must be mentioned</p> <p>ALLOW lone pair of electrons on O is (partially) drawn/ attracted/pulled into delocalised ring</p> <p>IGNORE 'activates the ring'</p> <p>DO NOT ALLOW charge density or electronegativity</p> <p>ALLOW <chem>Cl2</chem> is (more) attracted</p> <p>OR <chem>Cl2</chem> is not polarised by benzene</p> <p>OR induces dipoles (in chlorine/electrophile)</p>
Total			13	

Question		Expected Answers		Marks	Additional Guidance
3	a	Alternative approaches		4	<p>ALLOW ammoniacal AgNO₃/ Ag⁺(NH₃)₂ / Ag⁺(NH₃)</p> <p>ALLOW acidified dichromate OR Fehlings as an alternative to Tollens – observation ‘turn green’ OR ‘red precipitate’ respectively</p> <p>ALLOW acidified manganate(VII) and observation as either brown precipitate/decolourised/pale pink</p> <p>ALLOW Brady’s (reagent)</p> <p>ALLOW orange/red/yellow for colour of the 2,4-DNP(H) precipitate</p> <p>ALLOW solid/crystals in place of precipitate</p> <p>IGNORE any reference to melting points</p> <p>ALLOW PCl₅ as a test for the acid – observation would be ‘white fumes (of HCl)’</p> <p>ALLOW detection of (carboxylic) acid by reacting with an alcohol to make an ester but no mark for the observation.</p> <p>DO NOT ALLOW detection of (carboxylic) acid by pH or indicator</p> <p>Please annotate, use ticks to show where marks are awarded</p>
		<p>Tollens’ test AND ‘silver precipitate/mirror’ ✓ is the aldehyde ✓</p> <p>react with 2,4-DNP(H) and ‘orange precipitate’ ✓</p> <p>must be the ketone ✓</p>	<p>Tollens’ test AND ‘silver precipitate/mirror’ ✓ is the aldehyde ✓</p> <p>react with carbonate/hydrogencarbonate/Na/Mg and ‘fizzes/bubbles/ effervesces/ gas evolved’ ✓</p> <p>must be the (carboxylic) acid ✓</p>		
	b	<p>2,4-DNP(H) AND orange precipitate ✓ is either aldehyde OR ketone ALLOW carbonyl OR C=O ✓</p> <p>Tollens’ test & ‘silver ppt/mirror’ ✓ is the aldehyde ✓</p>	<p>2,4-DNP(H) and no orange precipitate ✓ is the (carboxylic) acid ✓</p> <p>Tollens’ test & ‘silver ppt/mirror’ ✓ is the aldehyde ✓</p>		
		<p>Peak in range 2500–3300 (cm⁻¹) or (around) 3000 shows O–H ✓ [need wavenumber (or range) and O–H bond]</p>		1	<p>DO NOT ALLOW single peak quoted within range 2500–3300 other than 3000 (cm⁻¹) for OH</p> <p>DO NOT ALLOW range 3200–3550 (cm⁻¹)</p> <p>IGNORE any reference to C–O or C=O</p>

Question		Expected Answers	Marks	Additional Guidance
c		<p>Alternative approaches depending on whether or not the aldehyde is correct</p> <p>Doublet indicates adjacent C is bonded to only 1H OR (relative) peak area indicates 2 x CH₃ (in the same environment) ✓</p> <p>If aldehyde is correct (CH₃)₂CH—CH₂—CHO ✓✓</p> <p><i>If aldehyde is correct only need to explain doublet OR peak areas</i></p>		<p>ALLOW 3-methylbutanal, any correct unambiguous structure ALLOW two marks for correct aldehyde with no explanation</p> <p>ALLOW doublet/peak at 0.9ppm due to R—CH ALLOW the splitting shows adjacent to CH/environment that contains 1 H/proton</p> <p>ALLOW 6 Hs/ protons in same environment DO NOT ALLOW 6 Hs in same environment next to CHO</p> <p>e.g. </p> <p>would score two marks if the doublet and the peak areas were correctly explained</p>
		<p>Doublet indicates adjacent C is bonded to only 1H ✓ AND (relative) peak area indicates 2 x CH₃ (in the same environment) ✓</p> <p>If aldehyde identified is incorrect ✗</p> <p><i>if aldehyde is incorrect must explain both doublet or peak areas</i></p>		
d	i	<p> ✓</p> <p>ketone 3</p>	1	ALLOW displayed/skeletal formulae
	ii	<p>There are 4 (different C) environments ✓ (therefore) it is ketone 2 /</p> <p> ✓</p> <p>(C responsible for peak at δ = 210 ppm) is C=O/carbonyl carbon ✓</p>	3	<p>ALLOW 2 Cs are in same environment/equivalent</p> <p>ALLOW 3-methylbutan(-2-)one/ any correct unambiguous structure</p> <p>ALLOW 2-methylbutan-3-one</p> <p>ALLOW</p> <p></p>
Total			12	

Question		er	Mark	Guidance	
4	(a)	$(\text{CH}_3\text{CO})_2\text{O} + \text{CH}_3\text{CH}(\text{OH})\text{CH}_3$ $\rightarrow \text{CH}_3\text{COOCH}(\text{CH}_3)_2 + \text{CH}_3\text{COOH}$ 1st mark Correct structure of ester: $\text{CH}_3\text{COOCH}(\text{CH}_3)_2$ ✓ 2nd mark Equation contains correct formulae for $(\text{CH}_3\text{CO})_2\text{O}$, $\text{CH}_3\text{CH}(\text{OH})\text{CH}_3$ AND CH_3COOH ✓	2	ALLOW correct structural OR displayed OR skeletal formula ALLOW combination of formulae as long as unambiguous DO NOT ALLOW molecular formulae ALLOW $(\text{CH}_3)_2\text{CHOOCCH}_3$ OR $(\text{CH}_3)_2\text{CHOCOCH}_3$	
	(b)	(i)	(relative) solubility ✓	1	IGNORE partition DO NOT ALLOW adsorption OR absorption
		(ii)	The esters would have similar retention times AND similar structures/molecules OR same functional groups OR similar polarities OR similar solubilities ✓ Alcohol would have short retention time AND alkane would have long retention time ✓	2	IGNORE similar properties

Question	Answer	Mark	Guidance
4 (c)	<p>Elemental analysis and molecular formula – 2 marks</p> <p>Use of percentages (to find EF) AND 144 ✓</p> <p>Molecular formula = C₈H₁₆O₂ ✓</p>	2 marks	<p>ANNOTATIONS MUST BE USED</p> <p><i>Working</i></p> $\text{C : H : O} = 66.63/12 : 11.18/1 : 22.19/16$ $5.5525 : 11.18 : 1.386875$ $4 : 8 : 1$ <p>Alternative method:</p> <p>carbon: (144 x 66.63/100)/12 = 8 hydrogen: (144 x 11.18/100)/1 = 16 oxygen: (144 x 22.19/100)/16 = 2</p>
	<p>ester structure – 4 marks</p> $\begin{array}{ccccccc} & \text{CH}_3 & & \text{O} & & & \\ & & & & & & \\ \text{H}_3\text{C} & - \text{C} & - \text{CH}_2 & - \text{C} & - \text{O} & - \text{CH}_2 & - \text{CH}_3 \\ & & & & & & \\ & \text{CH}_3 & & & & & \end{array}$ <p style="text-align: right;">✓✓✓✓</p>	4 marks	<p>ALLOW correct structural OR displayed OR skeletal formula ALLOW combination of formulae as long as unambiguous NO ECF from earlier structures</p> <p>If not fully correct award following marks:</p> <p>If structure an ester of formula C₈H₁₆O₂ OR the organic structure contains C(CH₃)₃ ✓</p> <p>If structure is an ester of formula C₈H₁₆O₂ AND ester contains C(CH₃)₃ ✓✓</p> <p>If structure is an ester of formula C₈H₁₆O₂ AND ester contains O–CH₂C(CH₃)₃ AND ester contains CH₃CH₂COO ✓✓✓ <i>i.e. If the ester link is reversed</i></p> $\begin{array}{ccccccc} & & & \text{O} & & & \text{CH}_3 \\ & & & & & & \\ \text{CH}_3 & - \text{CH}_2 & - \text{C} & - \text{O} & - \text{CH}_2 & - \text{C} & - \text{CH}_3 \\ & & & & & & \\ & & & & & & \text{CH}_3 \end{array}$ <p>IGNORE any name</p>

Question	Answer	Mark	Guidance
	<p>NMR analysis – 4 marks</p> <p>Triplet (at δ 1.3) shows an adjacent CH₂ OR triplet (at δ 1.3) shows (C with) 2 adjacent Hs/protons ✓ <i>(because of splitting: so triplet)</i></p> <p>Peak at (δ) 2.2 shows H adjacent to C=O AND adjacent to (C with) no hydrogens ✓ <i>(because of no splitting: so singlet)</i></p> <p>Peak at (δ) 4.2 shows H–C–O AND adjacent CH₃ OR 3 adjacent Hs/protons ✓ <i>(because of splitting: so quartet)</i></p> <p>Peak at (δ) 0.9 show 3 x CH₃ ✓ <i>(because of singlet and area 9)</i></p>	<p>4 marks</p>	<p>NOTE: Each peak can be identified from:</p> <ul style="list-style-type: none"> • its δ value: ± 0.2 ppm • a range, eg ‘the peak between 2 and 3’ • its relative peak area (CARE two peaks have an area of 2) • its splitting (CARE: two peaks are singlets) • labelling on the spectrum <hr/> <p>QWC: triplet must be spelled correctly ALLOW neighbouring Hs for adjacent to Hs</p> <p>For peak at (δ) 2.2 ALLOW singlet at (δ) 2.2 ALLOW singlet labelled 2</p> <p>For peak at (δ) 4.2 ALLOW quartet (labelled 2)</p> <hr/> <p>Check back for any responses added to spectra</p> <p>ADD ^ MARK TO THE SPECTRUM PAGE TO SHOW THAT IT HAS BEEN LOOKED AT</p>
	Total for 4(c)	10	
	Total	15	