| Question |  |  | Answer |  | Mark | Guidance |
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| 1 | (a) | (i) |   |  | 2 | ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous <br> DO NOT ALLOW peptide chains |
|  | (a) | (ii) | alanine at pH 6.0 <br> serine at pH 10.0 |  | 2 | ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous <br> ALLOW + charge on N or H : i.e. ${ }^{+} \mathrm{NH}_{3}$ or $\mathrm{NH}_{3}{ }^{+}$ <br> DO NOT ALLOW ‘-‘ charge on C i.e. ${ }^{-} \mathrm{COO}$ <br> DO NOT ALLOW if structure is incomplete |


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


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| (b) | ${ }^{1} \mathrm{H}$ NMR spectrum for serine |  |  |  | 2 | ALLOW $\delta$ values $\pm 0.2 \mathrm{ppm}$, as a range or a value within the range <br> ALLOW a response that implies a splitting into three for a triplet/into two for a doublet |
|  |  | chemical shift, $\delta / p p m$ | relative peak area | splitting pattern |  |  |
|  |  | 2.0 to 3.0 | 1 | triplet |  |  |
|  |  | 3.3 to 4.2 | 2 | doublet |  |  |
|  |  | One mark for each correct row $\quad \checkmark \checkmark$ |  |  |  |  |
| (c) | (i) |  |  |  | 1 | ALL correct for one mark |
| (c) | (ii) | any two from: <br> no/fewer side effects <br> increases the (pharmacological) activity/effectiveness <br> Reduces/stops the need for/cost/difficulty in separating stereoisomers/optical isomers |  |  | 2 | IGNORE toxic/harmful <br> IGNORE a response that implies a reduced dose IGNORE "it takes (less) time to separate" |


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| (c) | (iii) |  | 4 | ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous ALLOW + charge on H of $\mathrm{NH}_{2}$ groups, i.e. $\mathrm{NH}_{2}{ }^{+}$ IGNORE negative (counter) ions |
| (c) | (iv) | idea of separating (the components/compounds) <br> AND idea of (identifying compounds by) comparison with a (spectral) database | 1 | ALLOW (identifies compounds) using fragmentation (patterns)/fragment ions (but IGNORE molecular ions) <br> IGNORE retention times |
|  |  | Total | 15 |  |

Question

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| (b) | (i) | 2 <br> 1st mark: reactants, correctly balanced, $\checkmark$ ie $2 \mathrm{C}_{6} \mathrm{H}_{5} \mathrm{Cl}+\mathrm{Cl}_{3} \mathrm{CCHO}$ <br> 2nd mark: product, (correctly balanced) $\checkmark$ ie $\mathrm{H}_{2} \mathrm{O}$ | 2 | Each mark is independent of the other <br> ALLOW $\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{Cl}$ for chlorobenzene <br> ALLOW any unambiguous structure for $\mathrm{Cl}_{3} \mathrm{CCHO}$, e.g. $\mathrm{CCl}_{3} \mathrm{CHO}$ <br> BUT ..... DO NOT ALLOW CCl $\mathrm{COH}_{3}$ <br> Standalone mark <br> Standalone mark |
|  | (ii) | $6 \checkmark$ | 1 |  |
| (c) |  | substitution/nitration/ $\mathrm{NO}_{2}$ at different positions (on the ring) OR <br> forms different isomers <br> OR <br> multiple substitution/nitration $\checkmark$ | 1 | ALLOW examples, e.g. 1-chloro-2-nitrobenzene and 1-chloro-2-nitrobenzene <br> ALLOW 'it' for nitro group <br> ALLOW examples, e.g. 1-chloro-2,3-dinitrobenzene IGNORE nitrate/ $\mathrm{NO}_{3}$ |
| (d) |  | In phenol, <br> (lone) pair of electrons on O is (partially) delocalised into the ring $\checkmark$ <br> QWC: delocalised/delocalized/delocalise, etc must be spelt correctly in the correct context for benzene OR phenol at least once <br> electron density increases/is high $\checkmark$ ORA <br> $\mathrm{Cl}_{2} /$ electrophile is (more) polarised $\checkmark$ ORA | 3 | ANNOTATIONS MUST BE USED <br> ALLOW diagram to show movement of lone pair into ring but delocalised ring must be mentioned ALLOW lone pair of electrons on $O$ is (partially) drawn/ attracted/pulled into delocalised ring <br> IGNORE 'activates the ring' <br> DO NOT ALLOW charge density or electronegativity <br> ALLOW Cl ${ }_{2}$ is (more) attracted <br> $\mathrm{OR} \mathrm{Cl} \mathrm{I}_{2}$ is not polarised by benzene <br> OR induces dipoles (in chlorine/electrophile) |
|  |  | Total | 13 |  |




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| 4 | (a) |  | $\begin{aligned} & \text { (CH3 CO) })_{2} \mathrm{O}+\mathrm{CH}_{3} \mathrm{CH}(\mathrm{OH}) \mathrm{CH}_{3} \\ & \qquad \rightarrow \mathrm{CH}_{3} \mathrm{COOCH}\left(\mathrm{CH}_{3}\right)_{2}+\mathrm{CH}_{3} \mathrm{COOH} \\ & \text { 1st mark } \\ & \text { Correct structure of ester: } \mathrm{CH}_{3} \mathrm{COOCH}\left(\mathrm{CH}_{3}\right)_{2} \checkmark \\ & \text { 2nd mark } \\ & \text { Equation contains correct formulae for }\left(\mathrm{CH}_{3} \mathrm{CO}\right)_{2} \mathrm{O} \text {, } \\ & \mathrm{CH}_{3} \mathrm{CH}(\mathrm{OH}) \mathrm{CH}_{3} \text { AND } \mathrm{CH}_{3} \mathrm{COOH} \checkmark \end{aligned}$ | 2 | ALLOW correct structural OR displayed OR skeletal formula ALLOW combination of formulae as long as unambiguous DO NOT ALLOW molecular formulae <br> ALLOW $\left(\mathrm{CH}_{3}\right)_{2} \mathrm{CHOOCCH}_{3} \mathrm{OR}\left(\mathrm{CH}_{3}\right)_{2} \mathrm{CHOCOCH}_{3}$ |
|  | (b) | (i) | (relative) solubility $\checkmark$ | 1 | IGNORE partition <br> DO NOT ALLOW adsorption OR absorption |
|  |  | (ii) | The esters would have similar retention times <br> AND <br> similar structures/molecules OR same functional groups <br> OR similar polarities OR similar solubilities <br> Alcohol would have short retention time <br> AND <br> alkane would have long retention time $\checkmark$ | 2 | IGNORE similar properties |


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| 4 | (c) | Elemental analysis and molecular formula - 2 marks <br> Use of percentages (to find EF) AND $144 \checkmark$ <br> Molecular formula $=\mathrm{C}_{8} \mathrm{H}_{16} \mathrm{O}_{2} \checkmark$ | $\begin{gathered} 2 \\ \text { marks } \end{gathered}$ | ANNOTATIONS MUST BE USED <br> Working $\begin{array}{ccccc} \mathrm{C}: \mathrm{H}: \mathrm{O} & =66.63 / 12 & : & 11.18 / 1 & : \\ & 5.5525 & : 11.18 & 22.19 / 16 \\ & 4 & : & 8 & : \\ & & 1.386875 \\ & \end{array}$ <br> Alternative method: <br> carbon: $(144 \times 66.63 / 100) / 12=8$ <br> hydrogen: $(144 \times 11.18 / 100) / 1=16$ <br> oxygen: $(144 \times 22.19 / 100) / 16=2$ |
|  |  | ester structure - 4 marks | $\begin{gathered} 4 \\ \text { marks } \end{gathered}$ | ALLOW correct structural OR displayed OR skeletal formula ALLOW combination of formulae as long as unambiguous NO ECF from earlier structures <br> If not fully correct award following marks: <br> If structure an ester of formula $\mathrm{C}_{8} \mathrm{H}_{16} \mathrm{O}_{2}$ OR the organic structure contains $\mathrm{C}\left(\mathrm{CH}_{3}\right)_{3}$ <br> If structure is an ester of formula $\mathrm{C}_{8} \mathrm{H}_{16} \mathrm{O}_{2}$ <br> AND ester contains $\mathrm{C}\left(\mathrm{CH}_{3}\right)_{3}$ <br> If structure is an ester of formula $\mathrm{C}_{8} \mathrm{H}_{16} \mathrm{O}_{2}$ <br> AND ester contains $\mathrm{O}-\mathrm{CH}_{2} \mathrm{C}\left(\mathrm{CH}_{3}\right)_{3}$ <br> AND ester contains $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{COO} \checkmark \checkmark \checkmark$ <br> i.e. If the ester link is reversed <br> IGNORE any name |



