

# GCE

## **Chemistry A**

Unit H432A/02: Synthesis and analytical techniques

Advanced GCE

## Mark Scheme for June 2017

#### H432A/02

#### Mark Scheme

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This mark scheme is published as an aid to teachers and students, to indicate the requirements of the examination. It shows the basis on which marks were awarded by examiners. It does not indicate the details of the discussions which took place at an examiners' meeting before marking commenced.

All examiners are instructed that alternative correct answers and unexpected approaches in candidates' scripts must be given marks that fairly reflect the relevant knowledge and skills demonstrated.

Mark schemes should be read in conjunction with the published question papers and the report on the examination.

OCR will not enter into any discussion or correspondence in connection with this mark scheme.

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## Annotations available in RM Assessor

Annotation	Meaning
<b>V</b>	Correct response
X	Incorrect response
	Omission mark
BOD	Benefit of doubt given
CON	Contradiction
RE	Rounding error
SF	Error in number of significant figures
ECF	Error carried forward
L1	Level 1
L2	Level 2
L3	Level 3
NBOD	Benefit of doubt not given
SEEN	Noted but no credit given
I	Ignore

Abbreviations, annotations and conventions used in the detailed Mark Scheme (to include abbreviations and subject-specific conventions).

Annotation	Meaning
1	alternative and acceptable answers for the same marking point
✓	Separates marking points
DO NOT ALLOW	Answers which are not worthy of credit
IGNORE	Statements which are irrelevant
ALLOW	Answers that can be accepted
()	Words which are not essential to gain credit
	Underlined words must be present in answer to score a mark
ECF	Error carried forward
AW	Alternative wording
ORA	Or reverse argument

## H432/02 Subject-specific Marking Instructions

#### **Mark Scheme**

### INTRODUCTION

Your first task as an Examiner is to become thoroughly familiar with the material on which the examination depends. This material includes:

- the specification, especially the assessment objectives
- the question paper
- the mark scheme.

You should ensure that you have copies of these materials.

You should ensure also that you are familiar with the administrative procedures related to the marking process. These are set out in the OCR booklet **Instructions for Examiners**. If you are examining for the first time, please read carefully **Appendix 5 Introduction to Script Marking: Notes for New Examiners**.

Please ask for help or guidance whenever you need it. Your first point of contact is your Team Leader.

H432/02	
Question	

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Question	Кеу	Marks	Guidance
1	В	1	
2	D	1	
3	С	1	
4	В	1	
5	D	1	
6	С	1	ALLOW 3 (This is the trigonal planar atom)
7	A	1	
8	С	1	ALLOW 4 (This is the number of chiral centres)
9	С	1	
10	D	1	
11	С	1	ALLOW 3 (This is the number of peaks in the NMR spectrum)
12	A	1	
13	A	1	
14	С	1	
15	A	1	
	L	I	<u> </u>

H432/	02		Mark Sche	eme	
Que	stio	n	Answer	Marks	Guidance
16 (a	a)		Compound A (is branched so) has less points of contact / less surface interaction between molecules $\checkmark$	2	Both answers need to be comparisons ALLOW ORA throughout
					DO NOT ALLOW 'more contact between atoms'
					<b>IGNORE</b> van der Waals' forces/VDW for induced dipole–dipole interactions (ambiguous as this term refers to both permanent dipole – dipole and induced dipole–dipole forces)
			Induced dipole-dipole interactions / London (dispersion) forces are weaker.		ALLOW fewer induced dipole-dipole interactions.
			AND Require less energy to break (these interactions / forces) ✓		IGNORE it is easier to break the induced dipole- dipole / London forces. (reference to energy required) IGNORE less energy required to separate molecules IGNORE less energy is needed to break the bonds.
(k	<b>)</b>	(i)	Hex-3-en-1-ol ✓	1	ALLOW Hex-3-ene-1-ol
					<b>ALLOW</b> 1-hydroxyhex-3-ene as this is unambiguous
					Hex-3-enol is <b>not</b> sufficient
					IGNORE lack of hyphens, or addition of commas

1432/02	Mark Sch	eme		Jun
Question	Answer	Marks	Guidance	
(ii)	Same structural formula AND Different arrangement (of atoms) in space OR different spatial arrangement (of atoms) ✓	1	<ul> <li>ALLOW have the same structure/displayed formula/skeletal formula</li> <li>DO NOT ALLOW same empirical formula OR same general formula</li> <li>IGNORE same molecular formula</li> <li>Reference to <i>E/Z</i> isomerism or optical isomerism is <b>not</b> sufficient</li> </ul>	
	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	2	<ul> <li>ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous</li> <li>ALLOW one mark if both stereoisomers of compound C are shown but in the incorrect columns</li> <li>ALLOW one mark for correct stereoisomers of compound C in correct columns where – CH<sub>2</sub>CH<sub>2</sub>OH is represented as -C<sub>2</sub>H<sub>5</sub>O or – C<sub>2</sub>H<sub>4</sub>OH</li> <li>DO NOT ALLOW incorrect connectivity e.g. – CH<sub>3</sub>CH<sub>2</sub> on first occasion but allow ECF in second structure.</li> </ul>	

H432/02	Mark Sch	eme		June
Question	Answer	Marks	Guidance	
(c)	Image: point of the second secon	1	DO NOT ALLOW C=C in diagram DO NOT ALLOW overlapping p orbitals on left hand side in the diagram. DO NOT ALLOW a diagram that contains four lobes on the right hand side. e.g. IGNORE any atoms joined to the bonds Note: labels are not required ALLOW the following diagram to show the π- bond (π-bond)	
(d) (i)	(The H atom of HBr) accepts a pair of electrons $\checkmark$	1		
(ii)	$\begin{array}{ c c c c c c } \hline CH_3 & CH_2CH_3 & CH_3 & CH_2CH_3 \\ \hline H_3C & \hline C & \hline C & \hline C & \hline H & H_3C & \hline C & \hline C & \hline H & \hline H & Br & \checkmark & Br & H & \checkmark \end{array}$	2	ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous ALLOW in either order	

H432/02	Mark Scl	neme		June 2017
Question	Answer	Marks	Guidance	
(iii)	Curly arrow from C=C bond to H of H–Br $\checkmark$	3	ANNOTATE ANSWER WITH TICKS AND CROSSES	
	Correct dipole shown on H–Br <b>AND</b> curly arrow showing the breaking of H–Br bond $\checkmark$ CH <sub>3</sub> CH <sub>2</sub> CH <sub>3</sub>		<b>ALLOW</b> any combination of skeletal <b>OR</b> structural <b>OR</b> displayed formula as long as unambiguous	
	$CH_3$ $H^{\delta+}$		<b>DO NOT ALLOW</b> partial charges shown on C=C double bond ( the second marking point)	
	$\int_{Br}^{1} \delta^{-}$ Correct carbocation <b>AND</b> curly arrow from Br <sup>-</sup> to C <sup>+</sup> of carbocation $\checkmark$			
	$\begin{array}{c c} CH_3 & CH_2CH_3 \\ CH_3 - C - C - H \\ H \end{array} \begin{array}{c} CH_3 - C - C - H \\ H \end{array} \begin{array}{c} CH_3 - C - C - H \\ CH_3 - C - C - H \\ H \end{array} \begin{array}{c} CH_3 - C - C - H \\ CH_3 - C - C - H \\ H \end{array}$		<b>DO NOT ALLOW</b> $\delta$ + on C of carbocation Curly arrow must come from a lone pair on Br <sup>-</sup>	
	Br		<b>OR</b> from the negative sign of Br <sup>-</sup> ion (then lone pair on Br <sup>-</sup> ion does not need to be shown)	

H432/02	Mark Sch	neme	June 2
Question	Answer	Marks	Guidance
(iv)	$\begin{array}{c c} & CH_3 & CH_2CH_3 \\ H_3C & \hline C & \hline C & \hline H \\ & & \\ Br & H \\ \end{array}$ 2-bromo-2-methylpentane AND	1	<ul> <li>Note: the correct product and explanation are both required for the mark</li> <li>The major product may be identified by its <ul> <li>corresponding letter (E or F) from the table in (d)(ii)</li> <li>correct structure</li> <li>correct name</li> </ul> </li> </ul>
	(the) <b>carbocation</b> intermediate (in the formation of 2-bromo-2-methylpentane) is more stable (than the carbocation in the formation of the other product) $\checkmark$		<b>DO NOT ALLOW</b> product comes from the more stable <b>secondary or primary</b> carbocation <b>IGNORE</b> explanations based on Markownikoff's rule.
(e) (i)	$n(\text{myrcene}) = \frac{204 \times 10^{-3}}{136.0} = 1.5(0) \times 10^{-3} \text{ (mol)} \checkmark$	2	Correct working <b>required</b> for the first marking point.
	Volume of H <sub>2</sub> = $3 \times 1.5(0) \times 10^{-3} \times 24000$ = 108 (cm <sup>3</sup> ) $\checkmark$		ALLOW ECF from incorrect moles of myrcene i.e. <i>n</i> (myrcene) × 3 × 24000 Common incorrect answers
			108000 cm <sup>3</sup> = 1 mark (not converted to g) $12cm^{3} = 1 mark$ ( divided by 3) $36 cm^{3} = 1 mark$ ( not multiplied by 3)
			<b>IGNORE</b> Calculations based on $pV = nRT$

H432/02	Mark Sche	eme		June 201
Question	Answer	Marks	Guidance	
(ii)	Amount of hydrogen $n(H_2) = \frac{5.28}{24.0} = 0.22(0) \text{ (mol) } \checkmark$ Number of double bonds $= \frac{0.220}{0.0200} = 11 \checkmark$	4	<b>ALLOW</b> Evidence of $n(H_2) = \frac{5.28}{24.0}$ if 0.22 is not seen Evidence for 11 double bonds could come from	
	Formula of saturated product C <sub>40</sub> H <sub>78</sub> ✓ Equation		11H₂ in equation Formula could be shown as the product of an equation	
	$C_{40}H_{56} + 11H_2 \longrightarrow C_{40}H_{78} \checkmark$		ALLOW ECF from $C_{40}H_{82}$ and $C_{40}H_{80}$ only i.e. $C_{40}H_{60} + 11H_2 \longrightarrow C_{40}H_{82}$ $C_{40}H_{58} + 11H_2 \longrightarrow C_{40}H_{80}$	
	Total	20		

Answer	Marks	Guidance
Generation of electrophile	5	ANNOTATE ANSWER WITH TICKS AND CROSSES
$HNO_3 + H_2SO_4 \longrightarrow H_2O + HSO_4^- + NO_2^+ \checkmark$		<b>ALLOW</b> HNO <sub>3</sub> + $2H_2SO_4 \rightarrow H_3O^+ + 2HSO_4^- +$
Electrophilic substitution		NO <sub>2</sub> <sup>+</sup>
Curly arrow from $\pi$ -bond to NO <sub>2</sub> <sup>+</sup> $\checkmark$		<b>ALLOW</b> HNO <sub>3</sub> + H <sub>2</sub> SO <sub>4</sub> $\rightarrow$ H <sub>2</sub> NO <sub>3</sub> <sup>+</sup> + HSO <sub>4</sub> <sup>-</sup> then
СООН		$H_2NO_3^+ \rightarrow H_2O + NO_2^+$
		<b>ALLOW</b> $^+NO_2$ <b>OR</b> $NO_2^+$
NO <sub>2</sub> +		First curly arrow must come from the ring to $NO_2^+$
Correct intermediate ✓		<b>DO NOT ALLOW</b> the following intermediate:
Curly arrow back from C-H bond to reform $\pi$ -ring AND H <sup>+</sup> as product $\checkmark$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$		$\pi$ -ring should cover approximately 4 of the 6 sides of the benzene ring structure <b>AND</b> the correct orientation, <i>i.e.</i> gap towards C with NO <sub>2</sub> <b>ALLOW</b> + sign anywhere inside the 'hexagon' of
	HNO <sub>3</sub> + H <sub>2</sub> SO <sub>4</sub> $\longrightarrow$ H <sub>2</sub> O + HSO <sub>4</sub> <sup>-</sup> + NO <sub>2</sub> <sup>+</sup> $\checkmark$ Electrophilic substitution Curly arrow from $\pi$ -bond to NO <sub>2</sub> <sup>+</sup> $\checkmark$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$	HNO <sub>3</sub> + H <sub>2</sub> SO <sub>4</sub> $\longrightarrow$ H <sub>2</sub> O + HSO <sub>4</sub> <sup>-</sup> + NO <sub>2</sub> <sup>+</sup> $\checkmark$ Electrophilic substitution Curly arrow from $\pi$ -bond to NO <sub>2</sub> <sup>+</sup> $\checkmark$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$

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Question	Answer	Marks	Guidance	
(ii)*	<ul> <li>Please refer to the marking instructions on page 5 of this mark scheme for guidance on how to mark this question.</li> <li>Level 3 (5–6 marks)</li> <li>Outlines the main steps of recrystallisation to produce a pure sample of 3-nitrobenzoic acid from the impure solid.</li> <li>AND</li> <li>Calculates correct percentage yield of 3-nitrobenzoic acid.</li> </ul>	6	Guidance         Indicative scientific points, with bulleted elements, may include:         1. Purification         • Recrystallisation         • Dissolve impure solid in minimum volume of hot water/solvent         • Cool solution and filter solid         • Wash with cold water/solvent and dry         2. Percentage yield         • n(benzoic acid) used = $\frac{4.97}{122}$ = 0.0407 (mol)	
	<ul> <li>AND Method of checking purity to include comparison to relevant data.</li> <li>A well-structured response with the steps for recrystallisation and the determination of purity being given in the correct order. Correct use of terminology throughout.</li> </ul>		• <i>n</i> (benzoic acid) used = $\frac{122}{122} = 0.0407$ (mol) • <i>n</i> (3-nitrobenzoic acid) made = $\frac{4.85}{167} = 0.0290$ (mol) • percentage yield = $\frac{0.0290}{0.0407} \times 100 = 71.3$ (%) <b>ALLOW</b> 71 to calculator value of 71.29001554 correctly rounded.	
	<ul> <li>Level 2 (3–4 marks) Attempts all three scientific points but explanations may be incomplete. OR Explains two scientific points thoroughly with very few omissions.</li> <li>The description of checking for purity or recrystallisation is clear and any calculations structured. Key terminology used appropriately.</li> <li>Level 1 (1–2 marks) A simple explanation based on at least two of the main scientific points. OR</li> </ul>		<b>CHECK</b> for extent of errors by <b>ECF</b> Alternative correct calculation may calculate theoretical mass of 3-nitrobenzoic acid that can be produced as $0.0407 \times 167 = 6.80$ (g) followed by: percentage yield = $\frac{4.85}{6.80} \times 100 = 71.3$ (%) Calculation <b>must</b> attempt to calculate <i>n</i> (benzoic acid) in mol.	

		(Scheme		
uestion	Answer	Marks	Guidance	
	<ul> <li>Explains one scientific point thoroughly with few omissions.</li> <li>There is an attempt at a logical structure. The description of the practical techniques provides some detail but may not be in the correct order.</li> <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><b>0 marks</b></li> <li>No response or no response worthy of credit.</li> </ul>		<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> <li>ALLOW alternative approach based on spectroscopy or TLC</li> <li>Spectroscopy <ul> <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> </li> <li>TLC <ul> <li>Run a TLC</li> <li>Compare (<i>R</i><sub>f</sub> value) to known data</li> <li>Pure sample will have a very similar <i>R</i><sub>f</sub></li> </ul> </li> </ul>	
(b) (i)	Phenol is the most easily nitrated/ most reactive AND Benzoic acid is the least easily nitrated /least reactive ✓	1	Response <b>must</b> give rank order of reactivity e.g. nitration becomes more difficult from phenol (to benzene) to benzoic acid <b>OR</b> nitration becomes easier from right to left in the table	
(ii)	Reactivity of phenol <ul> <li>a (lone) pair of electrons on O is (partially)</li> <li>delocalised/donated into the π-system / ring ✓</li> </ul>	3	<b>ANNOTATE ANSWER WITH TICKS AND CROSSES</b> <b>ALLOW</b> the electron pair in the p orbitals of the O atom becomes part of the $\pi$ -system / ring <b>ALLOW</b> diagram to show movement of lone pair into ring	

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Question	Answer		Guidance	
	Reactivity of benzoic acid The –COOH group on benzoic acid is an electron withdrawing group ✓		ALLOW lone pair of electrons on O is (partially) drawn/attracted/pulled into π-system / ring IGNORE activating and deactivating.	
	<ul> <li>Links electron density in π-bond to reactivity</li> <li>In phenol electron density is higher AND</li> <li>The ring is more susceptible to attack</li> <li>OR</li> <li>In benzoic acid electron density is lower AND</li> <li>The ring is less susceptible to attack ✓</li> </ul>		<ul> <li>ALLOW the following alternatives for susceptibility to attack:</li> <li>phenol attracts electrophiles / NO<sub>2</sub><sup>+</sup> more</li> <li>phenol polarises electrophiles / NO<sub>2</sub><sup>+</sup> more</li> <li>benzoic acid attracts electrophiles / NO<sub>2</sub><sup>+</sup> less</li> <li>benzoic acid polarises electrophiles / NO<sub>2</sub><sup>+</sup> less</li> </ul>	
(c) (i)	Bromination: Br₂ AND A <i>t</i> Br₃/FeBr₃/Fe ✓ Intermediate	3	ALLOW any combination of skeletal OR structural O displayed formula as long as unambiguous ALLOW any suitable halogen carrier catalyst	PR

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Question		Marks	Guidance			
	Reduction: Sn AND (concentrated) HC1 ✓		ALLOW Kekulé structure IGNORE names <i>(question asks for formulae)</i> IGNORE reaction conditions even if incorrect IGNORE 'dilute' for HC <i>l</i> IGNORE H <sub>2</sub> IGNORE H <sub>2</sub> IGNORE NaOH if seen as a reagent to convert nitro group into amine e.g 'Sn/(concentrated) HCl then NaOH' scores the			
(ii)	NH <sub>2</sub> is 2,4 directing $\checkmark$ Products (1 mark for each): $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$	3	mark IGNORE references to electron donating/withdrawing groups ALLOW –NH <sub>2</sub> activates the ring causing the new group to join at positions 2 and 4. ALLOW ortho and para directing for 2,4 directing IGNORE 6-directing ALLOW Kekulé structure IGNORE names	p		
	Total	21				

Question	Answer	Marks	Guidance
Question 18 (a) (i)	Answer curly arrow from <sup>-</sup> CN to carbon atom of C-C <i>l</i> bond $\checkmark$ Dipole shown on C-C <i>l</i> bond, C <sup>δ+</sup> and C <i>l</i> <sup>δ-</sup> , AND curly arrow from C-C <i>l</i> bond to C <i>l</i> atom $\checkmark$ $C_2H_5 \longrightarrow C_1^{\delta_+} \longrightarrow C_1^{\delta} \longrightarrow C_1^{\delta$	Marks 3	Guidance ANNOTATE ANSWER WITH TICKS AND CROSSES Curly arrow must come from lone pair on C of $^{-}$ CN OR CN <sup>-</sup> OR from minus sign on C of $^{-}$ CN ion (then lone pair on CN <sup>-</sup> does not need to be shown) IGNORE NaCl ALLOW S <sub>N</sub> 1 mechanism: Dipole shown on C–Cl bond, C <sup>δ+</sup> and Cl <sup>δ-</sup> , AND curly arrow from C–Cl bond to Cl atom $\checkmark$ Correct carbocation AND curly arrow from $^{-}$ CN to carbocation. Curly arrow must come from lone pair on C of $^{-}$ CN OR CN <sup>-</sup> OR from minus sign on C of $^{-}$ CN ion (then lone pair on C of $^{-}$ CN OR CN <sup>-</sup> OR from minus sign on C of $^{-}$ CN ion (then lone pair on CN <sup>-</sup> does not need to be shown) $\checkmark$ correct organic product AND Cl <sup>-</sup> $\checkmark$ H H H C <sub>2</sub> H <sub>5</sub> $\stackrel{+}{\longrightarrow}$ $\stackrel{+}$

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Question	Answer Ma		Guidance
(ii)	Compound <b>G</b>	3	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous         IGNORE name(s)         ALLOW $OH$ $OH$ $H$ $OH$ $H$ $OH$ $H$ $OH$ $H$ $H$ $H$ $H$ $H$ $H$
	Reagents		
	Reaction 2: H₂ AND Ni ✓		ALLOW any suitable metal catalyst e.g. Pt ALLOW LiAlH <sub>4</sub> for reagent in reaction 2 DO NOT ALLOW NaBH <sub>4</sub> for reagent in reaction 2 IGNORE names (question asks for formulae) IGNORE references to temperature and/or pressure
	Reaction 3: Correct formula of an aqueous acid e.g. HC <i>l</i> (aq)/H₂SO₄(aq) ✓		ALLOW H <sup>+</sup> (aq) IGNORE dilute ALLOW formula of an acid AND water e.g. HC <i>l</i> AND H <sub>2</sub> O H <sub>2</sub> SO <sub>4</sub> AND H <sub>2</sub> O

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Question	Answer	Marks	Guidance
Question (iii)	Explanation         Nitrogen electron pair OR nitrogen lone pair         AND         accepts a proton/H <sup>+</sup> ✓         Structure of salt         OH       H         H       +	Marks 2	GuidanceIGNORE NH2 group donates electron pairALLOW nitrogen donates an electron pair to H*DO NOT ALLOW nitrogen donates lone pair to acidIGNORE comments about the O in the –OH groupCompound H is a base is not sufficient (role of lone pair required)DO NOT ALLOW nitrogen/N lone pair accepts hydrogen (proton/H* required)ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous
	H H AND CT ✓		ALLOW $H = - \begin{pmatrix} 0H & H \\ - & - &$

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Question	Answer		Guidance	
(iv) (iv) (b) (i)	$H = \begin{pmatrix} H & 0 \\ - & - & - \\ H & H \end{pmatrix}$ Ester link $\checkmark$ Rest of structure $\checkmark$ (polymer J is biodegradable because) the ester / ester bond / ester group / polyester can be hydrolysed $\checkmark$ $H = \begin{pmatrix} H \\ H$	Marks 3 2 1	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous DO NOT ALLOW more than two repeat units for second marking point. 'End bonds' MUST be shown (do not have to be dotted) IGNORE brackets IGNORE n Broken down by water is <b>not</b> sufficient IGNORE references to photodegradable ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous ALLOW ClCl MUST be a whole number. DO NOT ALLOW an answer that uses an incorrect molar mass in the working. ALLOW 96	
	Total	14		

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Question	Answer		Guidance	
19 (a)	$C_5H_{10}O + 7O_2 \longrightarrow 5CO_2 + 5H_2O \checkmark$	1	ALLOW multiples e.g. $2C_5H_{10}O + 14O_2 \longrightarrow 10CO_2 + 10H_2O$ ALLOW any equation involving an unsaturated alcohol with correct balancing e.g. $C_5H_8O + 6.5O_2 \longrightarrow 5CO_2 + 4H_2O$ $C_5H_6O + 6O_2 \longrightarrow 5CO_2 + 3H_2O$ $C_5H_4O + 5.5O_2 \longrightarrow 5CO_2 + 2H_2O$ $C_5H_2O + 5O_2 \longrightarrow 5CO_2 + H_2O$	
(b) (i)	Diagram showing a water molecule and an ethanol molecule with at least one H <sup><math>\delta</math>+</sup> and one O <sup><math>\delta</math>-</sup> on <b>BOTH</b> molecules $\checkmark$ Hydrogen bond between one lone pair on O atom in one of the molecules and the H atom of another. <b>AND</b> Hydrogen bonding stated or labelled on diagram $\checkmark$ e.g. Hydrogen bond $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$ $\downarrow$	2	IGNORE state symbols ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous DO NOT ALLOW $\delta$ + on H atoms of alkyl group DO NOT ALLOW any marks for a diagram containing O <sub>2</sub> H If more than one hydrogen bond is shown they must all be correct to award the mark.	

H432/02 Mark Scheme June 2017 Question Answer Marks Guidance Statements **MUST** be comparative (ii) Hexane-1,6-diol has more OH groups (than hexan-1-ol) 1 AND e.g. hexane-1,6-diol has two -OH groups and hexan-1-ol has one -OH group (hexane-1,6-diol) forms more hydrogen bonds with water **ALLOW** hydroxyl or hydroxy DO NOT ALLOW hydroxide/OH<sup>-</sup> **ALLOW ORA** Starting material from reduction reaction 5 ALLOW any combination of skeletal **OR** structural (C) (i) **OR** displayed formula as long as unambiguous Watch for missing methyl groups **Reagent for reduction IGNORE** H<sup>+</sup>/ acid or H<sub>2</sub>O or ethanol NaBH₄ ✓ ALLOW sodium borohydride **OR** sodium tetrahydridoborate Product from reaction with NaBr/H<sub>2</sub>SO<sub>4</sub> **ALLOW** LiAlH<sub>4</sub> Br **Structural isomers** 

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uestion	Answer	Marks	Guidance		
			ALLOW in either order		
(ii)	3-methylcyclohexanol ✓	1	ALLOW 3-methylcyclohexan-1-ol ALLOW 1-methylcyclohexan-3-ol IGNORE lack of hyphens, or addition of commas		
(d)	Structures of organic products	5	ANNOTATE WITH TICKS AND CROSSES		
			Use of any primary alcohol containing 3, 5 or more carbons can be awarded up to 4 marks.		
			ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous		
			IGNORE names		
			<b>DO NOT ALLOW</b> $CH_3CH_2CH_2COH$ for the structure of the aldehyde.		
			ALLOW $CH_3CH_2CH_2CO_2H$ for the structure of the carboxylic acid.		
	Equations		ALLOW marks for structures from equations as		
	$\begin{array}{c} CH_3CH_2CH_2CH_2OH + [O] \longrightarrow CH_3CH_2CH_2CHO + H_2O \\ \checkmark \end{array}$		long as unambiguous. <b>ALLOW</b> molecular formulae in equations e.g. $C_4H_{10}O + [O] \longrightarrow C_4H_8O + H_2O$		
	$\begin{array}{c} CH_{3}CH_{2}CH_{2}CH_{2}OH + 2[O] \longrightarrow CH_{3}CH_{2}CH_{2}COOH + \\ H_{2}O \checkmark \end{array}$		$C_{4}H_{10}O + 2[O] \longrightarrow C_{4}H_{8}O_{2} + H_{2}O$ $C_{4}H_{9}OH + [O] \longrightarrow C_{3}H_{7}CHO + H_{2}O$ $C_{4}H_{9}OH + 2[O] \longrightarrow C_{3}H_{7}CO_{2}H + H_{2}O$		
	Reaction conditions		<b>IGNORE</b> incorrect structures in equations i.e. $C_4H_{10}O + [O] \longrightarrow C_3H_7COH + H_2O$		

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Que	stion	Answer	Marks	Guidance			
		Distillation to produce aldehyde/CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> CHO AND Reflux to produce carboxylic acid/CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> COOH ✓		scores equation mark Conditions <b>must</b> be linked to aldehyde/carboxylic acid or correct products. Conditions may be written above arrow of equation.			
		Total	15				

Q	uestion	Answer	Marks	Guidance
20	(a)	Empirical formula         Mole Ratio C : H : O = 5.88 : 5.92 : 1.47         Empirical formula = $C_4H_4O \checkmark$ Molecular formula         Molecular formula = $C_8H_8O_2$ AND         Evidence of 136 in working or from labelled peak in spectrum $\checkmark$	3	ANNOTATE ANSWER WITH TICKS AND CROSSES         ALLOW $\frac{70.58}{12.0}$ : $\frac{5.92}{1.0}$ : $\frac{23.50}{16.0}$ ALLOW $\frac{44:1}{12.0}$ : $\frac{136 \times 70.58}{10.0}$ : $\frac{136 \times 70.58}{12.0}$ = 8         H: $\frac{136 \times 5.92/100}{1.0}$ = 8         O: $\frac{136 \times 23.50/100}{16.0}$ = 2
	(b)	Functional groups	3	
		Phenol AND ketone ✓ Explanation		<b>DO NOT ALLOW</b> any other functional groups for first marking point.

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Question	Answer	Marks	Guidance	
	Links phenol to (weak) acidity AND no reaction with $Na_2CO_3$ (so not carboxylic acid) $\checkmark$		<b>ALLOW</b> identity of functional groups in the explanation if not stated on functional group prompt line.	
	Links 2,4-DNP(H) or Brady's reagent observation to carbonyl AND Tollens' reagent observation (so not an aldehyde) ✓		ALLOW "aldehyde or ketone" in place of carbonyl	
(c)	Carbon NMR analysis	3	ALLOW peaks to be identified by:	
	Peaks between 110–160 ppm are the (four) aromatic (carbon environments) $\checkmark$		Peaks labelled on spectrum	
	Compound contains a C=O between 190 - 200 ppm		<ul> <li>Peaks indicated on a chemical structure</li> </ul>	
	Compound contains a C-C at 20-30 ppm ✓		<ul> <li>Peaks indicated from within text</li> </ul>	
	Structure OH V		<b>Note:</b> If identifying aromatic peaks from the spectrum all four peaks should be indicated.	
			<b>ALLOW</b> any combination of skeletal <b>OR</b> structural <b>OR</b> displayed formula as long as unambiguous	
	Total	9		

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Question	Answer	Marks	Guidance	
21*	Please refer to the marking instructions on page 5 of this mark scheme for guidance on how to mark this question. Level 3 (5–6 marks) Structure of L is CH <sub>3</sub> CH <sub>2</sub> COOCH <sub>2</sub> C(CH <sub>3</sub> ) <sub>3</sub> OR (CH <sub>3</sub> ) <sub>3</sub> CCH <sub>2</sub> COOCH <sub>2</sub> CH <sub>3</sub> AND A comprehensive explanation with most of the spectral data analysed and few omissions. There is a well-developed line of reasoning which is clear and logically structured. The information presented is relevant and substantiated. Splitting patterns used to deduce the correct structure of L. Level 2 (3–4 marks) Attempts all three scientific points but explanations may be incomplete and/or structure of L incorrect. OR Explains two scientific points thoroughly with few omissions. There is a line of reasoning presented with some structure. The information presented in the most part relevant and supported by some evidence. The analysis is clear and includes some interpretation of NMR/IR peaks. Level 1 (1–2 marks) A simple explanation based on at least two of the main scientific points. OR Explains one scientific point thoroughly with few omissions. There is an attempt at a logical structure with a line of reasoning. The information is in the most part relevant.	6	Indicative scientific points may include: 1. <sup>1</sup> H NMR spectrum • $\delta = 1.1$ ppm, triplet, 3H CH <sub>3</sub> CH <sub>2</sub> - • $\delta = 1.3$ ppm, singlet, 9H (CH <sub>3</sub> ) <sub>3</sub> C- • $\delta = 2.3$ ppm, quartet, 2H CH <sub>3</sub> CH <sub>2</sub> -C=O • $\delta = 4.0$ ppm, singlet, 2HCH <sub>2</sub> -O- ALLOW approximate values for chemical shifts. 2. Infrared spectra IR spectrum of M • peak at 2300-3700 (cm <sup>-1</sup> ) is O-H • peak at ~1720 (cm <sup>-1</sup> ) is C=O • M is a carboxylic acid IR spectrum of N • peak at 3100-3700 (cm <sup>-1</sup> ) is O-H • N is an alcohol ALLOW ranges from <i>Data Sheet</i> IGNORE references to C-O peaks 3. Structure of L • L is an ester (as it reacts with HC <i>l</i> (aq) to form carboxylic acid and alcohol)	

H432/02	Mark Scheme			
Question	Answer	Marks	Guidance	
	The analysis is communicated in an unstructured way and includes interpretation of a few peaks from the NMR/IR spectra. <b>0 marks</b> No response or no response worthy of credit.		• Correct structure $\begin{array}{c c} H & H & O & H & CH_3 \\ H & C & C & C & O & C & C & CH_3 \\ H & H & H & H & CH_3 \end{array}$ ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous	
	Total	6		

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