

GCE

Chemistry A

Unit H432/02: Synthesis and analytical techniques

Advanced GCE

Mark Scheme for June 2018

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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.

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Annotations available in RM Assessor

Annotation	Meaning
✓	Correct response
×	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
LI	Level 1
12	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
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

Mark Scheme

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Subject-specific Marking Instructions

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.

Question	Answer	Marks	Guidance
1	Α	1	
2	C	1	
3	В	1	
4	С	1	
5	В	1	
6	В	1	ALLOW 4 (This is the number of peaks in the NMR spectrum)
7	С	1	
8	D	1	
9	В	1	
10	C	1	
11	В	1	ALLOW 2 (This is the number of straight chain isomers with a chiral C atom)
12	С	1	
13	Α	1	
14	В	1	
15	В	1	
	Total	15	

Question		n	Answer	Marks	Guidance
16	(a)	(i)	3-methylbutan-2-ol ✓	1	IGNORE lack of hyphens or addition of commas ALLOW 3-methylbutane-2-ol DO NOT ALLOW 2-methylbutan-3-ol OR 3-methylbut-2-ol OR 3-methylbutan-2-ol OR 3-methybutan-2-ol OR 3-methylbutan-2-ol
		(ii)	(CH ₃) ₂ CHCHOHCH ₃ ✓	1	ALLOW brackets around OH e.g. (CH ₃) ₂ CHCH(OH)CH ₃ ALLOW any unambiguous structural formula e.g. CH ₃ CH(CH ₃)CHOHCH ₃ CH ₃ CH(CH ₃)CH(CH ₃)OH
		(iii)	One mark for each correct structure.	2	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous ALLOW in either order

Questio	on	Answer	Marks	Guidance
	(iv)	$\downarrow \downarrow \downarrow + NaCl + H_2SO_4 \rightarrow \downarrow \downarrow \downarrow + NaHSO_4 + H_2O$ Correct haloalkane \checkmark Correctly balanced equation \checkmark	2	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous ALLOW H ⁺ for H ₂ SO ₄ ALLOW equations forming Na ₂ SO ₄ $\xrightarrow{OH} + 2NaCl + H_2SO_4 \rightarrow 2 \rightarrow 4$ ALLOW equations with HCl $\xrightarrow{OH} + HCl \rightarrow + H_2O$ DO NOT ALLOW equations that form NaOH
(b)		$H_{3}C \xrightarrow{CH_{3}}_{OH} \xrightarrow{H}_{OH} \xrightarrow{H}_{OH} \xrightarrow{H}_{OH} \xrightarrow{H}_{S} [O] \xrightarrow{H}_{3}C \xrightarrow{CH_{3}}_{OH} \xrightarrow{O}_{OH} \xrightarrow{H}_{2H_{2}O} \xrightarrow{CH_{3}}_{OH} \xrightarrow{O}_{OH} \xrightarrow{H}_{2H_{2}O}$ Correct organic product \checkmark Rest of equation \checkmark	2	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous ALLOW any vertical bond to the tertiary OH group e.g. ALLOW $H_3C - CH_3 - CH$

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Question	Answer	Marks	Guidance
(c)	Product from excess CH_3OH/H_2SO_4 H_3COOC	3	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous e.g OOC IGNORE connectivity in each product ALLOW the <i>E</i> or <i>Z</i> isomer as product from excess CH ₃ OH/H ₂ SO ₄
	Product from steam, H ₃ PO ₄ HOOC COOH OH V		
	Repeat unit of polymer C		'End handa' MUST he about (de not have to he
	Н СООН НООС СООН С		dotted) IGNORE brackets IGNORE n ALLOW more than one repeat unit but has to be a whole number of repeat units
	Total	11	



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Q	uesti	on	Answer	Marks	Guidance
					OR CI^{-} $H_{3}N^{+}$ C_{-} C_{-} C_{-} OH $(CH_{2})_{4}$ OH CI^{-} + NH_{3} (H missing from α C atom)
	(c)		$HO - CH_2 - O^-(Na^+) - CH_2 - O^-(Na^+) - O^-(Na^+)$	3	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous IGNORE NH ₃ (question asks for organic products) ALLOW $-COO^-$ OR $-COONa$ DO NOT ALLOW negative charge on C atom DO NOT ALLOW $-COO-Na$ (covalent bond) BUT ALLOW ECF if seen in subsequent structures DO NOT ALLOW COOH in this structure DO NOT ALLOW (sodium) salt of alcohol group i.e. $O_{-}CH_{2}$ $H_{2}N$ $O_{-}(Na^{+})$

Mark Scheme



Qu	Question		Answer	Marks	Guidance
18 ((a) ((i)	Number of peaks 2 marks 2-nitrophenol AND 3-nitrophenol have six peaks/environments/types of carbon ✓ 4-nitrophenol has four peaks/environments/types of carbon ✓ 4-nitrophenol has four peaks/environments/types of carbon ✓ Statement 1 mark 4-nitrophenol can be distinguished OR 2-nitrophenol and 3-nitrophenol cannot be distinguished ✓	3	IGNORE any numbers shown on structures ALLOW 1 mark only IF a response identifies that all the compounds have 6 peaks/environments/types of C OR all the compounds have 4 peaks/environments/ types of carbon IGNORE chemical shifts
	((ii)	 (In phenol) a (lone) pair of electrons on O is(partially) delocalised/donated into the π-system / ring ✓ Electron density increases/is higher (than benzene) ✓ ORA (phenol) is more susceptible to electrophilic attack OR (phenol) attracts/accepts electrophile/HNO₃ more OR (phenol) polarises electrophile/HNO₃ more ✓ ORA 	3	DO NOT ALLOW ECF from an incorrect number of peaks/environments/types of carbon ALLOW the electron pair in the p-orbitals of the O atom becomes part of the π -system / ring ALLOW diagram to show movement of lone pair into ring ALLOW lone pair of electrons on O is (partially) drawn/attracted/pulled/ into π -system / ring IGNORE activating IGNORE charge density IGNORE charge density IGNORE phenol reacts more readily <i>(no reference to electrophile)</i> ALLOW NO ₂ ⁺ for electrophile

Question	Answer	Marks	Guidance
(b)		3	ANNOTATE WITH TICKS AND CROSSES
			NOTE : curly arrows can be straight, snake-like, etc. but NOT double headed or half headed arrows
	Curly arrow from π -bond to S in SO ₃ AND		 1st curly arrow must go to the S of SO₃
	CH ₃		 start from, OR close to circle of benzene ring
	$ \begin{array}{c} \circ \\ S \\ \delta^{\dagger} \\ C \\ O \\ \delta^{-} \end{array} $		2nd curly arrow must start from, OR be traced back to, any part of S=O bond and go to O
			ALLOW 2nd curly arrow from S=O to any O in SO $_3$
			Intermediate must have correct SO ₃ ⁻ structure fully displayed

Questic	on	Answer	Marks	Guidance
		Correct intermediate ✓		DO NOT ALLOW the following intermediate:
		Correct intermediate \checkmark Curly arrow from C-H bond to reform π -ring \checkmark		DO NOT ALLOW the following intermediate: Γ_{H_3} $\Gamma_{++++++++++++++++++++++++++++++++++++$
				H H H H
		Total	9	

Q	uestior	Answer	Marks	Guidance
19	(a)	Links rate of reaction to strength of bond/bond enthalpy ✓ e.g. the weaker the bond the faster the reaction stronger bond takes longer to break lower bond enthalpy reacts faster	2	Each marking point must be a comparison
		Correct comparison of rate of reaction for at least two C– Hal bonds e.g. C–F bond is hydrolysed slow est C–I bond is hydrolysed faster than C–Br C–Br has shorter reaction time than C–CI OR		IGNORE references to halogens as elements: <i>i.e.</i> chlorine is less reactive than bromine etc. DO NOT ALLOW chloride, bromide and iodide
		Correct comparison of C–Hal bond strength/enthalpy of at least two of C–Hal bonds e.g. C–I bond is the weak est C–I has lower bond enthalpy than C–Br C–Br is broken more easily/readily than C–CI C–Hal bond strength decreases down group (7) ✓		IGNORE references to bond length, polarity and electronegativity

Question	Answer	Marks	Guidance
Question (b)	Answer Curly arrow from HO ⁻ to carbon atom of C–Cl bond \checkmark Dipole shown on C–Cl bond, $C^{\delta+}$ and $Cl^{\delta-}$ AND curly arrow from C–Cl bond to Cl atom \checkmark $^{\delta+}$ $^{O+}$ $O+$ $^{O+}$ $^{O+}$ $O+$ $^{O+}$ $O+$ $^{O+}$ $O+$	Marks 3	Guidance ANNOTATE ANSWER TICKS AND CROSSES NOTE: curly arrows can be straight, snake-like, etc. but NOT double headed or half headed arrows 1st curly arrow must • go to the C of C-Cl AND • start from, OR be traced back to any point across width of lone pair on O of OH ⁻ • OR start from - charge on O of OH ⁻ • OR start from - charge on O of ⁻ OH ion • OH OH OH • OH OH CH • OH OH • OH OH • OH OH
	GNORE presence of Na ⁺ but OH [−] needed i.e. Na ⁺ OH [−] can be allowed if criteria met Correct organic product AND Cl [−] \checkmark \checkmark \checkmark \Box		• OR start from – charge on O of ⁻ OH ion :OH OH :OH (Lone pair NOT needed if curly arrow shown from O ⁻) 2nd curly arrow must start from, OR be traced back to, any part of C–CI bond and go to CI C–CC C C C C C C C

Image: state of the state	Question	Answer	Marks	Guidance
Second mark Correct carbocation AND curly arrow from HO ⁻ to carbocation Image: Correct carbocation and the second curly arrow must come from lone pair on O of HO ⁻ OR OH ⁻ OR from minus on O of HO ⁻ ion (no need to show lone pair if curly came from negative charge) ✓ Third mark Correct organic product AND Cl ⁻ ✓				ALLOW S _N 1 mechanism First mark Dipole shown on C–Cl bond, C ^{δ^+} and Cl ^{δ^-} , AND curly arrow from C–Cl bond to Cl atom \checkmark \checkmark \checkmark \checkmark \checkmark \checkmark \checkmark \checkmark

Question Answer M	Marks	Guidance
Question Answer M (c) (i) Diagram Diagram showing round bottom/pear shaped flask AND upright condenser ✓ Water out ✓ Water out ✓ Water out ✓ Water out ✓ Condenser Water in (Round-bottom /pear-shaped) Water in Iss Iss Iss Iss Iss Iss Heat Labels (Round-bottom/pear-shaped) flask AND condenser AND water in at bottom and out at top AND heat (source) ✓	2	Guidance DO NOT ALLOW conical flask, volumetric flask, beaker in place of round bottom/pear shaped flask DO NOT ALLOW distillation DO NOT ALLOW stopper/bung on top of condenser IGNORE a thermometer in condenser IGNORE a small gap between flask and condenser IGNORE a small gap between flask and condenser IGNORE a small gap between flask and condenser IGNORE a label

Question	Answer	Marks	Guidance
(c) (ii	Precipitate G 1 mark silver bromide/AgBr AND $M = 1.88/0.01 = 188 \text{ (g mol}^{-1})$ 188 - 107.9 = 80.1 (so halide is Br ⁻)	3	 ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous Note: working is required for first mark ALLOW use of 108 as A_r of Ag
	Alcohol F and Haloalkane E 2 marks		
	E and F clearly identified F/alcohol: butan-2-ol H OH H_3C C C CH ₃ H H		Note: E and F can be identified by correct name or structure BUT IGNORE incorrect names
	 E/haloalkane: E is haloalkane of C₄H₉X with same halogen as G AND same carbon chain as F √ 		
	To	tal 10	

C	Questio	n	Answer	Marks	Guidance
20	(a)		priority groups/atoms are on different/opposite sides \checkmark High(est) priority groups are C ₆ H ₅ AND CHO OR Lowest priority groups are H and CH ₃ \checkmark	2	 ALLOW suitable alternatives to 'priority' e.g. groups with highest atomic number or more important groups etc. ALLOW high priority groups are diagonal(ly across) IGNORE references to relative mass of groups, <i>A</i>_r, <i>M</i>_r, ALLOW identification by name e.g aldehyde for CHO phenyl/benzene group for C₆H₅ alkyl for CH₃ ALLOW response in terms that O has higher priority than H in context of –CH₃ and –CHO IF 'priority' is not mentioned ALLOW 1 mark for 'C₆H₅ and CHO are on different sides' OR H and CH₃ are on different sides
	(b)	(i)	Bromine/ Br₂ AND goes colourless/decolourised ✓	1	Note: both reagent and observation are required ALLOW bromine water/ Br ₂ (aq)
		(ii)	Tollens' (reagent) AND Silver (mirror/precipitate/ppt/solid) √	1	 Note: both reagent and observation are required for the mark. ALLOW ammoniacal silver nitrate OR Ag⁺/NH₃ ALLOW black ppt OR grey ppt

Question	Answer	Marks	Guidance
(iii)	(Add) 2,4-dinitrophenylhydrazine AND orange/yellow/red precipitate√	3	ALLOW errors in spelling ALLOW 2,4(-)DNP OR 2,4(-)DNPH ALLOW Brady's reagent or Brady's Test ALLOW solid OR crystals OR ppt as alternatives for precipitate
	Take melting point (of crystals) ✓ Compare to known values/database ✓		Mark second and third points independently of response for first marking point DO NOT ALLOW 2 nd and 3 rd marks for taking and comparing boiling points OR chromatograms

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Question	Answer	Marks	Guidance
Question (d)*	AnswerPlease refer to marking instructions on page 5 of mark scheme for guidance on how to mark this question.Level 3 (5–6 marks)An outline of the mechanism for the formation of either product which is mostly correct.ANDMajor and minor products identified with a correct explanation of which product is most/least likely to be formed.There is a well-developed line of reasoning which is clear and 	Marks 6	Guidance Please check all of page 23 which is included with this response. If this page is blank please annotate with SEEN Throughout: ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above if unambiguous Indicative scientific points: Mechanism for formation of either product. • Curly arrow from C=C to attack the 1 atom of the 1-C1 • Curly arrow from negative charge on c1 • Curly arrow from negative charge on C1 • Curly arrow from negative charge on C1 ⁻ or lone pair on C1 ⁻ to carbon atom with positive charge • Curly arrow from negative charge on C1 ⁻ or lone pair on C1 ⁻ to carbon atom with positive charge • Curly arrow from negative charge on C1 ⁻ or lone pair on C1 ⁻ to carbon atom with positive charge • Curly arrow from negative charge on C1 ⁻ or lone pair on C1 ⁻ to c4H5 to
	product is attempted. OR Basic explanation of which of the products is most/least likely to be formed. <i>There is an attempt at a logical structure with a line of reasoning.</i> <i>The information is in the most part relevant.</i>		$H \xrightarrow{CHO} H \xrightarrow{H} CHO \xrightarrow{H} CHO \xrightarrow{H} C_{e}H_{5} \xrightarrow{C} C_{e}H_{5} \xrightarrow{C} C_{e}H_{5} \xrightarrow{C} C_{e}H_{3} \xrightarrow{H} C_{e}H_{5} \xrightarrow{C} C_{e}H_{3}$

Question	Answer	Marks	Guidance
Question	Answer 0 marks No response or no response worthy of credit.	Marks	Guidance Organic products • Major/most likely product C_6H_5 C_6H_5 C_1 C_6H_5 C_1 <
			• Major/most likely product is formed from the most stable carbocation intermediate OR – Cl is attached to carbon atom with the least hydrogens attached OR the carbon with the most –C atoms attached OR the – I is attached to the carbon atom
	Total	18	

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Question	Answer	Marks	Guidance
(c) (i)	FIRST CHECK ANSWER ON ANSWER LINE IF answer = 7.5 × 10 ⁻⁴ award 2 marks	2	If there is an alternative answer, Apply ECF
Question (c) (i)	Answer FIRST CHECK ANSWER ON ANSWER LINE IF answer = 7.5×10^{-4} award 2 marks [K] in mol dm ⁻³ $\frac{9.13 \times 10^{-2}}{166}$ = 5.50×10^{-4} (mol dm ⁻³) \checkmark [L] from peak areas $5.50 \times 10^{-4} \times \frac{5.9}{4.3}$ OR $5.50 \times 10^{-4} \times 1.37$ = 7.5×10^{-4} (mol dm ⁻³) \checkmark 2 SF Required	2 2	Guidance If there is an alternative answer, Apply ECF Alternative method [K] in g dm ⁻³ with peak area of 5.9 9.13 × 10 ⁻² × $\frac{5.9}{4.3}$ OR 9.13 × 10 ⁻² × 1.37 = = 0.125 OR 0.13 (g dm ⁻³) ✓ Calculator: 0.125272093 [L] in mol dm ⁻³ $\frac{0.125}{166}$ = 7.5 × 10 ⁻⁴ OR $\frac{0.13}{166}$ = 7.8 × 10 ⁻⁴ (mol dm ⁻³) ✓ Common errors: Award 1 mark for: 0.099(from $\frac{9.13 × 10^{-2}}{166} × 180$) • $0.9 × 10^{-4}$ (from $\frac{0.125}{180}$) • $7.2 × 10^{-4}$ (from $\frac{0.13}{180}$) • $7.0 × 10^{-4}$ (from $\frac{0.13}{180}$)
			<u>0.25272093</u>) 180

Question	Answer	Marks	Guidance
Question (ii)	ester J HO H	Marks 3	Guidance ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous L and M can be identified either way round IGNORE 'C ₃ H ₇ ' in L and/or M as ambiguous
	HO HO HO HO HO HO HO HO HO HO HO HO HO H		IGNORE connectivity of phenol OH group (marks are for structures of alkyl groups)
	Total	12	

Qı	uestion	Answer	Marks	Guidance
22	(a)	$C_7H_{16} + 7^1/_2O_2 \rightarrow 7CO + 8H_2O$ OR $C_7H_{16} + 4O_2 \rightarrow 7C + 8H_2O \checkmark$	1	ALLOW multiples IGNORE state symbols ALLOW equations for incomplete combustion that give CO and/or C with CO ₂ e.g C ₇ H ₁₆ + 9O ₂ \rightarrow 4CO + 3CO ₂ + 8H ₂ O C ₇ H ₁₆ + 6O ₂ \rightarrow 4CO + 3C + 8H ₂ O
	(b)	Heptane compared to hexane heptane (has a longer chain so) has more points of contact / more surface interaction (between molecules) ✓	4	ANNOTATE WITH TICKS AND CROSSES ALLOW ORA throughout
		heptane has stronger/more induced dipole(–dipole) interactions ✓		ALLOW heptane has more electrons IGNORE IDID
		Pentan-1-ol compared to heptane and/or hexane pentan-1-ol has hydrogen bonds that are strong(er than induced dipole–dipole interactions) OR (alcohols have) hydrogen bonds and induced dipole(-dipole)		ALLOW stronger/more London forces IGNORE van der Waals' forces/VDW for induced dipole– dipole interactions (<i>ambiguous as this term refers to both</i> <i>permanent dipole–dipole interactions and induced</i> <i>dipole–dipole interactions</i>)
		interactions/London forces ✓ Energy required to break forces		IGNORE 'pentan-1-ol can form hydrogen bonds with water'
		interactions in heptane than hexane OR More energy is required to break hydrogen bonds √		 ALLOW 'more energy to break intermolecular forces' if intermolecular forces are not stated. IGNORE it is harder to break the intermolecular forces <i>no reference to energy</i>) IGNORE more energy needed to separate molecules IGNORE more energy is needed to break bonds

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Question	Answer	Marks	Guidance
(c) (i)		5	Consult your team leader if an alternative creditworthy approach is seen
	<i>n</i> (CO ₂) = 2.97/44 = 0.0675 (mol) ✓		
	$n(H_2O) = 1.62/18 = 0.0900 \text{ (mol)}$		IGNORE ratio of CO_2 to H_2O is 3:4
	3:8√		OR a correct structure if not shown in working
	Molecular formula C₃H ₈ O₂✓		DO NOT ALLOW an incorrect molecular formula
	Structure any correct structure of $C_3H_8O_2 \checkmark$		
	e.g.		Mark independently from molecular formula but structure MUST contain 3C, 8H and 2O
	носсон		ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous
			ALLOW any vertical bond to the OH group e.g. ALLOW
			ОР ОН НО
	н—с—о—с—о—с—н 		DO NOT ALLOW OH-

Question	Answer	Marks	Guidance
Question (C) (ii)	Answer HO	Marks 2	Guidance ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous

Question (d)*

Answer	Marks	Guidance
Please refer to the marking instructions on page 5 of this mark scheme for guidance on how to mark this question.	6	Indicative scientific points: Empirical and Molecular Formula
Level 3 (5–6 marks) Compound is a structure of $C_6H_{12}O_3$ that is consistent with splitting pattern and chemical shifts in NMR spectrum. AND		• $C: H: O = 54.54/12 : 9.10/1 : 36.36/16$ 4.545 : 9.10 : 2.273 2 : 4 : 1
Comprehensive reasoning with most of the data analysed.		• Empirical formula = C_2H_4O
There is a well-developed line of reasoning which is clear and logically structured. The information presented is relevant and substantiated.		• uses $m/z = 132.0$ to determine molecular formula as $C_6H_{12}O_3$
Level 2 (3–4 marks)		¹ H NMR analysis
Compound has a feasible chemical structure that is consistent with the splitting pattern in NMR spectrum but may have incorrect molecular formula. AND Reasoning provided with some of the data analysed.		Spectrum:• $\delta = 4.0$ ppm, quartet, 1H,CH ₃ -CH-O• $\delta = 1.3$ ppm, singlet, 6H,(CH ₃) ₂ -C• $\delta = 1.2$ ppm, doublet, 3H,CH ₃ -CH-Without D-O:
There is a line of reasoning presented with some structure. The information presented is relevant and supported by some evidence.		 Peak at 11.0 ppm COOH or OH peak at 3.6 ppm OH Note: Data Sheet shows O-H chemical shift can occur
Level 1 (1–2 marks) Correct determination of empirical formula and/or molecular formula. OR Analyses most of the NMR data		around 11.0 ppm
OR		Structure
Attempts to determine empirical and/or molecular formula AND analyses some of the NMR data.		ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous
There is an attempt at a logical structure with a line of reasoning. The information is in the most part relevant.		Contains

Question	Answer	Marks	Guidance
	0 marks No response or no response worthy of credit.		 region that gives doublet and quartet e.g. H - C - C - C - C - C - C - C - C - C -
			Examples of structures consistent with splitting and chemical shift in NMR $H_3C - C - C - C - C - C - C - C - C - C -$

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Ques	tion	Answer	Marks	Guidance
				$ \begin{array}{cccccccccccccccccccccccccccccccccccc$
				H_3C O C CH_3 H_3C OH H_3C OH H_3C OH OH OH OH OH OH OH OH
				СН ₃ ОН H ₃ CСОН OСН СН ₃
				Note: there may be other possible structures that are consistent with the splitting pattern and chemical shifts in NMR – if an alternative structure is seen, please contact your team leader
		Total	18	

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