

GCE

Chemistry A

H432/02: Synthesis and analytical techniques

Advanced GCE

Mark Scheme for Autumn 2021

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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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1. Annotations

Annotation	Meaning
\checkmark	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
L1	Level 1
L2	Level 2
L3	Level 3
NBOD	Benefit of doubt not given
SEEN	Noted but no credit given
I	Ignore

2. 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
WA	Alternative wording
ORA	Or reverse argument

Question	Answer	Marks	AO element	Guidance
1	С	1	AO2.1	ALLOW 4 (This is the number of structural isomers)
2	В	1	AO1.2	
3	С	1	AO2.2	
4	С	1	AO2.6	
5	D	1	AO2.1	
6	В	1	AO1.2	
7	Α	1	AO1.2	
8	С	1	AO2.1	
9	С	1	AO1.2	
10	Α	1	AO2.1	
11	D	1	AO2.5	
12	В	1	AO2.1	
13	В	1	AO2.1	
14	С	1	AO1.1	
15	Α	1	AO1.2	
	Total	15		

C	Question		Answer	Marks	AO element	Guidance
16	(a)	(i)	σ-bond: Overlap of orbitals between (bonding) atoms $✓π$ -bond: Sideways overlap of (adjacent) p-orbitals $✓$	2	AO1.1 ×2	ALLOW labelled diagrams IGNORE the type of orbital for σ-bond DO NOT ALLOW pi-orbital
		(ii)	$ σ$ -bonds: 9 \checkmark $ π$ -bonds: 2 \checkmark	2	AO1.2 ×2	
	(b)	(i)	$H \rightarrow CH_2CH_3$ $H \rightarrow H^{\delta^+}$ $H \rightarrow H^{\delta^+}$ $H \rightarrow H^{\delta^+}$ Curly arrow from C=C bond to H of H-Br \checkmark DO NOT ALLOW partial charge on C=C Correct dipole shown on H-Br AND curly arrow showing breaking of H-Br bond \checkmark	4	AO1.2 ×2 AO2.5 ×2	NOTE: curly arrows can be straight, snake like, etc. but NOT double headed or half headed arrows 1st curly arrow must • go to the H atom of H–Br AND • start from, OR be traced back to any point across width of C=C C = C

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October 2021

Question	Answer	Marks	AO element	Guidance
	Correct carbocation AND curly arrow from Br ⁻ to C ⁺ of carbocation \checkmark DO NOT ALLOW δ + on C of carbocation $H - CH_2CH_3$ $H - CH_2CH_3$ $H - CH_2CH_3$ $H - CH_2CH_3$ $H - CH_2CH_3$			 3rd curly arrow must go to the C+ of carbocation AND start from, OR be traced back to any point across width of lone pair on :Br⁻ OR start from – charge of Br⁻ ion C⁺ C⁺ C⁺ B⁺ B⁺ (Lone pair NOT needed if curly arrow shown from – charge of Br⁻ ion)
	Correct product \checkmark H $\begin{array}{c} H \\ H $			ALLOW ECF for product from incorrect carbocation, i.e. H CH_2CH_3 H $-C$ $-C$ $-HH$ H $-C$ $-C$ $-HH$ H $-C$ $-C$ $-HH$ $-C$ $-C$ $-H-H$ $-C$ $-C$ $-H-H$ $-C$ $-C$ $-H$ $-H-H$ $-C$ $-C$ $-H$ $-H$ $-H$ $-C$ $-C$ $-H$ $-H$ $-H$ $-H$ $-C$ $-H$ $-H$ $-H$ $-H$ $-H$ $-H$ $-H$ $-H$

Question		Answer	Marks	AO element	Guidance
	(ii)	(major product forms from) most/more stable intermediate/carbocation ✓	2	AO1.1	For carbocation, ALLOW carbonium ion or cation
		(major product forms from a) secondary carbocation OR carbocation bonded to more C atoms / more alkyl groups OR carbocation bonded to fewer H atoms ✓		AO1.2	IGNORE descriptions of the major/minor product in terms of Markownikoff's rule e.g. H atom joins to C with most H
					IGNORE references to stability of the product
					ALLOW ORA, i.e. (minor product forms from) least/less stable intermediate/carbocation ✓
					(minor product forms from a) primary carbocation OR carbocation bonded to less C atoms / less alkyl groups
	(:::)	3 ✓	1	AO1.2	OR carbocation bonded to more H atoms \checkmark
	(iii)		1	AUT.2	
(c)	(i)	Same structural formula AND Different arrangement (of atoms) in space OR different spatial arrangement (of atoms) ✓	1	AO1.1	ALLOW structure/displayed/skeletal formula DO NOT ALLOW same empirical formula OR same general formula
					IGNORE same molecular formula Reference to <i>E</i> / <i>Z</i> isomerism or optical isomerism is not sufficient
	(ii)	Student is not correct AND 2 groups on one carbon atom (of C=C) are the same OR C–C bond can rotate ✓	1	AO3.1	DO NOT ALLOW one side of C=C

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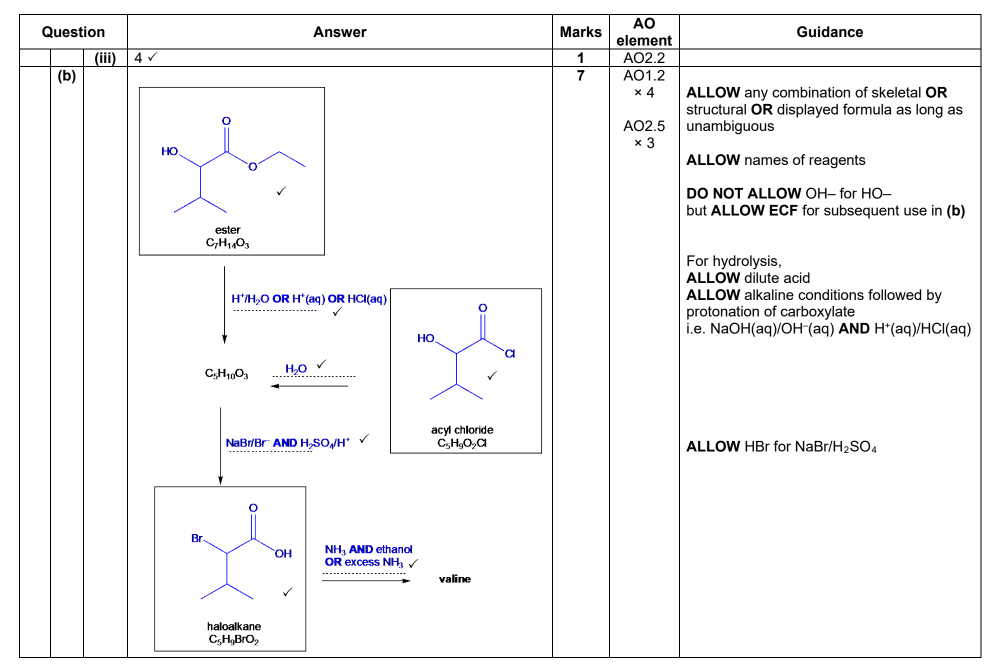
C	Question		Answer	Marks	AO element	Guidance
	(d)	(i)	1 mark for each curly arrow ✓✓	2	AO2.5 ×2	IGNORE any dipoles shown NOTE: curly arrows can be straight, snake- like, etc. but NOT half headed or double headed arrows Curly arrow from C=C bond must start from, OR be traced back to, Lower left: any part of C=C bond and go to C-C Upper left: any part of C=C bond and go to gap between C=C and C=C
		(ii)	OCH3 OCH3 OCH3 V	2	AO3.2 ×2	
			Total	17		

C	Question		Answer	Marks	AO element	Guidance	
17	(a)		Formation of Cl • $CClF_3 \rightarrow CF_3 \bullet + Cl \bullet \checkmark$	3	AO2.5	IGNORE dots for formation Cl^{\bullet} , i.e. ALLOW $CClF_3 \rightarrow CF_3 + Cl$	
			Breakdown of O_3 $Cl \cdot + O_3 \rightarrow \cdot ClO + O_2 \checkmark$		AO1.1 ×2	DO NOT ALLOW ECF Dots required in this equation	
			• $ClO + O \rightarrow Cl \cdot + O_2 \checkmark$			IGNORE O + O ₃ \rightarrow 2O ₂ ALLOW 1 mark if both equations are correct by atom but dot(s) missing or incorrect	
	(b)	(i)	F F C F CI ✓	1	AO2.5	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous 'End bonds' MUST be shown DO NOT ALLOW more than 1 repeat unit	
						IGNORE brackets	
		(ii)	More points of contact / more surface interaction (between molecules) AND Stronger/more dipole(–dipole) interactions √	2	AO2.1 ×2	Both answers need to be a comparison IGNORE surface area ALLOW more electrons ALLOW induced/permanent dipole interactions ALLOW London forces ALLOW van der Waals' forces (as permanent dipole-dipole and induced dipole-dipole interactions are present within this polymer) IGNORE IDID	
			More energy needed to break the intermolecular forces \checkmark				

Question	Answer	Marks	AO element	Guidance
Question (c)	Answer $ \begin{array}{c cccccccccccNH_2 \\ H_2N-C-C-C-C-N_{H_2} \\ H_H H_H H_H \\ \hline \\ \end{array} $ $ \begin{array}{c ccccccc} H_1 H_1 H_1 \\ \hline \\ H_1 H_1 H_1 \\ \hline \\ H_1 H_1 H_1 \\ \hline \\ H_1 \\$	Marks 4		For polymer, DO NOT ALLOW > 1 repeat unit 'End bonds' MUST be shown (do not have to be dotted) ALLOW -NH- at either end i.e. -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C -C
				IGNORE n
	Total	10		

Question	Answer	Marks	AO element	Guidance
18 (a) (i)	Non-superimposable mirror images (about a chiral centre) \checkmark	1	AO1.1	
	Correct groups attached to chiral C of alanine seen once e.g. $\begin{array}{c} $	2	AO2.1 × 2	Each structure must have four central bonds with at least two wedges . For bond into paper accept: ALLOW two 3D structures with 2 groups swapped e.g.

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Question	Answer	Marks	AO element	Guidance
(c) (i)	C ₁₃ H ₁₈ O ₂ ✓	1	AO2.1	ALLOW C, H and O in any order
(ii)	FIRST CHECK ANSWER ON THE ANSWER LINE If answer = 1.17×10^{21} award 3 marks	3	AO2.2 × 3	
	M (ibuprofen) = 206 \checkmark			ALLOW ECF from (c)(i)
	$n(\text{ibuprofen}) = \frac{400 \div 1000}{206} = 1.94 \times 10^{-3} \text{ (mol)} \checkmark$			Calculator: 1.941747573 × 10 ⁻³
	Number of molecules = $1.94 \times 10^{-3} \times 6.02 \times 10^{23}$ = 1.17×10^{21} to 3 SF \checkmark			ALLOW ECF from <i>n</i> (ibuprofen) 3 SF essential
(d) (i)		2	AO3.2 × 2	IGNORE small slip in carbon chains
	$ \begin{array}{c} $			ALLOW
(ii)	More soluble in water ✓	1	AO3.1	Answer must be a comparison ALLOW dissolve faster/quicker IGNORE absorbed more quickly (given in question)
	Total	18		

C	Questi	on	Answer	Marks	AO element	Guidance
19	(a)	(i)	3-methylbut-2-enal ✓	1	AO1.2	IGNORE lack of hyphens, or addition of commas
		(ii)			AO1.2 ×4 AO2.5 ×3	ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous ALLOW names of reagents and catalyst For oxidation, ALLOW K ₂ Cr ₂ O ₇ for Cr ₂ O ₇ ²⁻ ALLOW H ₂ SO ₄ for H ⁺ For left hand side esterification IGNORE C ₃ H ₇ OH IF esterification is given instead of hydrogenation contact your Team Leader

QuestionAnswerAO elementGuidance	
(b)*Refer to marking instructions on page 5 of mark scheme for guidance on marking this question.A02.4 $\times 2$ Indicative scientific points may incl Calculation of mass of C eH ₅ CH ₂ CI AND Planned synthesis to form the intermediate definited.A02.4 $\times 2$ Indicative scientific points may incl Calculation of mass of C eH ₅ CH ₂ CI AND Planned synthesis to form the intermediate $C_{eH_5}CH_2CI$ There is a well-developed line of reasoning which is clear and logically structured. The information presented is relevant and substantiated.A03.3 $\times 2$ A03.3 $\times 2$ Level 2 (3-4 marks) Correct calculation of the mass of CeH ₅ CH ₂ CI AND Planned synthesis to form the intermediate CeH ₅ CH ₂ CI is partly correct AND Planned synthesis includes formation of the intermediate CeH ₅ CH ₂ CI is partly correct AND Planned synthesis includes formation of the intermediate CeH ₅ CH ₂ CI followed by hydrolysis to form A with some of the reagents identified OR Attempts to calculate mass of CeH ₅ CH ₂ CI but makes little progress AND Planned synthesis includes formation of the intermediate CeH ₅ CH ₂ COI but makes little progress AND ALLOW small slip/rounding errors suc M_7 e.g. use of 137 instead of 136 for CeH ₅ CH ₂ COI but makes little progress ANDAND Planned synthesis includes formation of the intermediate C H ₂ CH ₂ CON AND ALLOW small slip/rounding errors suc M_7 e.g. use of 137 instead of 136 for CeH ₂ COH Examples of partly correct calculations Mass = 1.265 g from 0.0400 $\times \frac{25}{100} \times 12$	00 5

Question	Answer	Marks	AO element	Guidance
	There is a line of reasoning presented with some structure. The information presented is relevant and supported by some evidence.Level 1 (1-2 marks) Calculation of the mass of $C_6H_5CH_2Cl$ is partly correct 			Synthesis: reagents and conditionsStage 1: Formation of intermediate, $C_6H_5CH_2CN$ • Reagents: $CN^-(/ethanol)$ • Equation: $C_6H_5CH_2CI + CN^- \rightarrow C_6H_5CH_2CN + CI^-$ OR $C_6H_5CH_2CI + NaCN \rightarrow C_6H_5CH_2CN + NaCl$ (OR use of KCN)Stage 2: Formation of A, $C_6H_5CH_2COOH$ • Reagents: H^+/H_2O (ALLOW 'acid hydrolysis'• Equation: $C_6H_5CH_2CN + 2H_2O + H^+ \rightarrow C_6H_5CH_2COOH + NH_4^+$ OR $C_6H_5CH_2CN + 2H_2O + HCI \rightarrow C_6H_5CH_2COOH + NH_4CI$
	Total	18		

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C	Quest	ion	Answer	Marks	AO element	Guidance
20	(a)	(i)	Stage 1	6		ANNOTATE WITH TICKS AND CROSSES
			$(\bigcirc, \bigcirc, \downarrow, \bigcirc, \downarrow, \downarrow, \bigcirc, \downarrow, \downarrow,$		AO1.1 AO1.2 AO2.5	NOTE: curly arrows can be straight, snake-like, etc. but NOT double headed or half headed arrows Curly arrow from OH ⁻ must • go to the H of O-H AND • start from, OR be traced back to any point across width of lone pair on O of OH ⁻ \overrightarrow{OH} \overrightarrow{OH} \overrightarrow{OH} \overrightarrow{OH} • OR start from - charge ⁻ OH ion \overrightarrow{OH} \overrightarrow{OH} \overrightarrow{OH} Curly arrow from O-H bond must start from, OR be traced back to, any part of O-H bond and go to O IGNORE dipoles on O-H bond IGNORE Na ⁺

Mark Scheme

Question	Answer	Marks	AO element	Guidance
	Stage 2 Curly arrow from π -ring to C in CO ₂ AND curly arrow from the C=O bond to O atom \checkmark		AO2.5	 1st curly arrow must go to the C of CO₂ AND start from, OR close to circle of benzene ring Image: Image: Imag
	Correct intermediate ✓ Curly arrow from C–H bond to reform π-ring AND H ⁺ formed ✓		AO2.5 AO1.2	DO NOT ALLOW the following intermediate: $ \begin{array}{c} $

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Q	luest	on	Answer	Marks	AO element	Guidance
			$ \begin{array}{c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ $			DO NOT ALLOW mark for intermediate if phenolic O ⁻ is missing curly arrow must start from, OR be traced back to, any part of C-H bond and go inside the 'hexagon'
		(ii)	OH⁻: base ✓	2	AO2.1 ×2	ALLOW alkali IGNORE 'nucleophile', 'donates electron pair'
			CO_2 : electrophile OR electron pair acceptor \checkmark			IGNORE lone pair acceptor (No lone pair involved)
		(iii)	O	3	AO3.1	
			OH O		AO3.2	
			$2 \longrightarrow 0 + 2H_2O$		AO2.6	
			One ester link in organic product \checkmark			
			Correct structure of organic product \checkmark			
			Correct equation AND balanced ✓			

Questi	on	Answer	Marks	AO element	Guidance
(b)	(i)	 Dissolve in hot water/solvent ✓ Minimum amount of solvent ✓ Cool AND Filter AND (leave to) dry ✓ All three needed 	3	AO3.3 ×3	 ALLOW any solvent IGNORE Initial filtering hot filtration to remove insoluble impurities DO NOT ALLOW adding of a drying agent (e.g. MgSO₄)
	(ii)	C : H : N : O 31.44/12 : $1.31/1$: $18.34/14$: $48.91/16OR 2.62 : 1.31 : 1.31 : 3.06 \checkmark6:3:3:7ORC6H3N3O7 \checkmarkMolecular formula = C6H3N3O7AND use of M = 229.0 (directly linked to molecularformula) \checkmarkAny trisubstituted -NO2 substituted phenol that isconsistent with M = 229.0 \checkmarkEvidence for substitution2,4,6 OR 3,4,5 substituted phenolAND 4 peaks/ C environments from 13C NMR \checkmark2,4,6 substituted phenolAND directing effects of -OH \checkmark$	6	AO1.2 × 2 AO3.1 AO3.2 AO3.1 ×2	ALLOW alternative approach for empirical formula and evidence that 229 is equal to C ₆ H ₃ N ₃ O ₇ DO NOT ALLOW ECF from the empirical formula with the wrong molar ratio $O_{2N} + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (+++) + (++++) + (++++) + (++++) + (++++) + (++++) + (++++) + (++++) + (++++) + (++++) + (++++) + (++++) + (+++++) + (+++++) + (+++++) + (+++++) + (+++++) + (+++++) + (+++++) + (+++++) + (+++++) + (++++++) + (+++++) + (++++++) + (+++++) + (++++++) + (++++++) + (++++++) + (++++++) + (++++++) + (++++++) + (++++++) + (++++++) + (++++++) + (+++++++) + (+++++++) + (+++++++) + (++++++++$
		Total			

Question	Answer	Marks	AO element	Guidance
21*	Refer to marking instructions on page 5 of mark scheme for guidance on marking this question.	6	AO3.1 ×4	Indicative scientific points may include: Observations from Test-tube tests
	 Level 3 (5–6 marks) Compounds D, E AND F correctly identified AND Most of the observations and NMR data analysed. There is a well-developed line of reasoning which is clear and logically structured. The information presented is relevant and substantiated. Level 2 (3–4 marks) Most of compounds D, E AND F correctly identified AND Some of the observations and NMR data analysed. 		AO3.2 ×2	2,4 DNPD has no C=O E and F have C=O presentH+/Cr2O72-D is primary OR secondary alcohol E and F are ketones (negative test shows not aldehydes)Br2D, E and F have no C=C/are saturated $\frac{13}{C}$ NMR analysisD: • 3 carbon environments/types of C • $\delta = 24$, 36 ppmC-C • $\delta = 73$ ppm,C-O
	There is a line of reasoning presented with some structure. The information presented is relevant and supported by some evidence. Level 1 (1–2 marks) Most of compounds D, E AND F correctly identified OR Some of compounds D, E AND F correctly identified			¹ H NMR analysisE:• $\delta = 2.4$ ppm, quartetCH ₃ -CH ₂ -C=O• $\delta = 1.1$ ppm, tripletCH ₃ -CH ₂ -F:• $\delta = 2.6$ ppm, heptet/multiplet(CH ₃) ₂ -CH-C=O• $\delta = 2.1$ ppm, singlet,CH ₃ -C=O• $\delta = 1.1$ ppm, doubletCH ₃ -CH-
	AND Analyses some of the observations or NMR data OR Analyses most of the observations from the test-tube tests. OR Analyses most of the NMR data. OR Analyses some of the observations and NMR data			• $\sigma = 1.1$ ppm, doublet CH ₃ =CH= Structures ALLOW any combination of skeletal OR structural OR displayed formula as long as unambiguous OH D OH OH

Mark Scheme

Question	Answer	Marks	AO element	Guidance	
	There is an attempt at a logical structure with a line of reasoning. The information is in the most part relevant.				
	0 marks No response or no response worthy of credit.				
	Total	6			

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