

## **GCE**

# **Chemistry A**

Unit F322: Chains, Energy and Resources

Advanced Subsidiary GCE

Mark Scheme for June 2014

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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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## F322 Mark Scheme June 2014

Annotation	Meaning
BP	Blank Page – this annotation must be used on all blank pages within an answer booklet (structured or unstructured) and on each page of an additional object where there is no candidate response.
BOD	Benefit of doubt given
CON	Contradiction
×	Incorrect response
ECF	Error carried forward
I	Ignore
NAQ	Not answered question
NBOD	Benefit of doubt not given
POT	Power of 10 error
^	Omission mark
RE	Rounding error
SF	Error in number of significant figures
<b>✓</b>	Correct response

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

## **Subject-specific Marking Instructions**

The following questions should be annotated with ticks to show where marks have been awarded in the body of the text:

2(b), 3(a), 4(a), 4(b)(iii), 6(a)(i), 7(d), 8(a), 8(b)

All questions where an ECF has been applied.

#### **Checking additional pages**

All the Additional Pages in the examination script must be checked to see if any candidates include any answers.

- When you open question 1(a)(i) you will see a view of page 22 one of the Additional Pages.
- If the page is blank then, using the marking mode, annotate the page with an omission mark, ^.
- Scroll down to page 24 and annotate with a ^ if the page is blank.
- If pages 23 or 24 are not blank then use the paper clip icon to link the pages to the correct questions.

You may need to contact your Team Leader if you do not know how to do this.

#### **Generic comments**

#### **ORGANIC STRUCTURES**

For a 'structure' or 'structural formula',

• ALLOW correct structural OR displayed OR skeletal formula OR mixture of the above (as long as unambiguous)

For an alkyl group shown within a structure,

- **ALLOW** bond drawn to C or H, e.g. **ALLOW** CH<sub>3</sub>-,CH<sub>2</sub>-, C<sub>3</sub>H<sub>7</sub>-, etc.
- ALLOW vertical 'bond' to any part of an alkyl group

For an OH group shown within a structure,

- **DO NOT ALLOW** formula with horizontal —HO **OR** OH –
- ALLOW vertical 'bond' to any part of the OH group

For a CHO group shown within a structure,

DO NOT ALLOW COH

#### For a 3D structure.

For bond in the plane of paper, a solid line is expected:	
For bond out of plane of paper, a solid wedge is expected:	
For bond into plane of paper, ALLOW:	Minning Minning Minning
ALLOW a hollow wedge for 'in bond' OR an 'out bond', provided it is different from the other in or out wedge e.g.:	

#### **NAMES**

Names including alkyl groups:

- ALLOW alkanyl, e.g. ethanyl (i.e. IGNORE 'an')
- **DO NOT ALLOW** alkol, e.g. ethol (ie 'an' is essential)

#### Names of esters:

- Two words are expected, e.g. ethyl ethanoate
- ALLOW one word, e.g. ethylethanoate

Names with multiple numbers and hyphens:

Use of 'e'

- ALLOW superfluous 'e', e.g. propane-1-ol ('e' is kept if followed by consonant)
- **ALLOW** absence of 'e', e.g. propan-1,2-diol ('e' is omitted if followed by vowel)

Hyphens separate name from numbers:

ALLOW absence of hyphens, e.g. propane 1,2 diol

Multiple locant numbers must be clearly separated:

- ALLOW full stops: e.g. 1.2 OR spaces: 1 2
- DO NOT ALLOW e.g. 12

Locant numbers in formula must be correct

• **DO NOT ALLOW** propan-3-ol

Order of substituents should be alphabetical:

• ALLOW any order (as long as unambiguous), e.g. 2-chloro-3-bromobutane

#### **ABBREVIATIONS**

van der Waal's forces

**ALLOW** vdw forces **OR** VDW forces (and any combination of upper and lower cases)

C	uesti	ion	Answer	Mark	Guidance
1	(a)	(i)	(series of compounds with the) same functional group OR same/similar chemical properties OR same/similar chemical reactions ✓  each successive/subsequent member differing by CH₂ ✓	2	IGNORE references to physical properties IGNORE has same general formula (in question) DO NOT ALLOW have the same empirical formula OR have the same molecular formula
		(ii)	C <sub>n</sub> H <sub>2n</sub> ✓	1	
		(iii)	More carbons (in ring) OR more (surface area of) contact  AND more van der Waals forces OR stronger van der Waals forces ✓	2	Both answers need to be comparisons ALLOW ORA throughout  ALLOW has more electrons OR larger (carbon) ring OR higher molecular mass IGNORE bigger molecule IGNORE chain instead of ring DO NOT ALLOW 'more contact between atoms'  ALLOW 'VDW' for van der Waals 'More intermolecular forces' is not sufficient
			More energy needed to break the intermolecular forces ✓		ALLOW it is harder to overcome the intermolecular forces ALLOW intermolecular bonds / van der Waals bonds ALLOW more energy is needed to separate molecules IGNORE more energy is needed to break bonds

Qu	estion	Answer	Mark	Guidance
	(b)	tetrahedral ✓	2	Mark each point independently
		four <b>bonding</b> pairs repel <b>OR</b> four <b>bonds</b> repel ✓		IGNORE surrounded by four atoms IGNORE four areas of electron charge repel IGNORE four electron pairs repel (one could be lp) DO NOT ALLOW atoms repel
	(c)	Br — C — C — Br	1	ALLOW correct structural OR displayed OR skeletal formula OR mixture of the above (as long as unambiguous)  ALLOW structure of 1,2-isomer  H H H H H H H H H H H H H H H H H H H
	(d) (i)	$C_6H_{14} \rightarrow C_6H_{12} + H_2 \checkmark$	1	ALLOW correct structural OR displayed OR skeletal formula OR mixture of the above (as long as unambiguous)  ALLOW any correct multiple  IGNORE state symbols

Qı	uestic	on	Answer	Mark	Guidance
		(ii)	Cyclohexane will burn more efficiently ✓	1	KEY IDEA IS COMBUSTION OR BURNING
					Assume 'it' refers to cyclohexane ALLOW ORA for hexane
					ALLOW cyclohexane allows smoother burning OR promotes more efficient combustion OR increases octane number OR reduces knocking OR less likely to produce pre-ignition OR burns better OR easier to burn OR combusts more easily OR improves combustion OR burns more cleanly DO NOT ALLOW cyclohexane ignites more easily IGNORE cyclohexane increase volatility of fuel IGNORE reference to boiling points IGNORE cyclohexane gives a better fuel
	(e)	(i)	(Compounds with the) same structural formula but a different arrangement (of atoms) in space ✓	1	ALLOW different spatial arrangement of atoms.  DO NOT ALLOW different displayed formula.
		(ii)	$H_3C$ $CH_3$ $H_3C$ $CH_3$ $CH_3$ $CH_3$	2	ALLOW displayed OR skeletal formula OR mixture of the above.  ALLOW structures in either order  IGNORE molecular formula  IGNORE structural formula  IGNORE names  IGNORE E/Z and cis/trans labels  ALLOW 1 mark for a pair of E/Z isomers of an incorrect hydrocarbon structure with four C atoms e.g. C, or CH or CH <sub>2</sub> instead of CH <sub>3</sub> groups.

Question		Answer	Mark	Guidance
(f) (i)	Step Initiation (1 mark)  Propagation (2 marks)  Termination (2 marks)	Equation $Br_{2} \rightarrow 2Br^{\bullet} \checkmark$ $C_{6}H_{12} + Br^{\bullet} \rightarrow C_{6}H_{11}^{\bullet} + HBr \checkmark$ $C_{6}H_{11}^{\bullet} + Br_{2} \rightarrow C_{6}H_{11}Br + Br^{\bullet} \checkmark$ $C_{6}H_{11}^{\bullet} + Br^{\bullet} \rightarrow C_{6}H_{11}Br$ $C_{6}H_{11}^{\bullet} + C_{6}H_{11}^{\bullet} \rightarrow C_{12}H_{22}$ $Br^{\bullet} + Br^{\bullet} \rightarrow Br_{2}$ Two correct $\checkmark$ All three correct $\checkmark\checkmark$	5	IGNORE state symbols  IGNORE dots  If an incorrect hydrocarbon with six C atoms is used:  DO NOT ALLOW any marks for the propagation steps but ALLOW ECF for termination steps (i.e. 3 max)
(ii)	OR the breaking	ng of a (Br-Br) bond <b>AND</b> forms (two) radicals g of a (Br-Br) bond <b>AND</b> one electron (from the goes to each atom/bromine ✓	1	ALLOW the breaking of a covalent bond' ALLOW the splitting of the bond in bromine  ALLOW the breaking of a covalent bond where each atom keeps one of the bonding electrons IGNORE particle for atom ALLOW one electron goes to each product / species DO NOT ALLOW molecule or compound for atom IGNORE homolytic fission equations
(g) (i)	$C_6H_{12} + 2B$	$r_2 \rightarrow C_6H_{10}Br_2 + 2HBr \checkmark$	1	ALLOW molecular formula only.
(ii)	OR 1,2-dibi	omocyclohexane romocyclohexane romocyclohexane romocyclohexane ✓	1	Locant numbers MUST lowest possible e.g. DO NOT ALLOW 2,4-dibromocyclohexane etc.  IGNORE structures
		Total	21	

C	uestion	Answer	Mark	Guidance
2	(a)	It is an electron pair donor <b>OR</b> can donate a lone pair ✓	1	
	(b)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	3	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC  IGNORE connectivity to C <sub>3</sub> H <sub>7</sub> throughout
		CCH <sub>3</sub> Dipole shown on the C-Br bond, $C^{\delta +}$ and $Br^{\delta -}$ and curly arrow from the C-Br bond to the Br atom $\checkmark$		IGNORE alkyl group in first marking point. Curly arrow must start from C–Br bond and not from C atom. Dipole must be partial charge and not full charge
		Curly arrow from :OCH₃ to carbon atom in the C-Br bond ✓ Correct organic product ✓ S <sub>N</sub> 1 mechanism		CH <sub>3</sub> O <sup>-</sup> curly arrow must come from one lone pair on O of CH <sub>3</sub> O <sup>-</sup> ion <b>OR</b> from negative sign on O of the CH <sub>3</sub> O <sup>-</sup> ion <b>ALLOW</b> arrow from lone pair on O in OCH <sub>3</sub> <sup>-</sup> Lone pair not required <b>DO NOT ALLOW</b> CH <sub>3</sub> O <sup>δ</sup> -
		$C_3H_7$ $H \longrightarrow C \xrightarrow{\delta^+ \frown} Br^{\delta^-}$		DO NOT ALLOW incorrect connectivity of CH <sub>3</sub> O group in the final product –CH <sub>3</sub> O IGNORE Br <sup>δ-</sup> as a product
		$H$ $C_3H_7$ $H \longrightarrow C^+$ $\vdots$ $OCH_3$ $H_3CO \longrightarrow C_3H_7$ $\vdots$ $H_3CO \longrightarrow H$		ALLOW $S_N1$ mechanism Dipole shown on the C-Br bond, $C^{\delta^+}$ and $Br^{\delta^-}$ and curly arrow from C-Br bond to the Br atom $\checkmark$ curly arrow from $CH_3O^-$ to carbonium ion $\checkmark$ correct organic product $\checkmark$

Question	Answer	Mark	Guidance
(c)	1-lodobutane increases the rate  AND	1	
	C—I bonds are weaker (than C—Br)  OR C—I bond has a lower bond enthalpy  OR C—I bond needs a smaller amount of energy to break  OR C—I bond is easier to break ✓		All statements must be comparative ALLOW ORA IGNORE C—I bond is longer IGNORE polarity and references to electronegativity
(d)		2	ALLOW only skeletal formula
	butyl ethanoate ✓		DO NOT ALLOW ECF from incorrect structure. ALLOW butylethanoate ALLOW butanyl for butyl DO NOT ALLOW butly
(e) (i)	( <u>136.9</u> × 100) = 47% 291.1√	1	ALLOW 47 up to calculator value correctly rounded. 47.0 or 47.03 or 47.029 will be correct common answers
			IGNORE any working shown.
(e) (ii	NaBr <b>OR</b> LiBr ✓	1	ALLOW correct name or formula  DO NOT ALLOW HBr (it is an acid)
(e) (iii	Look at answer if 88.8% AWARD 3 marks if 88.75% AWARD 2 marks (not 3 sig. fig.)  Moles of butan-1-ol = 0.08(00) ✓	3	Answer MUST be to 3 significant figures.  ALLOW ECF but do not allow a yield >100%
	Moles of 1-bromobutane = 0.071(0) ✓		ALLOW Mass of 1-bromobutane expected = 10.952 g
	% yield = 88.8% ✓  Total	12	
	lotai	12	

Question		Mark	Guidance
3 (a)	There are 3 marking points required for 2 marks $H_2(g) + I_2(g)$ $\Delta H$ $2HI(g)$ $H_2 \text{ and } I_2 \text{ on LHS}$ $\mathbf{AND} \text{ 2HI on RHS}$ $\mathbf{AND} \text{ correctly labelled Ea} \checkmark$ $\Delta H \text{ labelled with product below reactant}$ $\mathbf{AND} \text{ arrow downwards} \checkmark$	2	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC  IGNORE state symbols.  E <sub>a</sub> :  ALLOW (+)173 only as an alternative label for Ea ALLOW no arrowhead or arrowheads at both ends of activation energy line  The E <sub>a</sub> line must point to maximum (or near to the maximum) on the curve OR span approximately 80% of the distance between reactants and maximum regardless of position  ALLOW AE or A <sub>E</sub> for E <sub>a</sub> ΔH:  IF there is no ΔH labelled ALLOW –9 as an alternative label for ΔH.  IF ΔH is labelled IGNORE any numerical value.  DO NOT ALLOW –ΔH.  ALLOW this arrow even if it has a small gap at the top and bottom i.e. does not quite reach reactant or product line
(b)	(+)182 ✓	1	This is the <b>ONLY</b> acceptable answer

Quest	ion	Answer	Mark	Guidance
(c)		Look at answer if +63 kJ AWARD 2 marks If 63 (no sign) OR-63 (incorrect sign) AWARD 1 mark  No of moles of HI = 14 moles ✓	2	ALLOW one mark for +126 kJ
		Enthalpy Change = +63 kJ ✓		Sign and value required.  ALLOW ECF from incorrect number of moles of HI
(d)	(i)	Rate of the forward reaction is equal to the rate of the reverse reaction ✓  OR	1	ALLOW both reactions occur at same rate
		concentrations do not change√		IGNORE conc. of reactants = conc. of products
	(ii)		2	Mark each point independently
		More H₂ and I₂ <b>OR</b> less HI ✓		ALLOW more reactants OR less products
		(equilibrium position shifts) to the left  AND  (Forward) reaction is exothermic  OR reverse reaction is endothermic  OR in the endothermic direction✓		Note: ALLOW suitable alternatives for to the left e.g. towards reactants $ \begin{array}{c} \text{OR towards H}_2  /  I_2 \\ \text{OR in reverse direction} \\ \text{OR favours the left.} \end{array} $
				ALLOW gives out heat for exothermic ALLOW takes in heat for endothermic
	(****	N. "		IGNORE responses in terms of rate
	(iii)	No effect AND	1	
		Same number of (gaseous) moles on both sides ✓		ALLOW same number of molecules on each side

C	Question	Answer	Mark	Guidance
	(e)	Look at answer if (+)298 AWARD 2 marks	2	
		If answer is -298 AWARD 1 mark (incorrect sign)		
		2 x H-I bond enthalpy correctly calculated		
		(436 +151-(-9) =) (+)596 ✓		<b>ALLOW</b> 1 mark for (+)293.5 kJ mol <sup>-1</sup> (bonds broken divided
		H-I bond enthalpy correctly calculated		by 2)
		(Bond energy for H-I <u>(+)596</u> =) (+)298 kJ mol <sup>-1</sup> ✓		<b>ALLOW</b> 1 mark for (+)289 kJ mol <sup>-1</sup> (incorrect expression i.e. [436 +151+(-9)])
		2		2
		Tota	11	

Q	uesti	on	Answer	Mark	Guidance
4	(a)		FIRST, CHECK THE ANSWER ON ANSWER LINE IF answer = -38.3 (kJ mol <sup>-1</sup> ) award 4 marks IF answer = (+)38.3 (kJ mol <sup>-1</sup> ) award 3 marks (incorrect sign) IF answer = -38,300 (kJ mol <sup>-1</sup> ) award 3 marks (used J instead of kJ).	4	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC
			Energy $q$ calculated correctly = 1149.5(J) $\checkmark$ OR 1.1495 (kJ) $\checkmark$		Note: $q = 50.0 \times 4.18 \times 5.5$ ALLOW 1149.5 OR correctly rounded to 3 sig figs (J) IGNORE sign IGNORE working ALLOW 53.18 × 4.18 × 5.5 OR 1222.6082 OR 1220 OR correctly rounded to 3 or more sig figs in J or kJ
			Moles Amount, $n$ , of Na <sub>2</sub> CO <sub>3</sub> calculated correctly= 0.03(00) $\checkmark$		IGNORE working IGNORE trailing zeros
			Calculating ΔH correctly calculates ΔH in kJ mol <sup>-1</sup> to 3 or more sig figs ✓		IONODE since at this integral distant
			Rounding and Sign calculated value of $\Delta H$ rounded to 3 sig. fig. with minus		IGNORE sign at this intermediate stage ALLOW ECF from incorrect q and/or incorrect n
			sign√		Final answer must have correct sign and three sig figs
					<b>ALLOW</b> –40.8 kJ mol <sup>-1</sup> if 53.18 used in calculation of q <b>ALLOW</b> –40.7 kJ mol <sup>-1</sup> if q is rounded to 1220 from 53.18 earlier
	(b)	(i)	(Enthalpy change) when one mole of a compound ✓ is formed from its elements ✓	3	ALLOW energy required OR energy released ALLOW one mole of substance OR one mole of product DO NOT ALLOW one mole of element
			298 K / 25 °C <b>AND</b> 1 atm / 100 kPa / 101 kPa / 1 bar ✓		IGNORE reference to concentration

Question	Answer	Mark	Guidance
(ii)	1/2N <sub>2</sub> (g) + 2H <sub>2</sub> (g) + $1/2$ Cl <sub>2</sub> (g) + 2O <sub>2</sub> (g) → NH <sub>4</sub> ClO <sub>4</sub> (s) correct species ✓ correct state symbols <b>and</b> balancing ✓	2	Second mark can only be awarded if all species in the equation are correct
	·		DO NOT ALLOW multiples of this equation
(iii)	FIRST, CHECK THE ANSWER ON ANSWER LINE IF answer = (+)90 award 3 marks IF answer = -90 award 2 marks IF answer = ±270 award 2 marks IF answer = ±2947 award 1 mark	3	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC
	Processing ∆H <sub>f</sub> values		
	±(3832 − 885) ±2947✓		<b>Note:</b> $\pm 2947 = \pm [-1676 + (-704) + (6 \times -242)] - (3 \times -295)]$
	OR		
	± (3832 – 885)		
	subtraction using ∆H reaction		
	±(2947-2677)= ±270 ✓		
	Calculation of $\Delta H$ formation NO $270/3 = (+)90 \checkmark$		<b>ALLOW ECF</b> for dividing by 3 from working that includes at least one $\Delta H_{\rm f}$ and one balancing number and $\Delta H$ (-2677) for 1 mark
	Total	12	

Q	uesti	on	Answer	Mark	Guidance
5	(a)		$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	2	Displayed formulae MUST be used to award each mark
			Correct polymer with side links ✓  Balanced equation for formation of correct polymer - correct use of <i>n</i> in the equation and brackets ✓		n on LHS can be at any height to the left of formula <b>AND</b> n on the RHS must be a subscript (essentially below the side link)
	(b)	(i)	$CH_2CHCl + 2O_2 \longrightarrow CO + CO_2 + HCl + H_2O \checkmark$	1	ALLOW any other correctly balanced equation with the same reactants and products ALLOW C <sub>2</sub> H <sub>3</sub> C <i>l</i> for CH <sub>2</sub> CHC <i>l</i>
		(ii)	Sodium hydrogencarbonate neutralises HC <i>l</i> ✓	1	Assume that 'it' refers to sodium hydrogencarbonate but DO NOT ALLOW other chemicals e.g. sodium  ALLOW NaHCO <sub>3</sub> is a base ALLOW forms a salt or sodium chloride or NaCl ALLOW equation to show formation of NaCl from NaHCO <sub>3</sub> and HCl even if not balanced.  IGNORE reacts

Question	Answer	Mark	Guidance
(c)	abundance (in atmosphere) OR amount (in atmosphere) OR (atmospheric) concentration OR percentage (in air) ✓ OR ability to absorb infrared/IR (radiation) ✓ OR	2	ALLOW absorption of infrared/IR
(d) (i)	residence time ✓  Any balanced equation between a metal oxide and carbon dioxide to form a carbonate e.g CaO + CO₂    CaCO₃ ✓	1	ALLOW MO for metal oxide
(ii)		1	Assume that 'it' refers to carbon dioxide but DO NOT ALLOW carbon  DO NOT ALLOW reacted with oxides or stored as carbonates.
	Total	8	

Question	Answer	Mark	Guidance
6 (a) (i)		4	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC
	Number of Molecules  T <sub>1</sub> T <sub>2</sub>		Candidates do not need Ea on graph
	axes labelled (number of) molecules and (kinetic) energy ✓		ALLOW particles instead of molecules on the y axis DO NOT ALLOW atoms instead of particles/molecules ALLOW ECF for the incorrect use of atoms (instead of molecules/particles) DO NOT ALLOW enthalpy on the x-axis
	Compat drawing a of a two Doltzmann a distributions		<b>DO NOT ALLOW</b> increase of more than one small square at high energy end of curve.
	Correct drawing of a two Boltzmann distributions i.e. both curves must start within the first small square nearest to the origin <b>AND</b> must not touch the x axis at high energy ✓		Maximum of curve for higher temperature to right  AND lower than maximum of lower temperature curve  AND above lower temp line at higher energy  Higher temp line should intersect lower temp line once
	Drawing of Boltzmann distribution at <b>two</b> different temperatures with higher and lower temperature clearly identified ( ie $T_2 > T_1$ ) $\checkmark$		DO NOT ALLOW lower activation energy QWC requires more molecules have or exceed activation energy/E <sub>a</sub> . IGNORE more molecules have enough energy to
	QWC - (At a higher temperature) more molecules have energy above activation energy OR greater area under the curve above the activation energy✓		react for the <b>QWC</b> mark (as not linked to E <sub>a</sub> ) <b>ORA</b> if states the effect when the temperature is lower <b>IGNORE</b> (more) successful collisions

Question	Answer	Mark	Guidance
(a) (ii)	(Decreasing the pressure) decreases the rate of reaction  AND	2	Correct effect on rate must be linked to reason for the first marking point.
	Decreased concentration of molecules  OR  Number of molecules remains the same but the volume increases  OR  Less molecules per (unit) volume ✓		ALLOW molecules are further apart IGNORE less crowded ALLOW particles or atoms for molecules ALLOW 'space' for volume DO NOT ALLOW area instead of volume
	Less <b>frequent</b> collisions ✓		ALLOW collisions occur less often OR decreased rate of collision IGNORE less chance of collisions
			'less collisions' alone is <b>not</b> sufficient <b>IGNORE</b> successful
(b) (i)	$C_6H_{12}O_6 \longrightarrow 2C_2H_5OH + 2CO_2 \checkmark$	2	ALLOW correct molecular OR structural OR displayed OR skeletal formula OR mixture of the above (as long as unambiguous) IGNORE state symbols
	Temperature: Between 20 °C and 45 °C inclusive AND Condition: Absence of oxygen OR anaerobic ✓		DO NOT ALLOW acidic or alkaline conditions If there is a contradiction or an incorrect answer in any condition given then do not award this mark. ALLOW conditions shown in the equation A limited supply of oxygen is not sufficient IGNORE pressure IGNORE yeast (in question) ALLOW Lack of oxygen
(b) (ii)	2NO + 2CO → 2CO <sub>2</sub> + N <sub>2</sub> ✓	1	ALLOW multiples IGNORE state symbols
	Total	9	

Question	Answer	Mark	Guidance
7 (a)	CH <sub>3</sub> CH <sub>3</sub>	1	ALLOW correct structural OR displayed OR skeletal formula OR mixture of the above  DO NOT ALLOW molecular formula  ALLOW dichloro or diiodo compound instead of the dibromo compound as the only alternatives.
(b)	Reagent <b>A</b> : correct halogen ✓ e.g. Br <sub>2</sub> / bromine	1	ALLOW Cl <sub>2</sub> if dichloro compound drawn ALLOW I <sub>2</sub> if diiodo compound drawn  IGNORE state symbols Answer must match box from (a) to score
(c) (i)	Steam <b>AND</b> acid catalyst ✓	1	ALLOW H <sup>+</sup> / named acid / H <sub>2</sub> SO <sub>4</sub> / H <sub>3</sub> PO <sub>4</sub> ALLOW H <sub>2</sub> O(g) ALLOW water only if a temperature of 100 °C or above is quoted.  IGNORE any temperature given with steam IGNORE pressure
(ii)	(compounds or molecules) having the same molecular formula but different structural formulae ✓	1	ALLOW different structure OR different displayed formula OR different skeletal formula for structure  Same formula is <b>not</b> sufficient  Different arrangement of atoms is <b>not</b> sufficient
(iii)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	2	ALLOW correct structural OR displayed OR skeletal formula OR mixture of the above ALLOW any vertical bond to OH DO NOT ALLOW OH—
(iv)	Does not contain OH group(s)  OR does not contain hydroxyl group(s)  OR is not an alcohol ✓	2	ALLOW ORA throughout DO NOT ALLOW OH (ions) / hydroxide (ions)
	Does not form hydrogen bonds with water ✓		'Does not form hydrogen bonds' is <b>not</b> sufficient

Question	Answer	Mark	Guidance
(d)	Reagents: Acid/H <sup>+</sup> and (potassium or sodium) dichromate/Cr <sub>2</sub> O <sub>7</sub> <sup>2-</sup> seen <b>once</b> ✓	6	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC
	Observations: Orange to Green <b>OR</b> Orange to Blue✓		ALLOW H <sub>2</sub> SO <sub>4</sub> and K <sub>2</sub> Cr <sub>2</sub> O <sub>7</sub>
	Distillation / Distil produces aldehyde/CH <sub>3</sub> CH <sub>2</sub> CHO: ✓ CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> OH + [O] → CH <sub>3</sub> CH <sub>2</sub> CHO + H <sub>2</sub> O ✓		ALLOW correct displayed formula OR correct structural formula OR skeletal formula OR a mixture of
			the above DO NOT ALLOW molecular formulae
			<b>ALLOW</b> C <sub>3</sub> H <sub>7</sub> OH for propan-1-ol in equations
			DO NOT ALLOW CH <sub>3</sub> CH <sub>2</sub> COH for aldehyde
	Reflux (of propan-1-ol) produces carboxylic acid/CH₃CH₂COOH✓		IGNORE further oxidation of aldehyde
	CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> OH + 2[O] → CH <sub>3</sub> CH <sub>2</sub> COOH + H <sub>2</sub> O ✓		<b>ALLOW</b> CH <sub>3</sub> CH <sub>2</sub> CO <sub>2</sub> H for carboxylic acid
	Total	14	

Question	Answer	Mark	Guidance
8 (a)	Molar mass of <b>B</b> = 74 ✓	6	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC
	B-F clearly identified		Check and annotate page 19 below this response
	B/alcohol:		Molar mass = <u>2.59</u> = 74 0.035
	H H OH H		For structure of <b>B</b> , <b>C</b> , <b>D</b> or <b>E/F ALLOW</b> correct displayed <b>OR</b> correct structural formula <b>OR</b> correct skeletal formula <b>OR</b> mixture of the above as long as unambiguous.
	C/ketone:		<b>DO NOT ALLOW</b> missing H atom(s) in a displayed formula for one structure but <b>ALLOW</b> missing H atoms in subsequent structures.
	H—C—C—C—H		IGNORE names of organic compounds
	D/carboxylic acid:		
	H—————————————————————————————————————		E and F can be identified either way round  ALLOW H <sub>2</sub> O or displayed formula for mark
	E and F:		For <b>E</b> and <b>F</b> – <b>ALLOW</b> the two optical isomers
	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		$C_2H_5$ $C_2H_5$ $C_2H_5$ $C_2H_5$ $C_2H_5$ $C_2H_5$ $C_2H_5$ $C_2H_5$ $C_2H_5$
	H₂O/water ✓		

Question	Answer	Mark	Guidance
8 (b)	Mole ratio C : H : O = $\frac{55.8}{12.0} : \frac{7.0}{1.0} : \frac{37.2}{16.0}$	7	ANNOTATE ANSWER WITH TICKS AND CROSSES ETC
	<b>OR</b> $4.65:7.0:2.33/2.325$ <b>OR</b> $2:3:1$ <b>OR</b> $C_2H_3O$ $\checkmark$ Molecular formula of <b>G</b> $C_4H_6O_2$ $\checkmark$		<b>ALLOW</b> mass of C = 0.558 x 86 <b>or</b> 48 <b>AND</b> mass of H = 0.07 x 86 <b>or</b> 6 <b>AND</b> mass of O = 0.372 x 86 = 32
	Mass spectrum for G: 2 marks		
	Peak <b>X or peak 41</b> indicates C <sub>3</sub> H <sub>5</sub> <sup>+</sup> ✓		
	Peak Y or peak 45 indicates COOH <sup>+</sup> ✓		+ charge required for each response <b>ALLOW</b> one mark if both formulae are correct but with no charge/incorrect charge
	Infrared for G: 1 mark		<b>ALLOW</b> any possible fragments that contain C, H and/or O that have the correct mass. E.g. Peak X indicates C <sub>2</sub> OH <sup>+</sup> , Peak Y indicates C <sub>2</sub> H <sub>5</sub> O <sup>+</sup> Unfeasible fragments are not allowed e.g. C <sub>3</sub> H <sub>9</sub> <sup>+</sup> (too many H atoms)
	Peak at 1640–1750 cm <sup>-1</sup> indicates presence of C=O <b>AND</b> Peak at 2500–3300 cm <sup>-1</sup> (indicates the presence of) –OH group linked carboxylic acid/COOH <b>QWC</b> ✓		LOOK ON THE SPECTRUM for labelled absorbance which can be given credit Candidates must link absorbance to bond in order to gain the mark
			<b>ALLOW</b> 1700 cm <sup>-1</sup>
			For 2500–3300 cm <sup>-1</sup> , <b>ALLOW</b> 2900 cm <sup>-1</sup> or any stated wavenumber with range 2500–3300 cm <sup>-1</sup> <b>ALLOW</b> wavenumber range up to 2400–3500 cm <sup>-1</sup>

Question	Answer	Mark	Guidance
Question	Answer  Structure of G:  Correct structure: $ \begin{array}{cccccccccccccccccccccccccccccccccc$	Mark	ALLOW structural, skeletal or displayed formula.  DO NOT ALLOW ECF from incorrect molecular formula
	OR ——COOH		
	Total	13	

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