1. From the evidence, candidates may have identified compound **F** as propanone, propanal or propanoic acid

If **F** is propanone or propanoic acid, then maximum score = 7; **but** if **F** is propanal then maximum score = 6

The mark scheme for \mathbf{F} = propanone and propanal is shown below.

mass spec of E- Remember to check the spectrum

Quality of Written Communication – mass spec gives M^+ or molecular ion of 60 OR mass spec gives parent ion of 60 OR highest m/z (ALLOW m/e) value is 60 \checkmark

m/z = 45 indicates loss of CH₃

OR m/z = 45 indicates presence of CH₃CHOH

OR CH₂CH₂OH OR C₂H₅O ✓

IR of F – Remember to check the spectrum

IR shows no broad absorption between 2500 to 3300 cm^{$^{-1}$} so no O—H bond **OR** no broad absorption between 2500 to 3300 cm^{$^{-1}$} so not a carboxylic acid \checkmark

IR shows absorption at 1700 cm⁻¹ due to a C=O bond

OR absorption at 1700 cm⁻¹ indicates a ketone **OR** aldehyde present

Identification and equation

F is CH₃COCH₃ **OR** propanone ✓

E is CH₃CHOHCH₃ **OR** propan-2-ol ✓

 $CH_3CHOHCH_3 + [O] \rightarrow CH_3COCH_3 + H_2O$

If \mathbf{F} has been incorrectly identified as propanal, mark identification and equation as ECF, so $\max = 2$

ALLOW E is CH₃CH₂CH₂OH ✓

ALLOW: $CH_3CH_2CH_2OH + [O] \rightarrow CH_3CH_2CHO + H_2O \checkmark$

The mark scheme for F = propanoic acid is shown below.

mass spec of E-Remember to check the spectrum

QWC – mass spec gives M⁺ or molecular ion of 60

OR mass spec gives parent ion of 60

OR highest m/z (**OR** m/e) value is 60 \checkmark

m/z = 45 indicates loss of CH₃

OR m/z = 45 indicates presence of CH₃CHOH

OR CH_2CH_2OH **OR** C_2H_5O \checkmark

IR of F- Remember to check the spectrum

IR shows (broad) absorption somewhere between 3500 and 2500 cm⁻¹ suggests carboxylic acid **OR** O−H bond ✓

IR shows absorption at 1700 cm⁻¹ due to C=O **OR** absorption at 1700 cm⁻¹ indicates a carboxylic acid ✓

Identification and equation

F is CH₃CH₂COOH **OR** propanoic acid ✓
E is CH₃CH₂CH₂OH **OR** propan-1-ol ✓
CH₃CH₂CH₂OH + 2[O] → CH₃CH₂COOH + H₂O ✓

Extra guidance for marking of question

If E has not been identified OR if F has been identified as a ketone or aldehyde, use the first mark scheme

If F has been identified as a carboxylic acid, use the second mark scheme

Mass spec

These two marking points stand as **independent** marks whichever compounds have been identified.

The positive sign for fragment ions is not required. **IGNORE** negative charge. The mass spec may well be on the actual spectrum.

IR mark

These stand as **independent** marks whichever compounds have been identified. The IR analysis may well be on the actual spectrum.

Identification marks

If both structure and name are given they must **both** be correct but allow 'propanol' drawn with the correct structure because the position number of the –OH has been clearly identified

ALLOW ECF for identification of **F** e.g. if **E** is pentan-2-ol \star then an answer of pentan-2-one for **F** will be given a mark \star as ECF

ALLOW identification marks for **E** and **F** from equation

Equation mark

ALLOW ECF for any correct equation showing the oxidation of **any** alcohol to the appropriate product.

ALLOW molecular formulae in equations,

i.e.
$$C_3H_7OH + [O] \rightarrow C_2H_5CHO + H_2O \checkmark$$
; $C_3H_8O + [O] \rightarrow C_3H_6O + H_2O \checkmark$; $C_3H_7OH + [O] \rightarrow C_2H_5COH + H_2O \checkmark$

[7]

2. (a) method 1:

fermentation of sugars or carbohydrates **OR** reaction with yeast with sugar or carbohydrates \checkmark $C_6H_{12}O_6 \rightarrow 2C_2H_5OH + 2CO_2 \checkmark$

method 2:

hydration of ethene **OR** reaction of ethene with water **OR** reaction of steam with ethene \checkmark

$$C_2H_4 + H_2O \rightarrow C_2H_5OH \checkmark$$

ALLOW sugar from equation

ALLOW sugar from equation

ALLOW C_2H_6O in equation

ALLOW correct multiples

IGNORE state symbols

ALLOW ethene from the equation

IGNORE mention of any catalyst

ALLOW C_2H_6O in equation **OR** H_2O over the arrow

ALLOW correct multiples

IGNORE state symbols

4

$$(CH_3)_2CHOH + [O] \rightarrow (CH_3)_2CO + H_2O \checkmark$$

If name and formula given both need to be correct

ALLOW propanone OR acetone

IGNORE propone

NOT incorrect named compound

$$ALLOW C_3H_8O + [O] \rightarrow C_3H_6O + H_2O$$

ALLOW O instead of [O]

ALLOW correct multiples

IGNORE state symbols

2

(ii) CH₃CH₂COOH **OR** propanoic acid ✓

Any number or range of numbers between 1750–1640 (cm $^{-1}$) for C=O \checkmark

Any number or range of numbers between 2500–3300 (cm $^{-1})$ for O–H \checkmark

ALLOW C=O and O—H marks independent of compound identified **i.e. stand alone marks**

ALLOW correct bonds shown by the appropriate absorption on the IR spectrum

IGNORE reference to C—O bond

3

(c) (i) 2-methylpropan-2-ol ✓

ALLOW methylpropan-2-ol OR tertiarybutanol

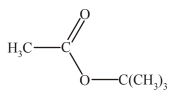
1

(ii) ester ✓

1

(iii) $CH_3CO_2C(CH_3)_3$ **OR** $CH_3COOC(CH_3)_3$

OR



ester group shown 🗸

rest of molecule ✓

ALLOW skeletal formula OR displayed formula

ALLOW ester linkage even if rest of structure is wrong

2

[13]

3. (a) (i)

$$C_2H_5$$
 C_2H_5
 C

C–I curly arrow from the bond not from carbon atom ✓

curly arrow from the OH⁻ ✓

correct partial charges on C—I ✓

no need to show any lone pairs on oxygen but must have a clear negative sign rather than partial negative charge

IGNORE lone pairs

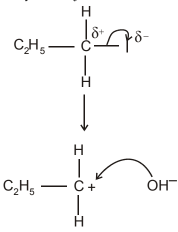
IGNORE products of this reaction

ALLOW curly arrow from a negative charge or from any part of hydroxide ion

If SNI mechanism is given then use the mark scheme below correct partial charges on C-I

C–I curly arrow from the bond not from carbon atom \checkmark

*curly arrow from the OH***-** *to the correct carbocation* ✓



(ii) nucleophilic substitution ✓

1

3

(b) C–I bonds broken more easily ✓ C–I bonds are weaker **OR** have less bond enthalpy **OR** C–I

bonds are longer ✓

ALLOW ora e.g. C—Br bonds are stronger OR broken less easily

2

[6]

4. from IR absorption, **J** contains O–H **OR** from IR **J** is an alcohol \checkmark

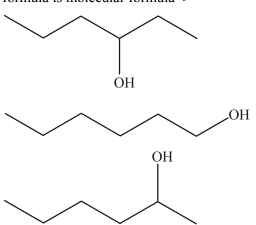
C: H: O =
$$\frac{70.59}{12.0}$$
: $\frac{13.72}{1.0}$: $\frac{15.69}{16.0}$

OR 5.8825 : 13.72 : 0.9806 ✓

empirical formula = $C_6H_{14}O$ \checkmark

(from mass spectrum), $M_{\rm r} = 102 \checkmark$

evidence that it has been shown that the empirical formula is the molecular formulae e.g. Mr of $C_6H_{14}O = 102$ so empirical formula is molecular formula \checkmark



One mark for each correct structure \checkmark \checkmark This is a QWC mark ALLOW two marks for correct empirical formula with no working out This is a QWC mark ALLOW structural or displayed formulae IGNORE incorrect names ALLOW one minor slip in drawing structures e.g. one missing hydrogen but ALLOW ecf for bigger slips such as showing just sticks and no hydrogen atoms ALLOW bond to H in OH **ALLOW one** mark for three isomers of $C_6H_{13}OH$ whether branched or unbranched as a catch mark if no other mark has been awarded for the structures If more than three isomers of $C_6H_{13}OH$ drawn • 1 branched and 3 unbranched award two marks • any other combination award one mark ALLOW one mark for hexan-1-ol, hexan-2-ol and hexan-3-ol if structures not drawn [8] (i) Any two realistic fragments, e.g. CH_3^+ : 15; $C_2H_5^+$: 29; $C_3H_7^+$: 43; $C_4H_9^+$: 57; OH^+ : 17, etc. (1) (1) Do not penalise missing charge. 2 (ii) breathalysers/monitoring of air pollution, MOT emission testing, etc. (1) [3] mole ratio = 88.89/12 : 11.1/1 = 7.41 : 11.1 (1) empirical formula = C_2H_3 (1) relative mass of $C_2H_3 = 27$. $M_{\rm r} = 2 \times 29$ so molecular formula = C4H6 (1) X reacts with 2 mol H₂ so there are 2 double bonds (1) Possible structure = 1,3-butadiene / [5] (a) (i) 1 1

5.

6.

7.

(ii)

(i)

(b)

Orange to green/black/blue

7

1

contains a C=O/aldehyde, ketone, carboxylic acid and ester/

carbonyl/carbonyl in an aldehyde

- (ii) does **not** contain a O–H/ (hydrogen bonded in a) carboxylic acid
- (iii) distillation (no mark) **because** distillation allows loss of volatile components /removes butanal from oxidising mixture prevents formation of RCOOH/ partial oxidation would be achieved or reverse argument for reflux not being used in that reflux prevents loss of volatile components hence complete oxidation would be achieved/RCOOH would be formed

[7]

8. (i) $H^+ \checkmark Cr_2O_7^{2-}$

(ii)

(iii) carboxylic acid would have an absorption between $1680 - 1750 \text{ cm}^{-1} / 1700 \text{ cm}^{-1} \text{ or } 2500 - 3300 \text{ cm}^{-1}$.

[6]

2

1

- 9. (a) (i) (volatile components) can escape/distil out 1
 ethanal is most volatile/b pt less than 60°C/partial oxidation 1
 - (ii) (volatile components) cannot escape/ refluxed 1 complete oxidation will be achieved/oxidised to the acid 1
 - (b) $C_2H_5OH + 2[O] \rightarrow CH_3COOH + H_2O$ $(CH_3COOH + H_2O \checkmark)$ 2
 - (c) spectrum C
 spectrum C only shows absorption at 1700 cm⁻¹ for the C=O
 the other two spectra contain the OH group absorption at approx 3000 cm⁻¹

10. acrylic acid 1
approx 1700 cm⁻¹ (range 1650 – 1750) indicates C=O 1

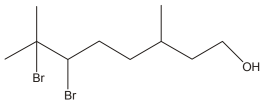
[9]

[3]

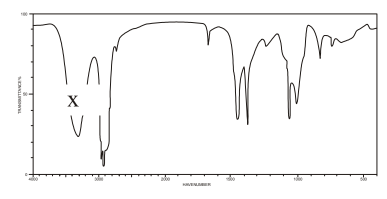
11. (a) (i) alkene ✓

alcohol/hydroxy/hydroxyl ✓ 1

- (b) (i) $I = \text{alkene \& II} = \text{alcohol... both are needed } \checkmark$
 - (ii) decolourised / colourless ✓
 - (iii) **✓**



(iv) X as shown below ✓



- (c) (i) Ni/Pt/Rh/Pd ✓
 - (ii) compound **B** is $C_{10}H_{22}O$ \checkmark
 - (iii) $C_{10}H_{20}O + H_2 \rightarrow C_{10}H_{22}O \checkmark$

12. (a) (i) Alkene/C=C ✓

Alcohol/ROH/hydroxy/hydroxyl/OH (not OH⁻ or hydroxide) ✓ 1

(ii) One of the C in both C=C is joined to two atoms or groups that are the same ✓ 1

[9]

(b)	Observation		decolourisation (of Br ₂) ✓	1	
	Mole	cular formula	$C_{10}H_{18}OBr_4 \checkmark \checkmark$	2	
			$C_{10}H_{18}OBr_2$ gets 1 mark		
(c)	reagent		CH₃COOH ✓	1	
	catalyst		$\mathrm{H_2SO_4/H}^+\mathrm{/HC}l$ (aq) or dilute loses the mark \checkmark	1	
(d)	(i)	(i) $C_{10}H_{18}O + 2[O] \rightarrow C_{10}H_{16}O_2 + H_2O \checkmark \checkmark$ 1 mark for H_2O and 1 mark for $2[O]$		2	
	(ii) The infra-red spectrum was of compound Y because absorption between 1680 − 1750 cm ⁻¹ indicates a C=O ✓				
				1	
		and the absence of a peak between $2500 - 3300 \text{ cm}^{-1}$ shows the absence			
		of the OH hydro	gen bonded in a carboxylic acid 🗸	1	[12]