

M1. (a) Functional group (isomerism)

1

(b)

M1 Tollens' (reagent)
(*Credit ammoniacal silver nitrate OR a description of making Tollens'*)
(*Ignore either AgNO₃ or [Ag(NH₃)₂]⁺ or "the silver mirror test" on their own, but mark M2 and M3*)

M1 Fehling's (solution) or Benedict's solution
(*Ignore Cu²⁺(aq) or CuSO₄ on their own, but mark on to M2 and M3*)

M2 silver mirror

M2 Red solid/precipitate
(*Credit orange or brown solid*)

OR

black solid/precipitate
(*NOT silver precipitate*)

M3 (stays) colourless
or no change or no reaction

M3 (stays) blue
or no change or no reaction

Mark on from an incomplete/incorrect attempt at the correct reagent, penalising M1

No reagent, CE=0

Allow the following alternatives

M1 (acidified) potassium dichromate(VI) (solution)

M2 (turns) green

M3 (stays) orange/no change

OR

M1 (acidified) potassium manganate(VII) (solution)

M2 (turns) colourless

M3 (stays) purple/no change

For M3

Ignore "nothing (happens)"

Ignore "no observation"

3

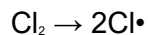
(c) (Both have) C=O **OR** a carbonyl (group)

1

(d) (i) (Free-) radical substitution ONLY
Penalise "(free) radical mechanism"

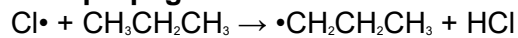
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(ii) **Initiation**



Penalise absence of dot once only.

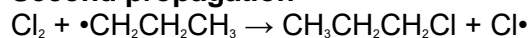
First propagation



Penalise incorrect position of dot on propyl radical once only.

Penalise $\text{C}_3\text{H}_7\cdot$ once only

Second propagation

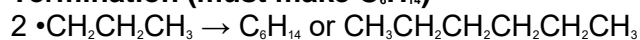


OR



Accept $\text{CH}_3\text{CH}_2\text{CH}_2\cdot$ with the radical dot above/below/to the side of the last carbon.

Termination (must make C_6H_{14})



Use of the secondary free radical might gain 3 of the four marks

4

- (e) $M_r = \underline{44.06352}$ (for propane)
 $M_r = \underline{43.98982}$ (for carbon dioxide)

Mark independently

M1 a correct value for both of these M_r values.

M2 a statement or idea that two peaks appear (in the mass spectrum)

OR

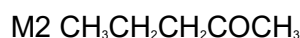
two molecular ions are seen (in the mass spectrum).

2

[12]

M2. (a) (i) M1 pentan-3-one only

1



(insist on C=O being drawn out)

(penalise use of C_5H_{10})

- 1
- (ii) aldehyde $(\text{CH}_3)_2\text{CHCH}_2\text{CHO}$ 1
- ketone $(\text{CH}_3)_2\text{CHCOCH}_3$ 1
- (insist on a clear structure for the C=O of the functional groups, but do not be too harsh on the vertical bonds between carbon atom on this occasion)*
- (If both structures correct, but wrong way around, award one mark)*
- (ignore names)*
- (b) (i) $\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{CHO} + [\text{O}] \rightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{COOH}$ 1
- (accept $\text{C}_4\text{H}_9\text{CHO}$ going to $\text{C}_4\text{H}_9\text{COOH}$)*
- (insist on a balanced equation – for example do not credit [O] over the arrow alone)*
- (ii) pentanoic acid 1
- (credit pentan–1–oic acid)*
- (c) (i) $\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{OH}$ OR pentan–1–ol 1
- (If both a structure and a formula are given, credit either correct one of these provided the other is a good, if imperfect, attempt)*
- (ii) Primary 1
- (credit 1° or 1)*

[8]

M3.(a) Eliminate / reduce fire risk;
Allow ethanol flammable / burns / combusts.

1

(b) Orange to green;

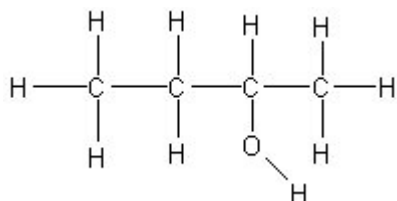
Need full colour change to score mark.

1

[2]

M4. (a) **M1**

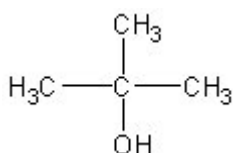
Displayed formula for butan-2-ol



M1 displayed formula must have all bonds drawn out, including the O—H but ignore angles

Penalise “sticks”

M2 Alcohol X is



M2 structure must be clearly identifiable as

2-methylpropan-2-ol and may be drawn in a variety of ways.

M3 Alcohol Y is named (2)-methylpropan-1-ol ONLY

M3 must be correct name, but ignore structures

3

(b) **M1** The infrared spectrum shows an absorption/peak in the range 3230 to 3550 (cm⁻¹)(which supports the idea that an alcohol is present)

In M1, allow the words “dip”, “spike”, “low transmittance” and “trough” as alternatives for absorption.

M2 Reference to the ‘fingerprint region’ or below 1500 (cm⁻¹)

M3 Match with or same as known sample/database spectra

Check the spectrum to see if alcohol OH is labelled and credit.

OR

M2 Run infrared spectra (of the alcohols)

M3 Find which one matches or is the same as this spectrum.

3

- (c) **M1** balanced equation

$$\text{C}_6\text{H}_{12}\text{O}_6 \rightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} + 2\text{CO}_2 + \text{H}_2\text{O}$$
 or $\text{C}_4\text{H}_9\text{OH}$
Or multiples for M1 and M3
In M1 and M3 penalise use of $\text{C}_4\text{H}_{10}\text{O}$ or butan-2-ol once only

M2 Any one from

- excess/adequate/sufficient/correct amount of/enough/plenty/ a good supply of oxygen or air
- good mixing of the fuel and air/oxygen
For M2, do not accept simply “oxygen” or “air” alone
Ignore reference to “temperature”

M3 $\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH} + 6\text{O}_2 \rightarrow 4\text{CO}_2 + 5\text{H}_2\text{O}$
 or $\text{C}_4\text{H}_9\text{OH}$

M4 A biofuel is a fuel produced from (renewable) biological (re)source(s)

OR

(renewable) (re)source(s) from (a specified) plant(s)/fruit(s)/tree(s)

In M4

Ignore references to “carbon neutral”

Ignore “sugar” and “glucose”

4

- (d) **M1** butan-1-ol is a primary or 1° (alcohol)

M2 Displayed formula (ONLY) for butanal $\text{CH}_3\text{CH}_2\text{CH}_2\text{CHO}$

M3 Displayed formula (ONLY) for butanoic acid $\text{CH}_3\text{CH}_2\text{CH}_2\text{COOH}$

M2 and M3 displayed formula must have all bonds drawn out including the O—H but ignore angles.

If butanal and butanoic acid formulae are both correctly given but not displayed, credit one mark out of two.

M4 Oxidation (oxidised) OR Redox

M5 orange to green

Both colours required for M5

Ignore states

5

[15]

M5. (a) Allow 1 mark each for any correctly drawn primary, secondary and tertiary alcohol of molecular formula C_4H_8O 3

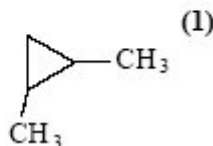
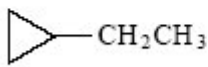
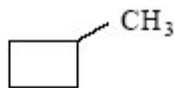
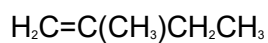
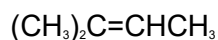
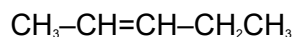
Tertiary alcohol cannot be oxidised 1

(b) Region $1500-400\text{ cm}^{-1}$ 1

exact match to spectrum of known compound 1

(c) **A** $CH_3CH_2CH_2OH$ **B** $CH_3CH_2-O-CH_3$ (1)
 or $CH_3CH(OH)CH_3$ (1)

C one alkene e.g. **D** one cycloalkane e.g.



(1)
etc

E CH_3CH_2CHO (1)

F CH_3COCH_3 (1)

6

[12]