

1

- (b) This question is marked using levels of response. Refer to the Mark Scheme Instructions for Examiners for guidance on how to mark this question.

All stages are covered and the explanation of each stage is generally correct and virtually complete.

Answer is communicated coherently and shows a logical progression from stage 1 to stage 2 then stage 3.

Level 3
5 – 6 marks

All stages are covered but the explanation of each stage may be incomplete or may contain inaccuracies OR two stages are covered and the explanations are generally correct and virtually complete.

Answer is mainly coherent and shows progression from stage 1 to stage 3.

Level 2
3 – 4 marks

Two stages are covered but the explanation of each stage may be incomplete or may contain inaccuracies, OR only one stage is covered but the explanation is generally correct and virtually complete.

Answer includes isolated statements but these are not presented in a logical order or show confused reasoning.

Level 1
1 – 2 marks

Insufficient correct chemistry to gain a mark.

Level 0
0 marks

Indicative Chemistry content

Stage 1: Formation of product

- Nucleophilic attack
- Planar carbonyl group
- H^- attacks from either side (stated or drawn)

Stage 2: Nature of product

- Product of step 1 shown
- This exists in two chiral forms (stated or drawn)
- Equal amounts of each enantiomer / racemic mixture formed

Stage 3: Optical activity

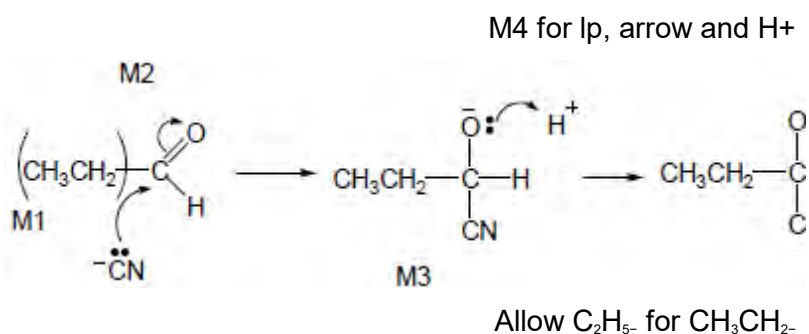
- Optical isomers / enantiomers rotate the plane of polarised light equally

- in
With a racemic / equal mixture the effects cancel

6
[7]

M2.(a) Nucleophilic addition

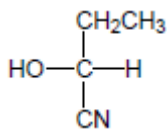
1



- M1 and M4 include lone pair and curly arrow.
- Allow: CN⁻ but arrow must start at lone pair on C.
- M2 not allowed independent of M1, but allow M1 for correct attack on C⁺.
- + rather than δ⁺ on C=O loses M2.
- Penalise incorrect partial charges.
- M3 is for correct structure including minus sign but lone pair is part of M4.
- Penalise extra curly arrows in M4.

4

(b) (i) M1



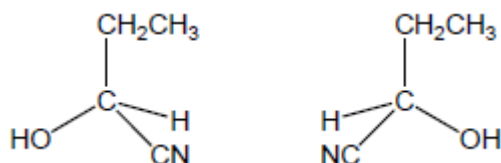
M1 for correct structure of product of part (a).

Allow C₂H₅- for CH₃CH₂-.

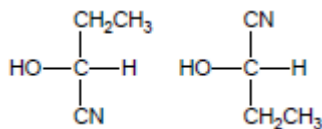
Penalise wrongly bonded, OH or CN or CH₂CH₃ once only in clip.

1

M2

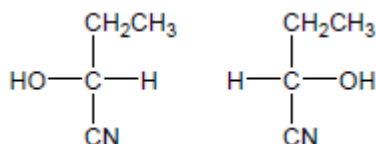


M2 cannot be gained by simply swapping two or more groups with no attempt to show a mirror image., e.g. do not allow M2 for



because these do not show the enantiomers as mirror images.

Students must show an attempt at mirror images, eg allow

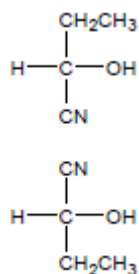


ie vertical groups same and horizontal swapped as if there was a mirror between them

No mirror need be shown

Do not penalize wedge bond when wedge comes into contact with both C & N

However these two could score M2 if placed as below as if with a "mirror" horizontally between them.



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- (ii) M1 (Plane) polarized light
M2 only scores following correct M1

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M2 Rotated in opposite directions (equally) (only allow if M1 correct or close)

Not just in different directions but allow one rotates light to the left and one to the right.

Not molecules rotate.

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(c) 2-hydroxybutane(-1-)nitrile

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(d) Weak acid / (acid) only slightly / partially dissociated / ionised
Ignore rate of dissociation.

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[CN⁻] very low

Allow (very) few cyanide ions.

Mark independently.

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(e) (i) $\text{H}_2\text{C}=\text{CH}-\text{CH}_3 + \text{NH}_3 + \frac{3}{2}\text{O}_2 \longrightarrow \text{H}_2\text{C}=\text{CH}-\text{CN} + 3\text{H}_2\text{O}$

OR

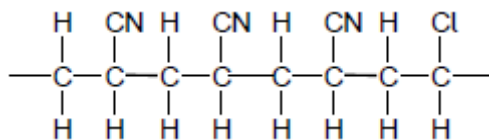
$\text{H}_2\text{C}=\text{CH}-\text{CH}_3 + \text{NH}_3 + 3\text{O}_2 \longrightarrow \text{H}_2\text{C}=\text{CH}-\text{CN} + 3\text{H}_2\text{O}_2$

OR doubled.

Allow C₃H₆ and CH₂CHCN or C₃H₃N on this occasion only.

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(ii)

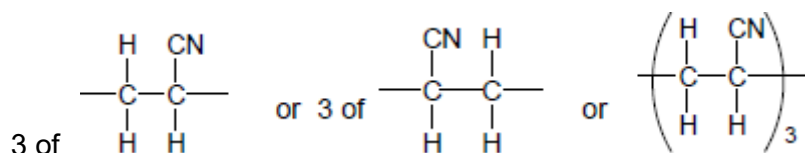


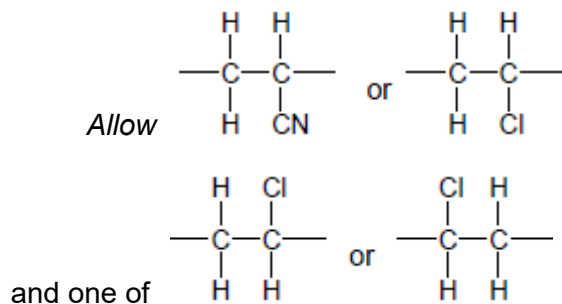
Ignore n.

Must show trailing bonds.

Do not penalise C–NC bond here on this occasion.

Must contain, in any order,





Allow $-\text{CH}_2\text{CH}(\text{CN})\text{CH}_2\text{CHCl}-$ etc.

1

(iii) Addition (polymerization)

Allow self-addition.

Do not allow additional.

1

[15]

M3. Dichromate(VI) will also oxidise / give a positive test with alcohols

Allow 'dichromate'.

Allow 'dichromate(VI) will oxidise other organic molecules / functional groups'.

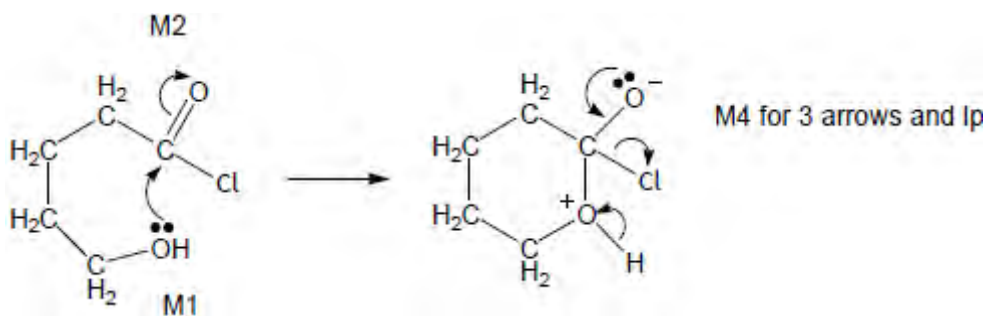
[1]

M4.(a) (i) (nucleophilic) addition-elimination

Not electrophilic addition-elimination

Ignore esterification

1



M3 for structure

- If wrong nucleophile used or O–H broken in first step, can only score M2.
- M2 not allowed independent of M1, but allow M1 for correct attack on C+
- + rather than $\delta+$ on C=O loses M2.
- If Cl lost with C=O breaking lose M2.
- M3 for correct structure with charges but lone pair on O is part of M4.
- Only allow M4 after correct / very close M3.
- Ignore HCl shown as a product.

4

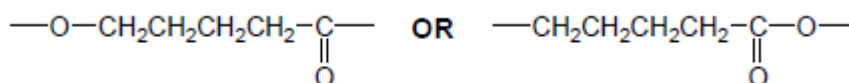
a 20-50 (ppm) or single value or range entirely within this range
If values not specified as a or b then assume first is a.

1

b 50-90 (ppm) or single value or range entirely within this range

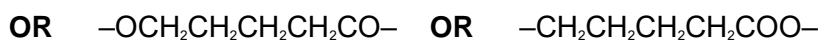
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(ii)

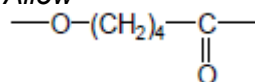


Must have trailing bonds, but ignore n.

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Allow



but not $\text{---C}_4\text{H}_8\text{---}$

one unit only

Condensation

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(b)

| | | | |
|--|----------|------------------------|--------------------------------|
| | Tollens' | Fehling's / Benedict's | Acidified potassium dichromate |
|--|----------|------------------------|--------------------------------|

Penalise wrong formula for Tollens or missing acid with potassium dichromate but mark on.

1

| | | | |
|----------|--|---|--|
| J | No reaction / no (visible) change / no silver mirror | No reaction / no (visible) change / stays blue / no red ppt | No reaction / no (visible) change / stays orange / does not turn green |
|----------|--|---|--|

Ignore 'clear', 'nothing'.
Penalise wrong starting colour for dichromate.

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| | | | |
|----------|--|---|----------------------|
| K | Silver <u>mirror</u> / grey <u>ppt</u> | Red <u>ppt</u> (allow brick red or red-orange) | (orange) turns green |
|----------|--|---|----------------------|

1

J Two (peaks)
Allow trough, peak, spike.

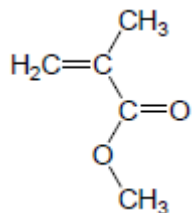
1

K Four (peaks)
Ignore details of splitting.
If values not specified as J or K then assume first is J.

1

(c) If all the structures are unlabelled, assume that the first drawn ester is L, the second ester is M; the first drawn acid is N, the second P. The cyclic compound should be obvious.

L
ester



OR $\text{H}_2\text{C}=\text{C}(\text{CH}_3)\text{COOCH}_3$

All $\text{C}_5\text{H}_8\text{O}_2$ L to P must have $\text{C}=\text{C}$.

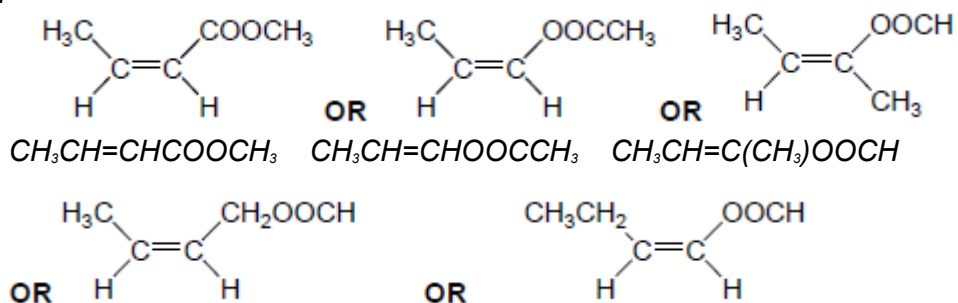
Allow CH_3^- .

Allow $-\text{CO}_2\text{CH}_3$ etc.

Allow $\text{CH}_2\text{C}(\text{CH}_3)\text{COOCH}_3$.

1

M
ester



$\text{CH}_3\text{CH}=\text{CHCH}_2\text{OOCH}$

$\text{CH}_3\text{CH}_2\text{CH}=\text{CHOOCH}$

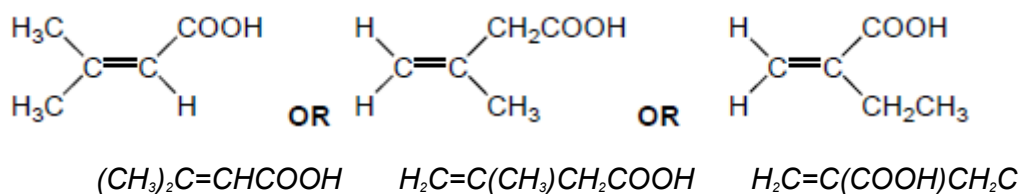
Allow either *E-Z* isomer.

Allow CH_3^- or C_2H_5^- but not CH_2CH_3^- .

Allow $\text{CH}_3\text{CHCHCOOCH}_3$ etc.

1

N
acid



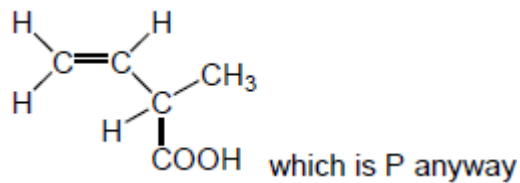
H_3

Allow CH_3^- or C_2H_5^- but not CH_2CH_3^- .

Allow $-\text{CO}_2\text{H}$.

Not cyclic isomers.

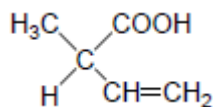
Not the optically active isomer.



Allow $(\text{CH}_3)_2\text{CCHCOOH}$ etc.

1

P
acid



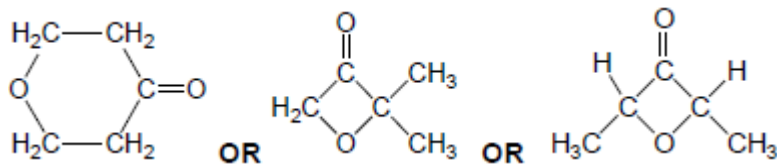
Allow $-\text{CO}_2\text{H}$.



Allow $\text{CH}_3\text{CH}(\text{CO}_2\text{H})\text{CHCH}_2$ or
 $\text{CH}_3\text{CH}(\text{CO}_2\text{H})\text{C}_2\text{H}_3$.

1

Q



Not cyclic esters.

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[19]

M5.(a) A mixture of liquids is heated to boiling point for a prolonged time

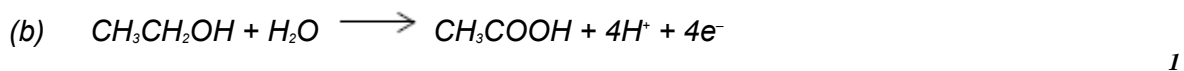
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Vapour is formed which escapes from the liquid mixture, is changed back into liquid and returned to the liquid mixture

1

Any ethanal and ethanol that initially evaporates can then be oxidised

1



(c) Mixture heated in a suitable flask / container
A labelled sketch illustrating these points scores the marks 1

With still head containing a thermometer 1

Water cooled condenser connected to the still head and suitable cooled collecting vessel 1

Collect sample at the boiling point of ethanal 1

Cooled collection vessel necessary to reduce evaporation of ethanal 1

(d) Hydrogen bonding in ethanol and ethanoic acid or no hydrogen bonding in ethanal 1

Intermolecular forces / dipole-dipole are weaker than hydrogen bonding 1

(e) Reagent to confirm the presence of ethanal:

Add Tollens' reagent / ammoniacal silver nitrate / aqueous silver nitrate followed by 1 drop of aqueous sodium hydroxide, then enough aqueous ammonia to dissolve the precipitate formed

OR

Add Fehling's solution

1

Warm

M2 and M3 can only be awarded if M1 is given correctly

1

Result with Tollen's reagent:

Silver mirror / black precipitate

OR

Result with Fehling's solution:

Red precipitate / orange-red precipitate

1

Reagent to confirm the absence of ethanoic acid

Add sodium hydrogencarbonate or sodium carbonate

1

Result; no effervescence observed; hence no acid present

1

M5 can only be awarded if M4 is given correctly

OR

Reagent; add ethanol and concentrated sulfuric acid and warm

Result; no sweet smell / no oily drops on the surface of the liquid,

hence no acid present

[16]

M6.B

[1]