$\boldsymbol{4.10,4.11}\ EXAM\ QUESTIONS\ mark\ scheme$

1.	(a)	electrophilic substitution;	1
		cone HN0 ₃ ;	1
		cone H ₂ SO ₄ either or both cone missing scores one for both acids;	1
	(b)	Sn or Fe/HCl (cone or dil or neither); (ignore extra NaOH)	1
		Sn or Fe/H ₂ SO ₄ (dil or neither) (not HNO ₃ at all)	
		or H_2/Ni (not $NaBH_4/LiAlH_4$ or Na/C_2H_5OH)	
		CH_3 CH_3	
		+ 6[H] + 2H ₂ O	
		NO_2 or $3H_2$ NH_2	
			1
	(c)	77 or 92;	1
	(d)	ÇH₃	
		$+\dot{ m N} m H_2$	1
		$(allow - NH_3^+)$	1
	(e)	\mathbf{G}	
		-CH ₂ NH ₂	
		Н	1
		NHCH ₃	

1

[9]

2. (a) $CH_3OH + CH_3CH_2COOH \rightarrow CH_3CH_2COOCH_3 + H_2O$

1

4

(b) (nucleophilic) addition-elimination NOT acylation

M3 for structure M4 for 3 arrows and lone pair ignore use of Cl^- to remove H^+

(c) $CH_3CH_2 - C$ $CH_3CH_2 - C$ $allow C_2H_5 and -CO_2 -$ $allow CH_3CH_2COOCOCH_2CH_3$

or $(CH_3CH_2CO)_2O$

- (d) (i) faster/not reversible/bigger yield/purer product/no(acid) (catalyst) required 1
 - (ii) anhydride less easily hydrolysed or reaction less violent/exothermic no (corrosive) (HCl) fumes formed or safer or less toxic/dangerous expense of acid chloride or anhydride cheaper 1 any one
- (e) (i) $C_8H_8O_2$
 - (ii) any two from

$$H = C$$

$$O = CH_{2}$$

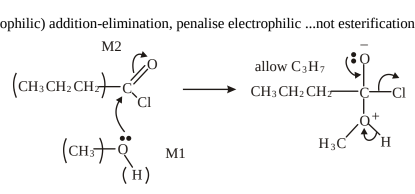
Allow $-CO_2$ - allow C_6H_5

[12]

2

- **3.** (a) butanoyl chloride
 - 25 27
 - (b) (i) \underline{Cl} has (two) isotopes or ^{35}Cl and ^{37}Cl 1 (ii) 106 and 108 1

(nucleophilic) addition-elimination, penalise electrophilic ...not esterification (c)



M3 for structure M4 for 3 arrows and lone pair (only allow for correct M3 or close)

M2 not allowed independent of Ml, but allow MI for correct attack on C+ if M2 shown as independent first.

[8]

4

1

1

- 4. (a) 5 (1)
 - (b) 2:2:2:3:3 (1) any order but not multiples

 $CH_3 - C - (R) (1)$ 01

 CH_3CH_2 or C_2H_5 or ethyl (1) (d) $\delta4.13$ (quartet): CH₂ peak <u>split by CH₃ / next to CH₃ (1)</u> $\delta 1.26$ (triplet): CH₃ peak split by CH₂ / next to CH₂ (1) 3

(e) CH_2CH_2 (1) 1

 $\begin{array}{c} (CO) & (CO) \\ CH_3 - C - CH_2CH_2 - C - OCH_2CH_3 \\ \parallel & \parallel \\ O & O \\ allow \textbf{(1)} \ for \ CH_3COCH_2OCOCH_2CH_3 \end{array}$ or CH₃COOCH₂COCH₂CH₃ Must be C₇H₁₂O₃

[9]

1

(b) (i) <u>C1</u> has (2) isotopes (1)

Allow ³⁵Cl and ³⁷Cl without word isotope – but must be correct isotopes

must have 3 different elements, i.e. not $C_3H_7^+$ but allow balanced equation including $C_3H_7^+$ for the equation mark

(ii) Fragmentation: $CH_3 - \overset{+}{C} = O$ (1) must be an ion (*)

Equation:
$$C_4H_7ClO^{+\bullet} \rightarrow CH_3 CO + C_2H_4Cl^{\bullet}$$
 (1) 3

(*) allow $C_2H_3O + or$ any form of it (i.e. CH_2CHO^{+} or CH_2COH^{+}) in equation, be generous with position of $+ or \tilde{N}$ if fragment ion completely wrong (not $m/z = 43$) no further marks

- (c) (i) $CDCl_3$ or CCl_4 (1) or D_2O , C_6D_6
 - (ii) $Si(CH_3)_4$ (1) or SiC_4H_{12}

2

(d)

	Peak 1	Peak 2	Peak
Integration value	3	3	1
Splitting pattern	doublet	singlet	quartet
Number of adjacent, non-equivalent protons	1	0	3

(1)

1

(f)

CH₃CH₂CH₂COCl

[9]

Name of mechanism: (nucleophilic) addition-elimination (1) (b) Mechanism: structure incl charges (1) 3 arrows **(1)** CH₃CH₂ 5 CH₃CH₂COOCH₂CH₃⁺ÑÈ CH₃CH₂C⁺=O (1)+ CH₃CH₂OÑ (c) equation (1) 2 CH₃CH₂CH₂COOCH₃ or (CH₃)₂CHCOOCH₃ (1) (d) Allow C₃H₇COOCH₃ 1 [9] 7. (a) (i) molecular formula (1) ¹³C isotope (1) (ii) 2 $(CH_3)_2CHCOCH_3^{+\bullet} \rightarrow (CH_3)_2CHCO^+ + CH_3$ (b) (i) **(1) (1) (1)** Structure 1 Structure 2 (ii) CH_3CO^+ $(CH_3)_2CH^+$ **(1)** 5 **(1)** two isotopes (1) (c) $C_3H_7^{35}Cl = 78$ $C_3H_7^{37}Cl = 80$ (1) relative abundances $^{35}Cl:^{37}Cl = 3:1$ (1) 3 [10] 8. (a) 2-chloropropanoic acid (1) 1 (b) δ 1.72 Doublet : next to CH (1) $\delta 4.44$ Quartet : next to CH₃ (1) 2 Two triplets (1) 1 (c)

(allow - COO -) (1)

1

6.

$$(d) \qquad \stackrel{\textbf{(1)}}{\overset{\text{Cl}}{\overset{\text{Cl}}{\overset{\text{CH}_{3}-\text{CH}-\text{COOH}}{\overset{\text{CH}_{3}-$$

(e) (i)
$$H_3^+$$
 (CH₂)₄ - C - COOH +NH₃ (Cl⁻)

(ii)
$$H_2N - (CH_2)_4 - C - COO^- (Na^+)$$
 (1) NH_2

Or anhydride

3 [12]

9. (a) (i)
$$CH_3 - C - CH_2CH_2CH_3 + 2$$
 [H] $\rightarrow CH_3CH(OH)CH_2CH_2CH_3$ (1) O or $C_5H_{10}O$ or $C_5H_{11}OH$

Name of mechanism: nucleophilic addition (1) (ii)

Mechanism:

(CH₃) (1) :H⁻ (1)
$$CH_3$$
 (1) CH_3 (1)

racemic (racemate) mixture formed (1) OR explained e.g. 2 enantiomers in equal amounts

Fragment 1: 43; 43 (b) Fragment 2: 71; 15 Any two \times (1)

2

7

[9]

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S: C=O or carbonyl (1)
                                                                                                       2
       (b)
              aldehyde (1)
                                                       -CHO or RCHO (1)
                                                                                                       1
       (c)
              (i)
                     Reason 1: TMS inert or non-toxic or volatile / easily removed
                     Reason 2: single (intense) peak
                                peak of 12 protons
                                has 12 equivalent protons
                                all protons in same environment
                     OR
                                peak / signal upfield of others
                                highly shielded
                                more shielded
                                peak away from others or \delta = 0 or low
                     not solvent, not cheap
                     any 2 reasons x (1)
              (ii)
                     Solvent: CDCl<sub>3</sub> or CCl<sub>4</sub> (NOT D<sub>2</sub>O)
                     Reason: proton free (1)
                              allow no hydrogens (atoms)
                     NOT H<sup>+</sup> / hydrogen ions
                                                                                                       4
                   CH_3 - C - (1)
O
       (d)
                    -OH (1)
              (ii)
              (iii) -CH<sub>2</sub>-CH<sub>2</sub>-(1)
                                                                                                       3
             CH_3 - C - CH_2 - CH_2 - OH  (1)
                                                                                                       1
                                                                                                                  [11]
                    HCN or KCN/HCl (1)
11.
       (a)
              (i)
                     nucleophilic addition (1)
                    C_4H_8O
                                           C<sub>5</sub>H<sub>9</sub>NO
              (ii)
                     Mr = 72 (1)
                                            Mr = 99 (1)
                     If MF shown lose 1 for wrong Mr.
                     If no MF shown max 2 if Mr wrong
                    5g \rightarrow \frac{5}{72} \times 99 \ (1) \ (= 6.88g)
                    64% yield = 0.64 \times \frac{5}{72} \times 99 = 4.40g (1)
                     (allow answer 4.36 - 4.42)
                                                                                                       6
```

10.

(a)

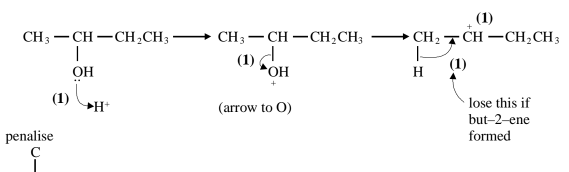
R: O-H (alcohols) (1)

- (b) (i) NaBH₄ or LiAlH₄ or H_2/Ni or Na/C₂H₅OH (1)
 - (ii) racemic mixture formed (1) or equal amounts of enantiomers
 - (iii) butanone has peak at $\sim 1700 \text{ cm}^{-1}$ (1) (but not at $\sim 3350 \text{ cm}^{-1}$) B has peak at $\sim 3350 \text{ cm}^{-1}$ (1)

(but not at $\sim 1700 \text{ cm}^{-1}$)

(c)

OH



(d) $\left(- \text{CH}_2 - \text{CH} \right)_{(n)}$ (1) $\begin{array}{c} \text{CH}_2 \text{CH}_3 \\ \text{Or } \text{C}_2 \text{H}_5 \end{array}$

[15]

3

4

12. (a) A is RCOOR' (1)

$$R + R' = 102 - 44 = 58$$
 (1) $\equiv C_4H_{10}$

$$\therefore C_5 H_{10} O_2 (1)$$

(b) 2:2:3:3(1)

(c) Two CH₂CH₃ groups present (1)

(d) $CH_3^a CH_2 COOCH_2^c CH_3^d$ (1)

(e)

Chemical shift, δ/ppm	1.09	1.33	2.32	4.13
Label of group	a (1)	d (1)	b (1)	c (1)

[10]

13. $(CH_3)_4$ Si or tetramethylsilane (1) (a)

1

(b) 4 (1) 1

(c) 2:1:6:3 (1) 1

- (d) $-CH_2CH_3$ (1)
 - CH₃ splits CH₂ to form a quartet (1)
 - CH₂ splits CH₃ to form a triplet (1)

3

(e) two equivalent CH₃ groups (1) 1

1

(f) $(CH_3)_2$ C CH_2CH_3 (1) ОН

[8]

- 14. CH₃COOCH₂CH₃ or CH₃COOC₂H₅ (1) (i)
 - 3 (1) (ii)
 - (iii) $3 + 4 \underline{\text{or}} \text{ triplet} + \text{quartet } (1)$

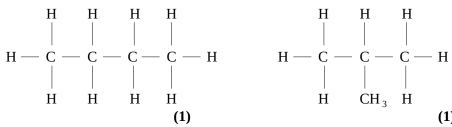
[3]

- **15.** same molecular formula / same number of each type of atom (1) (a) different arrangements of atoms (in the molecule) (1)
 - (not just same structural formula)

2

3

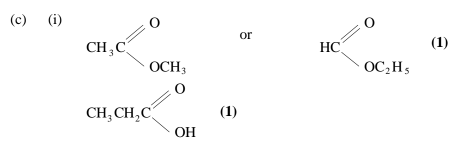
(b) (i)



- **(1)**
- 43: CH₃CH₂CH₂⁺/ CH₃C⁺HCH₃ / C₃H₇⁺ (1) (ii)
 - $CH_3CH_2^+/C_2H_5^+$ (1) 29:
 - $CH3^{+}$ (1)
 - (2 max if +ve sign omitted or -ve) (+ can be anywhere)
- 3

2

- (iii) Isomer 1 (dependent on candidate's order) (1)
 - Isomer 2 could not (easily) give peak at $29 / C_2 H_5^+$ (1)



(ii) ester (1) carboxylic acid (1)

names must be appropriate way round relative to (i) these marks dependent on correct answers in (i)

2 **[13]**

2

- - (b) C D

 CH₃CH₂CH₂OH CH₃CH₂-O-CH₃ (1)

 or CH₃CH(OH)CH₃ (1) 2

 - (d) **G H**CH₃CH₂CHO (1) CH₃COCH₃ (1) 2
 - CH₃CH₂CHO (1)

 CH₃COCH₃ (1)

 J

 CH₃COCH₃ (1)

 CH₃COCH₃ (1)

 CH₃COCH₃ (1)

 CH₃COCH₃ (1)

 CH₃COCH₃ (1)

 CH₃COCH₂ (1)

 CH₃COCH₃ (1)

 CH₃COCH₂ (1)

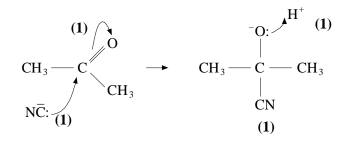
 CH₃CH₂ CH₂ CH

 $(CH_3)_2C = CHCH_3$ etc CH_2CH_3 CH_3 CH_3 CH_3

17. (a) *Name*

nucleophilic addition (1)

Mechanism



(b) (i) Equation

 $CH_3COCH_3 + 2[H] \rightarrow CH_3CH(OH)CH_3$ (1)

Reducing agent

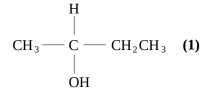
 $NaBH_4(1)$

(ii)

Carbonyl compound A

Alcohol C

(1)



Carbonyl compound **B**

 $Alcohol\, m{D}$

(1)
$$CH_3 \longrightarrow CH \longrightarrow CH_2OH$$
 (1) CH_3

6

5

[11]

- 1715 cm⁻¹ C=O group (**1**) 18. 3350 cm^{-1} O–H group В alcohol (1) A CH₃CH₂COCH₂CH₃ (1)

 † (1) q (1) two environments or two kinds of proton (1) CH₃CH₂ adjacent or coupled (1) ratio 2:3 or 4:6 (1) symmetric (1) $[CH_3CH_2COCH_2CH_3]^{+\bullet} \rightarrow CH_3CH_2CO^{+} + CH_3CH_2^{\bullet}$ (1) m/z = 86 (1)**(1)** or M_r for A $CH_3CH_2CHCH_2CH_3 \rightarrow CH_3CH = CHCH_2CH_3$ (1) В $\text{CH}_3\text{CH}_2\overset{^+\text{C}}{\text{C}}\text{HCH}_2\text{CH}_3$ (1) and $\text{CH}_3\overset{^+\text{C}}{\text{C}}\text{HCH}_2\text{CH}_2\text{CH}_3$ (1) both secondary (1) hydration gives ${\bf B}$ and ${\rm CH_3CHCH_2CH_2CH_3}$ (1) about 50% of each (1) $A \rightarrow B$ reduction $B \rightarrow C$ dehydration <u>or</u> elimination (1) C is an alkene (1) cis/trans isomers (1)

NOT acid

D is a racemate (1) or optical isomers

any 20

[20]

(b)
$$H_2C = C - CH_2CH_3$$
 $H_3C - C = CHCH_3$ $H_3C - CH - CH = CH_2$ CH_3 CH_3

Allow conseq dibromocompounds following incorrect unbranched alkenes

NOT allow dibromocompound consequent on a duplicate alkene NOT allow monobromocompounds if HBr added 3

1

6:3:1 either next to correct structure or to none

Allow a mark for identifying correct dibromocompound with three peaks even if integration ratio is wrong

if 6:3:1 missing or wrong, no marks for splitting

Only award a mark for splitting if it is clear which integration number it refers to 1

6 singlet or drawn 1

3 doublet or drawn 1

1 quartet/quadruplet or drawn 1

20. (a) (i) 3 peaks or shown in a list 1 $m/z = 126, 128 \text{ and } 130 (56 + 70/72/74) (all \ 3 \text{ scores } 2)$ 2

mz = 120, 128 and 130 (30 +70/72/74) (and 3 scores 2) (if 56 wrong allow (x + 70/72/74) for I(x cannot be zero) (any two scores 1)

(ii) 3

$$[C_4H_8Cl_2]^{+\bullet} \longrightarrow CH_3CH_2 \longrightarrow CH_3CH_4 \longrightarrow CH_2Cl$$

$$(1) \qquad (1) \qquad (1)$$

[16]

(b)	(1)	optical		1
		equal mixture of enantiomers		1
		(optically) inactive or effects cancel	· · · · · · · · · · · · · · · · · · ·	1
			stereospecific reagent (QoL)	1
		rotated in opposite/different directions (QoL) reac	ts with one isomer only	1
	(::)	carbocation	or with one isomer only	
	(ii)	planar – (must refer to carbocation or in	ntermediate)	1
		attack from either side equally likely –	uermeaiaie)	1
		(must refer to carbocation /intermediate	2)	1
				7 max
				,
(a)	(i)	2 pooles (if A peaks allow splitting only)		1
(c)	(i)	2 peaks (if 4 peaks allow splitting only)		1
		ratio 6:2 or 3:1		1
		doublet (6 or 3)		1
		quartet (2 or 1)		1
	(ii)	S		
	` '	Cl		
		H_3C — C — CH_2CH_3		
		Cl		1
		T		
		CH ₃		
		H ₂ C — CH ₂ Cl		
		CH_3		1

[19]

21. (a) HOOC COOH

NB The bonds shown in the structure must be correct

Isomerism: Geometric or cis-trans

If written answer is correct, ignore incorrect labelling of structures.

If no written answer, allow correctly labelled structures.

Both COOH groups must be on the same side/ close together/ cis .

No rotation about C=C axis

Structure

Allow 1

(b) $Br_2 / HBr / H_2SO_4 / H^+ / Br^+ / NO_2^+ (Mark M1)$

HOOC COOH H COOH HOOC COOH $C = C \xrightarrow{(1) M2} M4(1) + C - C - H \longrightarrow Br - C - C - H$ $H \xrightarrow{H} (1) M3$ $:Br^{-}$

NB If electrophile $H^+/Br^+/NO_2^+$ allow M1, M2 and M4 If the acid is incorrect, M2 and M3 can still be scored Allow M4 consequentially if repeat error from part (a)

1

1

1

1

(c) e.g. $2NaOH + HO_2CCHCHCO_2H \rightarrow NaO_2CCHCHCO_2Na + 2H_2O$

Both H replaced

Balanced for atoms and charges NB Allow ionic equations and $2NaOH + C_4H_4O_4 \stackrel{.}{\to} C_4H_2O_4Na_2 + 2H_2O$ Allow one if structure incorrect but molecular formula correct

Allow one for a correct equation showing one H replaced

(d)M1Two peaks1M2No splitting or singlets1M3(Two) non-equivalent protons or two proton environments1M4No adjacent protons1M5Same area under the two peaks or same relative intensity1

NB Doublet could score M1 and M3 or M5 (Max 2)

More than two peaks CE = 0

Apply the "list principle" to incorrect answers if more than 3 given

[15]

Max 3

1

1

3

22. (a) [CH₃CH₂CO]⁺

 $CH_3CH_2COCl + AlCl_3 \rightarrow [CH_3CH_2CO]^+ + AlCl_4^-$

(Penalise wrong arrows in the equation or lone pair on Al In the equation, the position of the + on the electrophile can be on O or C or outside square brackets,

Can score electrophile mark in mechanism if not previously

Can score electrophile mark in mechanism if not previously gained)

(Arrow for M1 must be to C or to the + on C penalize + in intermediate if too close to C1; horseshoe should extend from C2 to C6)

(b)		$m/z = 105 \text{ C}_6\text{H}_5\text{CO}^+$	1
		$m/z = 77 \text{ C}_6 \text{H}_5^+$ (not Wheland intermediate) (Penalise missing + once) Allow position of + on O or C of CO or outside [] for the fragment ion $[C_6 H_5 CO]^+$	1
		Allow position of + on H or C or outside [] for the fragment ion $[C_6H_5]^+$ $[C_6H_5COCH_2CH_3]^+$ \longrightarrow $C_6H_5CO^+ + CH_3CH_2^-$ ('must be on H or C of CH_2 or outside bracket)	
		(1) for molecular ion (1) for RHS Allow molecular formulae, i.e. $C_9H_{10}O^+$. \longrightarrow $C_7H_5O^+ + C_2H_5$.	2
	(c)	Nucleophilic addition	1
		 1 Q contains asymmetric carbon or chiral centre or are chiral molecules 2 with 4 different groups/atoms attached (stated) not molecules attached 	
		3 <u>planar</u> C=O 4 attack from each side 5 equally likely or equal amounts of each isomer formed 6 <u>Racemic</u> mixture or <u>racemate</u> (Q of L) 7 of mirror images or enantiomers or d/l or +/- or R/S or drawn max	6
	(d)	Conc H ₂ SO ₄ or conc H ₃ PO ₄ or Al ₂ O ₃ or iron oxides Not HCl or HBr	1
		Geometrical or cis-trans	1
		Double bond or C=C not just π cloud (stated not just drawn)	1
		2 Different atoms/groups on each C (not molecules) (stated not just drawn)	1
			[20]

X is methyl propanoate 1 → CH₃CH₂-M4 = 3 arrows M1 M1 for arrow and lone pair, 4 M2 for arrow addition-elimination 1 Spectrum 2 1 if thinks Spectrum 1 = X can only score for structure of YY is CH₃COOCH₂CH3 1 The two marks for explanation are awarded for discussing one or more of the 2 four peaks (not those for the CH₃ of the ethyl groups) for stated δ values the integration or the splitting should be related to the structure: e.g. structure of **X** shows that at $\delta 3.7 - 4.1$ (1) spectrum of X should have integration 3 / singlet (1) or at $\delta 2.1 - 2.6$ (1) spectrum of **X** should have integration 2 / quartet (1) Spectrum 2 has these [OR Spectrum 1 has at 3.7 - 4.1 (1) quartet / integration 2 (1) so not X at 2.1 - 2.6 (1) singlet / integration 3 (1) so not X [10] K₂Cr₂O₇/H₂SO₄ reuced by CH₃CH₂CH₂CH₂OH (1) oxidised to $CH_3(CH_2)_2CHO$ (1) and $CH_3(CH_2)_2COOH(1)$ CH₃CH₂CH₂CHO (1) oxidised to $CH_3(CH_2)_2COOH$ (1) $Cr_2O_7^{2-} + 14H^+ + 6e^- \rightarrow 2Cr^{3+} + 7H_2O$ (1) Equation: 6 Note: Deduct one if all three compounds given as reducing

23.

24.

agents.

(b) Tollens' reduced by $CH_3CH_2CH_2CHO (1)$ oxidised to $CH_3(CH_2)_2COOH (1)$ Equation $[Ag(NH_3)_2]^+ + e^- \rightarrow Ag + 2NH_3 (1)$ 3

(c) $CH_3CH_2CH_2CH_2OH$ (1)

Product CH₃CH₂CH₂CH₂OOCCH₃ (1)

(CH₃)₃COH

Product $(CH_3)_3COOCCH_3$ (1)

4

2

(d) CH₃CH₂CH₂OH has five peaks (1)

 $(CH_3)_3COH$ has two peaks (1)

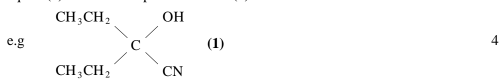
[15]

25. (a) 3 Ketones:

(b) 4 aldehydes:

(c) nucleophilic addition (1)

equal (1) mixture of optical isomers (1)



(d) Reagents are oxidizing agents (1)

Aldehydes can be (easily) oxidized (1)

Ketones are not (easily) oxidized (1)

3

7

[20]

26. Part (a) for each section:

A totally wrong reagent scores zero

An incomplete reagent such as silver nitrate for Tollens, loses the reagent mark, but can get both observation marks.

A wrong reagent such as $[Ag(NH_3)_2]^{2+}$ or bromide water loses the reagent mark and the next mark "gained", i.e. can only score 1/3 if both observations correct

If two test given and results given correctly for both compounds in both tests then full marks If one test on A and a different test on B with only these results given

if both results correct then score 2/3

if either result wrong then score 1/3

if either test would not work as a distinction, then score 0/3

If the candidate says A = ketone (or C = benzene), lose this mark.

If the candidate omits the letters when referring to the pair of compounds, e.g. says alkene decolourises / alkane no reaction penalise one mark only.

(a) (i) penalise observations which just say colour change occurs or only state starting colour

Tollens	[1]	Fehlings / Benedicts	[1]	Brady's or 2,4-dnph	[1]	sodium	[1]
No reaction A	[1]	no reaction A	[1]	no reaction A	[1]	bubbles or hydrogen A	[1]
silver mirror or grey or ppt B	[1]	red or ppt B	[1]	(Yellow / orange) Xtals or ppt	[1]	no reaction B	[1]
(not silver solution)	[1]	not red solution	[1]	not yellow / orange solution	[1]		

Carboxylic acid / H ₂ SO ₄	[1]	Schiff's	[1]	iodoform or I ₂ / NaOH	[1]	PCl ₅	[1]
(sweet) smell A	[1]	no reaction A	[1]	yellow (ppt) A	[1]	(misty) fumes A	[1]
no reaction B	[1]	goes pink B	[1]	no reaction B	[1]	no reaction B	[1]

(ii)

Bromine (water)	[1]	KMnO ₄	[1]	KMnO ₄ / H ₂ SO ₄	[1]	[1]
no reaction C	[1]	no reaction C	[1]	no reaction C	[1]	[1]
decolourised D	[1]	goes brown D	[1]	goes colourless D	[1]	[1]
not clear not discolour (is)ed						

(iii) not just smell for E

an identified (hydrogen) carbonate	[1]	correct metal	[1]	UI or stated indicator	[1]	PCl ₅	[1]
no reaction e	[1]	no reaction E	[1]	no change E	[1]	(misty) fumes E	[1]
bubbles or CO ₂ F	[1]	bubbles or H ₂ F	[1]	red or correct colour F	[1]	no reaction F	[1]

note MAX 8

(b) F has absorption at <u>2500 - 3000 cm⁻¹</u> (due to COOH) (1) **N.B.** Qu asks "How fingerprinting is used" i.e. no marks for simply stating fingerprint region unique.

Compare with (spectrum of) <u>known compound or database</u> (1) (exact) match

(c) major peak
$$[CH_3CO]^+$$
 (1)
m / z 43 (1)
 $CH_3COOCH_3^+ \rightarrow CH_3CO^+ + OCH_3^-$
(1 for molecular ion) (1 for correct other fragment)
Alternative:
major peak $[CH_3]^+$ (1)

m/z 15 (1)

 $CH_3COOCH_3^{+ \centerdot} \rightarrow CH_3^{+} + CH_3COO \cdot \text{ or COOCH}_3 \cdot \text{ or } C_2H_3O_2 \cdot \text{ or } C_3H_6O_2^{+ \centerdot}$

(1) (1 for radical)

If major peak wrong but possible e.g. CH_3OO^+ m/z = 59 no marks so far, but can score up to 2 for

 $CH_3COOCH_3^+ \rightarrow CH_3^+ + CH_3COO^+ \text{ or } ^+COOCH_3 + CH_3$

1 for correct other fragment]

[15]

4

27. (a) X contains > C = O(1)

if X and Y reversed lose this mark but allow remaining max 6/7

∴ X is CH₃CH₂COOH (1)

∴ Y is CH₃CH₂CH₂OH (1)

$$\therefore A \text{ is } CH_3CH_2C \bigcirc O$$

$$OCH_2CH_2CH_3$$
(1)

Propanol X reagent: acidified $K_2Cr_2O_7$ (1) Y reagent: NaBH₄ (1)

Conc H_2SO_4 : catalyst (1) 7

(b) 4

$$CH_{3}CH_{2}CH_{2} \stackrel{H}{\underset{C}{\leftarrow}} COOH \quad \textbf{(1)} \qquad \qquad \begin{matrix} H \\ I \\ CH_{3} \end{matrix} \qquad \begin{matrix} H \\ I \\ CH_{3}CH_{2} \stackrel{I}{\underset{C}{\leftarrow}} C-CH_{2}COOH \quad \textbf{(1)} \\ CH_{3} & \underline{C} \end{matrix}$$

(c)
$$-O\overset{a}{C}H_{2} - 3.1 - 3.9$$
 (1) $-\overset{b}{C}H_{2} - \overset{c}{C} - 2.1 - 2.6$ (1)

6

2

(d) 3269 cm^{-1} :: OH \bigcirc alcohol (1)

$$\therefore \underline{G} \text{ is } \overset{H}{\underset{HO}{\bigvee}} \overset{H}{\underset{OH}{\bigvee}} (1)$$

Notes

(a) first mark for C=O stated or shown in **X** *Ignore wrong names*

Y CH₃CH₂CH₂OH

allow C₃H₇ in **A** if **Y** correct or vice versa

Allow (1) for A if correct conseq to grong X and Y

other oxidising agents: acidified KMnO₄; Tollens; Fehlings

other reducing agents: LiAlH₄; Na/ethanol; NiH₂; Zn or Sn or Fe/HCl

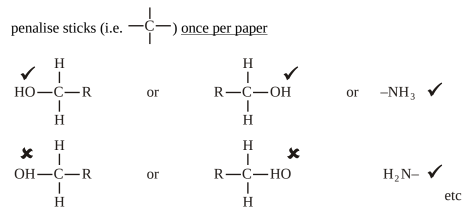
- (b) give (1) for carboxylic acid stated or COOH shown in each suggestion (1) for correct E
 any 2 out of 3 for B, C or D
 allow C₃H₇ for either the B or D shown on the mark schme
 i.e. a correct structure labelled B, C or D or E will gain 2.
- (c) protons a quartet must be correct to score 3 adjacent H mark. Same for b
- (d) allow (1) for any OH (alcohol) shown correctly in any structure ignore extra functional groups. Structure must be completely correct to gain second mark

[19]

Organic points

(1) <u>Curly arrows:</u> must show movement of a pair of electrons, i.e. from bond to atom or from lp to atom / space e.g.

(2) **Structures**



Penalise once per paper

allow
$$CH_3$$
- or $-CH_3$ or CH_3 or CH_3 or H_3C -

28. Identity of **X**; 2-methylpropene (1) (a)

Absorption at 1650 cm⁻¹ indicates an alkene present (1)

OR a chemical answer e.g. Br₂ (aq) brown to colourless

2

(b) Reagents

Step 1 KOH (allow NaOH) (1) alcoholic (1) warm (1) Only allow solvent and warm if reagent correct

Step 2 HBr (1)

Mechanism:

Mechanism:
$$A \rightarrow X$$

$$CH_{3} - CH_{3} - CH_{2} - CH_{2} - CH_{3} -$$

Or a carbocation mechanism

Mechanism
$$X \rightarrow B$$

$$CH_{3} C = C H H Br \rightarrow CH_{3} CH_{3} \rightarrow CH_{3} CH_{3} \rightarrow CH_{3} CH_{3} \rightarrow CH_{3} CH_{3}$$

11

A gives three peaks (1)

B gives one peak (1)

Allow one for "A has more peaks than B" when no number of peaks is given

29. B $1685 \text{ cm}^{-1} \rightarrow \text{C=O (1)}$

D

- C $3340 \text{ cm}^{-1} \rightarrow \text{OH or alcohol (1)}$
- D $1630 \text{ cm}^{-1} \rightarrow \text{C=C or}$ alkene (1) only 1,4-dimethylbenzene will give B as a single compound (1)

$$\begin{array}{c|c}
CH_3 & CH_3 \\
\hline
& electrophilic substitution \\
\hline
or acylation (1) & CH_3 \\
\hline
A & CH_3 & COCH_3
\end{array}$$

[1 mark if 1,2or 1,3- dimethyl]

$$CH_{3} CH = CH_{2} CH (OH) CH_{3}$$

$$CH_{3} CH = CH_{2} CH (OH) CH_{3}$$

$$CH_{3} CH (OH) CH_{3}$$

$$CH_{3} CH (OH) CH_{3}$$

$$CH_{3} CH (OH) CH_{3}$$

$$Ar \xrightarrow{CH} \stackrel{(1)}{\stackrel{}{\circ}} H^{+}$$

$$Ar \xrightarrow{CH} \stackrel{\dot{}{\circ}}{\stackrel{}{\circ}} H_{2}$$

$$CH_{3} \qquad CH_{3} \qquad (1)$$

$$Ar \xrightarrow{\dot{}} \stackrel{\dot{}{\circ}}{\stackrel{}{\circ}} H$$

$$(1) \xrightarrow{\dot{}} H \xrightarrow{\dot{}} CH_{2} \qquad (1)$$

chiral or asymetric C (1) optical isomers or two non-superimposable mirror-image forms possible (1)

nucleophilic

reduction (1)

addition or

[18]

- **30.** A: 4 peaks or 4 different environments (1)
 - 1:2:2:3 (1) OH singlet (1) CH₃ singlet (1)
 - 2 triplets (1) CH₂CH₂ coupled (1)
 - B: 4 peaks or 4 different environments (1)

1:2:2:3 (1) OH singlet (1) OCH₂O singlet (1)

quartet + triplet (1) CH₂CH₃ coupled (1)

- C: 2 peaks or 2 different environments (1)
 - 2:6 or 1:3 (1) CH₃ groups equivalent (1)
 - 2 singlets (1) no coupling (1)

[max 15]

31.	(a)	(i)	electrophile	lone pair acceptor/electron deficient species / electron seeking group / electron lover (not just positive group / species) (1)				
			substitution	replacement / swap / substitution of one atom / group (in a molecule) by another atom / group (1) not molecules replaced	2			
		(ii)	nitronium ion	NO ₂ ⁺ (1)				
			HNO_3++H_2	$SO_4 \rightarrow H_2NO_3^+ + HSO_4$ (1)				
			$H_2NO_3^+ + H_2SO_4 \rightarrow NO_2^+ + H_3O^+ + HSO_4^- $ (1)					
			allow 1 mark f	for $HNO_3 + H_2SO_4 \rightarrow NO_2^+ + HSO_4 + H_2O$				
			allow 2 marks 1 for species, 1	for HNO_3 $2H_2SO_4 \rightarrow NO_2^+ + H_3O^+ + 2HSO_4^-$ for balancing				
			mechanism sho	ows attack by NO ₂ ⁺ with curly arrow from ring (1)				
				ermediate with '+' charge in centre of ring, cle or 2 double bonds (1)				
			mark conseque	entially on electrophile given				
				om C–H bond to ring / deprotonation to give H ⁺ nust be nitrobenzene (1)	6			
		(iii)	above 60 °C li / nitration / (1	kelihood of multiple substitution	1			
			likely to carry	on reacting				
	(b)	,	chlorine (1)					
			AlCl ₃ / FeCl ₃ / Fe / other suitable halogen carrier (1)					
		absei	nce of sunlight /	room temp / anhydrous (1)	3			

		(ii)	compound D (1)		
			ume type of protons / hydrogen are all in same (chemical) conment / equivalent as they are all CH ₃ (1)	2	
		(iii)	compound C (1)		
			ratio = 6:2:1 (1)		
			2 CH ₃ groups have 6 equivalent protons, CH ₂ has 2 protons, CH 1 proton (1)	3	
			must say same type of proton / H		
			penalise first omission of 'same type'		
		(iv)	appropriate unambiguous formula for either but-1-ene or but-2-ene (1)		
			appropriate unambiguous formula for the remaining structural isomer		
			allow 1 mark if candidate draws cis and trans but-2-ene (1)	2	
		(v)	unambiguous structure for 2-methylpropan-1-ol – may be from mechanism (1)		
			curly arrow / attack by OH curly arrow from lone pair or charge only (1)		
			do not allow if Na -OH		
			curly arrow from bond to Cl / dipole shown on C-Cl bond / intermediate showing 3 full and 2 partial bonds to C (1)		
			loss of Cl ⁻ NaCl or Na ⁺ :Cl ⁻ (1)		
			not allowed	4	
			$\begin{array}{l} \mbox{if } S_N \mbox{1 mechanism given:} \\ \mbox{first mark as above - independent} \\ \mbox{second mark for correct carbocation formed including curly} \\ \mbox{arrow from } C \mbox{ to } Cl \mbox{ or } C^{S^+} - Cl^{S^-} \end{array}$		
			third mark for hydroxide attack as above final mark not available (wrong mechanism)		
			penalise missing proton once only		
					[24]
32.	(a)	(M–I	R) ^{+.} Is a radial-cation (1) covalent bond breaks (1)		
		to for	rm a cation (M^+) (1) and a radical (R^-) (1)	4	
	(b)	CH_3C	as a two isotopes (1) $CH_2^{35}Cl = 64$ and $CH_3CH_2^{37}Cl = 66$ (1) Eve abundances $^{35}Cl : ^{37}Cl = 3 : 1$ (1)		
			$CH_2Cl^{+\bullet} \rightarrow CH_3CH_2^+ + Cl^{\bullet}(1)$	4	
	(a)			7	
	(c)		$H_2CH_2Cl \text{ or } 3 \text{ isotopic combinations possible (1)}$ $H_3^{35}Cl_2 = 98 \text{ (1) } C_2H_4^{35}Cl^{37}Cl = 100 \text{ (1) } C_2H_4^{37}Cl_2 = 102 \text{ (1)}$	4	[12]
					['~]

(c) (i) 2-chloro(-2-)methylpropane / (2)methyl 2 chloropropane (1)

33. (a) A
$$C_6H_{14}$$
 (1) $CH_3 - C - C - CH_3$ Ratio 12:2 or 6:1 (1)

B/C C=O (1) C_5H_{10} O (1)

 $CH_3CH_2CCH_2$ CH₃ (1) ratio 6:4 or 3:2 (1)

 $CH_3 - C - C - C - CH_3$
 $CH_3 - C - C - CH_3$

(b) Tollens (1) silver mirror with aldehyde (1)

no reaction with ketone (1)

(or Fehlings red ppt with aldehyde, no reaction with ketone)

Fingerprint region (1)

Exact match with standard (1)

 $CH_3 - C - C - CH_3$
 $CH_3 - CH_3 - CH_3$
 $CH_3 - CH_3$
 CH_3

(c) 3300cm^{-1} :: OH group in both (1) 1650cm^{-1} :: C=C in D (1) :: D is CH₂ = CH- CH₂ CH₂ CH₂ OH (1) (or others) E is \bigcirc OH etc (1)

[18]

- 34. (a) Region 1500–400 cm⁻¹ (1) unique for each compound (1) compare spectrum with that of known compound (1) exact match (1)
 - (b) C₅ esters

Q is
$$H-C$$
 OC(CH₃)₃ (2)

R is CH_3-C OCH(CH₃)₂ (2)

S is (3 peaks) OCH₃ (2 CH_3) OCH₄ (2 CH_3) OCH

T (alcohol) is CH₃CH(OH)CH₃ (2)

(3 peaks)

U (acid) is CH₃COOH (2)

(2 peaks)

T absorption at 3250 cm⁻¹ confirms OH (alcohol) (1)

U absorption at 2900cm⁻¹ confirms OH (acid) or at 1700 cm⁻¹ confirms C=O (1) max 11

[15]

4

35. (a)

 $\mathbf{A} = butanal$

 $\mathbf{B} = \text{methylpropanal}$

C = butanone

 $\mathbf{D} = \text{ethyl ethanoate } (\mathbf{1})$

Ignore numbers in names unless they make them incorrect spellings must be correct accept alternative trivial names correctly spelled

(b) ethanol / correct formula (1)

ethanoic acid / ethanoyl chloride / ethanoic anhydride / correct formula (1)

temperature less than 100 °C / reflux heat / concentrated sulphuric acid (1) dilute sulphuric acid / acid conditions / H^+

(this mark dependent on sensible answers for first two marks)

for ethanoyl chloride, room temperature / dry / anhydrous

for ethanoic anhydride, heat / up to 100 °C

H CH₃ O

butanoic acid

(1)

methylpropanoic acid (1)

2

3

(d) heat with Fehling's solution / ammoniacal silver nitrate / Tollen's reagent / other suitable oxidising system eg acidified dichromate / Schiff's reagent

B gives red, green or brown (precipitate) / silver (mirror) or black/grey (1) precipitate / other, dependent on reagent

C shows no change (1)

B and **C** can be referred to as 'aldehyde' and 'ketone' only if names correct in (a) or if there is some other valid identification

3

(e) (i) **B**

two methyl groups / 6 Hs in identical chemical environments **or** (1) 2 Hs in unique environments

2

(ii) A

four different chemical environments (for protons) (1)

in (i) and (ii), second mark is dependent on first mark

2

(iii) same number of hydrocarbon groups with same number of protons in each 1

(f) nucleophilic addition (1)

(1) for intermediate

(1) for product

allow –ve charge on N but curly arrow must come from C

allow H from HCN or H₂O

[26]