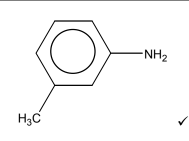
# F324 Rings, Polymers and Analysis

Qı	Question		Expected Answers	Marks	Additional Guidance		
1			+ Br <sub>2</sub> $+$ HBr	1	ALLOW $C_6H_6 + Br_2 \longrightarrow C_6H_5Br + HBr$ DO NOT ALLOW multiple substitution DO NOT ALLOW $Br^+$		
	(b)	(i)	White precipitate <b>OR</b> white solid <b>OR</b> white crystals ✓	2	DO NOT ALLOW colourless DO NOT ALLOW white ppt and bubbles  DO NOT ALLOW Br <sub>3</sub> C <sub>6</sub> H <sub>2</sub> OH OR 2,4,6-tribromophenol OR tribromophenol		
		(ii)	1,2-Dibromocyclohexane ✓	1	ALLOW 1,2dibromocyclohexane OR 1-2dibromocyclohexane OR 12dibromocyclohexane OR cyclo-1,2-dibromohexane DO NOT ALLOW dibromocyclohexane OR C <sub>6</sub> H <sub>10</sub> Br <sub>2</sub> OR structures		
		(iii)	MUST spell delocalised/delocalized or localised/localized correctly once in the answer to obtain all 5 marks  benzene electrons or π-bonds are delocalised ✓  phenol a lone or non-bonded pair of electrons on the oxygen or the OH group is (partially) delocalised into the ring ✓  cyclohexene electrons are localised OR delocalised between two carbons ✓  benzene has a lower electron density OR phenol has a higher electron density ✓ Cyclohexene has a dipole in Br₂ OR	5	ALLOW diagram to show overlap of all 6 p-orbitals for delocalisation  DO NOT ALLOW benzene has delocalised structure or ring  ALLOW diagram to show movement of lone pair into ring for phenol  ALLOW diagram or description of overlap of 2 adjacent p-orbitals for bonding in cyclohexene  DO NOT ALLOW cyclohexene has a C=C double bond IGNORE slip if cyclohexene is written as cyclohexane but π - bonding correctly described  DO NOT ALLOW charge density OR electronegativity instead of		
			phenol can polarise the $Br_2$ <b>OR</b> cyclohexene can polarise $Br_2$ or the $Br$ – $Br$ bond $\checkmark$		electron density <b>ALLOW</b> Br <sup>δ+</sup> <b>OR</b> electrophile Br <sup>+</sup> as alternate to polarise		





**ALLOW** ECF ✓✓ on incorrect amine

 $\rm HNO_2$  + HCl and temp < 10 °C OR NaNO $_2$  + HCl and temp < 10 °C  $\checkmark$ 

alkaline **AND** phenol (if temperature stated must be below 10  $^{\circ}$ C)  $\checkmark$ 

ALLOW

IGNORE Cl<sup>-</sup> ion

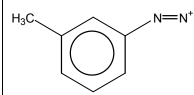
5

**DO NOT ALLOW** if ring is connected to the N triple bond in the diazonium or if diazonium has a negative charge

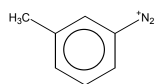
**ALLOW** one mark for correct displayed diazonium if alkyl group is not shown

**ALLOW** 



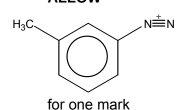


for both marks



for one mark

ALLOW



ALLOW  $N \equiv N^{\dagger}$ for one mark

**ALLOW** NaOH **OR** KOH &  $C_6H_5OH$  **OR** phenoxide ion **OR**  $C_6H_5O^-$  **ALLOW** reagents and conditions from the equations

Total

14

<u> </u>	1004	on	Expected Anguage		Additional Cuidanaa		
<u> </u>	uesti	_	Expected Answers	Marks 1	Additional Guidance		
2	(a)	(i)	silver mirror ✓		ALLOW Ag(s) OR Ag mirror OR precipitate OR ppt OR solid ALLOW brown OR black OR grey		
		(ii)	HOCH₂COOH ✓	1	ALLOW CH <sub>2</sub> OHCOOH OR CH <sub>2</sub> OHCO <sub>2</sub> H OR HOCH <sub>2</sub> CO <sub>2</sub> H OR displayed OR skeletal formula OR HOCH <sub>2</sub> COO <sup>-</sup> DO NOT ALLOW C <sub>2</sub> H <sub>4</sub> O OR 2-hydroxyethanoic acid		
	(b)				ALLOW displayed/skeletal formula/COOHCOOH ✓ ✓		
			HOCH <sub>2</sub> CHO + 3[O] $\rightarrow$ HOOCCOOH + H <sub>2</sub> O reagents $\checkmark$ both products $\checkmark$	2	if molecular formula used $C_2H_4O_2 + 3[O] \rightarrow C_2H_2O_4 + H_2O$ max = 1 $\checkmark$		
				1	Any correctly balanced equation for partial oxidation can score 1 mark $\checkmark$ HOCH <sub>2</sub> CHO + [O] $\rightarrow$ HOCH <sub>2</sub> COOH OR HOCH <sub>2</sub> CHO + 2[O] $\rightarrow$ OHCCOOH + H <sub>2</sub> O OR HOCH <sub>2</sub> CHO + [O] $\rightarrow$ OHCCHO + H <sub>2</sub> O OR HOCH <sub>2</sub> CHO + 2[O] $\rightarrow$ HOCCHO + H <sub>2</sub> O		
	(c)	(i)	HOCH <sub>2</sub> CH <sub>2</sub> OH ✓		ALLOW HO(CH <sub>2</sub> ) <sub>2</sub> OH OR (CH <sub>2</sub> OH) <sub>2</sub> OR skeletal formula OR displayed formula  DO NOT ALLOW molecular formula (C <sub>2</sub> H <sub>6</sub> O <sub>2</sub> )		
		(ii)	curly arrow from H $^-$ to C $^{\delta^+}$ $\checkmark$ dipoles <u>and</u> curly arrow from C=O bond to O $\checkmark$ intermediate $\checkmark$ curly arrow from intermediate to H $^{\delta^+}$ in H $_2$ O/ H $^+$ and if H $_2$ O is used it must show the curly arrow from the O–H bond to the O $\checkmark$	4	ALLOW curly arrow to C even if dipole missing or incorrect  ALLOW maximum of 3 marks if incorrect starting material is used		
			lone pairs are not essential		See page 36 for detailed mechanisms – <i>Alternative 3</i> scores all 4 marks even though the intermediate is not shown		

F324	Mark Scheme	January 2010
	Alternative 1 $HOH_2C$ $O$ $HOH_2C$ $O$ $HOH_2C$ $O$	δ <sup>+</sup> OH
	Alternative 2  HOH <sub>2</sub> C $ \begin{array}{c} \delta^{-} \\ \end{array} $ HOH <sub>2</sub> C $ \begin{array}{c} C \\ \end{array} $ HOH <sub>2</sub> C	products are not required
	H H	$OH_2C$ $O$ $H$ $H$
	Total 9	

F324		Mark Scheme		January 2010			
Quest	ion	Expected Answers	Marks				
3 (a)	(i)	adsorption ✓	1	ALLOW partition OR adsorbtion IGNORE solubility OR desorption DO NOT ALLOW absorption			
	(ii)	measure how far each spot travels relative to the solvent front or calculate the $R_{\rm f}$ value $\checkmark$ compare $R_{\rm f}$ values to those for known amino acids $\checkmark$	2	ALLOW compare R <sub>f</sub> values to database ALLOW compare to known amino acids DO NOT ALLOW retention times for first mark, but the 2nd mark would be available as ✓ ECF ALLOW alternative approach: on the same plate compare position of spots ✓ with known amino acids ✓			
	(iii)	(amino acids won't separate because) similar compounds have similar $R_{\rm f}$ (values) $\checkmark$	1	<b>ALLOW</b> spots often overlap <b>OR</b> don't (fully) separate <b>ALLOW</b> they have similar $R_f$ (values) or similar adsoptions or similar retention times ECF to <b>a(ii)</b>			
(b)	(i)	H 	1	ALLOW RCH(NH <sub>2</sub> )COOH any order for R, NH <sub>2</sub> and COOH but C must be next to H ' <u>CH'</u> must be shown ALLOW CO <sub>2</sub> H brackets around NH <sub>2</sub> are <b>not</b> essential ALLOW structure			
one	given  H <sub>2</sub> N.,,,,  H <sub>2</sub> C.,,,,  H both c	symbol in the "tools" to denote whether or not each chiral C is a reflection of the in the question  HOOC.  HOOC.	3	<ul> <li>each chiral C must have 2 — bonds, 1 wedge bond (IGNORE shading) &amp; 1 dash bond (IGNORE wedge) check the clockwise orientation of each C. For each C start with the H and if on the: <ul> <li>top C the H is followed by COOH it is not a mirror image. If it is a mirror image annotate using RE.</li> <li>bottom C the H is followed by CH<sub>3</sub> it is not a mirror image. If it is a mirror image annotate using RE.</li> </ul> the four groups can be attached in any order. If the molecule is drawn upside down – clockwise becomes anti-clockwise.</li> </ul> MUST check that the drawn structure is non-superimposable irrespective of the orientation or the way it has been drawn.			
				IGNORE bond linkage for all groups			

F324	Mark Scheme		January 2010
(c)	H <sub>3</sub> N — C — COO — H <sub>2</sub> N — C — COO — H <sub>3</sub> N — C — COOH — CH <sub>3</sub> N — C — COOH —		ALLOW CO <sub>2</sub> <sup>-</sup> ALLOW NH <sub>3</sub> <sup>+</sup> If NH <sub>3</sub> fully displayed ALLOW + charge on N or H  If COO fully displayed ALLOW <sup>-</sup> charge on O only
(d)	valine–glycine–leucine ✓	1	ALLOW val–gly–leu  DO NOT ALLOW structures
(e)	H <sub>2</sub> N(CH <sub>2</sub> ) <sub>6</sub> NH <sub>2</sub> ✓ HOOC(CH <sub>2</sub> ) <sub>8</sub> COOH ✓	2	ALLOW H <sub>2</sub> NCH <sub>2</sub> CH <sub>2</sub>

Total

Qι	ıestio	n Expected Answers	Marks	Additional Guidance		
4	(a)	infrared – 1 mark only shows (very broad) peak between 2500–3300 (cm <sup>-1</sup> ) (due to O–H bond) ✓	3	ALLOW (very broad) peak around 3000 (cm <sup>-1</sup> ) OR any stated value between 2500 and 3300 (cm <sup>-1</sup> ) for O–H DO NOT ALLOW peak in range 3200–3550 (cm <sup>-1</sup> )  IGNORE any reference to C=O or C–O as both are also present in an ester OR to fingerprint region		
		<sup>13</sup> C NMR – 2 marks (CH <sub>3</sub> ) <sub>2</sub> CHCH <sub>2</sub> COOH has 4 peaks (due to 4 different C environments) ✓ (CH <sub>3</sub> ) <sub>3</sub> CCOOH has 3 peaks (due to 3 different C environments) ✓		<b>ALLOW</b> <sup>13</sup> C NMR detects the number of/different C environments' for 1 ✓, suitable example for the 2nd mark		
	(b)	splitting pattern explains any two in terms of 'n + 1 rule' for two marks ✓✓ Explains any one peak for 1 mark ✓	6	mark for correct ester  if two splitting patterns are correctly analysed ignore the third		
		singlet therefore adjacent C (if any) has no Hs		ALLOW singlet because next or bonded to an O		
		multiplet <b>OR</b> split into 7 therefore adjacent Cs have lots of/6 Hs		ALLOW multiplet/heptet because next to 2 CH <sub>3</sub> s		
		doublet therefore adjacent C is bonded to 1H		ALLOW doublet because next to a CH		
		must spell <b>one</b> of multiplet / heptet, singlet, doublet correctly  max = 2 marks				
		chemical shifts		<b>ALLOW</b> tolerance on $\delta$ values; 3.6–3.8, 2.6–2.8 and 1.1–1.3		

F324	Ma	ark Sche	cheme		
	two marks if any two absorptions are identified correctly ✓✓ one mark if any one absorption is identified correctly ✓  • peak ~3.7 (ppm) – bonded to an O  • peak ~2.7 (ppm) – indicates it is next to a C=O  • peak ~1.2 (ppm) – bonded to other Cs OR part of a chain  max = 2 marks		(ppm)  ALLOW any two gets 2 marks, any one sco  HC—O  3.7 (ppm)  ALLOW peaks labelled on the spectrum  ALLOW singlet must be bonded to O, multiple doublet to CH or R for both chemical shift must be correctly identified	R——C <b>H</b> 1.2 (ppm)  plet to C=O and harks	
	compound identified as (CH <sub>3</sub> ) <sub>2</sub> CHCOOCH <sub>3</sub> ✓✓ 2 marks  compound identified as CH <sub>3</sub> COOCH(CH <sub>3</sub> ) <sub>2</sub> ✓ 1 mark				
	Total	9			

Question	Expected Answers	Marks	Additional Guidance
5 (a)	НО	1	ALLOW  HO  OF  OF
(b) (i	equation  (CH <sub>3</sub> CO) <sub>2</sub> O + H <sub>2</sub> N OH  reactants ✓  H <sub>3</sub> C OH  products ✓	2	ALLOW (CH <sub>3</sub> CO) <sub>2</sub> O + H <sub>2</sub> NC <sub>6</sub> H <sub>4</sub> OH → CH <sub>3</sub> CONHC <sub>6</sub> H <sub>4</sub> OH + CH <sub>3</sub> COOH  ALLOW H H H DO NOT ALLOW molecular formulae

1 324			•	Mark Scheme Sanda		
		(ii)	$C_{10}H_{11}NO_3$ is $O$ $H_3C$ $O$	1	ALLOW amide shown as either CH <sub>3</sub> CONH– OR H <sub>3</sub> CCONH– OR CH <sub>3</sub> COHN– OR H <sub>3</sub> CCOHN– ALLOW ester shown as either –OCOCH <sub>3</sub> OR –OOCCH <sub>3</sub>	
		(iii)	to ensure that there are no (harmful) side effects $\checkmark$	1	ALLOW impurities reduce effectiveness (of drug) OR might be toxic OR avoids litigation OR harmful OR hazardous ALLOW to ensure that the drug/active component is safe IGNORE dangerous OR nasty OR can kill OR increased dosage	
(	(c)		(aspirin contains) ester <b>AND</b> carboxylic acid ✓	2	IGNORE arene or benzene or aromatic or phenyl or methyl but any other group loses the mark ALLOW carboxyl group DO NOT ALLOW acid	
			(paracetamol contains) amide <b>AND</b> phenol ✓		IGNORE arene or benzene or aromatic or phenyl or methyl but any other group loses the mark ALLOW peptide ALLOW hydroxy(I) DO NOT ALLOW hydroxide or alcohol DO NOT ALLOW amine	
(	(d)	(i)	Both	3	<b>ALLOW</b> hydrolysis by H <sup>+</sup> (aq) or H <sup>+</sup> or HCl(aq) or HCl or H <sub>2</sub> SO <sub>4</sub> (aq)	

F324		Mark Scr	neme January 2010
	Na OR NaOH ✓  COO (Na <sup>+</sup> )  from aspirin  ✓  H <sub>3</sub> C O (Na <sup>+</sup> )  from paracetamol		or H₂SO₄ to give hydroxybenzoic acid + ethanoic acid with aspirin ✓ and ammonium salt of 4-aminophenol + ethanoic acid with paracetamol ✓  ALLOW hydrolysis by OH⁻(aq) or NaOH(aq) and other alkali leading to hydrolysis to give carboxylate salt and phenoxide salt on the ring + ethanoate with aspirin ✓ and 4-aminophenoxide ion + ethanoate ion with paracetamol ✓  ALLOW HNO₃ (and H₂SO₄) to give NO₂ in one or more positions on the ring in both aspirin and paracetamol ✓✓  DO NOT ALLOW NH₃ but correct ammonium salts can be awarded 2 marks ECF  DO NOT ALLOW H₂O but correct products can be awarded 2 marks ECF  if no reagent there cannot be any marks for the products If reagent selected is incorrect but would react with either aspirin or paracetamol ALLOW ✓ ECF for the correct organic product
(ii)	aspirin only NaHCO₃ OR Na₂CO₃ OR metal oxide ✓  H₃C  COO⁻ (Na⁺)  paracetamol only	2	ALLOW Mg, carbonates, NH <sub>3</sub> ALLOW alcohols ( <i>R</i> OH) to give ester if no reagent there cannot be any marks for the products  H <sub>3</sub> C  R  If reagent selected is incorrect but would react with BOTH aspirin and paracetamol ALLOW ✓ ECF for the correct organic product  ALLOW Br <sub>2</sub> water

		Tomo		
Br <sub>2</sub> ✓  H <sub>3</sub> C — C  N — OH  Br	2	ALLOW one or more Br at any position on the ring DO NOT ALLOW Br substitution of OH ALLOW acyl chloride or acid anhydride and corresponding ester ALLOW FeCl₃ to form a purple complex ion (structure not required) ALLOW diazonium and structure showing azo group substituting one of the Hs in the ring if no reagent there cannot be any marks for the products  If reagent selected is incorrect but would react with BOTH aspirin and paracetamol ALLOW ✓ ECF for the correct organic product		
	Total 14			

# **Grade Thresholds**

# Advanced GCE Chemistry A (H034/H434) January 2010 Examination Series

### **Unit Threshold Marks**

U	nit	Maximum Mark	а	b	С	d	е	u
F321	Raw	60	46	40	35	30	25	0
	UMS	90	72	63	54	45	36	0
F322	Raw	100	77	68	59	51	43	0
	UMS	150	120	105	90	75	60	0
F324	Raw	60	43	38	33	29	25	0
	UMS	90	72	63	54	45	36	0

## **Specification Aggregation Results**

Overall threshold marks in UMS (i.e. after conversion of raw marks to uniform marks)

	Maximum Mark	Α	В	С	D	E	U
H034	300	240	210	180	150	120	0

The cumulative percentage of candidates awarded each grade was as follows:

	A	В	C	D	Е	U	Total Number of Candidates
H034	12.9	37.5	62.7	83.1	96.2	100	1415

### 1415 candidates aggregated this series.

For a description of how UMS marks are calculated see: <a href="http://www.ocr.org.uk/learners/ums/index.html">http://www.ocr.org.uk/learners/ums/index.html</a>

Statistics are correct at the time of publication.

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