

## **GCE**

# **Chemistry A**

Unit F324: Rings, Polymers and Analysis

Advanced GCE

Mark Scheme for June 2016

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All examiners are instructed that alternative correct answers and unexpected approaches in candidates' scripts must be given marks that fairly reflect the relevant knowledge and skills demonstrated.

Mark schemes should be read in conjunction with the published question papers and the report on the examination.

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### Abbreviations, annotations and conventions

| Annotation   | Meaning  |  |  |  |
|--------------|--|--|--|--|
| DO NOT ALLOW | Answers which are not worthy of credit                     |  |  |  |
| IGNORE       | Statements which are irrelevant                            |  |  |  |
| ALLOW        | Answers that can be accepted                               |  |  |  |
| ()           | Words which are not essential to gain credit               |  |  |  |
| _            | Underlined words must be present in answer to score a mark |  |  |  |
| ECF          | Error carried forward                                      |  |  |  |
| AW           | Alternative wording  |  |  |  |
| ORA          | Or reverse argument  |  |  |  |

| Q | uesti | ion | Answer   | Mark | Guidance   |
|---|-------|-----|--|------|--|
| 1 | (a)   |     | Stearic acid/octadecanoic acid AND Saturated (fat)   | 1    | ALLOW stearic acid AND no C=C double bonds IGNORE comments about LDL and cholesterol DO NOT ALLOW stearic acid is a trans fatty acid |
|   | (b)   |     | $C_{17}H_{35}COOH + NaOH \rightarrow C_{17}H_{35}COO^-Na^+ + H_2O$                               | 1    | ALLOW C <sub>17</sub> H <sub>35</sub> COONa<br>IGNORE state symbols  |
|   | (c)   |     | At least one ester link fully displayed in a triglyceride structure   O  C  O  C  O              | 2    | $H - C - O$ $C - C_{17}H_{35}$          |
|   |       |     | Correct triglyceride structure ✓   |      | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above for the rest of the structure                   |
|   | (d)   | (i) | M1 Correct structure of a mono unsaturated fatty acid with 18 C                                  | 2    | Must be skeletal formula for M1  |
|   |       |     | M2 Correct position of double bond (12) in a mono unsaturated fatty acid AND trans arrangement ✓ |      | DO NOT ALLOW cis isomer for M2   |

| C | Question |      | Answer   | Mark | Guidance   |
|---|----------|------|--|------|--|
|   |          | (ii) | <b>Each</b> carbon atom <u>in the double bond</u> is attached to (two) different group <b>s</b> /atoms ✓ | 1    | ALLOW Each carbon atom of the double bond is attached to a H atom DO NOT ALLOW functional group for group DO NOT ALLOW the carbon atoms are attached to different groups IGNORE two of the substituent groups are the same |
|   |          |      | Total  | 7    |  |

| Q | uestio | n    | Answer   | Mark | Guidance   |
|---|--------|------|--|------|--|
| 2 | (a)    | (i)  | $H_2N(CH_2)_6NH_2$   | 2    | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  |
|   |        |      | HOOC(CH <sub>2</sub> ) <sub>4</sub> COOH   |      | ALLOW acid chloride, CIOC(CH <sub>2</sub> ) <sub>4</sub> COCI  |
|   |        | (ii) | Type of condensation polymer Polyamide  AND                                      | 1    | Both answers required for one mark  ALLOW nylon IGNORE numbers IGNORE polypeptide DO NOT ALLOW kevlar  |
|   |        |      | Use of condensation polymer Fibres in clothing ✓                                 |      | ALLOW any common use for nylon e.g. fibre, clothing, rope, fishing net, bristles, brushes, bags, cable ties etc. DO NOT ALLOW distinctive uses associated with kevlar or other polymers e.g. bullet-proof vests, crash helmets, bottles, cups IGNORE plastic |
|   | (b)    | (i)  | Ethanoic anhydride  O O O H <sub>3</sub> C O C C C C C C C C C C C C C C C C C C | 2    | ALLOW skeletal formula   |
|   |        |      | Other organic compound CH <sub>3</sub> COOH                                      |      | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  IGNORE names  |

| Question | Answer  | Mark | Guidance   |
|----------|---|------|--|
| (ii)     | FIRST CHECK THE ANSWER ON THE ANSWER LINE IF answer = 2.66 (g) award 3 marks IF answer = 4.36 (g) award 2 marks (% yield not used) IF answer = 7.14 (g) award 2 marks (% yield used | 3    | ANNOTATE WITH TICKS AND CROSSES ETC.   |
|          | incorrectly) n(phenylamine) (= 3.00/93.0) = 0.0323 mol  |      | ALLOW 3 SF: 0.0323 up to calculator value of 0.032258064 correctly rounded   |
|          | $n(compound A) = (0.0323 \times 0.61) = 0.0197 mol$   |      | ALLOW 3 SF up to calculator value  |
|          | Mr (compound A) = 135 AND   |      | Penalise rounding to 2 SF once ALLOW ECF on incorrectly rounded values   |
|          | Mass of compound A = (135)(0.0197) = 2.66 g   |      | Final answer must be expressed to 3 significant figures  |
|          | OR  |      | ALLOW ecf from incorrect Mr  |
|          | n(phenylamine) (= 3.00/93.0) = 0.0323 mol   |      |  |
|          | Mr (compound A) = 135 AND   |      |  |
|          | Theoretical mass of compound A = (0.0323 x 135) = 4.36  |      |  |
|          | Actual mass of compound A = (4.36 x 0.61) = 2.66 g  |      | IF answer = 2.65 (g) award 2 marks unless this alternative method is used (3 marks) 93 g gives 135 g 3.00 g gives 135/93 x 3.00 = 4.35 g |
|          |   |      | 4.35 x 0.61 = <b>2.65</b> g  |

| SWITH TICKS AND CROSSES ETC.<br>Show formation of the electrophile ${}_{2}SO_{4} + HNO_{3} \rightarrow 2HSO_{4}^{-} + H_{3}O^{+} + NO_{2}^{+}$ ${}_{3}O_{4} + HNO_{3} \rightarrow HSO_{4}^{-} + H_{2}NO_{3}^{+}$ ${}_{4}^{+} \rightarrow H_{2}O + NO_{2}^{+}$ |
|---|
| 7 112O + 11O2   |
| ssing or incorrect –NHCOCH <sub>3</sub> on intermediate   |
| <b>LOW</b> intermediate with the $\pi$ -system covering If the ring   |
| H NO₂   |
| harge anywhere inside the 'horseshoe' must have open end towards NO <sub>2</sub>  |
| kulé mechanism  |
| S S L<br>L  |

| Qı | Question |     | Answer  | Mark | Guidance  |
|----|----------|-----|---|------|---|
|    |          |     |   |      | NHCOCH <sub>3</sub> NHCOCH <sub>3</sub> NHCOCH <sub>3</sub> NO <sub>2</sub> |
|    |          |     |   |      | OR  NHCOCH3  NHCOCH3  NHCOCH3   |
|    |          |     | M5 Regeneration of the catalyst:<br>H <sup>+</sup> + HSO <sub>4</sub> <sup>-</sup> → H <sub>2</sub> SO <sub>4</sub> |      | <b>ALLOW</b> $H_3O^+ + HSO_4^- \rightarrow H_2SO_4 + H_2O$                  |
|    |          | (c) | reagents for step 1 Nitrous acid/HNO <sub>2</sub> (and HCl)   | 4    | ALLOW NaNO <sub>2</sub> + HCl   |
|    |          |     | conditions for step 1 ≤10 °C  |      | IGNORE reference to concentration   |
|    |          |     | compound C  |      | <b>ALLOW</b> –OH ionised as –O  |
|    |          |     | HO  |      | ALLOW KOH(aq)/NaOH(aq)/OH-(aq)  |

| Question | Answer                                  | Mark | Guidance   |
|----------|---|------|--|
|          | conditions for step 2 alkaline/alkali ✓ |      | ALLOW dilute NaOH or stated concentration IGNORE NaOH/KOH (must be aqueous) If temperature stated must be below 10°C DO NOT ALLOW heat/boil/warm |
|          | Total                                   | 17   |  |

| Qı | uestio | n   | Answer   | Mark | Guidance   |
|----|--------|-----|--|------|--|
| 3  | (a)    |     | V H − V − O H .  | 2    |  |
|    |        |     | Curly arrow from $OH^-$ to $C(\delta+)$ Dipole correct <b>AND</b> curly arrow from C=O bond to $O(\delta-)$                      |      | First curly arrow must come from either a lone pair on O or negative charge on O   |
|    | (b)    |     | Measure distance moved by spot / distance moved by solvent  Compare (R <sub>f</sub> ) value with data book values/known values   | 2    | <b>ALLOW</b> attempt at calculation of R <sub>f</sub> value using distances measured on the chromatogram <b>IGNORE</b> explanation of how chromatography works   |
|    |        |     | Two amino acids have the same/similar R <sub>f</sub> value <b>OR</b> similar adsorption <b>OR</b> move the same/similar distance | 1    | ALLOW One spot contains two amino acids ALLOW Two amino acids have not separated IGNORE relative solubility ALLOW two of the amino acids have similar structures |
|    | (c)    | (i) | The <b>pH</b> at which the amino acid exists as a <u>zwitterion</u>  | 1    | DO NOT ALLOW PH/ph  ALLOW zwitter ion  |
|    |        |     |  |      |  |

| Question | Answer   | Mark | Guidance  |
|----------|--|------|---|
| (ii)     | H <sub>2</sub> N — C — COO. CH <sub>2</sub>  | 1    | ALLOW correct structural OR displayed OR skeletal formulae OR combination of above as long as unambiguous  Two COO groups are required in the structure  ALLOW –COO Na+ OR -COONa  ALLOW delocalised carboxylate  ALLOW ONA OR  |
| (iii)    | $\begin{array}{c ccccccccccccccccccccccccccccccccccc$                                      | 2    | DO NOT ALLOW -COO-Na OR -O-Na (covalent bond)  ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  ALLOW tripeptide with the 3 amino acids in any order  ALLOW cyclic tripeptide  Isoleucine has two chiral centres, aspartic acid has one chiral centre and glycine has none.  ALL three correct for one mark |
|          | Н—С <sub>*</sub> —СН <sub>3</sub> Н Н Н СН <sub>2</sub> С <sub>2</sub> Н <sub>5</sub> СООН | 9    | ALLOW chiral centres correctly identified if the three amino acids are part of a polypeptide chain  |

| Qı | uestio | n      | Answer   | Mark | Guidance   |
|----|--------|--------|--|------|--|
| 4  | (a)    | 2      | 2(-)hydroxypropanoic acid  | 1    | DO NOT ALLOW 2-hydroxylpropanoic acid IGNORE other dashes, commas and spaces   |
|    | (b)    | (<br>( | Lactic acid synthesised in the laboratory will contain optical isomers/two optical isomers  OR  Lactic acid produced by bacteria will be present as one optical isomer | 1    | ALLOW enantiomer for optical isomer ALLOW racemic mixture IGNORE stereoisomer  |
|    | (c)    |        | CH <sub>3</sub> C C C C C C C C C C C C C C C C C C C  | 1    | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  |
|    | (d)    | (i)    | ——————————————————————————————————————   | 1    | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  DO NOT ALLOW more than one repeat unit  DO NOT ALLOW if structure has no end bonds  IGNORE brackets unless they are used to pick out the repeat unit from a polymer chain  IGNORE n |

| Question | Answer  | Mark | Guidance   |
|----------|---|------|--|
| (ii)     | (Ester links in PLA are) hydrolysed   | 3    | ANNOTATE WITH TICKS AND CROSSES ETC.  ALLOW (ester) hydrolysis/(ester) hydrolyses                                    |
|          | Any two from:   |      | IGNORE acid/alkaline (hydrolysis)  |
|          | Ester (links in the polymer) OR (PLA is a) polyester                              |      |  |
|          | Monomer/lactic acid/product (is soluble because it) forms hydrogen bonds to water |      | IGNORE PLA forms hydrogen bonds to water   |
|          | polymer is photodegradable  |      | IGNORE biodegradable   |
|          | the C=O bond absorbs radiation/uv/light   |      | IGNORE infrared radiation  |
|          | ✓ QWC: hydrolysed/hydrolysis/hydrolyses spelled correctly in the correct context  |      | Maximum of 2 marks if hydrolysed/hydrolysis/hydrolyses does not appear in the answer <b>ALLOW</b> (ester) hydrolyzed |
|          |   |      |  |
|          | Total   | 7    |  |

| Q | Question  |      | Answer  |                        |                                  | Mark | Guidance  |
|---|-----------|------|---|------------------------|----------------------------------|------|---|
| 5 | (a)       | (i)  | 111 1111  |                        |                                  | 3    | One mark for each correct row   |
|   |           |      | <sup>1</sup> H NMR spectrum for 2-aminopropan-1-ol  |                        |                                  | i    | <b>ALLOW</b> $\delta$ values as a range or a value within the specified |
|   |           |      | Chemical shift,   | Relative peak          | Splitting                        |      | range.  |
|   |           |      | δ/ppm   | area                   | pattern                          |      | <b>ALLOW</b> δ values +/- 0.2 ppm.                                      |
|   |           |      | 0.8 – 2.0   | 3                      | doublet                          |      | <b>ALLOW</b> a response that implies a splitting into two for a         |
|   |           |      | 2.3 – 3.0   | 1                      | multiplet                        |      | doublet etc.  |
|   |           |      | 3.3 – 4.2   | 2                      | doublet                          |      | ALLOW sextet/hextet/six (or more than 5) as alternative to              |
|   |           |      |   |                        |                                  |      | multiplet   |
|   |           |      |   |                        | $\checkmark\checkmark\checkmark$ |      | Relative peak area = CH <sub>3</sub> /3H etc. penalise once             |
|   |           | (ii) |   |                        |                                  | 2    | ALLOW correct structural OR displayed OR skeletal                       |
|   |           |      |   |                        |                                  |      | formulae <b>OR</b> a combination of above as long as                    |
|   |           |      | M <sup>+</sup> peak at 75 (peak 1)  |                        |                                  |      | unambiguous   |
|   |           |      | CH <sub>3</sub> CH(NH <sub>2</sub> )CH <sub>2</sub> OH <sup>+</sup> /C <sub>3</sub> H                                       | l <sub>9</sub> NO⁺     | ,                                |      |   |
|   |           |      |   |                        | ✓                                |      |   |
|   |           |      | _ , , , , , , ,   | I 0)                   |                                  |      |   |
|   |           |      | Fragment peak at 44 (pea<br>CH <sub>3</sub> CH(NH <sub>2</sub> ) <sup>+</sup> /C <sub>2</sub> H <sub>6</sub> N <sup>+</sup> | ak 2)                  |                                  |      | Positive charge is essential but ALLOW maximum of one                   |
|   |           |      | $CH_3CH(NH_2)^*/C_2H_6N^*$  |                        | ✓                                |      | mark if both formulae are correct <b>AND</b> neither species has        |
|   | (- )      |      |   |                        | V                                |      | a positive charge   |
| 5 | (b)       | (i)  | Ethanolic ammonia   |                        |                                  | 1    | ALLOW ammonia in a sealed tube  |
|   |           |      | OR ammonia/NH <sub>3</sub> AND ef   | thanol                 |                                  |      | ALLOW dilute ethanolic ammonia/NH <sub>3</sub>                          |
|   |           |      |   |                        | $\checkmark$                     |      | IGNORE heat   |
|   |           |      |   |                        |                                  |      | ALLOW alcohol for ethanol   |
|   |           |      | ,   |                        |                                  |      | DO NOT ALLOW any reference to water or hydroxide ions                   |
|   |           | (ii) | (compound D)  |                        |                                  | 1    | ALLOW correct structural OR displayed OR skeletal                       |
|   |           |      |   | $CH_3$                 |                                  |      | formulae <b>OR</b> a combination of above as long as                    |
|   |           |      |   |                        |                                  |      | unambiguous   |
|   | Н С—СН₂ОН |      |   |                        |                                  |      |   |
|   |           |      |   |                        |                                  |      |   |
|   |           |      |   | N H                    |                                  |      |   |
|   |           |      |   |                        |                                  |      |   |
|   |           |      | H <sub>3</sub> C  | C — CH <sub>2</sub> OH |                                  |      |   |
|   |           |      |   | 1.75                   |                                  |      |   |
|   |           |      |   | ù                      | ✓                                |      |   |
|   |           |      |   |                        |                                  |      |   |
|   |           |      |   |                        |                                  |      |   |

| Question | Answer                    |    | Guidance  |
|----------|---------------------------|----|---|
| (c) (i)  | Alcohol AND Amide/peptide | 1  | IGNORE phenol IGNORE hydroxyl/hydroxy IGNORE attempts to classify alcohol or amide as primary, secondary or tertiary DO NOT ALLOW hydroxide             |
| (ii)     | ОН                        | 2  | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above  ALLOW correct structural OR displayed OR skeletal                 |
|          | NH3*                      |    | formulae OR combination of above as long as unambiguous   |
|          | <b>√</b>                  |    | <b>ALLOW</b> + on N or H i.e. <sup>+</sup> NH <sub>3</sub> or NH <sub>3</sub> <sup>+</sup><br><b>ALLOW</b> NH <sub>3</sub> <sup>+</sup> Cl <sup>-</sup> |
|          | Total                     | 10 |   |

| Qı | uestion | Answer  |   | Guidance   |
|----|---------|---|---|--|
| 6  | (a)     | <u>Equation</u>   | 2 | <b>ALLOW</b> LiAlH <sub>4</sub> / lithium tetrahydridoaluminate(III)/lithium aluminium hydride   |
|    |         | CH <sub>3</sub> (CH <sub>2</sub> ) <sub>3</sub> CHO + 2[H] → CH <sub>3</sub> (CH <sub>2</sub> ) <sub>3</sub> CH <sub>2</sub> OH |   | <b>ALLOW</b> correct structural <b>OR</b> displayed <b>OR</b> skeletal formulae <b>OR</b> a combination of above <b>ALLOW</b> $C_4H_9CHO + 2[H] \rightarrow C_5H_{11}OH$ <b>ALLOW</b> molecular formulae: $C_5H_{10}O + 2[H] \rightarrow C_5H_{12}O$ <b>DO NOT ALLOW</b> —COH for aldehyde |
|    | (b)     |   | 7 | ANNOTATE WITH TICKS AND CROSSES ETC.   |
|    |         |   |   | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  |
|    |         |   |   | IGNORE names if structures are given   |
|    |         | <b>M1</b> Compound <b>F</b> structure is a secondary alcohol with the formula $C_5H_{11}OH$                                     |   | ALLOW 3-methylbutan-2-ol if structure not given  |
|    |         | M2 Compound $\mathbf{F} = CH_3CH(OH)CH(CH_3)CH_3$   | / | ALLOW ECF from an incorrect secondary alcohol for M3 e.g. pentan-2-ol → pentan-2-one e.g. pentan-3-ol → pentan-3-one   |
|    |         | M3 Compound $G = CH_3COCH(CH_3)CH_3$  | / | <b>ALLOW</b> (3-)methylbutanone if structure not given <b>IGNORE</b> any discussion of the reactions of compound <b>G</b> with 2,4-dinitrophenylhydrazine and/or Tollens' reagent.   |
|    |         |   |   | ALLOW 3 SF up to calculator value correctly rounded  |

| Question | Answer   | Mark | Guidance   |
|----------|--|------|--|
|          | M4 n(NaOH) = (0.125 x 22.8/1000) = 0.00285 (mol)  M5 M(compound H) = (0.211/0.00285 =) 74(.0) (g mol <sup>-1</sup> )  M6 Compound H = / CH <sub>3</sub> CH <sub>2</sub> COOH |      | IF M(compound H) = 74 award 2 marks (M4 + M5)  ALLOW ECF from incorrect calculation of amount of NaOH  ALLOW propanoic acid if structure not given   |
|          | M7 Compound I = $H = C = C = C = C = C = C = C = C = C = $   |      | ALLOW ECF from incorrect compound F (alcohol) and/or incorrect compound H (carboxylic acid) to form compound I (ester).  Compounds F, G, H and I must be placed in the correct box or correctly labelled for M2. M3, M6 and M7 |
| (c)      | The structural isomer is:  CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> CH <sub>3</sub>   | 1    | ALLOW correct structural OR displayed OR skeletal formulae OR a combination of above as long as unambiguous  ALLOW 2,2-dimethylpropan-1-ol   |
|          | Total  | 10   |  |

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