Surname				Centre Number	Candidate Number
Other Names					2
	GCE A LEVEL –	NEW			
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CHEMISTRY – A level component 2 Organic Chemistry and Analysis

MONDAY, 19 JUNE 2017 - MORNING

2 hours 30 minutes

	For Examiner's use only		e only
	Question	Maximum Mark	- Mark Awarded
Section A	1. to 7.	15	
Section B	8.	16	
	9.	18	
ADDITIONAL MATERIALS	10.	21	
 In addition to this examination paper, you will need a: calculator: 	11.	9	
Data Booklet supplied by WJEC.	12.	21	
	13.	20	
INSTRUCTIONS TO CANDIDATES	Total	120	

Use black ink or black ball-point pen.

Write your name, centre number and candidate number in the spaces at the top of this page.

Section A Answer all questions in the spaces provided.

Section B Answer **all** questions in the spaces provided.

Candidates are advised to allocate their time appropriately between **Section A (15 marks)** and **Section B (105 marks)**.

INFORMATION FOR CANDIDATES

The number of marks is given in brackets at the end of each question or part-question.

The maximum mark for this paper is 120.

Your answers must be relevant and must make full use of the information given to be awarded full marks for a question.

The assessment of the quality of extended response (QER) will take place in **Q.10**(*f*) and **Q.12**(*c*).

If you run out of space, use the additional page(s) at the back of the booklet, taking care to number the question(s) correctly.

PMT

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5. Draw the displayed formula of the compound that has the same molecular formula as ethanal, CH₃CHO. [1]

4.

H₃C

(a)

(b)

6.	Draw a dot and cross diagram of the hydroxide ion. [1]	Examiner only
7.	A sample of 1-butyl butanoate (boiling temperature 164 °C) is contaminated with around 10% of butanoic acid (boiling temperature 162 °C). Describe how you would obtain a pure sample of	o f
	1-butyl butanoate from this mixture. [3]	-
		·

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SECTION B

Answer all questions in the spaces provided.

8. (a) Ethanedioic acid, (COOH)₂, was first produced by the oxidation of carbohydrates with concentrated nitric acid more than 200 years ago. An equation showing the oxidation of sucrose by this method is shown below.

 $C_{12}H_{22}O_{11} + 12HNO_3 \longrightarrow 6(COOH)_2 + 12NO + 11H_2O$

In a laboratory experiment 25.0 g of sucrose $[M_r 342]$ were added to an excess of concentrated nitric acid, together with a suitable catalyst, in a 1 dm³ conical flask. The mixture was gently warmed to start the reaction and the rate of the exothermic reaction was controlled by appropriate cooling.

(i) Suggest a simple method of heating and subsequent cooling of the reaction mixture. [1]

.....

(ii) After 24 hours, the crystals of ethanedioic acid produced were filtered off, recrystallised and weighed. 18.0g of ethanedioic acid dihydrate, $(COOH)_2.2H_2O$ [M_r 126], were obtained. Calculate the percentage yield of the dihydrate. [2]

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Percentage yield =%

only

6 Examiner (iii) A by-product of the reaction was oxopropanedioic acid. COOH C = OCOOH oxopropanedioic acid State and explain how the ¹³C NMR spectrum of this compound would differ from that of ethanedioic acid. [2] (b) Another method to produce ethanedioic acid uses propene from the petrochemical industry as the starting material. Propene is reacted with nitric acid to give an intermediate, which is then treated with oxygen in a second stage. The nitrogen oxides are then oxidised and the product recycled. ► CH₃CHCOOH $CH_3CH = CH_2 + 3HNO_3$ $2H_2O$ 2NO + ONO_2 intermediate + $\frac{5}{2}O_2$ \rightarrow (COOH)₂ + CO₂ + HNO₃ CH₃CHCOOH + H₂O ONO₂ Combine these two equations to give an overall equation. [2]

(c) A summary of these two industrial processes to make ethanedioic acid is shown in the table.

Method	Organic starting materials	Reaction temperature/°C	Percentage yield/%
1	carbohydrates	< 100	65
2	propene	< 100	90

Give two advantages of using method 1.

1. 2.

(d) Many vegetables contain ethanedioic acid or its salts. In an experiment 140g of Brussels sprouts were treated to produce a solution containing ethanedioic acid. Aqueous calcium hydroxide was added to this solution to precipitate insoluble calcium ethanedioate, which was filtered off, washed and dried. It was then heated strongly to give 0.400g of calcium carbonate.

 $(COOH)_2.2H_2O \xrightarrow{Ca(OH)_2} (COO)_2Ca \xrightarrow{\text{strong heating}} CaCO_3$ $M_r 126 \qquad M_r 128 \qquad M_r 100$

Calculate the mass of ethanedioic acid dihydrate, $(COOH)_2.2H_2O$ in 100 g of Brussels sprouts, giving your answer to an appropriate number of significant figures. [3]

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[2]

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Examiner

(e) Group 2 salts of dicarboxylic acids undergo decarboxylation on heating. For example, strontium ethanedioate [M_r 176] produces strontium carbonate and carbon monoxide.

 $(COO)_2Sr \longrightarrow SrCO_3 + CO$

In an experiment 0.440 g of strontium ethanedioate were heated and produced 61.3 cm³ of carbon monoxide measured at 298 K and 1 atm pressure. Use these figures to confirm that the starting material was pure strontium ethanedioate. [1]

(f) A solution contains either compound **A**, compound **B** or compound **C**.



It is possible to identify the compound by using only aqueous bromine and sodium hydrogencarbonate. Complete the table by showing the observations that are made when the compounds are added separately to each reagent. Write 'no reaction' if no change is observed. [3]

Reagent	Compound A	Compound B	Compound C
aqueous bromine			
sodium hydrogencarbonate			

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9. (a) Derivatives of 2,4-dinitrophenylamine have been used as herbicides for about 80 years.



2,4-dinitrophenylamine

One method of preparing this compound is from phenylamine.



Examiner

(iii) By analogy with the nitration of benzene suggest how you could modify the nitration reaction in (ii) to ensure that the dinitro- product is formed. [1]

2,4-Dinitrophenylamine has a melting temperature of 180 °C and is recrystallised (iv) from a 1:1 mixture of ethanol and water. Outline the essential stages of this recrystallisation process so that pure dry crystals of 2,4-dinitrophenylamine are obtained. You should assume that a hot water bath is available for use. [4]

(b) Another use for 2,4-dinitrophenylamine is in the preparation of the azo dye 'Permanent Red 2G'.



Permanent Red 2G

This is prepared from the diazotisation of 2,4-dinitrophenylamine and then coupling the product with naphthalen-2-ol.

(i) State the reagents and conditions used to produce an aqueous solution of 2,4-dinitrobenzenediazonium chloride from 2,4-dinitrophenylamine. [2]

(ii)

(C)

Permanent Red 2G appears red in white light. State and explain the colour observed, if any, when Permanent Red 2G is viewed in

	I. 	blue light	[1]
	II.	red light	[1]
Ther 2,4-c o be carbo (i)	e are linitro more on do In ar Use	e environmental concerns about the use of herbicides phenylamine. Naturally-occurring herbicides, such as limonene, are de acceptable. Limonene is an unsaturated compound that contains to uble bonds in each molecule. In experiment 4.08 g of limonene just removed the colour from 9.58 g this information to calculate the relative molecular mass of limonene	based on considered wo carbon- of bromine. e. [3]
		<i>M</i> =	
(ii)	Whe men (i) to	en limonene reacts completely with hydrogen under suitable thane is produced. Use information from the question and your ans o find the relative molecular mass of menthane. Explain your answer	conditions, wer to part : [2]
		<i>M</i> _r =	
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Examiner

10. (a) Hydroxyproline is a cyclic α -amino acid that is produced in the liver. It is an essential part of the muscle protein collagen.



hydroxyproline

(i) Explain why the melting temperature of hydroxyproline is about 240 °C, which is much higher than expected from the formula given. [2]

(ii) Give the displayed formula of the compound formed when hydroxyproline has acted

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[1]

(b) The formula of a dipeptide formed between 2-aminopropanoic acid and another amino acid is shown below.



Give the displayed formula of the amino acid that forms this dipeptide with 2-aminopropanoic acid. [1]

as a base.

Examiner

(c) Secondary protein structure is concerned with hydrogen bonding between certain groups.
 State the groups present that take part in this hydrogen bonding and explain why it occurs.

(*d*) The graph shows the boiling temperatures of some unbranched primary alcohols, R–OH, and those of the corresponding unbranched primary thiols, R–SH.



Examiner only Explain why the boiling temperature of methanol is higher than would be suggested (i) by the trend in other alcohols. [3] (ii) Explain why the boiling temperatures of the thiols are lower than the boiling temperatures of the corresponding alcohols. [2]

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Turn over.

(e)	There react cons	e are two ed with eith idered as e	comm ner et quilibr	non methods hanoyl chloric rium reactions	of prod de or et	ucing 1-butyl ethan nanoic acid. Both of	oate. Butan- f these react	1-ol can be ions can be	Examiner only
CH₃CI	H ₂ CH	₂ CH ₂ OH(I)	+	CH ₃ COCI(I)		CH ₃ COOCH ₂ CH ₂ C	H ₂ CH ₃ (I) +	HCI(g)	
CH₃Cŀ	H₂CH₂	₂ CH ₂ OH(I)	+	CH ₃ COOH(I)		CH ₃ COOCH ₂ CH ₂ C	CH ₂ CH ₃ (I) +	H ₂ O(I)	
	(i)	Use the ec yield of 1-	quation outyl e	ns to suggest tethanoate than	why the n the me	method using ethano ethod using ethanoic	pyl chloride gi acid.	ves a highe [3]	r - -
	(ii)	Suggest o	ne pr	actical disadva	antage o	of the method that us	ses ethanoyl	chloride. [1]	

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(f) Compound **J** is a polyamide that is produced from two different monomers. This polyamide is hydrolysed by boiling it with aqueous sodium hydroxide. After acidifying the mixture obtained from hydrolysis, compound **K**, a dicarboxylic acid of general formula $HOOC-(CH_2)_n-COOH$ is isolated from the mixture. 1.24g of compound **K** is titrated against aqueous sodium hydroxide of concentration 0.500 mol dm⁻³. 37.55 cm³ are needed to just neutralise the acid.

Compound L is also formed from hydrolysis of the polyamide. It is a diamine, $H_2N-R-NH_2$, where R is a group containing only carbon and hydrogen. 0.500 g of this diamine reacts with nitric(III) acid to produce 278 cm³ of nitrogen gas, measured at 298 K and 1 atm.

 $H_2N-(C_x H_{2x})-NH_2 \longrightarrow 2N_2 + \text{ other products}$

The 13 C NMR spectrum of compound L shows that the R group contains three different carbon environments.

Use the information provided to deduce the displayed formulae of compounds **K** and **L**, and hence give the formula of the repeating section of polyamide **J**. [6 QER]

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11. (a) Furan, C_4H_4O , is a cyclic compound that contains one oxygen atom as part of the ring.

(i) The enthalpies of hydrogenation of dihydrofuran and furan are shown below.



Examiner

(b) But-2-enal can be used to prepare 2-oxopentanoic acid by a sequence of reactions.



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Examiner only

12. (a) When ethene is passed into aqueous chlorine one of the products is 2-chloroethanol.



(i) The presence of a C–Cl bond in this compound can be shown by a simple test tube reaction. Outline the practical steps that are used in this method. You can assume that 2-chloroethanol is soluble in water.



(ii) The oxidation of 2-chloroethanol (M_r 80.6) produces chloroethanoic acid together with a little chloroethanal.



5.80 g of chloroethanol were oxidised and the chloroethanoic acid in the resulting mixture reacted with 0.0600 mol of sodium hydroxide in a 1:1 stoichiometric ratio. Show that the percentage conversion to chloroethanoic acid was 83%. [3]

Examiner

(b) DDT is an effective insecticide but its use in recent years has become restricted because of its persistence in the environment. The DDT that is sold is a mixture of closely related compounds that includes DDT, DDE and DDD.







DDE



DDD

(i) Suggest why DDT is largely insoluble in water. [1]
(ii) Suggest suitable reagent(s) that can be used to convert DDE into DDD. [1]

Examiner only



(iii) Many areas around former DDT manufacturing sites remain contaminated. The soil from one contaminated area was analysed by gas chromatography.



Turn over.

Examiner only Over the last thirty years the production and use of CFCs has declined mainly because of their adverse environmental effects. Using fluorotrichloromethane as your example, outline how the compound causes a reduction in the amount of ozone present in the (C) upper atmosphere and state two problems that can be caused by this reduction. [6 QER]

Examiner only

(d) The formula of chloropentafluoroethane is shown below.



State and explain what you would expect to see in the molecular ion region of its mass spectrum and in its ¹³C spectrum. [4]

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13.	(a)	2,3-Dihydroxypropanal (glyceraldehyde), HOCH ₂ CH(OH)CHO, contains a chiral cer	ntre.	Examine only
		(i) State what is meant by a <i>chiral centre</i> .	[1]	
		(ii) Draw the two mirror image forms of 2,3-dihydroxypropanal.	[1]	
	(b)	Glucose, like 2,3-dihydroxypropanal, contains an aldehyde group. Fehling's / Benedict's solutions are used to show the presence of an aldehyde.		
		Describe another simple test to show the presence of this group. You should include the reagent(s) used and any observations made.	[2]	

Examiner

(c) Glucose occurs as two enantiomers. The concentration of an enantiomer can be found by using a polarimeter to measure the rotation of the plane of plane polarised light. In an experiment using an aqueous solution of D-glucose at a certain temperature, the initial rotation of the plane of plane polarised light was 20° using a cell of 1 dm length to hold the glucose solution.

Use the formula below to calculate the concentration of glucose in the solution in g dm⁻³. [2]

$$[\alpha_D] = \frac{r \times v}{m}$$

where specific rotation at this temperature [α _D] rotation of the plane of plane polarised light (r) volume of solution in 1 dm cell (v)	= 112° = 20° = 15 cm ³
m is the mass of glucose in 15 cm ³ of solution	
Concentration =	g dm ⁻³
Butan-2-ol (M_r 74) can be dehydrated to but-1-ene and but-2-	ene.
$CH_3CH_2CH(OH)CH_3 \longrightarrow CH_3CH_2CH=CH_2 + CH_3CH_3CH_2CH(OH)CH_3$	H ₃ CH=CHCH ₃ around 70%
(i) State a dehydrating agent for this reaction.	[1]
(ii) Both but-2-ene and penta-1,3-diene can exist as <i>E-Z</i> iso of isomerism occurs and give the displayed formula of <i>i</i>	omers. Explain why this type E-penta-1,3-diene. [2]

(d)

(e) Compound R is a neutral saturated aliphatic compound that contains only carbon, hydrogen and oxygen. The percentage of oxygen by mass is 24.6. This compound is hydrolysed in acid solution to give two new organic compounds, S and T. Further information about R, S and T is given below.

Compound R

- The ¹³C NMR spectrum shows that there are five different carbon environments
- The ¹H NMR spectrum shows the following signals

Signal δ/ppm	Relative peak area
1.20 singlet	9
1.25 triplet	3
4.11 quartet	2

- The infrared absorption spectrum shows a peak at 1731 $\rm cm^{-1}$ but no peak at 2500–3550 $\rm cm^{-1}$
- It is not an aldehyde or a ketone

$\text{Compound}\; \boldsymbol{S}$

- The ¹³C NMR spectrum shows that there are three different carbon environments
- The ¹H NMR spectrum shows the following signals

Signal δ/ppm	Relative peak area
1.20 singlet	9
around 11	1

• An aqueous solution is weakly acidic

Compound **T**

- The ¹³C NMR spectrum shows that there are two different carbon environments
- The ¹H NMR spectrum is shown below relative peak areas 1, 2 and 3



• The compound is neutral

Use this information to deduce the formulae of compounds S and T and hence the displayed formula of compound \mathbf{R} .	Examiner only
It is not necessary to name any of the compounds. [11]	

END OF PAPER

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For continuation only.	Examiner only

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