Questions

Q1.

A thermometric titration is a method for finding the end-point of a titration between aqueous solutions of ammonia and ethanoic acid.

A thermometric titration was carried out using the following steps:

- the temperatures of the aqueous ammonia and ethanoic acid solutions were measured and found to be 20.1 °C
- 30 cm³ of the aqueous ammonia was placed in a polystyrene cup
- a 10 cm³ portion of an ethanoic acid solution, concentration 1.10 mol dm⁻³ , was added to the polystyrene cup, the mixture stirred and the temperature measured
- further 10 cm³ portions of ethanoic acid solution were added, the mixture stirred, and the temperature measured immediately after each addition, until a total of 80 cm³ had been added.

Results for this experiment are shown in the table.

Volume of ethanoic acid added /cm³	0	10	20	30	40	50	60	70	80
Temperature /°C	20.1	21.8	23.5	25.1	26.4	25.8	24.9	24.1	23.3

(i) T	he	tem	perature of t	he reaction	n mixtu	re initia	ally inc	reased	l becau	ise the	reactio	on is	(4)
20 20 20 20 20 20 20 20 20 20 20 20 20 2] E		endothermic endothermic exothermic	so energy	y is rele is abso	eased b	by the by the v	water water					(1)
` '			e main reaso creased.	n why, afte	er the e	end-poi	int was	s reach	ed, the	tempe	rature	of the	(1)

(Total for question = 2 marks)

Q2.

A thermometric titration is a method for finding the end-point of a titration between aqueous solutions of ammonia and ethanoic acid.

A thermometric titration was carried out using the following steps:

- $\bullet~$ the temperatures of the aqueous ammonia and ethanoic acid solutions were measured and found to be 20.1 $^{\circ}\text{C}$
- 30 cm³ of the aqueous ammonia was placed in a polystyrene cup
- a 10 cm³ portion of an ethanoic acid solution, concentration 1.10 mol dm⁻³, was added to the polystyrene cup, the mixture stirred and the temperature measured
- further 10 cm³ portions of ethanoic acid solution were added, the mixture stirred, and the temperature measured immediately after each addition, until a total of 80 cm³ had been added.

Results for this experiment are shown in the table.

Volume of ethanoic acid added /cm³	0	10	20	30	40	50	60	70	80
Temperature /°C	20.1	21.8	23.5	25.1	26.4	25.8	24.9	24.1	23.3

(i) Plot the results using the axes provided.

Include two straight lines of best fit, extrapolated until they meet.

27 - 26 - 25 - 24 - 23 - 22 - 21 - 20 - 0 10 20 30 40 50 60 70 80

Volume of added ethanoic acid solution / cm3

(2)

(ii)	Determine the maximum temperature rise from your graph.	(1)
	Calculate the number of moles of ethanoic acid, with a concentration of 1.10 mol dm ⁻ ded at the end-point of the reaction.	
	The reaction that occurs is $NH_3(aq) \ + \ CH_3COOH(aq) \ \rightarrow \ NH_4^+(aq) \ + \ CH_3COO^-(aq)$ Calculate the enthalpy change per mole for this reaction. Include a sign and units in your answer. [Assume: specific heat capacity of the solution at the end-point = 4.18 J g $^{-1}$ °C $^{-1}$ 1.00 cm 3 of the solution at the end-point has a mass of 1.00 g]	(3)
	(Total for question = 8 mar	ks)

Q3.

This question is about the enthalpy change of combustion of methanol.

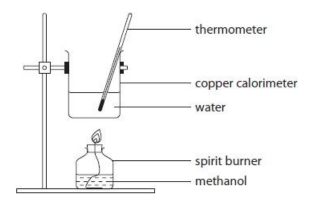
A teacher asked two students to carry out a practical task to determine the enthalpy change of combustion of methanol.

Both students were provided with the same apparatus and chemicals.

The following procedure was provided for the students.

Procedure

- Measure out 150 cm³ of distilled water, using a 250 cm³ measuring cylinder.
- Transfer the water to a copper calorimeter and note the initial temperature of the water (to the nearest 0.5°C) in **Table 1**.
- Weigh the spirit burner containing methanol and record its mass in **Table 1**.
- Place the spirit burner under the copper calorimeter, as shown in the diagram.
- Ignite the spirit burner and burn the methanol, whilst stirring the water with the thermometer.
- After heating the water for three minutes, extinguish the flame and immediately record the **highest** temperature reached by the water.
- As soon as possible, reweigh the spirit burner containing the methanol and record its mass in **Table 1**.



The results of Student 1 are recorded in **Table 1**.

Mass of spirit burner plus methanol before burning / g	213.47
Mass of spirit burner plus methanol after burning / g	211.87
Mass of methanol burned / g	
Highest temperature of the water / °C	64.5
Initial temperature of the water / °C	22.0
Temperature change of the water / °C	

Table 1

(a) Complete **Table 1**, giving the values to an appropriate number of decimal places.

(b) Write the equation that represents the reaction that occurs when the standard
enthalpy change of combustion of methanol, CH ₃ OH(I), is measured. Include
state symbols.

(2)

(c) Use Student 1's result to calculate the enthalpy change of combustion of methanol in kJ mol^{-1} .

Give your answer to an appropriate number of significant figures.

Specific heat capacity of water = 4.18 J g^{-1} °C⁻¹ Density of water = 1.00 g cm⁻³

(4)

(d) Student 1 compared the experimental value for the enthalpy change of combustion of methanol obtained in part (c) with the standard value given on the internet. The student's value was **less exothermic** than the standard value.

Student 1 decided to evaluate the uncertainty in the measurements made in this experiment.

(i) Student 1 used a 250 cm³ measuring cylinder to measure the volume of 150 cm³ distilled water. The uncertainty in this volume measurement is ±1 cm³. Calculate the percentage uncertainty in the volume of distilled water that Student 1 measured in the experiment.

(1)

	(ii) Compare and contrast the use of a 250 cm³ measuring cylinder to measure out the 150 cm³ distilled water with the use of a 25 cm³ measuring cylinder (uncertainty ±0.2 cr for each volume measurement) six times to measure the same volume.	
	(iii) Student 1 calculated the uncertainties in the remaining measurements. However, Student 1 realised that the measurement uncertainties did not explain the difference between the experimental value for the enthalpy change of combustion of methanol calculated in part (c) and the value obtained from the internet. Other than human error, give three reasons for the difference in the values.	(3)
••		
••		
••		

(e)	Student 1 decided to repeat the experiment.	
	Student 1 used the copper calorimeter and water from the first experiment and recorde the initial temperature as 60.0°C. Student 1 burned exactly the same mass of methanol as in the first experiment.	ed
	Explain, with a reason, how the value for the enthalpy change of combustion of metha from this experiment would differ, if at all, from the value obtained in the first experime	
•••		•
•••		•
(t)	Student 2 followed the existing instructions provided but extinguished the flowe ofter	•
	Student 2 followed the original instructions provided, but extinguished the flame after ur minutes rather than after three minutes.	
	Explain how the value calculated by Student 2 for the enthalpy change of combustion methanol compared with that obtained in Student 1's first experiment.	of
		(2)
en	Another student, Student 3, used the results from Student 1's first experiment to find thalpy change of combustion of methanol. Student 3 incorrectly used a value of 46.0 g ol ⁻¹ for the molar mass of methanol.	the
	State and justify how this mistake would affect the calculated value for the enthalpy change of combustion of methanol.	
	change of combustion of methanol.	(2)

(Total for question = 21 marks)

Q4.

This question is about enthalpy changes.

The equations for the combination of gaseous atoms of carbon and hydrogen to form methane, CH_4 , and propane, C_3H_8 , are

C(g) + 4H(g)
$$\rightarrow$$
 CH₄(g) $\Delta H = -1652 \text{ kJ mol}^{-1}$
3C(g) + 8H(g) \rightarrow C₃H₈(g) $\Delta H = -3998 \text{ kJ mol}^{-1}$

Calculate:

(i) the mean bond enthalpy of a C--H bond.

(1)

(ii) the mean bond enthalpy of a C--C bond.

(2)

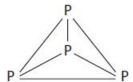
(Total for question = 3 marks)

Q5.

The halogens are elements in Group 7 of the Periodic Table.

Chlorine compounds have many uses, including water treatment.

(i) Chlorine and phosphorus (P₄) can react to form phosphorus(V) chloride. The structure of a molecule of phosphorus is



Some mean bond enthalpy values are shown in the table.

Bond	Mean bond enthalpy / kJ mol ⁻¹
P—P	+198
Cl—Cl	+243
P—Cl	+326

Calculate the enthalpy change for the reaction between chlorine and phosphorus to form phosphorus(V) chloride.

$$10Cl_2 + P_4 \rightarrow 4PCl_5$$

(3)

(Total for question = 4 marks)

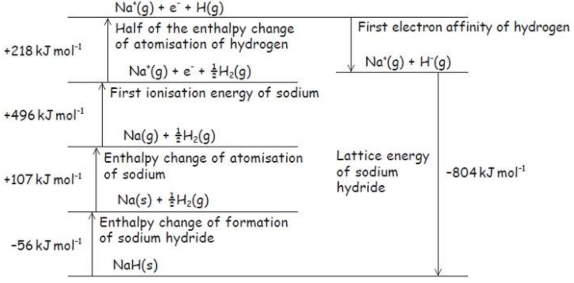
II)	Give a reason why bond enthalpy values are always positive.	
	(1)
• • •		

Q6.

Sodium hydride, NaH, can be used to generate hydrogen for fuel cells.

In order to calculate the first electron affinity of hydrogen, a student was asked to draw a Born-Haber cycle for sodium hydride.

The cycle had **two** errors but the numerical data were correct.



(i) Id	dentify and correct the two errors in this Born-Haber cycle.	
		(2)
(ii) C cycle	Calculate the first electron affinity, in kJ mol $^{-1}$, of hydrogen, using the values given in the.	ne

(1)

Q7.

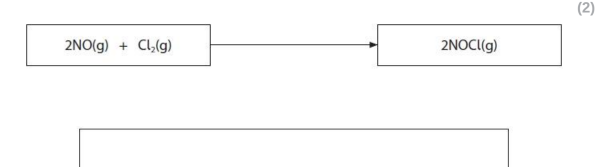
Nitrogen monoxide and chlorine gases react together to form a single product, nitrosyl chloride, NOCI.

Below 100 °C the yield of NOCI is almost 100 %, but as the temperature rises the yield of NOCI decreases as the equilibrium position shifts to the left.

$$2NO(g) + Cl_2(g) \rightleftharpoons 2NOCl(g)$$
 $\Delta_r H^{\Theta} = -75.6 \text{ kJ mol}^{-1}$

(i) Complete the Hess cycle to enable you to calculate the enthalpy change of formation, $\Delta_l H_{298}^{\Phi}$, of NOCI.

Include state symbols.



(ii) Calculate the enthalpy change of formation, $\Delta_f H_{298}^{\bullet}$, of NOCI given the data

$$2NO(g) + Cl_2(g) \rightleftharpoons 2NOCl(g)$$
 $\Delta_r H_{298}^{\Theta} = -75.6 \text{ kJ mol}^{-1}$ enthalpy change of formation of NO, $\Delta_f H_{298}^{\Theta}$, = $+90.3 \text{ kJ mol}^{-1}$

(2)

Q8.

Answer the question with a cross in the box you think is correct \boxtimes . If you change your mind about an answer, put a line through the box \boxtimes and then mark your new answer with a cross \boxtimes .

An equation for the formation of ammonia using the Haber process is shown.

$$N_2(g) + 3H_2(g) = 2NH_3(g)$$

(i) Calculate the enthalpy change for the forward reaction shown in the equation, selecting from the bond enthalpies in the table.

Include a sign in your answer.

(3)

Bond	Mean bond enthalpy / kJ mol ⁻¹
N—N	158
N=N	410
N≡N	945
N—H	391
Н—Н	436

(ii) A data book gives the standard enthalpy change of formation of ammonia as –46.1 kJ mol⁻¹.

Give two reasons for the difference between this value and the value that you calculated in (a)(i).

(2)

Reason 1	
Reason 2	

(iii) What is the percentage atom economy, by mass, for ammonia in the forw	
THE WINDS IS THE DESCENTAGE STOM ECONOMY BY MASS. FOR AMMONIA IN THE TORK	Mara reaction A
tiii) vyriat is trie bereeritade atom economy, by mass, for aminioma in trie forv	waru reaction:

$$N_2(g) + 3H_2(g) = 3(g)$$

- (1)
- □ A 17.6 %□ B 50.0 %□ C 82.4 %
- D 100 %
- (iv) What is the equilibrium expression for K_c ?

$$\square \quad \mathbf{A} \quad \mathcal{K}_{c} = \frac{[N_{2}][3H_{2}]}{[2NH_{3}]} \tag{1}$$

$$\square$$
 C $K_c = \frac{[NH_3]^2}{[N_2][H_2]^3}$

(Total for question = 7 marks)

Q9.

When solid calcium sulfate dihydrate, $CaSO_4 \cdot 2H_2O$, is heated in a crucible, it forms solid calcium sulfate hemihydrate, $CaSO_4 \cdot \frac{1}{2}H_2O$.

When water is added to calcium sulfate hemihydrate, there is a rise in temperature.

A student decided to investigate this reaction using the following procedure:

- Step 1 10 cm^3 of distilled water is measured using a measuring cylinder having an uncertainty of $\pm 0.5 \text{ cm}^3$, and is placed in an insulated cup with a lid.
- Step 2 A thermometer with an uncertainty of ± 0.5 °C is placed in the water.
- Step 3 Exactly 10.00 g of calcium sulfate hemihydrate is weighed out using a balance with an uncertainty of ± 0.005 g.
- Step 4 The weighed quantity of calcium sulfate hemihydrate is added to the water in the insulated cup.
- Step 5 The mixture in the insulated cup is stirred until no further temperature change is observed.

Results

Temperature of the water before adding the solid = 23.5 °C

Maximum temperature of the mixture after adding the solid = 26.3 °C

Other data

Molar mass of calcium sulfate hemihydrate, $CaSO_4$ ·½ H_2O = 145.2 g mol⁻¹ Density of water = 1.00 g cm⁻³

(i) Calculate the minimum volume of water needed to convert 10.00 g of CaSO₄· $\frac{1}{2}$ H₂O into CaSO₄·2H₂O.

 Calculate the enthalpy change, in kJ mol ⁻¹ , for this reaction. Include a sign in your answer and give your answer to an appropriate number of hificant figures.
Assume that the liquid has a mass of 10.00 g and a specific heat capacity of 4.18 J g ⁻¹
°C ⁻¹ . (4)
Deduce which measurement has the greatest uncertainty in this experiment. Justify your wer by calculating the percentage uncertainty of this piece of apparatus.
(2)
(Total for question = 8 marks)

Q10.

This question is about enthalpy changes and entropy changes.

Propan-1-ol is dehydrated to form propene.

The relevant mean bond enthalpies are given in the table.

Bond	Mean bond enthalpy / kJ mol ⁻¹
C— C	347
C — C	612
С—Н	413
0—Н	464

Calculate the C—O mean bond enthalpy, using the mean bond enthalpies given in the table and the enthalpy change of reaction.

(3)

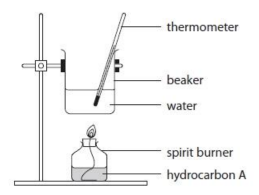
(Total for question = 3 marks)

Q11.

In an experiment, 1.000 g of a hydrocarbon, **A**, was burned completely in oxygen to produce 3.143 g of carbon dioxide and 1.284 g of water.

In a different experiment, the molar mass of the hydrocarbon, $\bf A$, was found to be 84.0 g mol⁻¹.

A spirit burner was filled with the liquid hydrocarbon, **A**. The burner was weighed, lit and then used to raise the temperature of a quantity of water in a beaker, as shown in the diagram. The burner was then reweighed.



Results

112.990 g
112.732 g
250 cm ³
21.3°C
29.5°C

Other data

Density of water	1.00 g cm ⁻³		
Specific heat capacity of water	4.18 J g ⁻¹ °C ⁻¹		
Molar mass of hydrocarbon A	84.0 g mol ⁻¹		

(i) Use these results to calculate the enthalpy change of combustion of hydrocarbon $\bf A$ in kJ mol⁻¹.

Give your answer to an appropriate number of significant figures and include a sign.

(3)

(ii) The beaker used in this experiment was made of copper rather than glass. Give a reason for this.	
	(1)
(Total for question = 4 mar	ks)

Q12.

In acid-base neutralisation reactions, there is a temperature change.

The enthalpy change when hydrochloric acid reacts with aqueous ammonia is -53.4 kJ mol⁻¹.

$$HCI(aq) + NH_3(aq) \rightarrow NH_4CI(aq)$$

Calculate the temperature change you would expect when $25.0~\text{cm}^3$ of $1.00~\text{mol}~\text{dm}^{-3}$ hydrochloric acid is mixed with $25.0~\text{cm}^3$ of $1.00~\text{mol}~\text{dm}^{-3}$ aqueous ammonia.

Give your answer to an appropriate number of significant figures.

Assume: the density of the solution is 1.00 g cm $^{-3}$ the specific heat capacity of the solution is 4.18 J g $^{-1}$ °C $^{-1}$

(3)

(Total for question = 3 marks)

Q13.

Ethanol burns completely in excess oxygen.

$$C_2H_5OH(I) + 3O_2(g) \rightarrow 2CO_2(g) + 3H_2O(I)$$

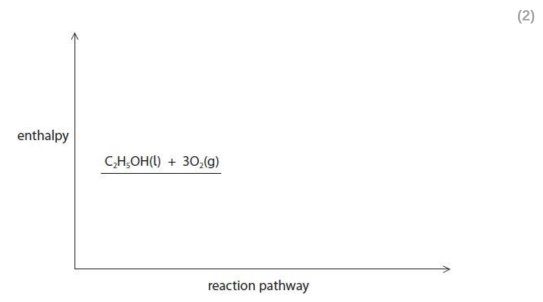
(i) The table shows some mean bond enthalpy data.

Bond	с—с	С—Н	с—о	о—н	0—0	c=0
Mean bond enthalpy / kJ mol ⁻¹	347	413	358	464	498	805

Calculate the enthalpy change, in kJ mol⁻¹, for the complete combustion of 1 mol of ethanol.

(3)

(ii) Complete the reaction profile diagram for the combustion of ethanol and fully label the diagram.



(iii) A data book value for the standard enthalpy change of combustion of ethanol is -1 kJ mol $^{-1}$.	1367.3
Give the main reason why the value you calculated in (i) is different from this data be value.	oook
	(1)
(Total for question = 6 n	narks)

Q14.

A student carries out two experiments to determine the enthalpy change that occurs when anhydrous sodium carbonate reacts to form hydrated sodium carbonate.

$$Na_2CO_3(s) + 10H_2O(l) \rightarrow Na_2CO_3.10H_2O(s)$$

(a) In the first experiment, the student determines the enthalpy change of solution for anhydrous sodium carbonate.

50.0g of distilled water is placed in a polystyrene cup and the temperature is recorded. A sample of anhydrous sodium carbonate is added to the water, the mixture is stirred and the final temperature recorded.

The results for this experiment are shown in the table.

mass used / g	5.09
initial temperature / °C	27.0
final temperature / °C	32.4

Calculate the enthalpy change of solution, in kJ mol⁻¹, for anhydrous sodium carbonate. Give your answer to an appropriate number of significant figures and include a sign. [Use 4.18 J g⁻¹ °C⁻¹ as the specific heat capacity of water]

$$Na_2CO_3(s) + aq \rightarrow Na_2CO_3(aq)$$

(4)

(b) I	n the second experiment,	the student	determines	the enthalpy	change of	solution for
hydra	ted sodium carbonate.					

$$Na_2CO_3.10H_2O(s) + aq \rightarrow Na_2CO_3(aq)$$
 $\Delta H = + 53.7 \text{ kJ mol}^{-1}$

Complete the Hess cycle and, together with your answer to (a) calculate the enthalpy change when anhydrous sodium carbonate reacts to form hydrated sodium carbonate. Include a sign in your answer.

 $Na_2CO_3(s) + 10H_2O(l)$ \rightarrow $Na_2CO_3.10H_2O(s)$

(c)	Hydrated sodium carbonate slowly loses some water of crystallisation when left in a	air.
	Explain how the enthalpy change in the second experiment would compare with the book value if an old sample of hydrated sodium carbonate had been used.	data
		(2)
•••		••••
•••		••••
•••		••••
•••		••••
•••		

(Total for question = 8 marks)

(2)

Q15.

An equation for the formation of ammonia using the Haber process is shown.

$$N_2(g) + 3H_2(g) = 2NH_3(g)$$

Ammonia is stable in air but can be oxidised on the surface of a copper catalyst.

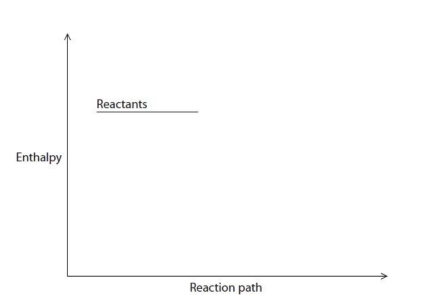
An equation for this reaction is

$$4NH_3(g) + 5O_2(g) \rightarrow 6H_2O(g) + 4NO(g)$$
 $\Delta_r H = -905.2 \text{ kJ mol}^{-1}$

The catalyst is usually warmed to approximately 300 °C to start the reaction, but after a short reaction time the copper catalyst often melts.

(i) Give a reason why the catalyst is warmed and a reason why the catalyst may melt.	
	(2)

(ii) Complete the reaction profile for this catalysed oxidation of ammonia, showing the enthalpy change, $\Delta_r H$.



(2)

iii) Describe the processes that occur on the surface of a heterogeneous catalyst during the xidation of ammonia in air.	е
	3)

(Total for question = 7 marks)

Q16.

The enthalpy change for the decomposition of sodium hydrogencarbonate can be determined indirectly using Hess's Law.

$$2NaHCO_3(s) \xrightarrow{\Delta_1 H} Na_2CO_3(s) + H_2O(l) + CO_2(g)$$

A student carried out two experiments.

Experiment 1 involved the reaction between sodium hydrogencarbonate and hydrochloric acid.

The student used the following procedure:

- use a measuring cylinder to measure 50 cm³ of 2.00 mol dm⁻³ hydrochloric acid and pour it into a polystyrene cup
- measure the initial temperature of the acid
- weigh the test tube containing sodium hydrogencarbonate
- tip the sodium hydrogencarbonate into the hydrochloric acid in the polystyrene cup, stir the mixture and record the lowest temperature reached
- weigh the empty test tube.

Results

Measurement	Value
Mass of test tube + NaHCO ₃ / g	21.23
Mass of empty test tube / g	15.61
Mass of NaHCO₃ used / g	
Initial temperature / °C	21.0
Final temperature / °C	14.4
Temperature fall / °C	

(i) Complete the table.

(1)

(ii) Show, by calculation, that the hydrochloric acid is in excess. You must show your working.

$$NaHCO_3(s) + HCI(aq) \rightarrow NaCI(aq) + H_2O(I) + CO_2(g)$$

(2)

(iii) Calculate the enthalpy change for the reaction between sodium hydrogencarbonate and hydrochloric acid, using the results of the experiment.

Include a sign and units in your answer.

Assume: mass of reaction mixture
$$= 50.0 \, \text{g}$$

specific heat capacity of the reaction mixture $= 4.18 \, \text{J g}^{-1} \, ^{\circ}\text{C}^{-1}$ (3)

(Total for question = 6 marks)

Q17.

Answer the question with a cross in the box you think is correct \boxtimes . If you change your mind about an answer, put a line through the box \boxtimes and then mark your new answer with a cross \boxtimes .

When solid calcium sulfate dihydrate, $CaSO_4 \cdot 2H_2O$, is heated in a crucible, it forms solid calcium sulfate hemihydrate, $CaSO_4 \cdot {}^1\!\!/_2H_2O$.

Which two terms could be used to describe this reaction?

(1)

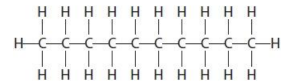
		Enthalpy change	Type of process
	Α	endothermic	hydration
Š	В	exothermic	hydration
×	c	exothermic	dehydration
	D	endothermic	dehydration

(Total for question = 1 mark)

Q18.

Decane, C₁₀H₂₂, is an alkane present in petrol and kerosene.

It has the displayed formula



The enthalpy change of combustion, $\Delta_c H^{\bullet}$, of decane can be estimated using mean bond enthalpy values and the equation shown.

$$C_{10}H_{22}(I) \ + \ 15.5O_2(g) \ \rightarrow \ 10CO_2(g) \ + \ 11H_2O(I)$$

(i) Calculate the enthalpy change of combustion of decane, using the mean bond enthalpy values in the table.

(3)

Bond	Mean bond enthalpy /kJ mol ⁻¹
с—с	347
С—Н	413
0-0	498
C=O	805
0—Н	464

(ii) A data book value for the enthalpy change of combustion of decane is −6 778 k			
	Give two reasons for the difference between your answer to (i) and this value.	(2)	

(Total for question = 5 marks)

Q19.

Methanol, CH₃OH, is a liquid fuel.

Methanol can be synthesised from methane and steam by a process that occurs in two steps.

Step 1
$$CH_4(g) + H_2O(g) \implies 3H_2(g) + CO(g)$$
 $\Delta H = +206 \, \text{kJ} \, \text{mol}^{-1}$
Step 2 $CO(g) + 2H_2(g) \implies CH_3OH(g)$ $\Delta H = -91 \, \text{kJ} \, \text{mol}^{-1}$

Calculate a value for the standard enthalpy change of combustion of gaseous methanol using the enthalpy change for Step **2** and the standard enthalpy change of combustion of gaseous carbon monoxide and of hydrogen.

Substance	Standard enthalpy change of combustion/kJ mol ⁻¹			
СО	-283			
H ₂	-286			

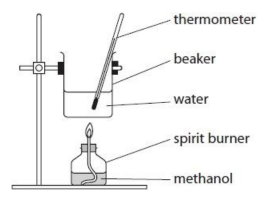
(3)

(Total for question = 3 marks)

Q20.

Methanol, CH₃OH, is a liquid fuel.

An experiment was carried out to determine the enthalpy change of combustion of liquid methanol.



The energy obtained from burning 2.08 g of methanol was used to heat 75.0 g of water.

The temperature of the water rose from 25.0 °C to 91.0 °C.

[Specific heat capacity of water = $4.18 \text{ J g}^{-1} \,^{\circ}\text{C}^{-1}$]

Use the data to calculate a value for the enthalpy change of combustion of one mole of methanol.

Give your answer to an appropriate number of significant figures and include a sign and units.

(4)

(Total for question = 4 marks)

Q21.

Prop-2-en-1-ol is an unsaturated alcohol with the structure shown.

A student planned to use bond enthalpy data to calculate a value for the enthalpy change of combustion of prop-2-en-1-ol.

(i) When researching the bond enthalpy data, the student claimed that it was not necessary to find the value for the C=C bond as they could use the value for a C-C bond and multiply it by two.

Explain why the student is incorrect.

(2)

(ii) Calculate a value for the enthalpy of combustion of prop-2-en-1-ol using the data shown.

$$C_3H_6O(g) + 4O_2(g) \rightarrow 3CO_2(g) + 3H_2O(g)$$

Bond	C–C	C=C	C-0	C=0	О-Н	C-H	0=0
Bond enthalpy / kJ mol ⁻¹	347	612	358	805	464	413	498

(3)

(iii) Explain, in terms of entropy, why the combustion of prop-2-en-1-ol is always feasible in the gaseous state.			
	(2)		

(Total for question = 7 marks)

Q22.

This question is about the oxidation of ammonia.

Nitric acid is made from ammonia. One of the stages in nitric acid production involves the oxidation of ammonia to produce nitrogen(II) oxide, NO. In this process, a mixture of ammonia and oxygen is passed over a platinum-rhodium catalyst. One manufacturer uses a pressure of 5 atm and a temperature of 850 °C.

$$4NH_3(g) + 5O_2(g) \rightarrow 4NO(g) + 6H_2O(g)$$
 $\Delta_t H = -904.8 \text{ kJ mol}^{-1}$

(i) Use this equation, and the enthalpy changes of formation of nitrogen(II) oxide and water, to calculate the enthalpy change of formation of ammonia in kJ mol⁻¹.

You may find it helpful to draw a Hess cycle first. You must show your working. $\Delta_t H(NO(g)) = +90.4 \, kJ \, mol^{-1}$

$$\Delta_f H (H_2O(g)) = -241.8 \text{ kJ mol}^{-1}$$

(3)

(ii) Calculate the atom economy by mass for the formation of NO in this reaction. Give your answer to an appropriate number of significant figures.

(2)

(Total for question = 5 marks)

Q23.

Nitric acid reacts with sodium hydroxide solution in a neutralisation reaction.

$$HNO_3(aq) + NaOH(aq) \rightarrow NaNO_3(aq) + H_2O(l)$$

In an experiment to determine the enthalpy change of neutralisation, the following results were obtained.

Volume of 1.00 mol dm^{-3} HNO₃ = 25.0 cm³

Volume of 1.05 mol dm^{-3} NaOH = 25.0 cm³

Temperature rise = 6.8 °C

Calculate the enthalpy change of neutralisation for the reaction between nitric acid and sodium hydroxide solution, using the results of the experiment.

Give your answer to an appropriate number of significant figures.

```
Assume: density of the reaction mixture = 1.0 \, \text{g cm}^{-3}
specific heat capacity of the reaction mixture = 4.18 \, \text{Jg}^{-1} \, \text{°C}^{-1}
```

(4)

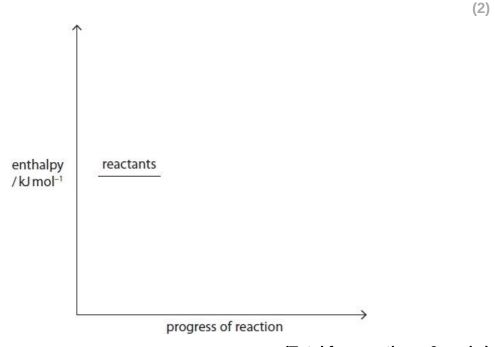
Q24.

This question is about enthalpy changes.

(i) State what is meant by the term 'standard enthalpy change of combustion'.		
		(2
(ii)	Write the equation, including state symbols, for the reaction that occurs when	

the standard enthalpy change of combustion of octane, C₈H₁₈(I), is measured.

(iii) The standard enthalpy change of combustion of octane is -5 470 kJ mol⁻¹. Complete the reaction profile diagram for the combustion of octane. Include labels showing the standard enthalpy change of combustion, $\Delta_c H^\theta$, and the activation energy, E_a .



(Total for question = 6 marks)

(2)

$\overline{}$	2E	
u	ZJ.	

This question is about enthalpy changes and energy changes.	
State what is meant by the standard enthalpy change of formation of aluminium oxide, $Al_2O_3(s)$. Include standard conditions.	
(3	3)
(Total for question = 3 marks	;)

Q26.

This question is about enthalpy changes and energy changes.

Magnesium carbonate reacts with dilute hydrochloric acid at room temperature.

$$MgCO_3(s) \ + \ 2HCI(aq) \ \rightarrow \ MgCI_2(aq) \ + \ CO_2(g) \ + \ H_2O(I)$$

When the reaction is carried out in a sealed container with a constant volume, the heat energy change is not the same as the enthalpy change for this reaction.

Give a reason why this is so.	
	(1)
	(Total for question = 1 mark)

Q27.

This question is about enthalpy changes.

Enthalpy changes of reactions which cannot be measured directly can be calculated using standard enthalpy changes of combustion. The table shows some of these values.

Substance	$\Delta_c H^{\Theta} / \text{kJ mol}^{-1}$
C(s)	-394
H ₂ (g)	-286
CH₄(g)	-890

Complete the Hess cycle and use it to calculate the standard enthalpy change for the following reaction.

 $C(s) + 2H_2(g) \rightarrow CH_4(g)$

(Total for question = 4 marks)

Q28.

Phosphorus(V) chloride, PCl_5 , can be thermally decomposed to phosphorus(III) chloride, PCl_3 , and chlorine, Cl_2 . The equation for this reaction is

$$PCI_5(g) \rightarrow PCI_3(g) + CI_2(g)$$

The enthalpy change for this reaction cannot be measured directly.

(i) Complete the Hess's Law cycle to include the enthalpy change of formation of both phosphorus chlorides.

Include the labels of the missing enthalpy changes.

 $\Delta_v H$ is the enthalpy change for the vaporisation of the substance from the state shown to the gaseous state.

 $\begin{array}{c} \Delta_{r}H \\ \\ PCl_{3}(g) + Cl_{2}(g) \\ \\ \Delta_{v}H\left[PCl_{3}(l)\right] \\ \\ PCl_{3}(l) + Cl_{2}(g) \\ \\ \end{array}$

(3)

(ii) Calculate the enthalpy change for the thermal decomposition of $PCl_5(g)$ to $PCl_3(g)$ and $Cl_2(g)$, using the data given in the table. Include a sign and units in your answer.

(2)

	Enthalpy change / kJ mol
$\Delta_{\rm f}H$ [PCl ₅ (s)]	-443.5
∆ _f H [PCl₃(l)]	-319.7
$\Delta_{v}H$ [PCl ₅ (s)]	+64.9
$\Delta_{v}H$ [PCl ₃ (l)]	+30.5

(Total for question = 5 marks)

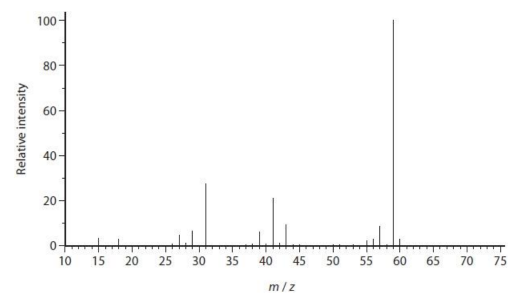
Q29.

This question is about 2-methylpropan-2-ol.

(a) Draw the fully **displayed** formula of 2-methylpropan-2-ol.

(1)

(b) The mass spectrum of 2-methylpropan-2-ol is shown.



(i) The relative molecular mass of 2-methylpropan-2-ol is 74.

Give a possible reason why there is no molecular ion peak in the mass spectrum of 2-methylpropan-2-ol.

.....

(ii) Write the formula for a species that could be responsible for the peak at m/z = 59.

(1)

(1)

(c)	The equation f	or the complete	combustion of	f 2-methylpropan-2-c	ol is

$$C_4H_{10}O(I) + 6O_2(g) \rightarrow 4CO_2(g) + 5H_2O(I)$$

(i) Using the bond enthalpies shown in the table, calculate a value for the enthalpy change, in kJ mol⁻¹, for the complete combustion of 2-methylpropan-2-ol.

(4)

Bond	Mean bond enthalpy / kJ mol ⁻¹
с—с	347
С—Н	413
с—о	358
О—Н	464
0=0	498
C =0	805

(ii) 2-methylpropan-2-ol burns in air with a smoky flame. Explain how burning with a smoky flame affects the value of the experimentally determined enthalpy change of combustion.	
	(2)
(''') A.D. (D. L. L. () (
 (iii) A Data Book value for the enthalpy change of combustion of 2-methylpropan-2-ol is -2643.8 kJ mol⁻¹. Give the main reason for the difference between this value and your answer to part (c)(i). 	;
	(1)

(Total for question = 10 marks)

Q30.

The enthalpy change for the decomposition of sodium hydrogencarbonate can be determined indirectly using Hess's Law.

$$2NaHCO_3(s) \xrightarrow{\Delta_1 H} Na_2CO_3(s) + H_2O(l) + CO_2(g)$$

A student carried out two experiments.

Experiment 2 involved the reaction between sodium carbonate and hydrochloric acid.

The student repeated the procedure for **Experiment 1** but used sodium carbonate instead of sodium hydrogencarbonate and measured the maximum temperature rise.

$$Na_2CO_3(s) + 2HCl(aq) \rightarrow 2NaCl(aq) + H_2O(l) + CO_2(q)$$

The student calculated the enthalpy change for this reaction as −29.4 kJ mol⁻¹.

(i) Complete the Hess cycle with appropriate formulae and labelled arrows.



(ii) Calculate the enthalpy change for the decomposition of sodium hydrogencarbonate. Include a sign and units in your answer.

$$2NaHCO3(s) \xrightarrow{\Delta_r H} Na2CO3(s) + H2O(I) + CO2(g)$$
(3)

(Total for question = 5 marks)

(2)

Q31.

Hess's law can be used to determine enthalpy changes for reactions which cannot be obtained directly.

An example is the reaction of anhydrous copper(II) sulfate with water to form hydrated copper(II) sulfate, $CuSO_4.5H_2O$.

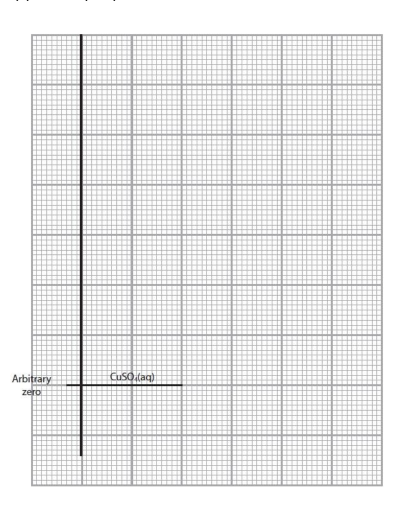
The following outline procedure was carried out.

- Step **1** 42.75 g of deionised water was weighed out in a polystyrene cup and the temperature measured.
- Step **2** 0.0250 mol of hydrated copper(II) sulfate was added to the water in the polystyrene cup with stirring, making a total of 45.00 g of water.
- Step **3** The temperature change was recorded.
- Step **4** Steps **1** to **3** were repeated using 45.00 g of deionised water and 0.0250 mol of anhydrous copper(II) sulfate.

The reaction of anhydrous copper(II) sulfate with water is shown.

$$CuSO_4(s) + aq \rightarrow CuSO_4(aq)$$
 $\Delta H_2 = -84.5 \text{ kJ mol}^{-1}$

(i) Draw to scale, on the graph paper, a labelled enthalpy level diagram which shows the enthalpy changes for the reactions of water with hydrated copper(II) sulfate (ΔH_1) and anhydrous copper(II) sulfate (ΔH_2) .



(3)

				(Total for	question = 4	marks)
	Δ.Η					(1)
	You must show	` '	O(I) → CuSO₄ gram.	5H₂O(s)		(1)
rea	action					

(ii) Use your enthalpy level diagram in (i) to determine the enthalpy change, $\Delta_r H$, for the

Q32.

Hess's law can be used to determine enthalpy changes for reactions which cannot be obtained directly.

An example is the reaction of anhydrous copper(II) sulfate with water to form hydrated copper(II) sulfate, $CuSO_4.5H_2O$.

The following outline procedure was carried out.

- Step **1** 42.75 g of deionised water was weighed out in a polystyrene cup and the temperature measured.
- Step **2** 0.0250 mol of hydrated copper(II) sulfate was added to the water in the polystyrene cup with stirring, making a total of 45.00 g of water.
- Step **3** The temperature change was recorded.
- Step **4** Steps **1** to **3** were repeated using 45.00 g of deionised water and 0.0250 mol of anhydrous copper(II) sulfate.

The reaction of hydrated copper(II) sulfate with water is shown.

CuSO₄.5H₂O(s) + aq
$$\rightarrow$$
 CuSO₄(aq) $\Delta H_1 = +18.2 \text{ kJ mol}^{-1}$

Calculate the temperature change that would have given this enthalpy change for the stated experimental procedure.

Give your answer to a measurable number of significant figures and state whether the temperature increases or decreases.

[Specific heat capacity of the solution = $4.18 \text{ J g}^{-1} \,^{\circ}\text{C}^{-1}$]

(3)

(Total for question = 3 marks)

Q33.

Hess's law can be used to determine enthalpy changes for reactions which cannot be obtained directly.

An example is the reaction of anhydrous copper(II) sulfate with water to form hydrated copper(II) sulfate, $CuSO_4.5H_2O$.

The following outline procedure was carried out.

- Step **1** 42.75 g of deionised water was weighed out in a polystyrene cup and the temperature measured.
- Step **2** 0.0250 mol of hydrated copper(II) sulfate was added to the water in the polystyrene cup with stirring, making a total of 45.00 g of water.
- Step **3** The temperature change was recorded.
- Step **4** Steps **1** to **3** were repeated using 45.00 g of deionised water and 0.0250 mol of anhydrous copper(II) sulfate.

State why the enthalpy change for the reaction of one mole of anhydrous copper(II) sulfate with five moles of water to form hydrated copper(II) sulfate, CuSO₄.5H₂O, cannot be measured directly.

(1

(Total for question = 1 mark)

Q34.

This question concerns iodine monochloride, ICI, a red-brown solid which melts at 27 °C to form a red-brown liquid.

lodine monochloride is used in measuring unsaturation in organic compounds.

lodine monochloride gas can be produced by the reaction between iodine vapour and chlorine gas. The reaction is exothermic.

$$I_2(g) + CI_2(g) \rightarrow 2ICI(g) \Delta_r H = -30 \text{ kJ mol}^{-1}$$

The table shows bond energy values for the bonds in iodine and chlorine.

Calculate the value of the bond energy of the I—Cl bond using these data and the equation.

Bond	Energy/kJ mol ⁻¹
I—I	151
cı—cı	243

(2)

(Total for question = 2 marks)

Q35.

Nitric acid reacts with sodium hydroxide solution in a neutralisation reaction.

$$HNO_3(aq) + NaOH(aq) \rightarrow NaNO_3(aq) + H_2O(l)$$

In an experiment to determine the enthalpy change of neutralisation, the following results were obtained.

Volume of 1.00 mol dm^{-3} HNO₃ = 25.0 cm³

Volume of 1.05 mol dm^{-3} NaOH = 25.0 cm³

Temperature rise = 6.8 °C

Give a reason why excess sodium hydroxide was used.	
	(1)

(Total for question = 1 mark)

Q36.

This question is about enthalpy changes and entropy changes.

Which is the equation for the standard enthalpy change of formation, $\Delta_t H^{\bullet}$, of aluminium oxide?

(1)

- \square **B** 4Al(s) + 6O(g) \rightarrow 2Al₂O₃(s)
- $\label{eq:decomposition} \square \quad \textbf{D} \quad \ 2\text{Al}(s) + 3O(g) \rightarrow \text{Al}_2O_3(s)$

(Total for question = 1 mark)

Q37.

One of the stages in the production of sulfuric acid from sulfide ores involves the oxidation of sulfur dioxide to sulfur trioxide. The equation for the reaction is

$$2SO_2(g) + O_2(g) = 2SO_3(g)$$
 $\Delta rH = -197 \text{ kJ mol}^{-1}$

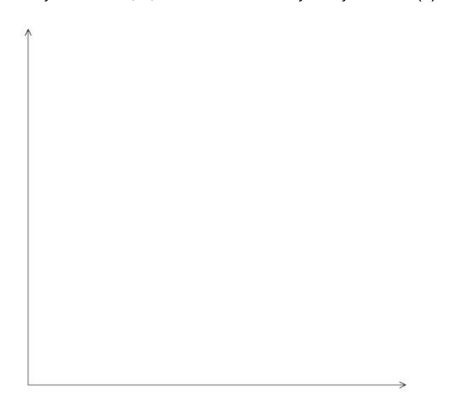
The conditions used in one industrial process are: 420°C and a pressure of 1.7 atm together with a vanadium(V) oxide catalyst.

It is proposed to change the conditions to 600°C and 10 atm pressure, while still using the same catalyst.

(i) On the axes provided, sketch the reaction profiles for the uncatalysed and catalysed reaction.

$$2SO_2(g) + O_2(g) = 2SO_3(g)$$
 $\Delta rH = -197 \text{ kJ mol}^{-1}$

Label the uncatalysed reaction, A, and the reaction catalysed by vanadium(V) oxide, B.



(ii) On your reaction profile, identify and label both the enthalpy change and the activation energy for the catalysed reaction.

(2)

(3)

(Total for question = 5 marks)

Q38.

Answer the question with a cross in the box you think is correct \boxtimes . If you change your mind about an answer, put a line through the box \boxtimes and then mark your new answer with a cross \boxtimes .

The standard molar enthalpy change of neutralisation is the enthalpy change when an acid and an alkali react under standard conditions to form one mole of water.

An experiment was carried out to determine the enthalpy change of neutralisation for the reaction between propanoic acid and sodium hydroxide.

The equation for this reaction is

$$CH_3CH_2COOH(aq) + NaOH(aq) \rightarrow CH_3CH_2COO^-Na^+(aq) + H_2O(I)$$

50.0 cm³ of sodium hydroxide solution, of concentration 1.00 mol dm⁻³, was placed in a polystyrene cup. The initial temperature was measured.

(i) Which piece of equipment has the **smallest** measurement uncertainty for the measurement of 50.0 cm³ of sodium hydroxide solution?

Equipment Measurement uncertainty for each reading 1 A burette $\pm 0.05 \, \text{cm}^3$ 50 cm³ measuring cylinder $\pm 1 \, \text{cm}^3$ 1 B C 25 cm³ pipette ±0.06 cm3 1 $\pm 0.08 \, \text{cm}^3$ 50 cm³ pipette D

(ii) 50.0 cm³ of propanoic acid solution, of concentration 1.00 mol dm⁻³, was added and thoroughly mixed with the sodium hydroxide solution in the polystyrene cup.

The maximum temperature rise was 6.5 °C.

Calculate the enthalpy change of neutralisation for propanoic acid, in kJ mol⁻¹, giving your answer to the **nearest whole number**.

[Assume density of the mixture = 1.00 g cm $^{-3}$, specific heat capacity of the mixture = 4.18 J g $^{-1}$ °C $^{-1}$]

(3)

(1)

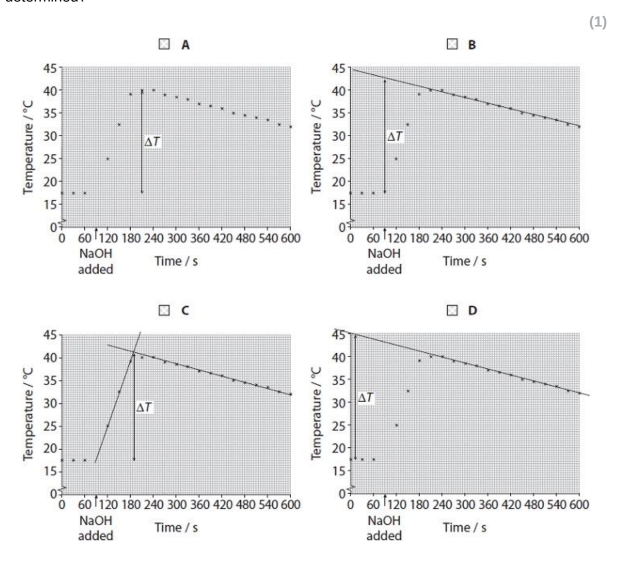
Q39.

Answer the question with a cross in the box you think is correct \boxtimes . If you change your mind about an answer, put a line through the box \boxtimes and then mark your new answer with a cross \boxtimes .

The standard molar enthalpy change of neutralisation is the enthalpy change when an acid and an alkali react under standard conditions to form one mole of water.

An experiment was carried out with a solution of ethanoic acid and sodium hydroxide solution of the same concentration.

(i) Which graph shows the correct way that the maximum temperature rise should be determined?



(ii) Explain why the data book value for the standard enthalpy change of neutralisation of ethanoic acid with sodium hydroxide is –55.2 kJ mol ⁻¹ but the value for hydrochloric acid is –57.1 kJ mol ⁻¹ .	
	(2)
	••••
	••••
	••••
(Total for question = 3 m	arks)



Write the equation to represent the standard enthalpy change of formation of ethanol. Include state symbols.

(2)

(Total for question = 2 marks)

Mark Scheme

Q1.

Question Number	Answer		Mark	
(i)	The only correct answer is C (exothermic so energy is absorbed by the water) A is not correct because the reaction is exothermic not endothermic B is not correct because the reaction is exothermic not endothermic and energy is absorbed not released by the water D is not correct because energy is absorbed not released by the			
Question Number	Answer Additional Guidance			
(ii)	An answer that makes reference to the following point: • (no further release of energy so colder) solution being added cools the reaction mixture OR Added ethanoic acid is at a lower temperature than the reaction mixture	Allow the heat energy is shared over a larger volume Ignore the reaction has stopped so no more energy is released Ignore heat loss	(1)	

Q2.

Question Number	Answer	Additional Guidance	Mark
(i)	An answer that makes reference to the following points: • points (1) plotted correctly • two (1) suitable straight lines of best fit drawn	26 25 24 22 23 21 20 0 10 20 30 40 50 60 70 Do not award dot to dot lines	(2)

Question Number	Answer	Additional Guidance	Mark
(ii)	temperature rise read from graph	26.7 – 20.1 = 6.6 (°C) Allow maximum temperature shown by graph – 20.1 or temperature from line of best fit at 0 cm³ added when these are not the same BUT do not award temperature rises which include subtraction of 20.0 unless the lines of best fit	(1)
		indicate this.	

Question Number	Answer	,,,	Additional Guidance	Mark
(iii)	 gives correct volume added at end-point from the graph (accurate to half a square) 	(1)	Example of calculation = 39 cm ³ Do not award 40 cm ³ unless the lines of best fit indicate this value	(2)
	finds moles of acid added	(1)	= 39 x 1.10 = 1000 0.0429 / 4.29 x 10 ⁻² (mol) Ignore units, even if incorrect Allow TE on first volume given, e.g. Use of 80cm ³ as volume giving 0.088 moles scores (1)	

Question Number	Answer	Additional Guidance	Mark
(iv)		Example of calculation	(3)
	• use of energy change = m x c x ΔT (1)	(30 + 39) x 4.18 x 6.6 = 1 903.6 / 1.9036 x 10 ³ (J)	
	• calculation of energy change per mole (1)	= <u>1 903.6</u> = 44 372 (J mol ⁻¹) 0.0429	
	• final answer with correct sign and units (1)	– 44 372 J mol ⁻¹ / – 44 400 J mol ⁻¹ / – 44.372 kJ mol ⁻¹ / – 44.4 kJ mol ⁻¹	
		Allow TE throughout from the graph in (a)(i) and calculations in (a)(ii) and (a)(iii) Ignore SF except 1 SF	

Q3.

Question Number	Acceptable Answers		Additional Guidance	Mark
(a)	• 1.60	(1)	Do not award MP1 for "1.6" (must be to 2 D.P.)	(2)
	• (+) 42.5	(1)	Do not award MP2 for "42.50" (must be to 1 D.P.)	
			Penalise D.P. error once only	

Question Number	Acceptable Answers	Additional Guidance	Mark
(b)	$CH_3OH(I) + 1.5O_2(g) \rightarrow CO_2(g) + 2H_2O(I)$		(2)
	Balanced equation (1)	Do not award multiples (enthalpy change is for the complete combustion of one mole) for MP1	
	State symbols all correct (1)	MP2 depends on the award of MP1 or correct species	

Question Number	Acceptable Answers	Additional Guidance	Mark
(c)	Calculation of energy change (1)	Example of calculation $(= mc\Delta T = 150 \times 4.18 \times 42.5 =)$ 26647.5 (J)	(4)
	Calculation of moles of CH₃OH	Moles CH ₃ OH = 1.60/32 (= 0.05(00))	
	(1)		
	Calculation of energy ÷ moles CH₃OH	26647.5 = 532950 (J mol ⁻¹) 0.05(00) Ignore any signs at this stage	
	ΔH final answer in kJ mol ⁻¹ and negative sign included and ΔH final answer to 2 or 3 S.F.	-533 (kJ mol ⁻¹) Or -530 (kJ mol ⁻¹)	
	(1)	Correct answer with no working gains full marks Penalise incorrect units for MP4 only Allow TE at each stage Allow correct rounding to 2SF or more at each stage	

Question Number	Acceptable Answers	Additional Guidance	Mark
(d)(i)	(±)0.7 (%)	Allow from 1 SF up to calculator value correctly rounded where (% uncertainty =) (\pm) \pm x 100 = 0.666667 (%) 150	(1)
		Allow 0.6 or $\frac{2}{3}$ Do not award 0.66/0.6	

Question Number	Acceptable Answers	Additional Guidance	Mark
(d)(ii)	An answer that makes reference to the following points:	Needs to show combined error in using the 25 cm³ six times is greater than using 250 cm³ measuring cylinder once only	(3)
	Calculation of the % uncertainty using the 25 cm³ measuring cylinder (1)	Award MP1 EITHER if multiplies errors: 100 x (0.2 /25) x 6 = 4.8% OR If adds errors 100 x (1.2 /150) = 0.8% Do not award (0.2 / 25) x 100 = 0.8 %	
	Then any two from:		
	% uncertainty with use of 25 cm³ measuring cylinder is greater (1)		
	Repeated use of the small measuring cylinder will lead to greater transfer losses (1)	r	
	Repeated use will take more time (1)	Do not award 'easier' to use larger measuring cylinder	

Question Number	Acceptable Answers	Additional Guidance	Mark
(d)(iii)	An answer that makes reference to any three of the following points: • heat/energy loss (to the surroundings) (1)	Ignore experiment carried out under non-standard conditions Ignore just 'no lid'	(3)
	evaporation of methanol / water from the calorimeter (1)		
	incomplete combustion (of methanol) (1)		
	(specific) heat capacity of the calorimeter/apparatus has been ignored (1)	Allow calorimeter has not been calibrated	

Question Number	Acceptable Answers	Additional Guidance	Mark	
(e)	An explanation that makes reference to the following points:		(2)	
	The second value will be less exothermic / less negative (1)	Allow 'more positive' or 'smaller in magnitude' Do not accept 'greater' or 'smaller' for 'less negative'		
	Some energy will be used to boil the water/boiling water is endothermic Water can only be heated to 100°C/ Temperature rise (measured) can only be (a maximum) of 40°C	Do not award just "the water boils"		
	Greater heat losses in the 60°C to 100°C range (1)	Mark points M1 and M2 independently		

Question Number	Acceptable Answers	Additional Guidance	Mark
(f)	An explanation that makes reference to the following points: Either • student 2's value will be similar / the same (1) • (As) both the energy change and moles/mass (of methanol) burned will be higher/ Ratio of energy change to moles/mass (of methanol) burned will be the same/ The energy change is proportional to the moles/mass (of methanol) burned (1)	Allow 'temperature change' for 'energy change'	(2)
	Or • student 2's value will be less negative/ less exothermic (1)	Allow 'more positive' or 'smaller in magnitude' or 'smaller' for 'less negative'	
	greater heat loss because higher temperature/heated for longer (1)	Mark points MP1 and MP2 independently within each route	

Question Number	Acceptable Answers	Additional Guidance	Mark
(g)	An answer that makes reference to the following points:		(2)
	(Calculated) value of moles (of methanol) burned will be less / too small (1)	Allow both marks for a calculation using M_r of 46.0 (instead of 32.0), giving a final ΔH value (approx.) of -766 (kJ mol ⁻¹)	
	The calculated value will be more exothermic / more negative (1)	Allow 'increase' or 'greater' for 'more negative' MP2 depends on MP1	

Q4.

Question Number	Acceptable Answer	Additional Guidance	Mark
(i)	calculation of mean C-H bond enthalpy	Example of calculation -1652 = -413 (kJ mol ⁻¹) 4 Therefore bond enthalpy is (+)413 (kJ mol ⁻¹)	(1)
		Correct answer with no working scores 1	

Question Number	Acceptable Answer	Additional Guidance	Mark
(ii)	 calculation of energy released when 8(C-H) bonds are formed in the formation of C₃H₈ (1) 	Example of calculation 8 x -413 = -3304 (kJ mol ⁻¹)	(2)
	calculation of mean C-C bond enthalpy (1)	-39983304 = -694 = -347 (kJ mol ⁻¹)	
		Therefore bond enthalpy is +347 (kJ mol ⁻¹)	
		Allow -347 (kJ mol ⁻¹) if -413 given as answer in (i) for 2 marks	
		Allow TE from (c)(i)	

Q5.

Question Number	Answer	Additional Guidance	Mark
	sum of bonds broken (1) and sum of bonds made (1) answer and with negative sign (1)	Example of calculation bonds broken = (6 x 198) + (10 x 243) = 3618 (kJ mol ⁻¹) bonds made = (20 x 326) = (-)6520 (kJ mol ⁻¹) enthalpy change = Bonds broken – bonds made = (3618 – 6520) = -2902 (kJ mol ⁻¹) Correct answer with no working	(3)
		TE on bonds broken and made	

Question Number	Answer	Additional Guidance	Mark
(ii)	bond breaking requires energy or by convention bond enthalpies refer to dissociation and so are endothermic	ALLOW bond breaking is endothermic ALLOW bond making is exothermic Ignore just 'bonds are broken' / 'it is endothermic'	(1)

Q6.

Question Number	Answer	Additional Guidance	Mark
(i)	An answer that makes reference to the following points:	Allow corrections to be made on the diagram	(2)
	identification and correction of the first error (1)	Error 1 – arrow for enthalpy change of formation should go down/be reversed	
	identification and correction of the second error (1)	Error 2 – the word 'half' should be deleted from the enthalpy change of atomisation of hydrogen	

Question Number	Answer	Additional Guidance	Mark
(ii)		Example of calculation	(1)
	calculation of first electron affinity of hydrogen	1st EA= -(218+496+107)-56 +804 = -73 (kJ mol ⁻¹)	
		Allow a TE 1st EA = +39(kJ mol-1) if the first arrow reversed direction is not identified	

Q7.

Question Number	Answer	Additional Guidance	Mark
(i)	correct species with state symbols in bottom box (1)	Example of Hess cycle $2NO(g) + Cl_2(g) \rightarrow 2NOCl(g)$ $N_2(g) + O_2(g) + Cl_2(g)$	(2)
	arrows in correct direction (1)		

Question Number	Answer	Additional Guidance	Mark
(ii)	 multiplies enthalpy change of formation of NO, Δ_fH^e₂₉₈(NO) by 2 or divides Δ_rH^e₂₉₈ by 2 (1) 	Example of calculation (2 x + 90.3) = 180.6 / 181 (kJ) or - <u>75.6</u> = -37.8 (kJ) 2	(2)
	calculates enthalpy of formation of NOCI (1)	$2\Delta_{\rm f} H_{298}^{\rm e}$ NOCI = $\underline{180.6\text{-}75.6}$ = 52.5 (kJ mol ⁻¹) 2 or $\Delta_{\rm f} H_{298}^{\rm e}$ NOCI = $90.3-37.8$ = 52.5 (kJ mol ⁻¹) Unit, if given, must be correct. Correct answer with no working scores (2) -52.5 (kJ mol ⁻¹) scores (1) 14.7 (kJ mol ⁻¹) scores (1) $+7.35$ (kJ mol ⁻¹) scores (1) -14.7 (kJ mol ⁻¹) scores (0) Ignore presence of absence of 298 Ignore SF except 1 SF M2 no TE other than the answers above No TE on an incorrect cycle	

Q8.

Question Number	Answer	Additional Guidance	Mark
(i)	 sum of bond energies of all reactants sum of bond energies of all products calculation of Δ_cH	Example of calculation 945 + (3 × 436) = (+)2253 (kJ mol ⁻¹) 6(N-H) = 6 × 391 = (-)2346 (kJ mol ⁻¹) -2346 + 2253 = -93 (kJ mol ⁻¹) TE from either/both M1 and M2 Correct answer with no working scores 3	(3)

Question Number	Answer	Additional Guidance	Mark
(ii)	An answer that makes reference to the following points: the equation in 9(a)(i) is for the formation of two moles of ammonia (1) the bond energies in the table are mean / not specific to ammonia (1)	Ignore any references to differing conditions for the Haber process Ignore heat losses	(2)

Question Number	Answer	Mark
(iii)	The only correct answer is D (100 %)	(1)
	A is not correct because this is the percentage of hydrogen	
	B is not correct because this is half the atom economy for making ammonia	
	C is not correct because this is the percentage of nitrogen	

Question Number	Answer	Mark
(iv)	The only correct answer is C $K_c = \frac{[NH_3]^2}{[N_2][H_2]^3}$ A is not correct because this expression shows molar quantities, not powers and is inverted B is not correct because this expression shows molar quantities, not powers D is not correct because this expression is for the reverse equation	(1)

Q9.

Question Number	Answer	Additional Guidance	Mark
Number (i)	calculation of moles of CaSO ₄ ·½H ₂ O (1) calculation of volume (or mass) of water required (1)	Example of calculation 10.00 g CaSO ₄ ·½H ₂ O = 10.00 ÷ 145.2 mol = 0.06887 mol Allow 0.069 (moles of water required = 0.06887 x 1.5 = 0.1033 mol) volume of water required = 0.1033 x 18 ÷ 1.00 = 1.86 cm ³ Allow 1.86 g Ignore SF except 1 SF Correct answer with no working scores (2) Allow calculation using multiples of these moles (still gets same final answer scores 2) Allow alternative correct calculations: e.g. comparison of moles of	(2)
		CaSO ₄ ·½H ₂ O with moles of water in 10.00 g.	

Question Number	Answer	Additional Guidance	Mark
(ii)	• calculation of ΔT (1) • use of $\operatorname{mc}\Delta T$ to find Q (1)	Example of calculation $\Delta T = 2.8$ °C m = 10.00 g, c = 4.18 J g ⁻¹ °C ⁻¹ Q = mc ΔT = 117.04 J / 0.11704 kJ Allow M1 and M2 if figure of 117.04 J is seen Ignore units unless converted to kJ	(4)
	 calculation of Δ_rH (1) correct final answer, with sign and 2 or 3 SF (1) 	$117.04 \div 0.06887 = -1699.4 \text{ (J mol}^{-1}\text{)}$ $-1.70 / -1.7 \text{ (kJ mol}^{-1}\text{)}$ Correct answer with no working scores (4) Allow TE throughout and from 4ci (for moles CaSO ₄ -½H ₂ O)	

Question Number	Answer	Additional Guidance	Mark
(iii)	selection of thermometer (1) calculation of percentage uncertainty (1)	Example of calculation 2 x 0.5 x 100 = 35.7 / 36 / 40 (%) 2.8 Allow selection of measuring cylinder and percentage uncertainty is 5%, scores (1) mark Do not award selection of balance Ignore SF	(2)

Q10.

Question Number	Answer	Additional Guidance	Mark
	calculation of energy needed to break bonds (1)	Example of calculation Energy to break bonds: (C-C) + (C-H) + (C-O) = 347 + 413 + (C-O) = (C-O) + 760 (kJ)	(3)
	calculation of energy released when bonds are made (1)	Energy released in forming bonds: (C=C) + (O—H) = 612 + 464 = (-)1076 (kJ)	
	calculation of mean bond enthalpy of C—O (1)	(C-O) + 760 - 1076 = 42 $(C-O) = (+)358 (kJ mol^{-1})$ TE on M1 and M2	
		If all bonds broken: Energy to break bonds = (C-O) + 4049 (kJ) Energy released in forming bonds = (-)4365 (kJ)	
		Ignore units	
		Correct answer with no working scores (3)	
		Allow correct working in M1 and M2 if answers not evaluated	

Q11.

Question Number	Acceptable Answer	Additional Guidance	Mark
(i)		example of calculation use of $Q = m c \Delta T$	(3)
	calculation of Q (1)	Q = 250 x 4.18 x 8.2 = 8569 (J) / 8.569 kJ ignore any sign at this stage	
	• mass of hydrocarbon burnt and value of $\Delta_c H$	= 112.990 - 112.732 = 0.258 g	
		$\Delta_c H = (-) 8569 \times 84/0.258$ = (-) 2789907 (J mol ⁻¹) /(-) 2789.907 (kJ mol ⁻¹)	
	sign and significant figures (1)	TE on incorrect value from M1	
		= -2790/-2800 (kJ mol ⁻¹) allow -2790000/-2800000 J mol ⁻¹ final answer to 2 or 3 sig figs only	
		Do not award M3 for incorrect method used in M2	
		correct final answer without working scores 3	

Question Number	Acceptable Answer	Additional Guidance	Mark
(ii)	an answer that makes reference to the following point: improved/better (thermal/heat) conduction	Allow copper is a good conductor (of heat) Allow reverse argument in terms of (thermal) insulators Ignore references to heat capacity/ heat lost to surroundings/ heat absorbed by container. Ignore any mention of glass breakage	(1)

Q12.

Question Number	Answer	Additional Guidance	Mark
	calculation of moles used (1)	Example of calculation moles used = 25.0 x 1.00/1000 = 0.0250	(3)
	calculation of energy for that number of moles (1)	energy released = 0.025 x 53.4 = 1.335 (kJ) / 1335 (J) TE on moles used Ignore sign	
	calculation of temperature change and gives answer to 2 SF (because a school	temperature change = 1335/(50.0 x 4.18) = 6.3876 = 6.4 (°C / K) TE on moles and energy	
	thermometer cannot measure to 3 SF) (1)	Allow final answer to 3 SF 6.39 (°C / K) Ignore units	
		Correct answer with no working scores (3)	

Q13.

Acceptable Answers	Additional Guidance	Mark
calculation or working of energy needed to break bonds (1) calculation or working of energy released when bonds made (1) calculation of enthalpy change of combustion with sign (1)	Example of calculation energy to break bonds = 347 + (5 x 413) + 358 + 464 + (3 x 498) = 4728 (kJ) energy released in making bonds = (4 x 805) + (6 x 464) = 6004 (kJ) enthalpy change of combustion = 4728 - 6004 = -1276 (kJ mol ⁻¹) or energy to break bonds = 347 + (5 x 413) + 358 + (3 x 498) = 4264 (kJ) energy released in making bonds = (4 x 805) + (5 x 464) = 5540 (kJ) enthalpy change of combustion = 4464 - 5540 = -1276 (kJ mol ⁻¹) TE on energies calculated to break and form bonds Correct answer with sign but no working scores 3 Ignore SF except 1SF Ignore missing units but do not allow incorrect units in M3 e.g. kJ mol ⁻	(3)
	calculation or working of energy needed to break bonds (1) calculation or working of energy released when bonds made (1) calculation of enthalpy change of combustion with sign	 calculation or working of energy needed to break bonds

Question Number	Acceptable Answers	Additional Guidance	Mark
(ii)	Enthalpy $C_2H_3OH(I) + 3O_2(g)$ Δ_cH $2CO_2(q) + 3H_2O(I)$	M1 is conditional on exothermic or endothermic value calculated in (c)(i) but if no value is calculated, award mark for exothermic reaction only	(2)
	Reaction pathway	Allow double headed arrows / lines, but penalise arrows pointing in wrong direction once only	
		Allow 'products'/ unbalanced formulae / missing state symbols as labels for product line	
	 products to the right of reactants <u>and</u> at a lower enthalpy <u>and</u> arrow labelled Δ_cH (1) 	$(-)\Delta H/(-)\Delta H_c$ /enthalpy	
	 curve and arrow labelled E_a (1 	Allow value calculated for energy needed to break bonds in (c)(i)	
S		Ignore any transition state	
0 8		Do not allow straight lines instead of Ea curve	
		If no other marks awarded, allow 1 mark for the correct labelled product line and activation energy curve if both arrows missing	

Question Number	Acceptable Answers	Additional Guidance	Mark
(iii)	standard enthalpy change of combustion refers to ethanol / water as liquid(s) but bond energies are calculated for gases or change of state data is not included	Ignore bond energies are mean values and the actual values in these compounds/ethanol may be different Ignore any reference to heat loss	(1)
	or ethanol / water are not in standard states for bond enthalpy calculation	Ignore any reference to incomplete combustion	

Q14.

Question Number	Acceptable Answers	Additional Guidance	Mark
(a)	calculation or working of heat evolved during reaction (1)	Example of calculation heat evolved = 50 x 4.18 x 5.4 = 1128.6 J or 1.1286 kJ Ignore any sign	(4)
	 calculation or working of mol Na₂CO₃ used (1) 	mol Na ₂ CO ₃ used = 5.09/106 = 0.04802	
	calculation of enthalpy change of solution (1)	enthalpy of solution = 1.1286/0.04802 = 23.5 TE on heat evolved and mol Na ₂ CO ₃	
	negative sign and answer to 2 or 3 SF (1)	-23.5/-24 (kJ mol ⁻¹) TE on enthalpy change in M3 Correct answer with – sign but no	
	(1)	working scores 4 Ignore missing units but penalise incorrect units once only in (a) or (b)	

Question Number	Acceptable Answers	Additional Guidance	Mark
(b)	both arrows in correct direction and Na ₂ CO ₃ (aq) (+ 10H ₂ O(I)) / 2Na ⁺ (aq) + CO ₃ ²⁻ (aq) (+ 10H ₂ O(I)) (1)	Na ₂ CO ₃ (s) + 10H ₂ O(l) → Na ₂ CO ₃ .10H ₂ O(s) (aq) Na ₂ CO ₃ (aq) Allow aq omitted from arrows Allow both arrows pointing upwards provided labelled as opposite signs	(2)
	answer to (a) – 53.7 with correct sign (1)	Example of calculation -23.5 - 53.7 = -77.2 (kJ mol ⁻¹) TE on answers to (a) but not on incorrect cycle Allow -77200 J mol ⁻¹ Ignore SF except 1SF Ignore missing units but penalise incorrect units	

Question Number	Acceptable Answers	Additional Guidance	Mark
(c)	An explanation that makes reference to the following points: • enthalpy change of solution will be lower/ less endothermic / less positive (than data book value) • because anhydrous sodium carbonate releases energy/reacts exothermically	Allow smaller / requires less energy Allow more exothermic / negative Conditional on M1	(2)
	with water or because less energy is needed to separate the (fewer) water molecules from the ions (in the crystal structure) (1)	Allow because there is (less water so) more Na ₂ CO ₃ (in the sample) Allow because less energy is needed to break the bonds between water and sodium carbonate	

Q15.

Question Number	Answer	Additional Guidance	Mark
(i)	An answer that makes reference to the following points: • provide / overcome the activation energy or (is slow at room temperature but) accelerates as temperature rises (1) • (sufficiently / very) exothermic enough to melt the copper / break bonds in copper (1)	Do not allow 'to lower the activation energy' Allow answers that link rise in temperature to rising rate	(2)

Question Number	Answer	Additional Guidance	Mark
(ii)	Intermediate Reactants A _r H Products	Allow transition state for intermediate	(2)
	Reaction path $ \begin{tabular}{ll} \hline & intermediate energy level/transition state \\ \hline & (1) \\ \end{tabular} $ $ \begin{tabular}{ll} \hline & & \\ $	Ignore type of arrows to and from intermediate Allow any diagram with a hump shown,	
	shown on down/ vertical arrow (1)	with / without intermediate / transition state label Do not penalise missing 'Products' label Allow use of $\Delta_r H/$ - 905.2 (kJ mol ⁻¹)	

Question Number	Answer	Additional Guidance	Mark
(iii)	An answer that makes reference to any three of the following points:		(3)
	100 Maria 100 Ma	Do not allow absorb	
	reactants adsorb onto catalyst/surface (1)		
	(there are) active sites on catalyst (surface) (1)		
	bonds in reactants weakened / broken		
	or		
	reaction takes place		
	(1)		
	products desorb from the catalyst/active		
<u>.</u>	site (1)		

Q16.

Question Number	Answer	Answer Additional Guidance	
(i)	mass and temperature fall correct	Mass of NaHCO₃ 5.62 used / g	(1)
	Programme 2017	Temperature fall / (-)6.6 °C	

Question Number	2	Answer		Additional Guidance	Mark
(ii)	•	calculation of amount of NaHCO ₃ and calculation of amount of hydrochloric acid	(1)	Example of calculation amount NaHCO ₃ = 5.62	(2)
	•	0.0669 mol NaHCO3 needs 0.0699 mol HC1 for reaction so HCl is in excess	(1)	23 + 1 + 12 + (3 x 16) = 0.0669 (mol) TE on mass of NaHCO ₃ in (a)(i) and amount HC1 = 50 x 2.00 = 0.10 (mol) 1000 Ignore SF including 1SF	
				Allow mol ratio = 1: 1 so HCl is in excess Allow just more moles of HCl used Allow 0.10 > 0.0669 (mol) Allow HCl is in excess by 0.033 (mol)	

Question Number				Additional Guidance	Mark
(iii)				Example of calculation	(3)
	•	calculation of heat absorbed	(1)	heat absorbed = 50.0 x 4.18 x 6.6 = 1379.4 (J) / 1.3794 (kJ)	
				Ignore sign	
	•	calculation of enthalpy	(1)		
		change	200	enthalpy change = <u>1379.4</u> 0.0669	
				$= 20619 (J \text{ mol}^{-1})$	
				or = 1.3794 0.0669	
				$= 20.619 \text{ (kJ mol}^{-1}\text{)}$	
				TE on heat absorbed and amount	
			(1)	NaHCO ₃ in (a)(ii)	
	•	positive sign and units			
	3300	P		Final answer +20.6(19) kJ mol ⁻¹ or +20619 J mol ⁻¹	
				TE on enthalpy change	
				Allow +19.7(06) kJ mol ⁻¹ from 0.07 mol in (a)(ii)	
				Allow kJ mol ⁻ / J mol ⁻	
				Ignore SF except 1 SF	
				Ignore incorrect / missing units in M1 and M2	
				Correct answer with sign and units scores (3)	

Q17.

Question Number	Answer	
	The only correct answer is D (endothermic, dehydration)	(1)
	A is not correct because hydration involves adding water	
	B is not correct because a reaction that requires heat is unlikely to be exothermic and hydration involves adding water	
9	C is not correct because a reaction that requires heat is unlikely to be exothermic	/4

Q18.

Question Number	Answer	Additional Guidance	Mark
(i)	calculation of energy associated with bond breaking (1)	Example of calculation = (22 x 413) + (9 x 347) + (15.5 x 498) = 19928 (kJ) (ans 1)	(3)
	calculation of energy associated with bond formation (1)	= (20 x 805) + (22 x 464) = 26308 (kJ) (ans 2) Ignore minus sign	
	calculation of the enthalpy change of combustion by subtraction and a negative sign (1)	= (ans 1) - (ans 2) = 19928 - 26308 = -6380 (kJ mol ⁻¹) Ignore units even if incorrect Allow TE throughout but for M3 do not award positive values Ignore SF except 1 SF Correct answer with no working scores (3) (+)6380 (kJ mol ⁻¹) with no working scores (2)	

Question Number	Answer	-00	Additional Guidance	Mark
(ii)	An answer that makes reference to the following points:			(2)
	 use of mean bond enthalpy values rather than actual values for the molecules involved 	(1)	Ignore just 'mean bond enthalpies are not accurate' without qualification	
	 substances in the wrong state for bond energy calculations 	(1)	Allow water / decane is a liquid / not a gas	

Q19.

Question Number	Answer	Additional Guidance Mark
	An answer that makes reference to the following points: • gives an equation linking the three values or processes together / constructs a Hess's Law cycle	Example of calculation $ \Delta_{c}H (CH_{3}OH) = -\Delta H (Step 2) + \Delta_{c}H(CO) + 2\Delta_{c}H(H_{2}) $ or $ CO(g) + 2H_{2}(g) \stackrel{-91}{\rightleftharpoons} CH_{3}OH(g) $ $ (+1.50_{2}) -283 \stackrel{-286}{\times} 2 $ $ CO_{2} + 2H_{2}O $
	 uses of numerical values in equation or on cycle, including use of 2 x Δ_cH(H₂) 	Do not penalise lack of 2 in $2H_2O$ in cycle or in $2\Delta_cH(H_2)$ if M2 not scored. Δ_cH (CH ₃ OH) = 91 + -283 + 2(-286)
	calculation of final value with correct sign	(1) = -764 (kJ mol ⁻¹) Correct answer with no working scores (3)
		Possible incorrect answers include: Award 2 marks for -478, - 1424, (+)946, -855, (+)764 Award 1 mark for -946, (+)478, -946, (+)1424

Q20.

Question Number	Answer		Additional Guidance	Mark
	An answer that makes reference to the following points: • calculation of the energy absorbed by water	(1)	Example of calculation $Q = m \times c \times \Delta T$ $= 75.0 \times 4.18 \times 66.0$ $= 20691 (J)$	(4)
	calculation of the number of moles of methanol	(1)	$= \frac{2.08}{32.0} = \frac{0.0650 / 0.065 /}{6.50 \times 10^{-2} \text{ (mol)}}$	
	 calculation of the energy absorbed per mole of methanol 	(1)	= 20 691 = 318323 (J mol ⁻¹)	
	 gives enthalpy change of combustion to 2 or 3 SF and correct sign and units (either J 	(1)	= -320 / -318 kJ mol ⁻¹ -320 000 / -318 000 J mol ⁻¹	
	mol ⁻¹ or kJ mol ⁻¹)		Do not award J/mol ⁻¹ Ignore sign until final answer when must be negative	
			Ignore significant figures until final answer	
			Allow TE throughout Correct answer with units and no working scores (4)	

Q21.

Question Number	Answer	Additional guidance	Mark
(i)	An explanation that makes reference to the following points • C=C bond is weaker than 2 x C-C bond (1) • as it consists of a pi and a sigma bond (rather than 2 sigma bonds) (1)	Ignore pi bond formed by sideways / less effective orbital overlap	(2)

Question number	Answer	Additional guidance	Mark
(ii)	calculation of energy required to break reactant bonds (1) calculation of energy release when product bonds form (1) calculation of enthalpy change (1)	Example of calculation: 5(C-H) + (C=C)+(C-C)+(C-O)+(O-H)+4(O=O) 5(413) + (612) + (347) + (358) + (464) + (4x498) = 5838 (kJ mol ⁻¹) 6(C=O) + 6(O-H) (6x805) + (6x464) = 7614 (kJ mol ⁻¹) 5838 - 7614 = -1776 (kJ mol ⁻¹) Ignore SF except 1 SF Allow TE from M1 and M2 Correct answer no working scores 3	(3)

Question Number	Answer	Additional Guidance	Mark
(iii)	An explanation that makes reference to one of the following points EITHER ΔS _{total} is always positive (1) (1) As both ΔS _{surroundings} and ΔS _{system} are positive (1) OR ΔG is always negative (1) as ΔH is negative and ΔS _(system) is positive (1)	If no marking points awarded allow 1 mark for idea that ΔS _{system} / ΔS _{surroundings} / entropy increases with correct explanation	(2)

Q22.

Question Number	Acceptable Answer	Additional Guidance	Mark
Number (i)	 calculates ΣΔ_fH(products) (1) ΣΔ_fH(products) - Δ_rH (1) calculates Δ_fH_(NH3) for 1 mol ammonia 	Example of calculation (+90.4 x 4) + (-241.8 x 6) = -1089.2 -1089.2 - (-904.8) = -184.4 -184.4/4 = -46.1 (kJ mol ⁻¹) TE from M1 to M2 M3 can be awarded for an	Mark
	(1)	incorrect answer to M2 divided by 4 correct answer with no working scores 3 marks	(3)

Question Number		Answer Acceptable	Additional Guidance	Mark
(ii)	•	correct expression (1)	Example of calculation 4NO 4NO + 6H₂O	
			OR 4NO 4NH ₃ + 5O ₂	
			may be shown as numbers only	
	•	correct evaluation of atom economy (1)	$\frac{4(14+16)}{4(14+16)+6(16+2)} \times 100$ OR	
			$\frac{4(14+16)}{4(14+3)+5(16\times2)} \times 100$	
			= 53/52.6(316)(%) allow answer to 2 or 3 SF only correct answer with no working scores 2 marks 0.53/0.526 scores M1 only	(2)

Q23.

Question Number	Acceptable Answer	Additional Guidance	Mark
	calculation of heat produced (1)	Example of calculation heat produced = 50.0 x 4.18 x 6.8 = 1421.2(J) / 1.4212 (kJ)	(4)
	• calculation of amount (mol) of HNO ₃ (1)	amount HNO ₃ used = $25.0 \times 1.00/1000$ = $0.025 / 2.5 \times 10^{-2}$ (mol)	
		Ignore moles NaOH and total	
		moles calculated	
	calculation of enthalpy change (1)	enthalpy change = $\frac{1421.2}{0.025}$ = 56848 (J mol ⁻¹)	
		or = 1.4212 = 56.848 0.025	
		(kJ mol ⁻¹)	
		TE on heat produced and amount HNO ₃	
	negative sign and units and answer to 2 / 1 SF	final answer -57 / -60 kJ mol ⁻¹ or -57 000 / -60 000 J mol ⁻¹ TE on enthalpy change Do not award 3 SF	
	(1)	Correct final answer with sign, units and 2 or 1 SF but no working scores (4)	
		Ignore units and sign of enthalpy change in M1 and M3	

Q24.

Question Number	Acceptable Answer	Additional Guidance	Mark
(i)	(standard enthalpy change of combustion is the enthalpy change when) one mole of a substance burns completely (in oxygen) / burns in excess oxygen / fully combusts	Accept energy released Ignore air Do not award one mole of atoms burns Do not award energy required / needed	(2)
	under standard conditions of 100 kPa and a stated temperature (1)	e.g 25°C / 298 K / 273 K / 293 K Allow 101 kPa / 1 atm Do not award just 'under standard conditions' / rtp	

Question Number	Acceptable Answer	Additional Guidance	Mark
(ii)	C ₈ H ₁₈ (I) + 12½O ₂ (g) → 8CO ₂ (g) + 9H ₂ O(I) • correct species (1) • balancing and state symbols (1)	Allow multiples only if one mole is not stated in (a)(i)	(2)

Question Number	Acceptable Answer	Additional Guidance	Mark
(iii)	enthalpy / kJ mol ⁻¹ reactants reactants (products) progress of reaction	¥	(2)
	Line rising to a maximum then falling to products lower than reactants (1)	Do not award double headed arrows	
	Labelled arrows for E_a and $\Delta_c H^a$ /-5 470 which touch or almost touch the maximum and be level or almost level with the product and reactant lines (1)	Do not award -∆ _c H* Do not award lines with no arrow heads Allow TE on an endothermic diagram	

Q25.

Question Number	Answer	Additional Guidance	Mark
	An answer that makes reference to the following points: • (the enthalpy/energy change when) 1 mol of aluminium oxide (1) • is formed from its elements in their standard states (1)	Allow $ 2\text{Al(s)} + 1\%\text{O}_2(g) \rightarrow \text{Al}_2\text{O}_3(s) \text{ for M1 and M2} $ If state symbols are missing or incorrect only M1 can be awarded $ \text{Allow M2 for multiples in equation provided state } $ symbols for the elements are correct	(3)
	at 100 kPa and a 'specified' / 'stated' temperature (1)	Allow 1 atm / 1 x 10 ⁵ Pa / 101 kPa / 1.01 x 10 ⁵ Pa for pressure Allow a value for the temperature of 298K / 25 ^o C Ignore 273K	
		Ignore other standard conditions e.g. 1 mol dm ⁻³	
		Do not allow °K	

Q26.

Question Number	Answer	Additional Guidance	Mark
	An answer that makes reference to the following point:		(1)
	(the system / it) is not at constant pressure or	Allow a gas / carbon dioxide is produced and this increases the pressure	
	enthalpy change is the heat change at a constant pressure	Allow the pressure is increased / increases Ignore reference to temperature	

Q27.

Question Number	Acceptable Answer	Additional Guidance	Mark
	• species and balanced (1)	Ignore state symbols even if incorrect Ignore absence of oxygen alongside arrows	(4)
	arrows pointing downwards		
	(1)	Example of calculation	
	• calculation of $\Delta_c H$ of reactants and show $\Delta_c H$ of product (1)	$\Delta_c H_{reactants} = -394 + (2 \text{ x} - 286) \text{ (kJ mol}^{-1})$ = -966 (kJ mol}^-1) $\Delta_c H_{products} = -890 \text{ (kJ mol}^{-1})$	
	• calculation of $\Delta_r H$ (1)	$\Delta_r H = -394 + (2 \times -286)890 = -76 \text{ (kJ mol}^{-1})$	
		Correct answer with no working scores final 2	
		Units not required, but if given must be correct	
		Ignore SF	
		Do not award kJ / mol ⁻¹ Allow TE on incorrect enthalpy of combustion calculation	

Q28.

Question Number	Answer	Additional Guidance	Mark
(i)	An answer that makes reference to the following points:	Penalise missing states only once (M1)	(3)
	correct elements in the lower box (P(s), Cl ₂ (g)) (1)	States are required Allow P ₄ (s) Ignore balancing numbers for M1	
	correct moles of each element, P(s) and 2½Cl₂(g) (1)	Allow ¹ / ₄ P ₄ (s) Ignore state symbols (if given)	
	 arrows correctly labelled (Δ_zH [PCl₅], Δ_zH [PCl₃]) 	Ignore state symbols (if given) on arrows	
PCl _s (g) + Cl ₂ (g)	

Question Number	Answer	Additional Guidance	Mark
(ii)	 use of ∑(Δ_tH[products] - ∑(Δ_tH[reactants] (1) correct answer with sign and units (1) 	Example of calculation (-319.7 + 30.5) - (-443.5 + 64.9) Allow correct sums $(-289.2 \text{ and } -378.6)$ but must be negative = +89.4 kJ mol ⁻¹ Sign and units must be shown Allow TE from M1(for omission of $\Delta_v H$ data (+123.8 kJ mol ⁻¹) Correct answer with no working scores (2)	(2)

Q29.

Question Number	Acceptable Answer	Additional Guidance	Mark
(a)	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	display all three methyl groups allow -OH do not award C-H-O	Antonoporo
			(1)

Question Number	Acceptable Answer	Additional Guidance	Mark
(b)(i)	An answer that makes reference to one of the following:		
	molecular ion/molecule fragments/is unstable		(1)

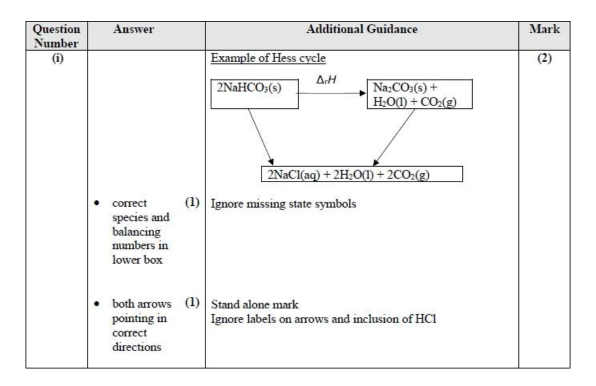
Question Number	Acceptable Answer	Additional Guidance	Mark
(ii)	СН ₃ — ст — СН ₃	allow + charge on any part of the ion/outside the structure but + must be shown	
		allow displayed/structural/skeletal/ molecular formulae or any combination of these.	

Question Number	Acceptable Answer	Additional Guidance	Mark
(c)(i)	calculation for bonds broken in the alcohol (*)	3(C-C) + 9(C-H) + (C-O) + (O-H)	
	calculation for bonds broken in oxygenand	=(3x347) + (9x413) + 358 + 464 = (+)5580 (kJ mol ⁻¹) 6(0=0) = (6 x 498) = (+)2988 (kJ mol ⁻¹)	
	total energy for bonds broken(**) (1) • calculation for bonds made(***) (1)	total = + 5580 + 2988 = (+)8568 (kJ mol ⁻¹) TE from ans * M1 + 2988 = 8(C=O) + 10(O-H) = (8x805) + (10x464) = -	
	 calculation of ∆_cH (2-methylpropan-2-ol) with sign (1) 	11080 (kJ mol ⁻¹)	
		answer(***) units not required but if given they must be correct correct final answer with no working scores 4 marks	(4)

Question Number	Acceptable Answer	Additional Guidance	Mark
(ii)	An explanation that makes reference to the following points:	mark independently	
	incomplete combustion (1)	do not award just	
	 Δ_cH (2-methylpropan- 2-ol) will be less negative /less exothermic than data book value 	lower/smaller/decreases/ more positive allow reduce the magnitude (of the value)	
	(1)		(2)

Question Number	Acceptable Answer	Additional Guidance	Mark
(iii)	An answer that makes reference to the following points:		
	Δ _c H figures are at 298 K /data book bond energies refer to gaseous state and	allow just liquid involved	
	water and/or 2-methylpropan- 2-ol are/is (both) liquid(s) (at 298 K)	do not award data book bond energies are mean (values)/not specific to 2-methylpropan- 2-ol	(1)

Q30.



Question Number		Answer		Additional Guidance	Mark
(ii)	•	expression for $\Delta_z H$	(1)	Example of calculation $\Delta_{r}H = 2x\Delta H_{1} - \Delta H_{2}$	(3)
	•	substitution of values into expression with both values in same units	(1)	$\Delta_r H = 2 \times 20.619 - (-29.4)$ or $\Delta_r H = 2 \times 20619 - (-29.400)$ M1 can be scored from values substituted into correct expression in M2 TE on ΔH_1 in (a)(iii) and expression in M1 No TE on incorrect arrows in	
	•	calculation of $\Delta_t H$ and sign and units		cycle $\Delta_r H = +70.638 \text{ kJ mol}^{-1}$ or $\Delta_r H = +70638 \text{ J mol}^{-1}$ TE on ΔH_1 in (a)(iii) and expression in M1 provided it is a +ve answer Ignore SF except 1 SF Correct answer with sign and units scores (3)	

Q31.

Question Number	Answer	Additional Guidance	Mark
(i)	 labelled y axis, including units, with appropriate scale (1) direction and placement of enthalpy changes, +18.2 and -84.5 (1) Entities with state symbols (1) 	Allow energy for 'enthalpy' Ignore horizontal axis Do not award enthalpy change/ΔH for y axis Allow ΔH ₁ and ΔH ₂ for respective values Arrows must be shown and in the correct direction Ignore activation energy 'curves' Do not award double-headed arrows Ignore inclusion of '+ aq' Example of diagram	(3)
		CuSO4 (s) Follula:	
		H 40- kjmel-1 20-	
		Arbitrary CuSO ₄ (aq) 4 2e(c) 2+18-2 CuSO ₄ .54 ₂ O ₆) 4	

Question Number	Answer	Additional Guidance	Mark
(ii)	• Use of Hess's law to calculate $\Delta r H$ shown on the diagram	Value from diagram = -102.7 (kJ mol ⁻¹)	(1)
		Allow $\Delta_r H = \Delta H_2 - \Delta H_1 = -84.5 - (+18.2) = -102.7 \text{ (kJ mol}^{-1}\text{)}$	
		Allow -103 (kJ mol ⁻¹)	
		Do not award if no working shown on the diagram	

Q32.

Question Number	Answer	Additional Guidance	Mark
	evaluation of Q (1)	Example of calculation Q = (ΔH × n)=18.2 × 0.025 = 0.455(kJ) or 455 J	(3)
	 rearrangement to give ΔT (1) 	Δ <i>T</i> =Q ÷(m x c) = 455 ÷ (45.00 x 4.18) = 2.4189(°C)	
	 Answer to 1 or 2SF and temperature change (1) 	ΔT = 2/2.4 °C/ K and decrease Allow -2/2.4 °C/ K Correct final answer without working scores (3)	
		TE throughout	

Q33.

Question Number	Answer	Additional Guidance	Mark
Number	An answer that makes reference to • Cannot react exactly 5 mol of water with 1 mol of anhydrous copper(II) sulfate	Cannot measure the temperature (change) for a solid Description that states more (than 5) water molecules will attach to some CuSO ₄ while less (than 5) water molecules will attach to other CuSO ₄ ACCEPT reasonable ideas such as some water may evaporate (due to exothermic reaction) Ignore heat loss to surroundings if given as an alternative reason Do not award heat is needed to start the reaction	(1)

Q34.

Question Number	Answer		Additional Guidance	Mark
	 calculation of energy required for breaking the bond in Cl₂ and I₂ 	(1)	= 151 + 243 = 394 (kJ mol ⁻¹)	(2)
	 calculation of energy in 2 moles of I-Cl bonds and divides by 2. 	(1)	$= \frac{394 + 30}{2} = (+)212 \text{ (kJ mol}^{-1})$	

Q35.

Question Number	Acceptable Answer	Additional Guidance	Mark		
	An answer that makes reference to the following point:				(1)
	 to make sure that (all) the (nitric) acid / HNO₃ / H⁺ has reacted / been neutralised / is used up 	Allow (nitric acid) / HNO3 is the limiting reagent			
	1000	Allow so that 0.025 mol of water / H ₂ O forms			
		Ignore to make sure that 1 mol of water / H ₂ O forms			
		Ignore just 'to ensure that reaction is complete'			

Q36.

Question Number	Answer	Mark
	The only correct answer is C	
	A is not correct because standard enthalpy of formation is for making 1 moles of a compound	
	B is not correct because standard enthalpy of formation is for making 1 moles of a compound	
	D is not correct because oxygen must be O₂	3

Q37.

Question Number	Acceptable Answer	Additional Guidance	Mark
Number (i)	Answer	Energy/Enthalpy Reaction Profile/ Progress of reaction	(3)
	 vertical axis labelled: H/enthalpy/e nergy/E (1) 	Do not award ΔH Ignore horizontal axis label Ignore units if given	
	level of reactants / 2SO ₂ + O ₂ above level of products / 2SO ₃ (1)	ignore state symbols even if incorrect	
	correct profile for uncatalysed reaction labelled A	allow vertical lines for catalysed and uncatalysed reactions to run together allow double hump profile	
	and peak lower for catalysed reaction labelled B (1)	reactants Ea1 Ea2 Intermediate (s)	

Question Number	Acceptable Answer	Additional Guidance	Mark
(ii)	enthalpy change, $\Delta_r H/\Delta H/(-)$ 197(kJ mol ⁻¹), shown correctly (1)	Ignore presence/absence of arrowheads Allow a degree of imprecision in the start/finish points of the lines for ΔH and E_a	(2)
	activation energy, E_a , shown correctly (upper diagram) (1)	Ea shown on double hump profile - shown in this diagram as Ea ₁ Ignore Ea2 if also shown	

Q38.

Question Number	Answer	Mark
(i)	The only correct answer is D	(1)
	A is not correct because the measurement uncertainty is doubled as there are two burette readings	
	B is not correct because this gives the largest measurement uncertainty	
	C is not correct because the measurement uncertainty is doubled as the pipette is used twice	

Question Number	Answer		Additional Guidance	Mark
(ii)			Example of calculation	(3)
	calculation of Q	(1)	Q=(100 x 4.18 x 6.5=) = 2717 (J) / 2.717 kJ	
	calculation of enthalpy change	(1)	ΔH= 2.717 ÷ 0.05= (-)54.340	
	answer to nearest whole number and w	rith negative sign (1)	= -54 (kJ mol ⁻¹)	
			Allow TE at each stage Correct answer with no working scores (3)	

Question Number	Answer	Mark
(i)	The only correct answer is B	(1)
	A is not correct because there is no extrapolation to the largest temperature increase carried out	
	C is not correct because the extrapolation is at the wrong time	
	D is not correct because the extrapolation extends beyond the time of addition of alkali	

Question Number	Answer	Additional Guidance	Mark
(ii)	An explanation that makes reference to		(2)
	 ethanoic acid is a weak(er) acid / only partially ionised/dissociated (1) 	Allow hydrochloric acid is a strong(er) acid/fully ionised	
	 (some) energy is used to fully/completely ionise the ethanoic acid (1) 	Do not award 'more NaOH will react so more energy given off'	

Q40.

Question Number	Acceptable Answers	Additional Guidance	Mark
	balanced equation (1)	$2C(s, graphite) + 3H_2(g) + 1/2O_2(g) \rightarrow C_2H_5OH(I)$ Allow C_2H_6O Do not allow multiples	(2)
	all state symbols (1)	Conditional on all species correct Allow C(s) / C(graphite)	