Surname	Centre Number	Candidate Number
First name(s)		2



GCE A LEVEL

A410U10-1





MONDAY, 12 JUNE 2023 – MORNING

CHEMISTRY – A level component 1

Physical and Inorganic Chemistry

2 hours 30 minutes

For Examiner's use only									
Question	Maximum Mark	Mark Awarded							
1. to 6.	15								
7.	19								
8.	17								
9.	13								
10.	24								
11.	18								
12.	14								
	Question 1. to 6. 7. 8. 9. 10.	Question Maximum Mark 1. to 6. 15 7. 19 8. 17 9. 13 10. 24 11. 18							

120

Section A

Section B

ADDITIONAL MATERIALS

- A calculator
- Data Booklet supplied by WJEC.

INSTRUCTIONS TO CANDIDATES

Use black ink or black ball-point pen. Do not use gel pen or correction fluid.

You may use a pencil for graphs and diagrams only. Write your name, centre number and candidate number in the spaces at the top of this page.

Section A Answer **all** questions. **Section B** Answer **all** questions.

Write your answers in the spaces provided in this booklet. 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.

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.7(a) and Q.11(a).



Total

SECTION A

Answer all questions.

1. The electronegativity values of some elements are listed in the table.

Element	Si	F	CI
Electronegativity value	1.8	4.0	3.5

	(a)	State what is m	neant by the term elect	ronegativity.		[1]
	(b)	Label the bond	s below with δ + and δ	– to show any	dipoles that are prese	nt. [[1]
			Si — CI	F-	- CI		
2.		e the feature of t ation energy of tl	he atomic emission sp ne atom.	ectrum of hydi	rogen that can be used	_	[1]



Element symbol Mass number



4.	An iodine	clock	reaction	was	studied	usina	different	concent	rations o	of reactar	nts.
	,	0.00.0			0100.00	~~9		001100110			

The time taken for the colour to change was measured.

Concentration of I ⁻ / mol dm ⁻³	Concentration of H ₂ O ₂ /mol dm ⁻³	Time for colour change/s	Rate
0.20	0.10	49.7	
0.20	0.20	24.9	
0.40	0.20	12.4	

(a)	(i)	Complete the table with the rate values for these experiments.	[1]
	(ii)	Give the unit of rate for these experiments.	[1]
		Unit	
(b)		cribe the relationship between the concentration of hydrogen peroxide and the relation.	ate [1]



[1]	

[1]

- Hydrogen chloride forms the strong acid hydrochloric acid when dissolved in water. 5. In one experiment the pH of the hydrochloric acid formed was 0.32.
 - Draw a dot and cross diagram for hydrogen chloride. (a)

State what is meant by the term strong acid. (b)

Calculate the concentration of hydrochloric acid of pH 0.32.

[1]

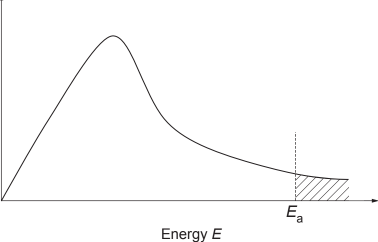
Concentration = mol dm⁻³

The graphs below show the energy distribution of particles at 298 K. 6.

Sketch the distribution at a lower temperature on the same axes.

[1]

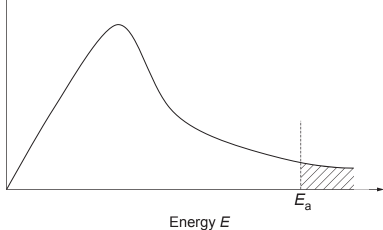
Fraction of molecules with energy *E*



Label the graph to show how catalysts increase the rate of chemical reactions. (b)

[1]

Fraction of molecules with energy E



15

A410U101

SECTION B

		Answer all questions.						
7.	The elements at the top and bottom of Group 4 have very different physical and chemic properties.							
	(a)	Describe the bonding and structures present in lead and one common elemental form of carbon.						
		You should explain how the bonding and structures affect the melting temperatures and electrical conductivity of these elements. [6 QEF						
			•••					
	•••••							



[2]

(b) A student is provided with a mixture of lead(II) oxide, PbO, and lead(IV) oxide, PbO₂.

To find the percentage by mass of each oxide in the sample the student treats the mixture with iodide ions in acid solution.

• Lead(IV) oxide oxidises the iodide ions in acid solution. This produces a precipitate of lead(II) iodide and an aqueous solution of iodine.

$$PbO_2 + 4I^- + 4H^+ \longrightarrow PbI_2 + I_2 + 2H_2O$$

Lead(II) oxide reacts with the iodide ions in acid solution. This produces a
precipitate of lead(II) iodide.

$$PbO + 2l^- + 2H^+ \longrightarrow Pbl_2 + H_2O$$

The solution and solid are separated by filtration. The solid is washed in cold water and the washings added to the solution. The solution is titrated using aqueous sodium thiosulfate of concentration $0.100\,\mathrm{mol\,dm^{-3}}$ to find the amount of iodine present. The solid sample is heated to constant mass.

The results are given below.

Mass of precipitate = 2.425 g

Volume of aqueous sodium thiosulfate required = 33.45 cm³

- (i) Give the colour of the lead(II) iodide precipitate. [1]
- (ii) Calculate the number of moles of lead(II) iodide formed.

Number of moles = mol

(iii) The thiosulfate ions act as reducing agents.

$$2S_2O_3^{2-} \longrightarrow S_4O_6^{2-} + 2e^{-}$$

Write the equation for the reduction of iodine by thiosulfate ions. [1]

		_
(iv)	Suggest a suitable indicator for this titration, giving the colour change expected at the endpoint. [2]	E
(v)	Calculate the number of moles of lead(IV) oxide present. [3]	
(vi)	Number of moles =mol Calculate the percentage by mass of lead(IV) oxide in the initial sample. [4]	

Percentage by mass = %

19



. Tran (a)		metals form a range of ions that can combine with ligands to form complex ions. e the electron arrangement for the Mn ²⁺ ion.	[1]
(b)	(i)	Cobalt(II) ions form complexes including $[{\rm CoCl_4}]^{2-}$ and $[{\rm Co(H_2O)_6}]^{2+}$. State the colours of these ions in aqueous solution.	[2]
		$[CoCl_4]^{2-}$	
	(ii)	Explain why octahedral transition metal complexes such as $[\mathrm{Co(H_2O)_6}]^{2^+}$ are coloured.	[3]

(c) Transition metal compounds are used in heterogeneous catalysts for a range of industrial processes.

(i) State what is meant by the term heterogeneous catalyst.

[1]

(ii) One industrial process that uses a heterogeneous catalyst is the carbonylation of methanol (Method 1).

 $CH_3OH + CO \rightleftharpoons CH_3COOH$

This reaction is usually carried out in the gas phase.

An equimolar gas phase mixture of CH $_3$ OH and CO is allowed to come to equilibrium and the resultant mixture contains 10.8% CH $_3$ COOH at a total pressure of 3.21 \times 10 6 Pa.

Calculate the value of K_p for this equilibrium giving its unit.

[4]

(A410U10-1)

ng ol to
or your [3]

Examiner only

(d) Most metals react with acids to produce a salt and hydrogen gas. Copper metal does not react in this way but it does react with nitric acid, HNO₃, in a different reaction.

Use the values of standard electrode potentials given in the table to explain these observations, writing an equation for the reaction you expect to occur between copper metal and nitric acid. [3]

	Standard electrode potential, E^{θ}/V
2H ⁺ + 2e [−] ← H ₂	+0.00
Cu ²⁺ + 2e [−]	+0.33
$NO_3^- + 2H^+ + e^- \rightleftharpoons NO_2 + H_2O$	+0.81

	Equation	
•••••		
		Γ

17



- **9.** Most Group 2 elements form oxides with similar properties.
 - (a) Draw a dot and cross diagram for the formation of magnesium oxide from atoms of its elements. [2]

- (b) The enthalpy change of formation of magnesium oxide can be found experimentally by measuring the temperature rise in two reactions and calculating their respective enthalpy changes.
 - · Adding magnesium metal to an excess of dilute acid
 - Adding magnesium oxide to an excess of dilute acid

Two students performed the experiment using the following method.

- Measure 50 cm³ of hydrochloric acid of concentration 1.0 mol dm⁻³ using a burette and place in a polystyrene cup.
- Place a lid on the cup and fit a thermometer through this. The thermometer measures to the nearest 0.2°C.
- Measure the temperature every 30 seconds for 3 minutes.
- Measure precisely a known amount of magnesium metal.
- Add the solid at precisely 3 minutes, stirring well.
- Measure the temperature every 30 seconds for another 4 minutes.
- Repeat these steps using magnesium oxide.
- Plot the results and calculate the enthalpy changes.



Examiner only

Tom decides to use the same mass of magnesium metal and magnesium oxide in his two experiments. He measures precisely $0.200\,\mathrm{g}$ of each and obtains the following results.

Results with magnesium metal

Time/s	0	30	60	90	120	150	180	210	240	270	300	330	360	390	420
Temperature /°C	19.2	19.4	19.4	19.4	19.4	19.4	XX	37.2	38.8	38.8	38.8	38.6	38.6	38.4	38.4

Results with magnesium oxide

Time/s	0	30	60	90	120	150	180	210	240	270	300	330	360	390	420
Temperature /°C	19.2	19.4	19.4	19.4	19.4	19.4	XX	22.0	22.4	22.8	22.8	22.6	22.6	22.6	22.4

Tom's teacher told him that his choice of mass of magnesium oxide has not given him suitable results and that he should repeat his experiment with a different mass.

I.	Suggest why Tom's mass of magnesium oxide did not give suitable resul	ts. [1]
II.	Tom decides to repeat both experiments using a mass of 1.00 g of each solid.	
	Suggest two reasons why 1.00 g would not be a suitable mass of magnesium metal to use.	[2]
		<u>.</u>



Turn over.

(ii) Cerys carried out the same experiments and obtained the following enthalpy change values. All substances were in their standard states.

Mg + 2HCl
$$\longrightarrow$$
 MgCl₂ + H₂ $\Delta H = -521 \text{ kJ mol}^{-1}$
MgO + 2HCl \longrightarrow MgCl₂ + H₂O $\Delta H = -142 \text{ kJ mol}^{-1}$

The enthalpy change of formation of H_2O is $-286 \, kJ \, mol^{-1}$.

Use this data to calculate the enthalpy change of formation of magnesium oxide.

. [3]



Barium oxide can be formed by heating barium carbonate. (c)

BaCO₃(s) \longrightarrow BaO(s) + CO₂(g) $\Delta H = 243 \text{ kJ mol}^{-1}$

	BaCO ₃ (s)	BaO(s)	CO ₂ (g)
Standard entropy/JK ⁻¹ mol ⁻¹	112	70	214

Give a reason why the standard entropy values of ${\rm BaCO}_3$ and ${\rm BaO}$ are lower than that of ${\rm CO}_2.$

Calculate the minimum temperature required for the thermal decomposition of barium carbonate. [4]

T =	k
, –	 ľ

13

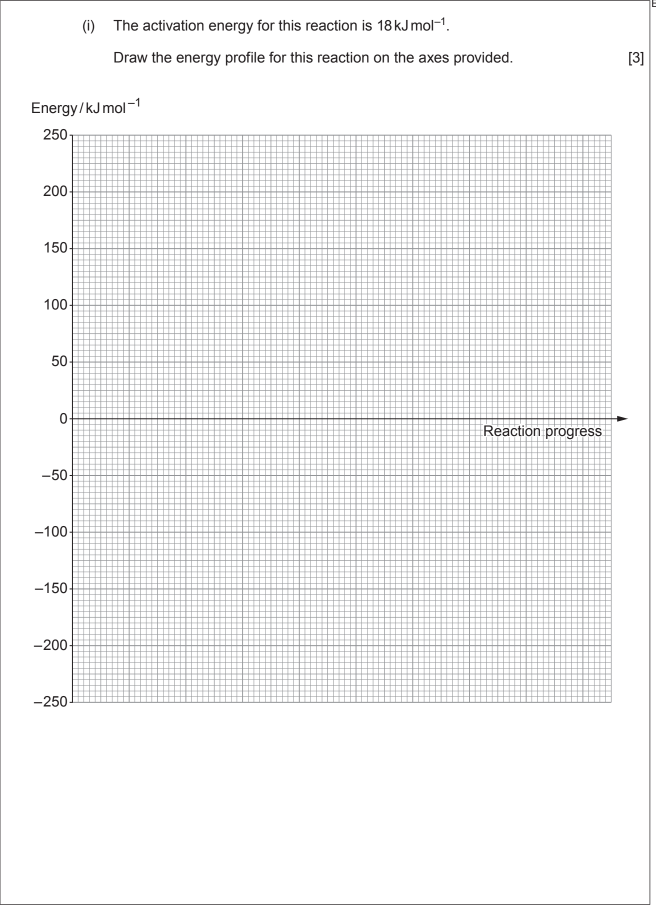
- **10.** Chlorine forms a range of oxides and oxyanions. Many of these are reactive species that can oxidise or chlorinate a range of elements and compounds.
 - Dichlorine heptoxide, $\mathrm{Cl_2O_7}$, is a colourless liquid that can be formed by reaction of (a) ozone with chlorine in the presence of ultraviolet light.

$$3Cl_2(g) + 7O_3(g) \longrightarrow 3Cl_2O_7(g)$$
 $\Delta H^{\theta} = -199 \text{ kJ mol}^{-1}$

$$\Delta H^{\theta} = -199 \,\mathrm{kJ} \,\mathrm{mol}^{-1}$$

Substance	Standard enthalpy change of formation, $\Delta_{\rm f} H^{ \theta} / {\rm kJ mol^{-1}}$
Cl ₂ (g)	0
O ₃ (g)	142

Examiner only





Calculate the temperature required to double the rate of reaction. (iii) The rate equation for this reaction is first order with respect to each reactant and the rate is measured in units of mol dm ⁻³ s ⁻¹ . I. Write the rate equation. [1] II. Give the unit of the rate constant. [1] (iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that of ozone is not. [2]	(iii) The rate equation for this reaction is first order with respect to each reactant and the rate is measured in units of mol dm ⁻³ s ⁻¹ . I. Write the rate equation. II. Give the unit of the rate constant. (iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that of the constant is the constant is the constant in the constant is	d.
 (iii) The rate equation for this reaction is first order with respect to each reactant and the rate is measured in units of mol dm⁻³ s⁻¹. I. Write the rate equation. [1] II. Give the unit of the rate constant. [1] (iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that of 	 (iii) The rate equation for this reaction is first order with respect to each reactant and the rate is measured in units of moldm⁻³ s⁻¹. I. Write the rate equation. II. Give the unit of the rate constant. (iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that ozone is not. 	[4]
II. Give the unit of the rate constant. [1] (iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that of	II. Give the unit of the rate constant. (iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that o ozone is not.	
(iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that of	(iv) Explain why the standard enthalpy of formation for chlorine gas is zero but that o ozone is not.	[1]
	ozone is not.	[1]



(v) Dichlorine heptoxide decomposes readily to form $Cl_2(g)$ and $O_2(g)$.

$$2Cl_2O_7(g) \longrightarrow 2Cl_2(g) + 7O_2(g)$$

Calculate the volume of gas produced when 2.70 g of ${\rm Cl_2O_7}$ decomposes at a temperature of $-12^{\circ}{\rm C}$ and 1 atm pressure. [4]

Volume = dm^3

(b)	solu	prine dioxide, CIO_2 , is an unstable oxide of chlorine. It is often stored in aqueous tion and in some countries solutions can only be transported if the concentration or than 0.30 g in $100\mathrm{cm}^3$ of water.	is
	Calo	culate the concentration of this solution in moldm ⁻³ .	[2]
		Concentration = mold	m ^{–3}
(c)	Chlo	prine perchlorate, ${\rm CIOCIO_3}$, is an oxide with the two chlorine atoms in different ation states.	
	(i)	If one chlorine atom has an oxidation state of +1, find the oxidation state of the other.	[1]
	(ii)	Predict the appearance of the molecular ion peaks seen in the mass spectrum chlorine perchlorate. Give reasons for your answer. You should refer to both the positions and heights of the peaks.	of [4]
	······		



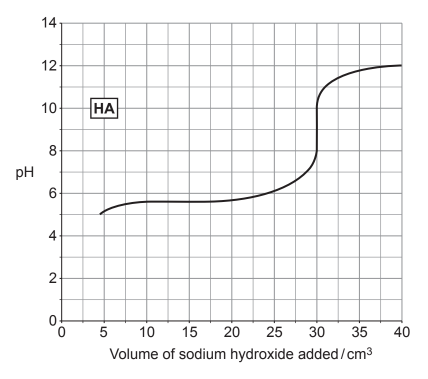
(d)	The chlorite ion, $O = CI - O^-$, has a non-linear shape.		Exa
	Explain why this ion is not linear.	[2]	

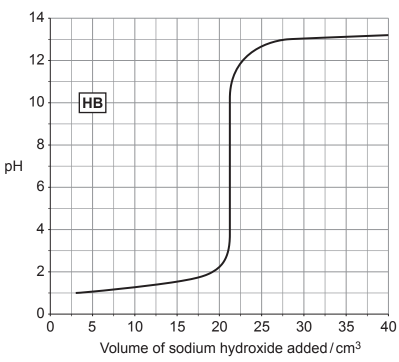


11. A student carried out two acid-base titrations using two acids, HA and HB.

A $25.0\,\mathrm{cm^3}$ sample of each acid was titrated against a sodium hydroxide solution of concentration $0.150\,\mathrm{mol\,dm^{-3}}$ giving the titration curves shown. The initial pH values are missing from both graphs.

One acid is a strong acid and one is a weak acid.







(a)	Desc	cribe and explain the differences between the two curves.	
	•	You should identify which acid is strong and which is weak, explaining fu you have reached your conclusion.	lly how
	•	You should identify which acid is more concentrated and which is more cexplaining how you have reached your conclusion.	dilute,
			[6 QER]
(b)	Calc	ulate $K_{ m a}$ for the weak acid.	[2]
(6)	Outo	ulate Na for the weak dold.	[4]
		K _a =	moldm ⁻³
		/\a =	. IIIOIUIII



(c)	simil	these titrations can be performed using appropriate acid-base indicators, ar experiments using a weak acid and a weak base cannot use acid-base essfully. Explain this difference.	indicat
(d)	The 0.15	students made $250\mathrm{cm}^3$ of aqueous sodium hydroxide of concentration 0 mol dm $^{-3}$ for these experiments.	
	(i)	Calculate the mass of NaOH required to make this solution.	
		Mass =	
	(ii)	Mass = Outline how this solution could be made.	
	(ii)		
	(ii) 		
	(ii)		
		Outline how this solution could be made.	
		Outline how this solution could be made.	
		Outline how this solution could be made.	
		Outline how this solution could be made.	



		27		
(i	iii)	Calculate the pH of this aqueous sodium hydroxide at 298K.	[3]	Examine only
		pH =		
				18



Turn over.

				20	
2.	(a)	Five ior	nisation energies are represen	ted by the letters A – E as shown belo	DW.
		A 1st ionisation energy of helium			
		В	final ionisation energy of nitro		
		С	final ionisation energy of oxy		
		D	1st ionisation energy of sodiu	um	
		E	2nd ionisation energy of mag	gnesium	
		The va	lues of these five ionisation en	ergies are given in the table.	
		V	/alue of ionisation energy /kJ mol ⁻¹	Letter representing the ionisation energy	
			84 078		
			64360		
			2372		
			1450		
			496		
	Complete the table using letters A – E to show which ionisation energy correach value.				
		Give ie	easons for your choices.		[5]
	•••••				
	•••••				
	•••••	•••••			
	•••••				



(b)	Three boiling temperatures are	listed below.		
	–33°C	–111°C	–132°C	
	These are the boiling temperat corresponds to each compound	ures of NH ₃ , PH d.	$_{\rm 3}$ and AsH $_{\rm 3}$. Identify which	n value
	Give reasons for your choices.			[3]
	Boiling temperature of NH ₃		°C	
	Boiling temperature of PH ₃		°C	
	Boiling temperature of AsH ₃		°C	
	Ç			

•••••				

(c) A student is provided with four solutions labelled **W**, **X**, **Y** and **Z** and is told that these contain common cations and anions that they have studied.

He observes that one solution is pale blue and the others are colourless. Flame tests on the solutions give apple-green and golden yellow flames with two solutions and unfamiliar colours with the others.

The student mixes pairs of solutions together and obtains the following results. He did not complete all the experiments.

Solution 1	Solution 2	Observation(s)	
W	х	white precipitate that dissolves when excess solution W is added	
W	Y	nixture of pale blue precipitate and white precipitate in a olourless solution	
W Z no visible change			
X Y		white precipitate in a pale blue solution	
X	Z		
Y	Z	white precipitate in a brown solution	

Compound X	Identify the four compounds and give reasons for your decisions.	[6]	Exam on
Compound Y Compound Z	Compound W		
Compound Z	Compound X		
	Compound Y		
	Compound Z		
END OF PAPER			1
	END OF PAPER		'



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Question number	Additional page, if required. Write the question number(s) in the left-hand margin.	Examine only
	3	1





GCE A LEVEL







MONDAY, 12 JUNE 2023 - MORNING

CHEMISTRY – A level component 1 Data Booklet

Avogadro constant
molar gas constant
molar gas volume at 273 K and 1 atm
molar gas volume at 298 K and 1 atm
Planck constant
speed of light
density of water
specific heat capacity of water
ionic product of water at 298 K
fundamental electronic charge

 $R = 8.31 \,\mathrm{J}\,\mathrm{mol}^{-1}\,\mathrm{K}^{-1}$ $V_m = 22.4 \,\mathrm{dm}^3 \,\mathrm{mol}^{-1}$ $V_m = 24.5 \,\mathrm{dm}^3 \,\mathrm{mol}^{-1}$ $h = 6.63 \times 10^{-34} \,\mathrm{J}\,\mathrm{s}$ $c = 3.00 \times 10^8 \,\mathrm{m}\,\mathrm{s}^{-1}$ $d = 1.00 \,\mathrm{g}\,\mathrm{cm}^{-3}$ $c = 4.18 \,\mathrm{J}\,\mathrm{g}^{-1}\,\mathrm{K}^{-1}$ $K_w = 1.00 \times 10^{-14} \,\mathrm{mol}^2 \,\mathrm{dm}^{-6}$ $e = 1.60 \times 10^{-19} \,\mathrm{C}$

 $N_A = 6.02 \times 10^{23} \,\mathrm{mol}^{-1}$

temperature (K) = temperature (°C) + 273

 $1 \,dm^3 = 1000 \,cm^3$ $1 \,m^3 = 1000 \,dm^3$ $1 \,tonne = 1000 \,kg$ $1 \,atm = 1.01 \times 10^5 \,Pa$

Multiple	Prefix	Symbol
10 ⁻⁹	nano	n
10 ⁻⁶	micro	μ
10 ⁻³	milli	m

Multiple	Prefix	Symbol
10 ³	kilo	k
10 ⁶	mega	M
10 ⁹	giga	G

2

Infrared absorption values

Bond	Wavenumber/cm ⁻¹
C — Br	500 to 600
C-CI	650 to 800
C - O	1000 to 1300
C = C	1620 to 1670
C = O	1650 to 1750
$C \equiv N$	2100 to 2250
C-H	2800 to 3100
O — H (carboxylic acid)	2500 to 3200 (very broad)
O — H (alcohol / phenol)	3200 to 3550 (broad)
N-H	3300 to 3500

13 C NMR chemical shifts relative to TMS = 0

Type of carbon	Chemical shift, δ (ppm)
Type of Carbon	Chemical Shift, 0 (ppin)
-c-c-	5 to 40
R — C — CI or Br	10 to 70
R-c-c- 0	20 to 50
R-C-N	25 to 60
-c-o-	50 to 90
c=c	90 to 150
$R-C \equiv N$	110 to 125
	110 to 160
R — C — (carboxylic acid / e	ster) 160 to 185
R — C — (aldehyde / ketone O) 190 to 220

¹H NMR chemical shifts relative to TMS = 0

Type of proton	Chemical shift, δ (ppm)
$-CH_3$	0.1 to 2.0
$R-CH_3$	0.9
$R-CH_2-R$	1.3
CH ₃ −C≡N	2.0
CH ₃ -C	2.0 to 2.5
$-CH_2-C$	2.0 to 3.0
\bigcirc CH ₃	2.2 to 2.3
HC-Cl or HC-Br	3.1 to 4.3
HC-O	3.3 to 4.3
R-OH	4.5 *
-C = CH	4.5 to 6.3
-c = cH - co	5.8 to 6.5
\leftarrow CH=C	6.5 to 7.5
◯ ⊢H	6.5 to 8.0
ОН	7.0 *
R-C H $R-C$ O OH	9.8 *
R-COH	11.0 *

^{*}variable figure dependent on concentration and solvent

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Krypton 36 Xenon 54 Radon 86 Helium 20.2 **Ne** 10 40.0 **Ar** Argon 18 83.8 **K** X 33 (222) **Rn** N Bromine 35 Chlorine Fluorine 9 Astatine 85 Lawrendum 103 lodine 53 79.9 **Br** (210) At 127 Lutetium 71 (257) Lr Oxygen 8 Polonium 84 Selenium Tellurium 79.0 Se **a** 158 (210) **Po** Ytterbium 70 Nobelium 102 (254) No ထ p block Arsenic 33 Bismuth 83 Nitrogen 7 Mendelevium 101 **Antimony** Thulium 69 74.9 **As** 122 **Sb** 203 **B**i (256) Md S Germanium Fermium 100 Silicon 72.6 **Ge** Erbium 68 Lead 82 30 Sn (253) Fm 28.1 Si 207 **Pb** 167 Er 4 Aluminium 13 Gallium Indium Einsteinium 99 10.8 **B** Boron 5 Holmium 67 69.7 **Ga** 204 1 (254) **Es** 27.0 **A** 132 3 Dysprosium 66 Californium 98 Cadmium Mercury 80 65.4 Zn Zinc 30 142 Cd 142 (251) Cf 163 THE PERIODIC TABLE Berkelium 97 Ag Ag Silver Terbium 65 Gold 79 (245) **BK** 197 **Au** f block Nickel 28 Palladium 195 Pt Platinum 78 Gadolinium 64 Curium 96 106 Pd (247) Cm 46 157 Gd Cobalt 27 Rhodium Iridium 77 Europium 63 Americium 95 ₽ ¹⁰³ (243) **Am** (153) Eu 45 Osmium 76 Plutonium 94 Samarium 62 Ruthenium 55.8 Fe 190 **Os** Iron 26 (242) Pu ₹ 2007 atomic number Group relative atomic mass d block Key Manganese 25 Rhenium Neptunium 93 echnetium 54.9 **Mn** (147) Pm (237) **Np** 98.9 7 186 Re A_r Symbol 9 Name Z / Chromium 24 Uranium 92 Molybdenum Tungsten 74 95.9 **Mo** 52.0 Cr ₹ ≥ 238 **○** 9 42 Protactinium 91 Praseodymium 59 Tantalum 73 /anadium Niobium 92.9 **Nb** (231) **Pa** <u>™</u> 2 <u>₹</u> ₽ Zirconium 40 Fitanium Hafnium 72 Thorium 90 Cerium 179 **H** (227) Ac •• Lanthanum 57 Actinium 89 Yttrium ► Lanthanoid Scandium elements ▶▶ Actinoid elements 139 **La** Radium 88 Magnesium 12 Calcium Strontium Barium Beryllium Ca 40.1 (226) **Ra** 87.6 Sr 137 **Ba** 26 38 s block Caesium 55 Rubidium 37 Hydrogen Sodium (223) Fr Francium 87 Lithium 3 Potassium 23.0 **Na** 85.5 **Rb** 133 Cs 6.94 Li 39.1 5. **⊥** 9 Period 2 S 9 / 3

(A410U10-1A)

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