## TOPICAL PAST PAPER QUESTIONS WORKSHEETS

# AS & A Level Chemistry (9701) Paper 2

Exam Series: Feb/Mar 2017 - Oct/Nov 2023

Format Type A:
Answers to all questions are provided as an appendix



## Introduction

Each Topical Past Paper Questions Compilation contains a comprehensive collection of hundreds of questions and corresponding answer schemes, presented in worksheet format. The questions are carefully arranged according to their respective chapters and topics, which align with the latest IGCSE or AS/A Level subject content. Here are the key features of these resources:

- 1. The workbook covers a wide range of topics, which are organized according to the latest syllabus content for Cambridge IGCSE or AS/A Level exams.
- 2. Each topic includes numerous questions, allowing students to practice and reinforce their understanding of key concepts and skills.
- 3. The questions are accompanied by detailed answer schemes, which provide clear explanations and guidance for students to improve their performance.
- 4. The workbook's format is user-friendly, with worksheets that are easy to read and navigate.
- 5. This workbook is an ideal resource for students who want to familiarize themselves with the types of questions that may appear in their exams and to develop their problem-solving and analytical skills.

Overall, Topical Past Paper Questions Workbooks are a valuable tool for students preparing for IGCSE or AS/A level exams, providing them with the opportunity to practice and refine their knowledge and skills in a structured and comprehensive manner. To provide a clearer description of this book's specifications, here are some key details:

- Title: Cambridge AS & A Level Chemistry (9701) Paper 2 Topical Past Papers
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# Chapter 1

# Atomic structure

 $1.\ 9701\_m22\_qp\_22\ Q\!:\, 1$ 

Fig. 1.1 shows how **first** ionisation energies vary across Period 2.

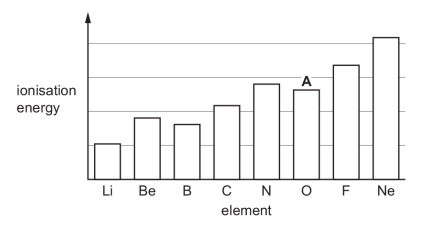


Fig. 1.1

(a)	Construct an equation to represent the first ionisation energy of oxygen.
	Include state symbols.

		[1]
(b)	(i)	State and explain the general trend in first ionisation energies across Period 2.
		[3]
	(ii)	Explain why ionisation energy <b>A</b> in Fig. 1.1 does <b>not</b> follow the general trend in first ionisation energies across Period 2.

(c) Element E is in Period 3 of the Periodic Table.

The first eight ionisation energy values of E are shown in Table 1.1.

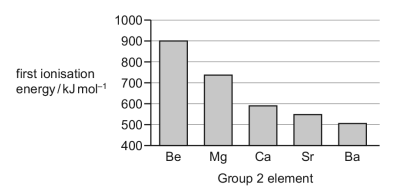
Table 1.1

ionisation	1st	2nd	3rd	4th	5th	6th	7th	8th
ionisation energy/kJ mol <sup>-1</sup>	577	1820	2740	11 600	14800	18400	23 400	27500

Deduce the full electronic configuration of <b>E</b> . Explain your answer.	
full electronic configuration of E =	
explanation	
	[3]
	[Total: 9]

[Total: 6]

The graph shows the first ionisation energies of some of the elements in Group 2.



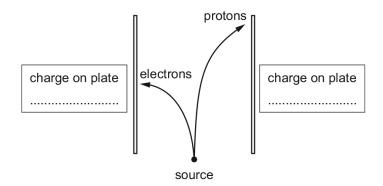
(a) Write an equation for the first ionisation energy of Mg.

	Include state symbols.
	[1]
(b)	Explain the observed trend in first ionisation energies down Group 2.
	[3]
(c)	The second ionisation energy of Be is 1757 kJ mol <sup>-1</sup> .
	Explain why the second ionisation energy of Be is higher than the first ionisation energy of Be.
	[2]

$$3.\ 9701\_w20\_qp\_22\ Q:\ 1$$

Atoms contain the subatomic particles electrons, protons and neutrons. Protons and electrons were discovered by observations of their behaviours in electric fields.

(a) The diagram shows the behaviour of separate beams of electrons and protons in an electric field.



- (i) Complete the diagram with the relative charge of each of the electrically charged plates.
- (ii) On the diagram, draw a line to show how a separate beam of neutrons from the same source behaves in the same electric field. [1]
- (b) Electrons in atoms up to  $_{36}$ Kr are distributed in s, p and d orbitals.
  - (i) State the number of occupied orbitals in an isolated atom of  $_{36} \rm Kr.$

type of orbital	S	р	d
number of orbitals			

[3]

	(ii)	Complete the diagram to show the number and relative energies of the electrons in an isolated atom of $_{\rm 14}{\rm Si}.$	
		4s	
		3р	
		3s	
		2p	
		2s	
		1s 1	
		[2]	
	(iii)	The diagram shows a type of orbital.	
		State the total number of electrons that exist in all orbitals of this type in an atom of ${}_9F.$	
		[1]	
	(iv)	The first ionisation energies of elements in the first row of the d block ( $_{21}$ Sc to $_{29}$ Cu) are very similar. For all these elements, it is a 4s electron that is lost during the first ionisation.	
		Suggest why the first ionisation energies of these elements are very similar.	
		[3]	
(c)	Нус	dron is a general term used to represent the ions ${}^1_1H^+$ , ${}^2_1H^+$ and ${}^3_1H^+$ .	
		te, in terms of subatomic particles in the nucleus, what is the same about each of these ions what is different.	
	san	ne	
	diffe	erent[1]	
		[Total: 12]	

# Chapter 2

Atoms, molecules and stoichiometry

$$4. \ 9701 \_w22 \_qp \_21 \ \ Q: 1$$

Atoms with nuclei containing an odd number of protons tend to have fewer isotopes than those with an even number of protons.

- (a) Gallium has two stable isotopes, <sup>69</sup>Ga and <sup>71</sup>Ga.
  - (i) Complete Table 1.1 to show the numbers of protons, neutrons and electrons in the two stable isotopes of gallium.

Table 1.1

isotope	number of protons	number of neutrons	number of electrons
<sup>69</sup> Ga			
<sup>71</sup> Ga			

(ii) Define relative atomic mass.

[2]

(iii) Define relative atomic mass.

[2]

(iii) The relative atomic mass of gallium, *A<sub>r</sub>*, is 69.723. The relative isotopic masses of <sup>69</sup>Ga and <sup>71</sup>Ga are:

<sup>69</sup>Ga, 68.926; <sup>71</sup>Ga, 70.925.

Use this information to calculate the percentage abundance of <sup>69</sup>Ga in elemental gallium. Show your working.

Assume that the element contains only the <sup>69</sup>Ga and <sup>71</sup>Ga isotopes.

Give your answer to four significant figures.

percentage abundance of <sup>69</sup>Ga = ..... %
[2]

<b>o</b> )	Pot	assium also has two stable isotopes. Both isotopes have the same chemical properties.
	(i)	Explain why both isotopes of potassium have the same chemical properties.
		[1]
	(ii)	State the full electronic configuration of an atom of potassium.
		[1]
(	iii)	The first, second and third ionisation energies of potassium are 418, 3070 and $4600\mathrm{kJmol^{-1}}$ , respectively.
		Use this information to explain why potassium is in Group 1.
		[2]
		[2]
		[Total: 10]

The composition of atoms and ions can be determined from knowledge of atomic number, nucleon number and charge.

(a) Complete the table.

 $5.\ 9701\_s17\_qp\_22\ Q:\ 1$ 

atomic number	nucleon number	number of electrons	number of protons	number of neutrons	symbol
3		2			<sup>6</sup> Li⁺
		23	26	32	

[2]

(b) Boron occurs naturally as a mixture of two stable isotopes, <sup>10</sup>B and <sup>11</sup>B. The relative isotopic masses and percentage abundances are shown.

isotope	relative isotopic mass	abundance/%
<sup>10</sup> B	10.0129	19.78
<sup>11</sup> B	to be calculated	80.22

(i)	Define the term relative isotopic mass.	
		[2]
(ii)	Calculate the relative isotopic mass of <sup>11</sup> B.	
	Give your answer to six significant figures. Show your working	

[2]

[Total: 6]

# Chapter 3

# Chemical bonding

6. 9701\_s23\_qp\_22 Q: 1

The melting points of some solids are shown in Table 1.1.

Table 1.1

solid	melting point/K
magnesium	923
phosphorus	317
sodium chloride	1074
sulfur	392

(a) (i)	State the type of bonding present in magnesium and in sodium chloride.
	bonding in magnesium
	bonding in sodium chloride
	[1]
(ii)	Explain the difference in the melting points of magnesium and sodium chloride.
	[1]
(iii)	Explain the difference in the melting points of phosphorus and sulfur in terms of structure and bonding.
	[2]
(b) (i)	Define electronegativity.
	[1]
(ii)	Explain why electronegativity increases across a period.

(iii)	Name the strongest intermolecular force that exists between $\mathrm{NH_3}(\mathrm{I})$ molecules.					
					[1]	
(iv)	Draw a diagram two molecules of		ation of the stronge	est intermolecular force be	etween	
	Include any releva	ant lone pairs of e	lectrons and dipoles			
					[2]	
(v)	The melting points	s of ice and ammo	onia are shown in Ta	ible 1.2.	[-]	
(-,	J P		ble 1.2			
		solid	melting point/K			
		ice	273			
		ammonia	195			
	Suggest <b>two</b> reas	ons for the differe	ence in the melting p	oints of ice and ammonia.		
					[2]	
				[To	tal: 12]	

The Pauling electronegativity values of elements can be used to predict the chemical properties of compounds.

Use the information in Table 1.1 to answer the following questions.

Table 1.1

element	Н	Li	С	0	S
Pauling electronegativity value	2.1	1.0	2.5	3.5	2.6
first ionisation energy/kJ mol <sup>-1</sup>	1310	519	1090	1310	1000
second ionisation energy/kJ mol <sup>-1</sup>	_	7300	2350	3390	2260

a)	(i)	Define electronegativity.	
			[1]
	(ii)	O and S are in Group 16.	
		Explain the difference in the Pauling electronegativity values of O and S.	
			[2]
b)	(i)	LiH is an ionic compound.	
		Draw a dot-and-cross diagram of LiH.	
		Include all electrons.	
			[2]
	(ii)	Suggest the shape of a molecule of H <sub>2</sub> S.	
			[1]

(c)	(i)	Write an equation that represents the first ionisation energy of H.	
	(ii)	Explain why there is no information given in Table 1.1 for the second ionisation energy	
		of H.	[1]
	(iii)	Give the full electronic configuration of S <sup>2+</sup> (g).	[1]
(d)	CO2	<sub>2</sub> and SO <sub>2</sub> are acidic gases.	
	(i)	Write an equation for the reaction of SO <sub>2</sub> with H <sub>2</sub> O.	[1]
	(ii)	Write an equation for the reaction of SO <sub>2</sub> with NaOH.	[1]
	(iii)	Construct an equation for the reaction of CO <sub>2</sub> with Mg(OH) <sub>2</sub> .	[.]
			[1]

(e) (i) Complete Table 1.2 by placing a tick (✓) to show which of the compounds have molecules with an overall dipole moment.

Table 1.2

compound	O=C=O	O=S=O	S=C=S	S=C=O
overall dipole moment				

[2]

(ii) At 150 °C and 103 kPa, all of the compounds listed in Table 1.2 are gases.

Under these conditions, 0.284g of one of the compounds occupies a volume of 127 cm<sup>3</sup>.

Use this information to calculate the  $M_{\rm r}$  of the compound. Hence, identify the compound from those given in Table 1.2.

Show your working.

$$M_{\rm r}$$
 = ...... identity of compound = ..... [3]

[Total: 17]

# Chapter 4

# States of matter

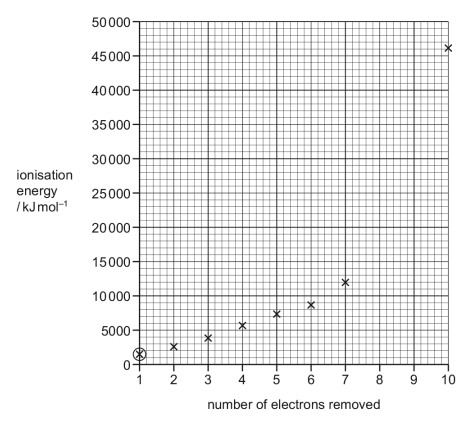
8. 9701\_s19\_qp\_21 Q: 3

(a) Construct an equation for the second ionisation energy of argon.

......[1]

(b) The graph shows successive ionisation energies for the element argon.

Complete the graph with predictions for the eighth and ninth ionisation energies of argon. Use a cross (x) for each data point. [2]



(c) The energy value required to remove the first electron from an atom of argon is circled on the graph.

Sketch the shape of the orbital that contains this electron.

(d)	Chlorine exists as a diatomic gas, $Cl_2(g)$ . A sample of $Cl_2(g)$ was made during a chemical reaction. When measured at 404 kPa and 25 °C the sample occupied a volume of 20.0 cm <sup>3</sup> .						
	(i)	Calculate the mass, in grams, of $Cl_2(g)$ formed.					
		For this calculation, assume that chlorine behaves as an ideal gas under these conditions.					
		mass of $Cl_2(g) =g$ [3]					
	(ii)	Calculate the number of chlorine atoms in this sample of ${\rm C}l_2({\rm g})$ . You may find it helpful to use your answer to (d)(i).					
		If you are unable to calculate an answer to <b>(d)(i)</b> , use 0.36g of $Cl_2$ . This is <b>not</b> the correct answer.					
		number of chlorine atoms = [2]					
	(iii)	$Cl_2(g)$ does <b>not</b> behave as an ideal gas under these conditions.					
		Explain why $Cl_2(g)$ behaves even less ideally at:					
		very high pressures					
		very low temperatures.					
		[2]					
		[Total: 11]					

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a) Define first io	nisation e	energy.						
) Successive id	onisation	energies f	or elemen	it <b>A</b> are sl	nown in Ta	ble 1.1.		
			Tabl	e 1.1				
ionisation	1st	2nd	3rd	4th	5th	6th	7th	8th
ionisation nergy/kJ mol <sup>-1</sup>	1310	3390	5320	7450	11 000	13 300	71 000	84 100
Use Table 1.1	I to deduc	e the grou	ıp of the P	eriodic Ta	ble that <b>A</b>	belongs to	o. Explain	your ans
Group								
Across Perio							to increas	e due to
increase in at								
Explain why phosphorus.	the first	ionisation	energy c	of sulfur i	s less tha	in the firs	t ionisatio	on energ
I) In an A $l^{2+}$ ion	the nucle	ear attract	ion for the	outer ele	ctron is st	ronger tha	an in an at	tom of Na
Compare the								nird ionisa
energy of alu	minium is	greater th	han the firs	st ionisati	on energy	of sodiun	n.	

(e) An isotope of copper has a relative isotopic mass of 65.

Complete Table 1.2 for an atom of copper-65.

Table 1.2

	atomic number	nucleon number	number of neutrons	electronic arrangement
copper-65				

			[3]
f)	(i)	The element copper has a relative atomic mass of 63.5.	
		Calculate how many atoms are present in 1.05g of copper.	
		atoms of copper present =	[1]
	(ii)	Copper has a melting point of 1085°C and a high electrical conductivity.	
		Explain these properties of copper by referring to its structure and bonding.	

.....[2]

[Total: 13]

10. 9701\_s21\_qp\_22 Q: 2

The strength of interaction between particles determines whether the substance is a solid, liquid or gas at room temperature.

(a) Lithium sulfide, Li<sub>2</sub>S, is a crystalline solid with a melting point of 938 °C. It conducts electricity when it is molten.

(i) Give the formulae of the particles present in solid lithium sulfide.

(ii) Explain, in terms of the structure of the crystalline solid, why lithium sulfide has a high melting point.

ici

- (b) Carbon monoxide, CO, is a gas at room temperature and pressure. It contains a coordinate bond.
  - (i) Explain what is meant by coordinate bond.

[1]

(ii) Draw a 'dot-and-cross' diagram to show the arrangement of outer electrons in CO.

Show the electrons belonging to the C atom as  $\times$ .

Show the electrons belonging to the O atom as •.

		. Neither CO nor $N_2$ is	an ideai
i) State two assumptions that are ma	ade about the behaviour	of particles in an ideal	gas.
1			
2			
			[2]
) Explain why N <sub>2</sub> does not behave a	s an ideal gas at very hi	ah pressures	
Explain why N <sub>2</sub> does not behave a	s an ideal gas at very ni	gn pressures.	
			[2]
i) Complete the table by naming all	the types of intermolec		
	the types of intermolec		
i) Complete the table by naming all	the types of intermolection (g).	cular forces (van der W	
ii) Complete the table by naming <b>all</b> separate samples of N <sub>2</sub> (g) and CO	the types of intermolec		
ii) Complete the table by naming <b>all</b> separate samples of N <sub>2</sub> (g) and CO	the types of intermolectics: $N_2(g)$	cular forces (van der W	
ii) Complete the table by naming all	the types of intermolecticg). $N_2(g)$	CO(g)	
i) Complete the table by naming all separate samples of N <sub>2</sub> (g) and CO umber of electrons per molecule resence of a dipole moment	the types of intermolecting).  N <sub>2</sub> (g)  14	CO(g) 14	
complete the table by naming all separate samples of N <sub>2</sub> (g) and CO separate samples o	the types of intermolecting).  N <sub>2</sub> (g)  14	CO(g) 14	
complete the table by naming all separate samples of N <sub>2</sub> (g) and CO separate samples o	the types of intermolecting).  N <sub>2</sub> (g)  14	CO(g) 14	
i) Complete the table by naming all separate samples of N <sub>2</sub> (g) and CO umber of electrons per molecule resence of a dipole moment	the types of intermolecting).  N <sub>2</sub> (g)  14	CO(g) 14	Vaals') in
Complete the table by naming all separate samples of N <sub>2</sub> (g) and CO mber of electrons per molecule esence of a dipole moment biling point/°C	the types of intermolecting).  N <sub>2</sub> (g)  14	CO(g) 14	
Complete the table by naming all separate samples of N <sub>2</sub> (g) and CO separate samples o	the types of intermolecting).  N <sub>2</sub> (g)  14  X  -195.8	CO(g) 14 ✓ -191.5	Vaals') in
complete the table by naming all separate samples of N <sub>2</sub> (g) and CO sumber of electrons per molecule resence of a dipole moment biling point/°C termolecular forces (van der Waals')	the types of intermolecticg).  N <sub>2</sub> (g)  14  X  -195.8	CO(g) 14  -191.5  Die moment.	Vaals') in

[Total: 13]

11.	9701	$_{ m s20}$ $_{ m qp}$ $_{ m 23}$ $_{ m Q:}$ $_{ m 2}$	
(a)	Exp	plain what is meant by the term <i>relative isotopic mass</i> .	
			[2]
(b)		ample of copper contains two isotopes, $^{63}$ Cu and $^{65}$ Cu. The relative atomic mass of oper in this sample is 63.55.	the
	Cal	culate the percentage abundance of each of these isotopes. Show your working.	
		percentage abundance of <sup>63</sup> Cu =	. %
		percentage abundance of <sup>65</sup> Cu =	
			[2]
(c)	(i)	Name the type of bonding within a sample of solid copper.	
` ,	` '		[1]
	/::\		1.1
	(ii)	Draw a labelled diagram to show the bonding within a sample of solid copper.	
			[2]

1s<sup>2</sup> .......[1]

(iii) State the electronic configuration of a copper atom.

(d) A student is provided with a sample of hydrated copper(II) sulfate, CuSO<sub>4</sub>•xH<sub>2</sub>O, and is asked to determine the value of x.

The student dissolves a sample of the hydrated copper(II) sulfate in water and adds it to an excess of aqueous potassium iodide to make a total volume of 250.0 cm<sup>3</sup> of solution.

$$2CuSO_4 + 4KI \rightarrow 2CuI + I_2 + K_2SO_4$$

The amount of iodine produced during this reaction is found by titrating a sample of this solution with sodium thiosulfate solution.

25.0 cm³ of the iodine-containing solution requires 20.0 cm³ of 0.10 mol dm⁻³ sodium thiosulfate solution.

$$I_2 + 2S_2O_3^{2-} \rightarrow S_4O_6^{2-} + 2I^{-}$$

(i) Calculate the amount, in mol, of copper(II) sulfate present in the original sample of hydrated copper(II) sulfate.

Show your working.

amount of copper(II) sulfate = ..... mol [2]

(ii) A total of 7.98 g of CuSO<sub>4</sub> is present in 10.68 g of CuSO<sub>4</sub>•xH<sub>2</sub>O.

Complete each row of the table to calculate the value of x, where x is an integer.

[M<sub>c</sub>: CuSO<sub>4</sub>,159.6]

amount of CuSO <sub>4</sub> in 10.68 g of CuSO <sub>4</sub> •xH <sub>2</sub> O	mol
amount of H <sub>2</sub> O in 10.68 g of CuSO <sub>4</sub> •xH <sub>2</sub> O	mol
value of x	x =

[3]

[Total: 13]

12. 9701\_w17\_qp\_22 Q: 1

The elements sodium to sulfur react with chlorine. The melting points of some of the chlorides formed are shown.

chloride	NaC1	MgCl <sub>2</sub>	AlCl <sub>3</sub>	SiC1 <sub>4</sub>	$PCl_3$	SCl <sub>2</sub>
melting point/K	1074	987	463	203	161	195

(a) Predict the shapes of  $AlCl_3$  and  $PCl_3$ .

Draw diagrams to show the shapes, name the shapes and state the bond angles.

$AlCl_3$
shape
angle

$PCl_3$	
shape	
angle	

[4]

(b)	(i)	Explain, in terms of structure and bonding, why the melting point of $SiCl_4$ is much low than that of NaC $l$ .	vei
			,
			[3]
	(ii)	Explain why the melting point of $\mathrm{SiC}l_4$ is higher than that of $\mathrm{PC}l_3$ .	
			••••

[1]

[Total: 10]

# Chapter 5

# Chemical energetics

Structure and bonding can be used to explain many of the properties of substances.

(a) Copper, ice, silicon(IV) oxide, iodine and sodium chloride are all crystalline solids.

Complete the table with:

- the name of a type of bonding found in each crystalline solid,
- the type of lattice structure for each crystalline solid.

crystalline solid	type of bonding	type of lattice structure
copper		
ice		
silicon(IV) oxide		
iodine		
sodium chloride		

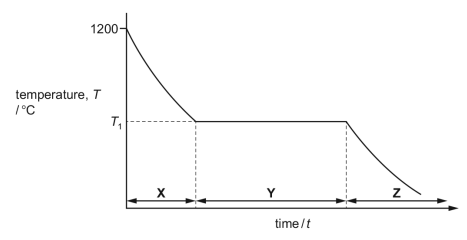
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п		١	ı

							_	
(b)	(i)	Name th	e stronge	est type (	of intermo	lecular	force	in ice

	-
יו	

(ii) Draw a fully labelled diagram of two water molecules in ice, showing the force in (i) and how it forms.

(c) The graph represents how the temperature of a sample of copper (melting point 1085°C) changes as it is gradually cooled from 1200°C.



(i) Identify the state(s) of matter present during each stage of the process shown in the graph.

X	
Υ	
Z	
	[2]

(ii) State what is happening to the energy and movement of the particles in the copper during stage X.

.....[2]

(iii) Explain why the temperature stays constant at  $T_1$  during stage  $\mathbf{Y}$ .

	101

.....[2]

[Total: 15]

# Appendix A

## Answers

### $1.\ 9701\_m22\_ms\_22\ Q\!:\, 1$

Question	Answer	Marks
(a)	$O(g) \rightarrow O^{+}(g) + e^{-}$	1
(b)(i)	increase across period <b>AND</b> increased nuclear attraction <b>for</b> (valence / outer) <b>electrons</b> [1] increase in (positive) nuclear charge / number of protons (in the nucleus) [1] similar shielding (of outer electrons) [1]	3
(b)(ii)	spin-pair repulsion (of electrons) in (2)p orbital [1] outweighs increased nuclear charge [1]	2
(c)	1s <sup>2</sup> 2s <sup>2</sup> 2p <sup>6</sup> 3s <sup>2</sup> 3p <sup>1</sup> [1] greatest jump between 3rd and 4th ionisations [1] indicates three electrons in outer shell [1]	3

## 2. 9701\_w20\_ms\_21 Q: 1

(a)	$Mg(g) \rightarrow Mg^{\dagger}(g) + e^{(-)}$	1
(b)	M1: distance between nucleus and outer e⁻ increases OR outer electron removed from higher energy shell	3
	M2: increased shielding	
	M3: decreased nuclear attraction	
(c)	M1: greater nuclear attraction	2
	M2: (2nd / 2s) electron being removed from smaller (ion)	

3.  $9701_{w20_{ms}_{2}}$  Q: 1

(a)(i)	positive / + on left AND negative / – on right  charge on plate  charge on plate	1
	source	
(a)(ii)	straight line vertically upwards from the source  tharge on plate  charge on plate  source	1
(b)(i)	type of orbital s p d	3
	number of orbitals 4 9 5	
	number of orbitals 4 3 3	
(b)(ii)	4s	2
(b)(iii)	5	1
(b)(iv)	Award one mark for each correct bullet point – max 3 marks  • nuclear charge increases  • extra electron(s) in inner shell / n=3 /d-subshell / d- orbital  • increased shielding (of 4s electrons by electrons in n=3 / 3 <sup>rd</sup> shell / 3d)  • (overall) similar nuclear attraction (for outer electron)	3
(c)	answer in terms of subatomic particles in the nucleus same (number of) protons AND different (number of) neutrons	1

## 4. 9701\_w22\_ms\_21 Q: 1

Question					Answer	Marks	
(a)(i)	columns 1	columns 1 & 3 identical					
	isotope	No of p's	No of n's	No of e's			
	<sup>69</sup> Ga	31	38	31			
	<sup>71</sup> Ga	31	40	31			
		•	√	√		1	
(a)(ii)	M1 (weight	M1 (weighted) average / mean mass of the isotopes / average mass of the atom(s) (of an element)					
	M2 compared to (the mass of) the unified atomic mass unit						
(a)(iii)	$69.723 = 68.926x + 70.925(1-x) : x = 0.6013$ $/69.723 = \frac{68.926x + 70.925(100 - x)}{100}$						
	60.13%					1	
(b)(i)	they have t	he same ele	ectron arran	gement / ele	tronic configuration	1	
(b)(ii)	1s <sup>2</sup> 2s <sup>2</sup> 2p <sup>6</sup>	3s <sup>2</sup> 3p <sup>6</sup> 4s <sup>1</sup>				1	
(b)(iii)	M2 second OR second (an OR	d third) elec	electron(s) is	s removed for moved from	om inner shell a shell closer to the nucleus clear attraction	2	

### $5.\ 9701\_s17\_ms\_22\ Q:\ 1$

(a)	atomic number	nucleon number	number of electrons	number of protons	number of neutrons	symbol		2
		6		3	3		1	
						<sup>58</sup> <sub>26</sub> Fe <sup>3+</sup>	1	
(b)(i)	EITHER mass of an atom / isotope relative / compared to 1/12 (the mass) of (an atom of) C-12 OR on a scale in which a C-12 (atom / isotope) has (a mass of exactly) 12 (units)  OR mass of one mol (of atoms) of an isotope relative / compared to 1/12 (the mass) of 1 mol of C-12 OR							2
(b)(ii)	on a scale in which o	`	sotope) has a mass	of (exactly) 12 g				
(~)(")	)(ii) $\frac{(10.0129 \times 19.78) + (80.22x)}{100} = 10.8$							
	x = 10.9941							1
						Total:		6

Question	Answer	Marks
(a)(i)	bonding in magnesium – metallic AND bonding in sodium chloride – ionic	1
(a)(ii)	bonds in NaCl are stronger than bonds in Mg	1
(a)(iii)	M1 S <sub>8</sub> / molecules of sulfur have more electrons (than P <sub>4</sub> / molecules of phosphorus) M2 S has stronger instantaneous dipole–induced dipole forces (than phosphorus / P)	2
(b)(i)	power of an atom to attract electrons to itself	1
(b)(ii)	(across a period)  increase in nuclear charge  similar shielding  (so) increase in nuclear attraction for bonding / outer / valence electrons  OR bonding / outer / valence electron(s) are more strongly attracted to nucleus  Two correct for one mark, three correct for two marks	2
(b)(iii)	hydrogen bond	1
(b)(iv)	M1 link shown as a dashed line between the lone pair of electrons from N of one NH <sub>3</sub> to one H on other NH <sub>3</sub> M2 minimum 3 correct partial charges (on adjacent atoms) over two NH <sub>3</sub> molecules EITHER <sup>5</sup> -N— <sup>5+</sup> H <sup>5-</sup> N OR <sup>5+</sup> H <sup>5-</sup> N— <sup>5+</sup> H	2
(b)(v)	M1 O is more electronegative than N M2 two H-bonds per water molecule : 1 per ammonia molecule.	2

## 7. 9701\_m23\_ms\_22 Q: 1

Question	Answer M.							
(a)(i)	power of an atom to attract electrons to itself							
(a)(ii)	O lower nuclear charge / lower proton number O has (one) fewer shell than S / less shielding greater attraction (for nucleus) in O	2						
(b)(i)	Li H —							
(b)(ii)	non-linear	1						
(c)(i)	$H(g) \rightarrow H^{\dagger}(g) + e^{-}$	1						
(c)(ii)	H (cannot undergo second ionisation because it only) has one electron / H <sup>+</sup> has no electron	1						
(c)(iii)	1s <sup>2</sup> 2s <sup>2</sup> 2p <sup>6</sup> 3s <sup>2</sup> 3p <sup>2</sup>	1						
(d)(i)	$SO_2 + H_2O \rightarrow H_2SO_3$	1						
(d)(ii)	$SO_2 + 2NaOH \rightarrow Na_2SO_3 + H_2O$	1						
(d)(iii)	$CO_2 + Mg(OH)_2 \rightarrow MgCO_3 + H_2O$							
(e)(i)	compound O=C=O O=S=O S=C=S S=C=O							
	overall dipole   moment   ✓							

Question		Answer					
(e)(ii)	conversion of units	103000 Pa	1				
	Use of $pV = (m/M_t)RT$	$M_{\rm r} = \frac{0.284 \times 8.31 \times 423}{103000 \times 127 \times 10^{-6}}$	1				
		$M_r$ = 76.3 <b>AND</b> compound = CS <sub>2</sub>	1				

## 8. 9701\_s19\_ms\_21 Q: 3

(a)	$Ar^{+}(g)  o Ar^{2+}(g) + e^{(-)} OR Ar^{+}(g) - e^{(-)}  o Ar^{2+}(g)$		1	
(b)	at x = 8, within range 13000–20000		1	
	at x = 9, within range 35000–45000		1	
(c)	OR S		1	
(d)(i)	M1 correct conversions of data to SI/consistent units $p = 404000$ ; $V = 20\times10^{-6}$ ; $T = 298$		1	
	M2 calculation of $n$ (= $pV/RT$ ) from M1 values $n = \frac{404000 \times 20 \times 10^{-6}}{8.31 \times 298} = 3.263 \times 10^{-3} \text{ mol of } Cl_2$			
	<b>M3</b> finding the mass of $Cl_2$ = 3.263 × 10 <sup>-3</sup> × 71.0 = 0.23 (g)			
(d)(ii)	Method 1 $ M1 = 3.263 \times 10^{-3} \times 2 $ Method 2 $ M1 = \frac{0.23}{71.0} \times 2 \text{ OR } 6.53 \times 10^{-3} $		1	
	M2 = $6.02 \times 10^{23} \times M1$ M2 = $6.02 \times 10^{23} \times M1$ = $3.93 \times 10^{21}$ atoms of Cl       = $3.90 \times 10^{21}$ atoms of Cl		1	
(d)(iii)	M1 size / volume of molecule / particle becomes significant / non-negligible OR IMFs become significant / non-negligible			
	M2 IMFs becomes significant / non-negligible / collisions are not elastic		1	

## 9. 9701\_s22\_ms\_23 Q: 1

Question	Answer			Ma	arks	
(a)	<ul> <li>energy required</li> <li>when one electron is removed</li> <li>from each atom in one mole of</li> <li>gaseous atoms</li> <li>two or three points for one mark, four points for two marks</li> </ul>				2	
(b)	Group VI / 16 AND large increase	(in IE) after 6th				1
(c)	M1 reference to spin pair repulsion in (3)p orbital (in S) OR due to repulsion of two electrons in a (3)p orbital (in S)  M2 outweighs increased nuclear charge (in S)					2
(d)	M1 similarity in electronic structure / shielding of Ale and Na both remove electron from (3)s1 / single electron in (3)s (sub-level / orbital)  OR Ale and Na have same electronic configuration  OR shielding (of outer electron) is the same  M2 greater nuclear charge / number of protons  Ale has greater nuclear charge  OR 13p compared to 11p					2
(e)	ato	mic no. nucleon	no. no.of neutrons	electronic arrangement	1	3
	copper –65 29	65	65 – 29 = 36	1s <sup>2</sup> 2s <sup>2</sup> 2p <sup>6</sup> 3s <sup>2</sup> 3p <sup>6</sup> 3d <sup>10</sup> 4s <sup>1</sup>		
	M1 29 AND 65					
	M2 nucleon no – atomic no ALLOW ecf from M1					
	M3 electronic arrangement					

Question	Answer	Marks
(f)(i)	$M_{\rm r} = 63.5$ (1.05 / 63.5) × 6.022 × 10 <sup>23</sup> = 9.958 × 10 <sup>21</sup> <b>OR</b> 9.96 × 10 <sup>21</sup>	1
(f)(ii)	M1 comment explaining high melting point of Cu many strong metallic bonds OR many strong (electrostatic) attractions between cations and delocalised electrons OR strong bonds in giant metallic structure.	2
	M2 comment explaining electrical conductivity of Cu delocalised electrons are free are to move through the structure (owtte)	

#### 10. 9701\_s21\_ms\_22 Q: 2

Question	Answer	Marks
(a)(i)	Li <sup>+</sup> AND S <sup>2-</sup>	1
(a)(ii)	M1 giant	1
	M2 (many) strong force(s) of attraction between oppositely charged ions OR (many) strong ionic bond(s)	1
(b)(i)	(covalent) bond with both electrons are provided from the same / one species OR shared pair (of electrons) are provided from the same species / one atom owtte	1
(b)(ii)	3 bonding pairs between C and O, 4 •'s AND 2×'s 1 lone pair on C, ××, AND 1 lone pair on O, ••.	2

Question	Answer		Mark		
(c)(i)	Any two assumptions about the behaviour of particles in an ideal gas from			2	
	(particles / molecules have mass but) negligible size / volume (compared to total volume of gas / container)				
	no / negligible forces / interactions (between particles / molecules)				
	collisions are elastic				
(c)(ii)	(c)(ii) M1 IMF become larger / more significant				
	M2 volume of molecules / particles becomes significant / no longer negligible			1	
(c)(iii)	N <sub>2</sub> (g)	CO(g)		2	
	instantaneous dipole-induced dipole 🗸	instantaneous dipole–induced dipole (and) permanent dipole–permanent dipole ✓			
(c)(iv)	O is more electronegative than C			1	

## 11. 9701\_s20\_ms\_23 Q: 2

(a)	EITHER			2	
	M1 mass of an atom / isoto	оре			
	M2 relative / compared to	1/12 (the mass) of (an atom of) C-12 OR			
	on a scale in which a C-12	(atom / isotope) has (a mass of exactly) 12 (	(units)		
	OR				
	M1 mass of one mol (of at	oms) of an isotope			
	,	1/12 (the mass) of 1 mol of C-12 OR			
	in which one mol C-12 (atom / isotope) has a mass of (exactly) 12 g				
(b)	% abundance of <sup>63</sup> Cu = 72			2	
	% abundance of <sup>65</sup> Cu = 27	$^{\prime}$ .5% ession AND correct calculation of $x$ for one is:	otope		
	% ab of $^{63}$ Cu = $x$ ( $x/100$	$\times$ 63) + ((1- x)/100 $\times$ 65) = 63.55 so x =72.5	5.600		
	OR % ab of $^{65}$ Cu = $x$ (1- $x$ )/100 × 63) + $x$ /100 × 65) = 63.55 so $x$ = 27.5				
	% ab of ${}^{\infty}Cu = x$ $(1-x)/100 \times 65) + x/100 \times 65) = 65.55$ so $x = 27.5$				
	M2 calculation of abundan	ce of other isotope by 100- x			
(c)(i)	metallic			1	
(c)(ii)	diagram showing the bond	ling in a sample of copper		3	
	$\odot$ $\odot$ $\odot$ $\odot$				
	$\bigcirc.\bigcirc.\bigcirc.\bigcirc.\bigcirc.\bigcirc.\bigcirc$				
	M1 diagram shows regular arrangement of spheres labelled as positively charged ions / +2 or +1 / cations M2 diagram shows surrounded by electrons and clearly labelled as 'delocalised electrons'				
		<u> </u>	ocalised electrons		
(c)(iii)	(1s²) 2s² 2p6 3s² 3p6 3d¹0 4s¹ <b>OR</b> (1s²) 2s² 2p6 3s² 3p6 4s¹ 3d¹0		1		
(d)(i)	M1 calculate the number n			2	
	20/1000 x 0.10 = $2x10^{-3}$ = 0.002 (mol $S_2O_3^{-2}$ .) <b>M2</b> calculate number mol CuSO <sub>4</sub> in 250cm <sup>3</sup>				
	(1mol S <sub>2</sub> O <sub>3</sub> <sup>2</sup> ·: 1 mol CuSO <sub>4</sub> ) = 0.002 mol CuSO <sub>4</sub> in 25cm <sup>3</sup>				
	so 0.02 mol CuSO <sub>4</sub> in 250cm <sup>3</sup>				
(d)(ii)	M1 amount of CuSO <sub>4</sub> in	7.98 / (159.6) = 0.05 (mol)	]	3	
	10.68 g of CuSO <sub>4</sub> ·xH <sub>2</sub> O	<u>5.55</u> (1151)			
	M2 amount of H₂O in	(10.68 – 7.98) / 18 = 2.7 / 18 =) 0.15 (mol)			
	10.68 g of CuSO <sub>4</sub> ·xH <sub>2</sub> O	, , , , , , , , , , , , , , , , , , , ,			
			1	l	

 $12.\ 9701\_w17\_ms\_22\ Q:\ 1$ 

(a)	3 marking points for each box: diagram, name and shape. for each box: all three correct = 2 marks two correct = 1 mark	4
(b)(i)	SiC L <sub>4</sub> simple / molecular AND Van der Waals' / id-id forces / London / dispersion forces / IMFs	1
	NaCl ionic OR giant	1
	bonding (in NaC <i>l</i> ) strong <u>er</u> (than forces in SiC <i>l</i> <sub>4</sub> ) owtte	1
(b)(ii)	SiCI <sub>4</sub> has more electrons ORA	1
	stronger Van der Waals' / id-id forces / London / dispersion forces / IMFs	1
(b)(iii)	:çi: :çi: :çi:	1

13. 9701\_s17\_ms\_21 Q: 2

(a)	substance	type of bonding	type of lattice structure	
	copper	metallic	giant/metallic	1
		1		
	ice	covalent OR hydrogen(-bonding) / H(-bonding)	hydrogen-bonded / simple / molecular	1
	silicon(IV) oxide	covalent	giant (molecular) / macromolecular	1
	iodine	covalent	simple / molecular	1
	sodium chloride	ionic	giant / ionic	1
(b)(i)	hydrogen bonding			1
(b)(ii)	H-bond between O and H of different molecules			1
	minimum three partial charges (in a row) over two $H_2O$ molecules, i.e.: either ${}^{\delta-}O$ — $H^{\delta^+}$ ${}^{\delta-}O$ or $H^{\delta^+}$ ${}^{\delta-}O$ — $H^{\delta^+}$			
	lone pair of electrons on O of H-bond,	in line with H-bond		1
(c)(i)	X = liquid AND Z = solid			1
	Y = liquid and solid OR 'liquid / solid' C	R 'liquid OR solid'		1
(c)(ii)	(kinetic) energy reducing			1
	motion slowing		owtte	1
(c)(iii)	energy given out / released forming bo	onds / forming bonds exothermic		1
	compensates for / counteracts heat los	ss / cooling	owtte	1
			Total:	15