

Boost your performance and confidence with these topic-based exam questions

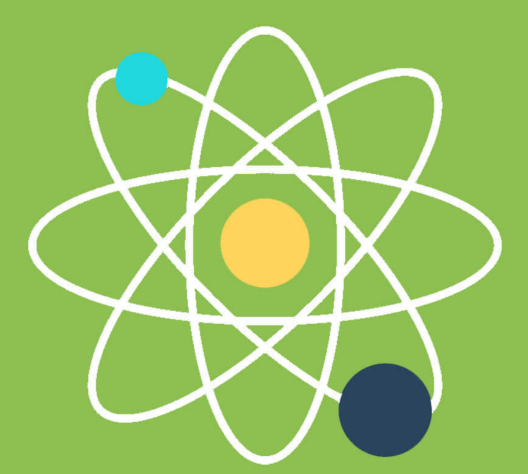
Practice questions created by actual examiners and assessment experts

Detailed mark schemes

Suitable for all boards

Designed to test your ability and thoroughly prepare you

3.1 The Periodic Table & Periodic Trends



IB Chemistry - Revision Notes

www.exampaperspractice.co.uk



3.1.1 The Periodic Table

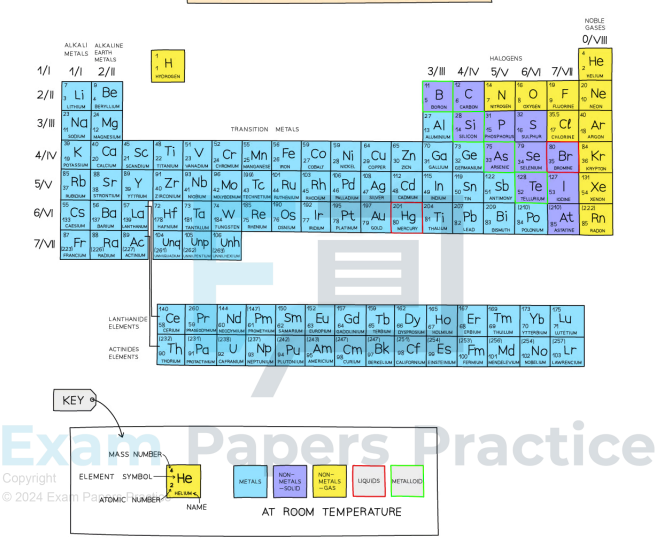
Structure of the Periodic Table

- The **periodic table** is a list of all known elements arranged in order of **increasing atomic number**, from 1 to 118.
- In addition to this, the elements are arranged in such a way that atoms with the same number of shells are placed together, and atoms with similar electronic configurations in the outer shell are also placed together. This is achieved as follows:
 - The elements are arranged in rows and columns.
 - Elements with **one** shell are placed in the **first row** (i.e. H and He)
 - Elements with **two** shells are placed in the **second row** (Lito Ne) and so on.
- A row of elements thus arranged is called a **period**. The period number, **n**, is the outer energy level that is occupied by electrons.
- In addition, the elements are aligned vertically (in columns) with other elements in different rows, if they share a similar outer-shell electronic configuration
 - The outer electrons are known as the **valence** electrons.
- A column of elements thus arranged is called a **group**

Exam Papers Practice

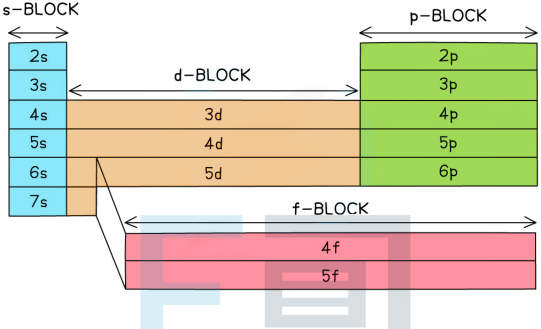


PERIODIC TABLE OF THE ELEMENTS



- Since the electronic configurations of **H** and **He** are unusual, they do not fit comfortably into any group. They are thus allocated a group based on similarities in physical and chemical properties with other members of the group
- He is placed in group 0 on this basis, but hydrogen does not behave like any other element and so is placed in a group of its own





The blocks of the periodic table

- All elements belong to one of four main blocks: the s-block, the p-block, the d-block and the fblock
 - The s-block elements are all those with only s electrons in the outer shell
 - The p-block elements are all those with at least one p-electron in the outer shell
 - The d-block elements are all those with at least one d-electron and at least one s-
 - electron but no for pelectrons in the outer shell (up to 5d)
 - The f-block elements are all those with at least one f-electron and at least one s-electron but no d or p electrons in the outer shell

Copyrig

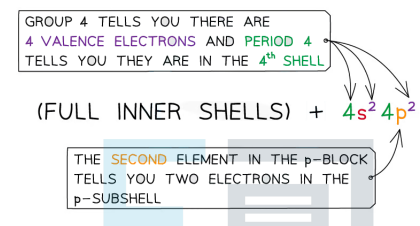
© 2024 The physical and chemical properties of elements in the periodic table show clear patterns related to the position of each element in the table

- Elements in the same group show similar properties, and properties change gradually as you go across a period
- As atomic number increases, the properties of the elements show trends which repeat themselves in each period of the periodic table
 - These trends are known as periodic trends and the study of these trends in known as periodicity



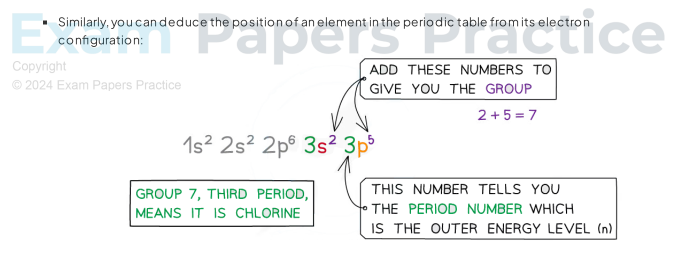
Deducing Electron Configurations

- The electron configuration of any element can be deduced from its position in the periodic table
- It is like an 'address' that tells you exactly where an element is found
- Using the element germanium as an example to illustrate how it works:



Deducing the electron configuration of germanium

- Germanium is in p block, in group 4 (using the simplify numbering system) and the second element across in period 4
- Group 4 tells you there are **four valence electrons** and period 4 tells you the **valence electrons** are in the **fourth shell**
- The **second** position in **p block** tell you that two electrons are in the **p subshell**



 $Deducing \ information \ from \ the \ electron \ configuration \ of \ chlorine$



• Test your understanding in the following example:

Worked example

Element Z is in period 4 and group 5 of the periodic table. Which statement is correct?

- A. Z has 5 occupied energy levels.
- **B.** Z can form ions with 3 charge.
- **C**. Z is a transition element.
- D. Z has 4 valence electrons.

Answer:

- The correct option is **B**. A group 5 element could form a 3 ion
- It cannot be **A** because 5 occupied energy levels would place it in period 5
- Transition elements would not be found in group 5, so C is also incorrect
- 4 valence electrons would match an element in group 4, so **D** must be wrong

Exam Papers Practice

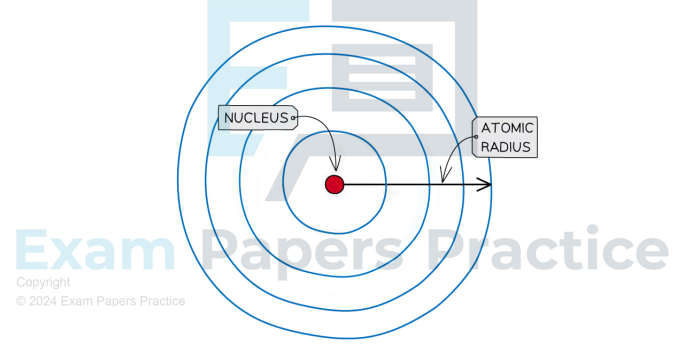


3.1.2 Periodic Trends: Physical – Atomic & Ionic Radius

Atomic & Ionic Radius

Atomic radius

- The atomic radius of an element is a measure of the size of an atom
- It is the distance between the **nucleus** of an atom and the outermost electron shell
- It can be quite hard to determine exactly where the boundary of an atom lies, so a variety of approches are taken such as half the mean distance between two adjacent atoms
- This will vary depending on the type of structure and bonding, but it gives a comparative value for atoms



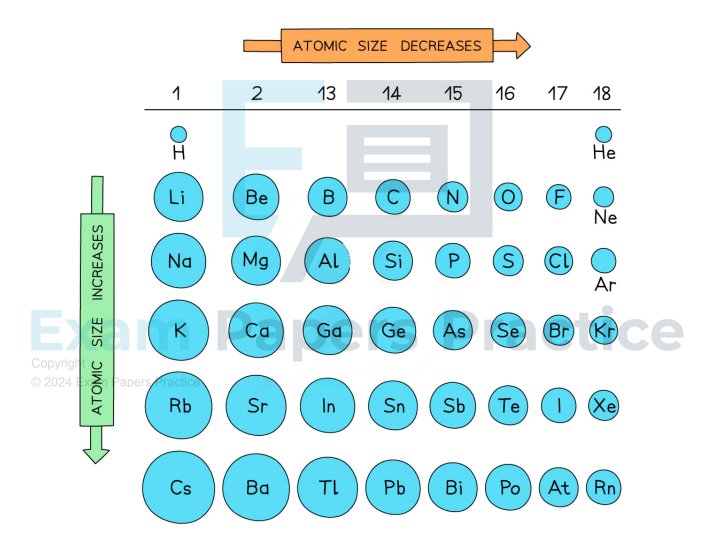
The atomic radius of an atom is the typical distance between the nucleus and the outermost electron shell

- Atomic radii show predictable patterns across the periodic table
 - They generally **decrease** across each period
 - They generally increase down each group
- These trends can be explained by the **electron shell theory**
 - Atomic radii decrease as you move across a period as the atomic number increases (increased positive **nuclear charge**) but at the same time extra electrons are added to the



same principal quantum shell

- The larger the nuclear charge, the greater the pull of the nuclei on the electrons which results in smaller atoms
- Atomic radii increase moving down a group as there is an increased number of shells going down the group
- The electrons in the **inner shells** repel the electrons in the **outermost shells**, **shielding** them from the positive nuclear charge
- This weakens the pull of the nuclei on the electrons resulting in larger atoms



Trends in the atomic radii across a period and down a group



- The diagram shows that the atomic radius increases sharply between the **noble gas** at the end of each period and the **alkali metal** at the beginning of the next period
- This is because the **alkali metals** at the beginning of the next period have **one** extra principal quantum shell
 - This increases shielding of the outermost electrons and therefore increases the atomic radius

lonic radius

- The ionic radius of an element is a measure of the size of an ion
- Ionic radii show predictable patterns
 - The trend down a group is the same as atomic radius it increases as the number of shells increases
 - The trend across a period is not so straightforward as it depends on whether it is positive or negative ions being considered
 - Ionic radii increase with increasing negative charge
 - Ionic radii decrease with increasing positive charge
- These trends can also be explained by the electron shell theory
 - lons with negative charges are formed by atoms accepting extra electrons while the nuclear charge remains the same
 - The extra electrons experience repulsion with the other valence electrons which increases the ionic radius
 - The greater the negative charge, the larger the ionic radius
 - Positively charged ions are formed by atoms losing electrons
 - The nuclear charge remains the same but there are now fewer electrons which undergo a greater electrostatic force of attraction to wards the nucleus which decreases the ionic

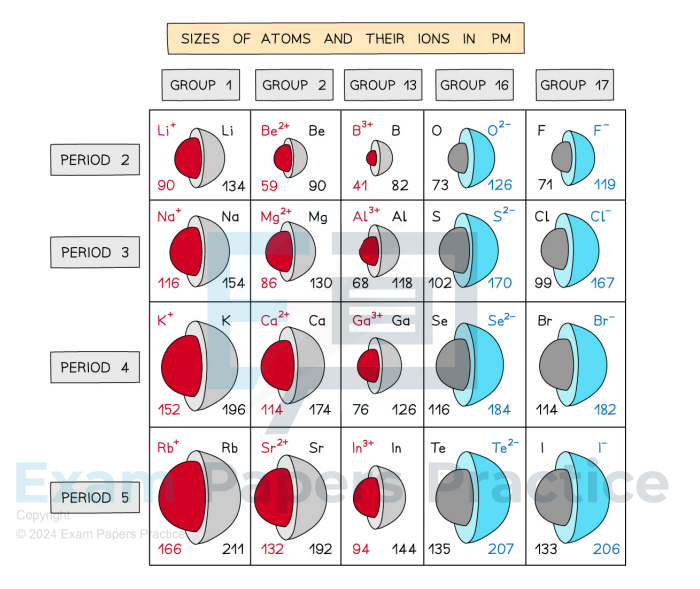




The greater the positive charge, the smaller the ionic radius

© 2024 Exam Papers Practice





Trends in the ionic radii across a period and down a group



Worked example

Which option shows atoms in order of decreasing atomic radius?

A. N > C > Be > Mg
B. Mg > N > C > Be
C. Be > C > N > Mg
D. Mg > Be > C > N

Answer:

Option **D** is the correct answer

- First, you need to identify that Be, C and N are all in period 2, but Mg is in period 3, so Mg will have the biggest radius.
- Secondly, atomic radius decreases across the period so Be, C and N decrease in that order as they belong to groups 2, 14 and 15, respectively

3.1.3 Periodic Trends: Physical – Ionisation Energy

First Ionisation Energy

The ionisation energy (IE) of an element is the amount of energy required to remove one mole of Copyrig electrons from one mole of atoms of an element in the gaseous state to form one mole of @ 2024 gaseous ions ractice

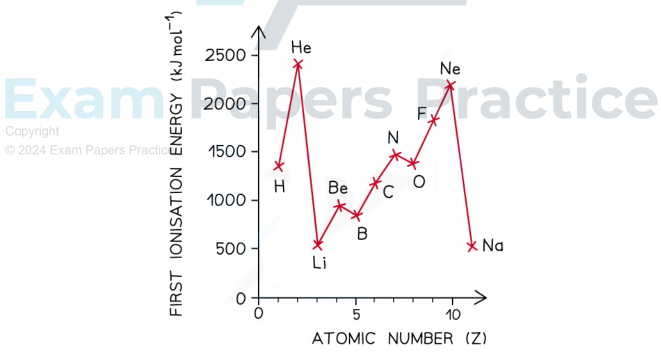
- Ionisation energies are measured under **standard conditions** which are 298 K and 100 kPa
- The units of IE are kilojoules per mole (kJ mol⁻¹)
- The **first ionisation energy** is the energy required to remove the one mole of electrons from one mole the gaseous atoms
 - E.g. the first ionisation energy of calcium:

 $Ca(g) \rightarrow Ca^+(g) + e^-$ lst $\Delta HI.E. = +590 \text{ kJ mol}^{-1}$



Ionisation Energies: Trends

- Ionisation energies show periodicity
- As could be expected from their electronic configuration, the group I metals show low IE whereas the noble gases have very high IEs
- The first ionisation energy increases across a period and decreases down a group and is caused by four factors that influence the ionisation energy:
 - Size of the nuclear charge: the nuclear charge increases with increasing atomic number, which means that there are greater attractive forces between the nucleus and outer electrons, so more energy is required to overcome these attractive forces when removing an electron
 - Distance of outer electrons from the nucleus: electrons in shells that are further away from the nucleus are less attracted to the nucleus so the further the outer electron shell is from the nucleus, the lower the ionisation energy
 - Shielding effect of inner electrons: the shielding effect is when the electrons in full inner shells repel electrons in outer shells preventing them to feel the full nuclear charge so the greater the shielding of outer electrons by inner electron shells, the lower the ionisation energy
 - **Spin-pair repulsion**: paired electrons in the same atomic orbital in a subshell repel each other more than electrons in different atomic orbitals; this makes it easier to remove an electron (which is why the first ionization energy is always the lowest)



A graph showing the ionisation energies of the elements hydrogen to sodium



lonisation energy across a period

- The ionisation energy across a period increases due to the following factors:
 - Across a period the **nuclear charge** increases
 - The distance between the nucleus and outer electron remains reasonably constant
 - The **shielding** by inner shell electrons remains the same
- There is a rapid **decrease** in ionisation energy between the **last** element in one period and the **first** element in the next period caused by:
 - The increased **distance** between the nucleus and the outer electrons
 - The increased **shielding** by inner electrons
 - These two factors outweigh the increased nuclear charge
- There is a slight **decrease** in 1st I.E. between **beryllium** and **boron** as the fifth electron in boron is in the 2p subshell which is further away from the nucleus than the 2s subshell of beryllium
 - Beryllium has a first ionisation energy of 900 kJ mol⁻¹ as its electron configuration is 1s² 2s²
 - Boron has a first ionisation energy of 801 kJ mol⁻¹ as its electron configuration is 1s² 2s² 2p¹
- There is a slight decrease in 1st I.E. between nitrogen and oxygen due to spin-pair repulsion in the 2p subshell of oxygen
 - Nitrogen has a first ionisation energy of 1402 kJ mol⁻¹ as its electron configuration is 1s² 2s²
 2p³
 - Oxygen has a first ionisation energy of 1314 kJ mol⁻¹ as its electron configuration is 1s² 2s²
 2p⁴

lonisation energy down a group

- Although going down a group the nuclear charge increases, the ionisation energy down a group decreases and it is due to the following factors:
- The distance between the nucleus and outer electron increases
- Copyright The shielding by inner shell electrons increases
- © 2024 Examine effective nuclear charge is decreasing as shielding increases

Ionisation Energy Trends across a Period & going down a Group Table



Across a Period: Ionisation Energy Increases	Down a Group: Ionisation Energy Decreases
Increase in nuclear charae	Increase in nuclear charge
Shell number is the same Distance of outer electron to nucleus is the same	Increase in shells Distance of outer electron to nucleus increases Shielding effect increases, therefore, the attraction of valence electrons to the nucleus decreases
Shielding remains reasonably constant	Increased shielding
Decreased atomic/ionic radius	Increased atomic/ionic radius
The outer electron is held more tightly to the nucleus so it gets harder to remove it	The outer electron is held more loosely to the nucleus so it gets easier to remove it

Successive ionisation energies of an element

Copyright The successive ionisation energies of an element increase as removing an electron from a © 2024 Exam Papers Practice positive ion is more difficult than from a neutral atom

• As more electrons are removed the attractive forces increase due to **decreasing shielding** and an increase in the proton to electron ratio

- The increase in ionisation energy, however, is not constant and is dependent on the atom's electronic configuration
- Taking calcium as an example:

Ionisation Energies of Calcium Table



Electronic Configuration	1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ²	1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ¹	1s ² 2s ² 2p ⁶ 3s ² 3p ⁶	1s ² 2s ² 2p ⁶ 3s ² 3p ⁵
IE	First	Second	Third	Fourth
IE (kJ mol ⁻¹)	590	1150	4940	6480

- The values become very large and difficult to represent meaningfully, so it is more convenient to show the logarithm of the ionisation energies
- 1st SHELL 6 ENERGY (kJ mol⁻¹) 2nd 5.5 SHELL 5 2s 3rd 2p SHELL 4.5 BIG JUMPS INDICATE LOG IONISATION A CHANGE OF SHELL 4 MINI JUMPS SHOW 3.5 4^{th} CHANGE OF SUBSHELL SHELL 3 2.5 2 10 11 3 4 5 6 7 8 9 12 13 14 15 16 17 18 19 20 1 NUMBER OF ELECTRONS REMOVED
- This helps us to see significant jumps in I.E.

Successive ionisation energies for the element calcium

• The **first** electron removed has a low ionisation energy as it is easily removed from the atom due to the spin-pair repulsion of the electrons in the 4s orbital



- The **second** electron is a little more difficult to remove than the first electron as you are removing an electron from a positively charged ion
- The **third** electron is much more difficult to remove than the second one corresponding to the fact that the third electron is in a **principal quantum** shell which is closer to the nucleus (3p)
- The graph shows there is a large increase in successive ionisation energy as the electrons are being removed from an increasingly positive ion
- The big jumps on the graph show the change of shell and the small jumps are the change of subshell

💽 Exam Tip

Be careful with how you interpret successive ionisation energy graphs as it is common for students to read them the wrong way around and count outer electrons from right to left instead of left to right so they get the jumps in the wrong place. This happens particularly when you are given only a partial successive ionisation energy graph and have to deduce which group the element comes from. It's a good idea if you see an ionisation energy graph in an exam question to label the shells and subshells so you are less likely to make this mistake!

Exam Papers Practice

© 2024 Exam Papers Practice



3.1.4 Periodic Trends: Physical – Electron Affinity

Electron Affinity

- When atoms gain electrons they become negative ions or **anions**
- Electron affinity (EA) can be thought of as the opposite process of ionisation energy and is defined as
 - The amount of energy released when **one mole** of electrons is gained by **one mole** of atoms of an element in the gaseous state to form **one mole** of gaseous ions
- Electron affinities are measured under standard conditions which are 298 K and 100 kPa
- The units of EA are kilojoules per mole (kJ mol⁻¹)
- The first electron affinity is always exothermic
 - E.g. the first electron affinity of chlorine is:

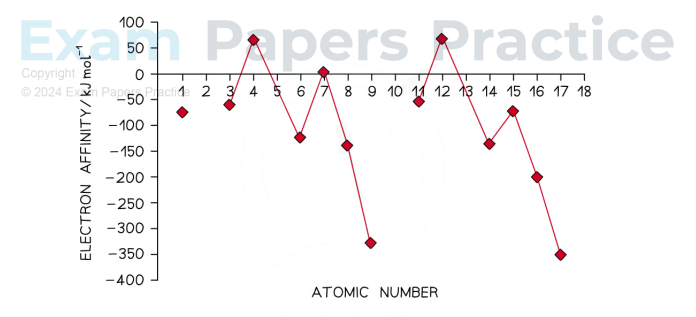
$$CI(g) + e^{-} \rightarrow CI^{-}(g)$$
 $\Delta H = -349 \text{ kJ mol}^{-1}$

However, the second electron affinity can be an endothermic process

$$O^{-}(g) + e^{-} \rightarrow O^{2-}(g)$$
 $\Delta H = +753 \text{ kJ mol}^{-1}$

• This is due to the fact that you are overcoming repulsion between the electron and a negative ion, so energy is required making the process endothermic overall

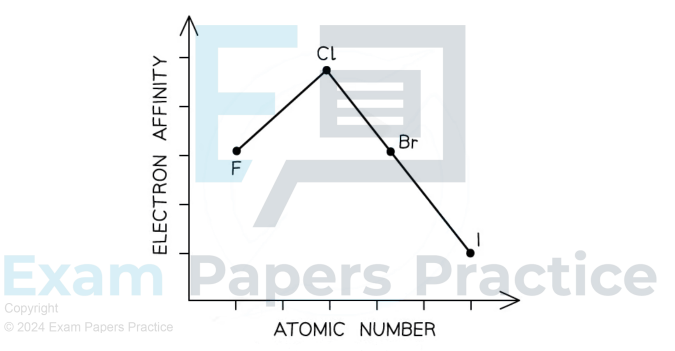
Trends in electron affinity





Electron affinities across a period

- Electron affinities show periodicity
- The pattern is very similar to ionisation energies, except that it is inverted and the minimum points are displaced one element to the right
- As might be expected, the most exothermic electron affinities are for group 17 elements which also have the highest electron egativities
- The strongest pull on electrons correlates with the greater amount of energy released when negative ions are formed
- Noble gases do not form negative ions, so they don't appear in this chart
- The electron affinities reach a peak for group 2 and group 5 elements



Electron affinities down a group

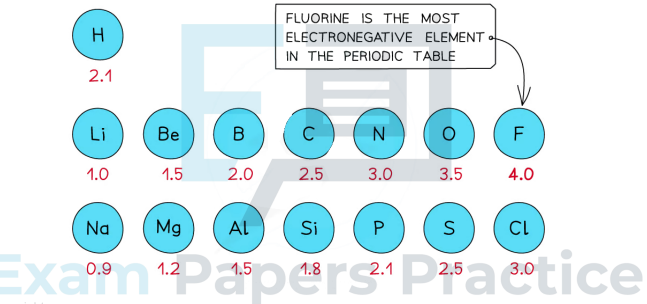
- Electron affinities generally decrease down a group
- As the atoms become larger the attraction for an additional electron is less, since the **effective nuclear charge** is reduced due to increased **shielding**
- Electron affinity become less exothermic going down the group
- An exception to this is fluorine whose **electron affinity** is smaller than expected
- This is because fluorine is such a small atom and an additional electron in the 2p subshell experiences considerable repulsion with the other valence electrons



3.1.5 Periodic Trends: Physical - Electronegativity

Electronegativity: Definition

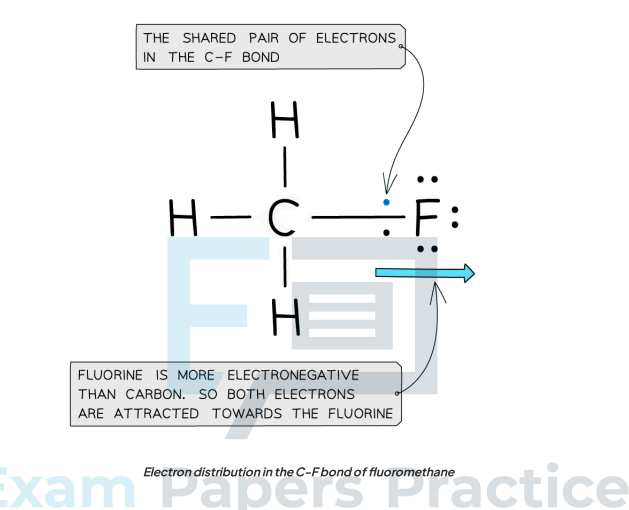
- Electronegativity is the ability of an atom to attract a pair of electrons towards itself in a covalent bond
- This phenomenon arises from the positive nucleus's ability to attract the negatively charged electrons, in the outer shells, towards itself
- The Pauling scale is used to assign a value of electronegativity for each atom



© 2024 Exam Papers First three rows of the periodic table showing electronegativity values

- Fluorine is the most electronegative atom on the Periodic Table, with a value of 4.0 on the **Pauling** Scale
- It is best at attracting electron density towards itself when covalently bonded to another atom





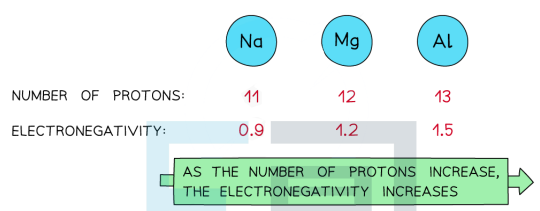
Copyright © 2024 Exam Papers Practice



Electronegativity: Affecting Factors

Nuclear charge

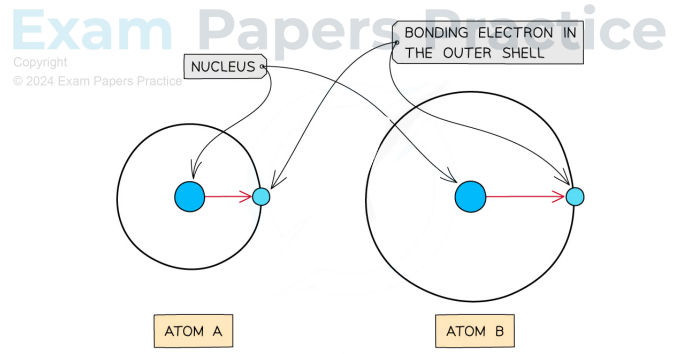
- Attraction exists between the positively charged protons in the nucleus and negatively charged electrons found in the energy levels of an atom
- An increase in the number of protons leads to an increase in nuclear attraction for the electrons in the outer shells
- Therefore, an increased nuclear charge results in an increased electronegativity



As the nuclear charge increases, the electronegativity of an element increases as well

Atomic radius

- The atomic radius is the distance between the nucleus and electrons in the outermost shell
- Electrons closer to the nucleus are more strongly attracted towards its positive nucleus
- Those electrons further away from the nucleus are less strongly attracted towards the nucleus
- Therefore, an increased atomic radius results in a decreased electronegativity



As the atomic radius increases, the nucleus has less of an attraction for the bonding electrons causing atom A to have a higher electronegativity than atom B

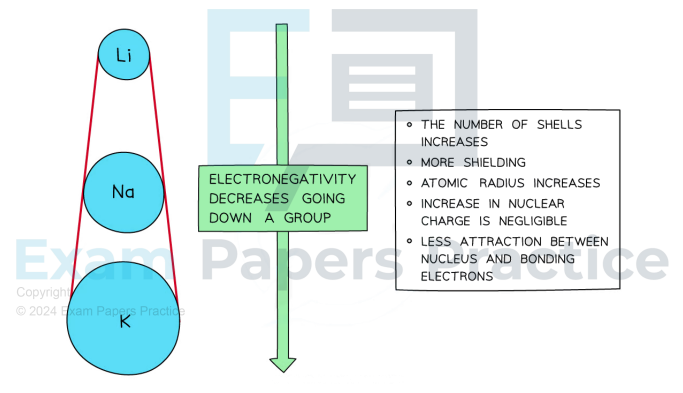


Electronegativity: Trends

• Electronegativity varies across periods and down the groups of the periodic table

Down a group

- There is a **decrease** in **electronegativity** going down the group
- The nuclear charge increases as more protons are being added to the nucleus
- However, each element has an extra filled electron shell, which increases shielding
- The addition of the extra shells increases the distance between the nucleus and the outer electrons resulting in **larger atomic radii**
- Overall, there is decrease in attraction between the nucleus and outer bonding electrons
- We say the **effective nuclear charge** has decreased down the group



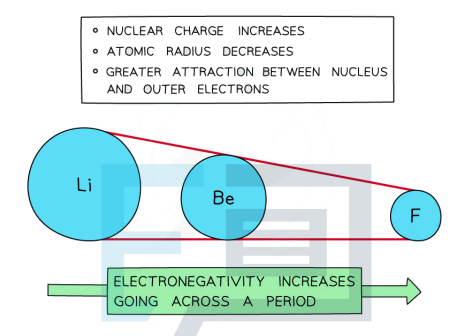
Electronegativity decreases going down the groups of the periodic table

Across a period

- Electronegativity increases across a period
- The nuclear charge increases with the addition of protons to the nucleus
- Shielding remains the same across the period as no new shells are being added to the atoms



- The nucleus has an increasingly strong attraction for the bonding pair of electrons of atoms across the period
- This results in **smaller atomic radii**



Electronegativity increases going across the periods of the periodic table

Trends down a Group & across a Period Table

EX	am Da	apers	Pract	ice
Copyright © 2024 Exan	Papers Practice	Down a Group	Across a Period	
	Nuclear charge	Increases	Increases	
	Shielding	Increases	Reasonably constant	
	Atomic radius	Increases	Decreases	
	Electronegativity	Decreases	Increases	



😧 Exam Tip

Make sure you learn the definition of **electronegativity** and can distinguish it from **electron affinity** as the two are often confused. **Electronegativity** is about chemical character and only applies to considerations of covalent bonds whereas **electron affinity** is a thermodynamic value that is measurable and applies to the formation of negative ions. You may come across something called **electropositivity** - this is a term used to describe the character of elements to form positive ions and is useful when talking about metal atoms and metal ions

Exam Papers Practice

© 2024 Exam Papers Practice



3.1.6 Periodic Trends: Chemical

Metallic & Non-metallic

 Periodic trends can also be seen in the typical properties of metals and non-metals which are summarised below:

Property	Metals	Non-metals
Electron arrangement	1–3 (more in Periods 5 & 6) outer shell electrons	4–7 electrons in the outer shell
Bonding	Metallic due to loss of outer shell electrons	Covalent by sharing of outer shell electrons
Electrical conductivity	Good conductors of electricity	Poor conductors of electricity
Type of oxide	Basic oxides (a few are amphoteric)	Acidic oxides (some are neutral)
Reaction with acids	Many react with acids	Do not react with acids
Physical characteristics	Malleable, can be bent and shaped High melting and boiling point	Flaky, brittle Low melting and boiling point

Copyright © 2024 Exam Papers Practice

The typical properties of metals and non-metals

- The typically properties of metals and non-metals can be explained by reference to their trends in atomic radius, ionic radius, ionisation energy, electron affinity and electron egativity
- The low **ionisation energies** and low **electronegativities** of metals can account for the ability of their **valence electrons** to move away from the nucleus
 - This is known as '**delocalisation**' of the electrons
- These properties increase from left to right as you transition from metal to metalloid to nonmetal
- The high **electronegativity** and **electron affinity** of non-metals can be related to their tendency to share electrons and form covalent bonds, either with themselves or other non-metal elements



• The similarities in **electronegativities** of the diagonal band of **metalloids** which divides the metals from the non-metals explains the behaviour of **metalloids**

Test yourself on the trends in the periodic table:

🖉 Worked exa	🖉 Worked example		
Which increase across a period from left to right?			
	А	lonic radius	Electronegativity
	В	Atomic radius	lonic radius
	С	1st ionization energy	Atomic radius
	D	1st ionization energy	Electronegativity

Answer:

The correct option is **D**

 Across a period atomic radius decreases, electronegativity and ionisation energy increase and ionic radius increases then decreases
 Comparison of the provide structure of the providest term of the provide structure of the provide structure of th

