COURSE TITLE: INTRODUCTORY INORGANIC CHEMISTRY
## MAIN COURSE

### CONTENTS

<table>
<thead>
<tr>
<th>Module 1</th>
<th>..................................................</th>
<th>1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit 1</td>
<td>The Periodic Table..........................</td>
<td>1</td>
</tr>
<tr>
<td>Unit 2</td>
<td>Modern Periodic Law..........................</td>
<td>9</td>
</tr>
<tr>
<td>Unit 3</td>
<td>Electronic Configuration.......................</td>
<td>16</td>
</tr>
<tr>
<td>Unit 4</td>
<td>Atomic Radii....................................</td>
<td>27</td>
</tr>
<tr>
<td>Unit 5</td>
<td>Ionisation Energy...............................</td>
<td>40</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Module 2</th>
<th>..................................................</th>
<th>46</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit 1</td>
<td>Electron Affinity............................</td>
<td>46</td>
</tr>
<tr>
<td>Unit 2</td>
<td>Electronegativity............................</td>
<td>51</td>
</tr>
<tr>
<td>Unit 3</td>
<td>Hydrogen........................................</td>
<td>57</td>
</tr>
<tr>
<td>Unit 4</td>
<td>Manufacture of Hydrogen........................</td>
<td>65</td>
</tr>
<tr>
<td>Unit 5</td>
<td>Ionic or Salt - Like Hydrides.....................</td>
<td>75</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Module 3</th>
<th>..................................................</th>
<th>80</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit 1</td>
<td>Hydrogen Bonding..............................</td>
<td>80</td>
</tr>
<tr>
<td>Unit 2</td>
<td>General Physical and Chemical Characteristics of the Alkali Metals...............</td>
<td>87</td>
</tr>
<tr>
<td>Unit 3</td>
<td>Compounds Alkali Metals.....................</td>
<td>98</td>
</tr>
<tr>
<td>Unit 4</td>
<td>Solvation of Alkali Metal Ions................</td>
<td>105</td>
</tr>
<tr>
<td>Unit 5</td>
<td>Alkaline Earth Metals........................</td>
<td>111</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Module 4</th>
<th>..................................................</th>
<th>118</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unit 1</td>
<td>Reactivity of Alkaline Earth Metals...........................................</td>
<td>118</td>
</tr>
<tr>
<td>Unit 2</td>
<td>Complexing Behaviour of Alkaline Earth Metals.....................................</td>
<td>126</td>
</tr>
</tbody>
</table>
1.0 INTRODUCTION
Scientists, from the very beginning have attempted to systematise the knowledge they gain through their observations and experiments. Development of the periodic law and the periodic table of the elements is one of such attempt. This has brought order in the study of the vast chemistry of more than a hundred elements known now.

It is therefore quite natural that you should begin your study of inorganic chemistry with the study of the periodic table. In this unit, you will be starting from the very beginning, that is, with the very first attempt made at classification of the elements.

By the mid-19th century, more than 60 elements were known and many more were being discovered. The rate of discovery of the new elements was so fast that the chemists started wondering "where it would all lead to"? Has nature provided a limit to the number of elements? And if so, how would one know about it? During this period, it was also realised that certain groups of elements exhibited similar physical and chemical
properties. Was it a mere coincidence or did a relationship exist among the properties of the elements? Attempts to reply such probing questions ultimately resulted in the development of the periodic table.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- list accurately at least two scientists who attempted to classify the elements into periods
- write with at least 70% accuracy, brief accounts of the attempts made by the two scientists and the result of these attempts
- state Mendeleev's periodic law
- state the property used by Mendeleev to classify the elements in his periodic table
- demonstrate an understanding of Mendeleev's law by applying it to predict properties of undiscovered elements.

3.0 MAIN CONTENT

3.1 The Beginning of Classification

One of the earliest attempts to classify elements was to divide them into metals and non metals. Metallic elements we all know have certain properties which include:

- having lustrous shining appearance, such as iron
- malleability, meaning they can be beaten into thin sheets such as is done when buckets are being produced
- metallic elements can also be drawn into wire such as is done when making electric wire. This property is known as ductility
- they can also conduct heat and electricity. If you hold a piece of metal in your hand and put one end into fire or in contact with any hot object, your hand will feel the heat as it travel from the point of contact with heat through the metal to your hand. Similarly if you hold one end of the metal through which an electric current is passed, you will be jolted by the current which travel through the metal to your hand.
- metallic elements also form basic oxides.

In contrast to metallic elements, non metallic elements have no characteristic appearance. They are brittle that is they break easily. They are poor conductors of electricity and heat. They form acidic oxides.
As more elements were discovered and knowledge of physical and chemical properties was refined, it became clear that within these two divisions of elements, there existed families of elements whose properties varied systematically from each other. Furthermore, certain elements, the metalloids possessed properties intermediate between the two divisions. Therefore, attempts were made to search for other classifications.

### 3.2 Attempts Made by J W Dobereiner

In 1829, J W Dobereiner observed that there exist certain groups of three elements which he called TRIADS. He also observed that elements in triad not only had similar properties, but also the atomic weight of the middle element was approximately an average of the atomic weights of the other two elements of the triad.

A few examples cited by him were: Li, Na, K, Ca, Sr, Ba, S, Se, Te and Cl, Br, I Although, Dobereineer's relationship seems to work only for a few elements, He was the first to point out a systematic relationship among the elements.

### 3.3 Attempts Made by A. de Chancourtois

In 1862, A. de Chanourtois arranged the elements that were known at that time in order of increasing atomic weight on a line which spiralled around a cylinder from bottom to top.

### 3.4 Attempts Made By John Newlands

In 1864, John Newlands, an English Chemist reported his "Law of Octaves". He suggested that if the elements were arranged in order of increasing atomic weight, every eighth element would have properties similar to the first element. For example, he arranged the elements in the following manner.

**Table 1.1: Arrangement of Elements according to John Newlands**
Thus, we see K resembles Na and Li, Ca resembles Mg and Be, Al resembles B, Si resembles C and so on. He called it the "Law of octaves" because he says the cycle of repetition shown by the elements is like that shown by octaves of music where every eight note resembles the first in octaves of music.

Newlands "Law of octaves" was rejected for two reasons. Firstly, it did not hold good for elements heavier than Ca. Secondly, he believed that there existed some mystical connection between music and chemistry.

3.5 The Work of Lothar Meyer

Lothar Meyer and Dmitri Mendeleev whom you will read about next played key role in the development of the periodic law as it is known today.

In 1869, Lothar Meyer reported that when physical properties like atomic volume, boiling point etc. were plotted against atomic weight, a periodically repeating curve was obtained in each case. Figure 1.1 is a graph showing the variation in atomic volume with atomic number. (Lothar Meyer also obtained semi curve by plotting atomic volume versus atomic weight)

The atomic volume behaviour is periodic. It goes through circles, dropping from a sharp maximum to a minimum and then sharply rising again. Each of the cycles is called a period. The location of element on the peak or in the troughs has an important correlation with their chemical reactivity. The elements of the peaks (example alkali metals) are the most reactive. Those in the troughs (example noble metals) are characteristically less reactive.

Fig. 1.1 Periodic Dependence of Atomic Volume on Atomic Number
3.6 Mendeleev's Periodic Law

In contrast to Lothar Meyer, Mendeleev used chemical properties like valence and formulae of hydrides, chloride, and oxides of the elements to illustrate his periodic law.

According to Mendeleev's periodic law, if the elements are arranged sequentially in the order of increasing atomic weight, a periodic repetition, that is, periodicity in properties is observed.

Mendeleev arranged elements in horizontal rows and vertical columns in order of increasing atomic weight so that the elements having similar properties were kept in the same vertical column.

![Mendeleev Periodic Table of 1871 against Each Element is the Value of Atomic Weight](image)

Though Newlands and Lothar Meyer also contributed in developing the periodic laws, the main credit goes to Mendeleev because of the following reasons:

- He included along with his table, a detailed analysis of the properties of all known elements and correlated a broad range of physical and chemical properties with atomic weights.
- He kept his primary goal of arranging similar elements in the same group quite clear. Therefore he was bold enough in reversing the order of certain elements. For example, iodine with lower atomic weight than that of tellurium (group VI) was placed in group VII along with fluorine, chlorine and bromine because of similarities in properties.
He also corrected the atomic weight of certain elements to include them in proper groups. For example, he corrected the atomic weight of beryllium (from 13.5 to 9) and indium (from 76 to 114) without doing any actual measurement. His competence was proved correct as Be and la with equivalent weight of 4.5 and 38 respectively are actually bivalent and trivalent.

Keeping to his primary goal of arranging similar elements in the same vertical column (group), he realised that some of the elements were still undiscovered and therefore left their places vacant in the table and predicted their properties. He predicted the existence in nature of over ten new elements and predicted properties of three of them, example eka-boron (scandium), eka aluminium (gallium) and eka silicon (germanium) from the properties of known elements surrounding them. When these elements were eventually discovered, Mendeleev prediction proved to be amazingly accurate.

This you can see by comparing the predicted and observed properties of eka-aluminium (gallium) and eka-silicon (germanium) given in Table 1.2. The validity of Mendeleev periodic law was dramatically and conclusively proven by the discovery of three out of the more than ten elements predicted by Mendeleev. The first to be discovered was eka-aluminium which was discovered by Lecoq de Boisbaudran in 1875.

**Table 1.2: Comparison of Predicted and Observed Properties of eka-aluminium (gallium) and eka silicon (germanium)**

<table>
<thead>
<tr>
<th>Property</th>
<th>Predicted by Mendeleev for eka-aluminium</th>
<th>Predicted by Mendeleev for eka silicon</th>
<th>Observed for gallium</th>
<th>Observed for germanium</th>
</tr>
</thead>
<tbody>
<tr>
<td>At weight</td>
<td>68</td>
<td>72</td>
<td>69.72</td>
<td>72.59</td>
</tr>
<tr>
<td>Density (kgm⁻³)</td>
<td>6.0 x 10¹ Low</td>
<td>5.9 x 10¹</td>
<td>302.8</td>
<td>5.3 x 10¹ 1220k</td>
</tr>
<tr>
<td>Melting point/k</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reaction with acids &amp; alkalines</td>
<td>Slow</td>
<td>Slow</td>
<td>Slow</td>
<td>Reacts with concentrated acids &amp; alkaline</td>
</tr>
<tr>
<td>Formula of oxide</td>
<td>E₂O₃</td>
<td>Ga₂O₃</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Density of oxide (kgm⁻³)</td>
<td>5.5 x 10¹</td>
<td>5.8 x 10¹</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Formula for chloride</td>
<td>ECl₃</td>
<td>GaCl₃</td>
<td>E₅Cl₄</td>
<td>GeCl₄</td>
</tr>
<tr>
<td>Boiling point of chloride (k)</td>
<td>Volatile</td>
<td>474</td>
<td>373</td>
<td>357</td>
</tr>
</tbody>
</table>

Lecoq de Boisbaudran called the element gallium and said its density was 4.7x10³. Mendeleev on hearing this wrote to Lecoq de Boisbaudran telling him that everything he said about the new element was correct...
except its density. On further position of the metal, Lecoq de Biosbaudran discovered that Mendeleev was right that the density of gallium was $5.8 \times 10^3$ kg just like it had been predicted by Mendeleev (Table 1.2).

Further proof of the law came via the works of Lars Fredrick Nilson who discovered scandium and Winkler who discovered germanium. Both elements were found to have properties corresponding to those of earlier predicted for them by Mendeleev.

The development of the periodic law is an excellent example where careful observation, critical analysis of available data without any preconceived notions and scientific foresight led to the discovery of a fundamental law of nature. Thus, when Mendeleev arranged elements in order of increasing atomic weights, he critically analysed the properties of the then known elements. He discovered that the properties of any element are an average of the properties of its neighbours in the periodic table. On this basis, he predicted the properties of undiscovered elements representing the gaps in his table.

4.0 CONCLUSION

In conclusion, the scientists’ tradition of recording and systemising knowledge gained through observations and experiments has enabled us to learn about the fundamental laws governing the arrangement of elements.

5.0 SUMMARY

In summary, you have learnt the following in this unit:

- scientists have always tried to systemise the knowledge they gain
- the effort to reveal the secrets of the periodic table were led by the following scientists J. W. Dobreiner, A de Chanourtois, John Newlands, Lothar Meyer and Dmitri Mendeleev
- the works of Dmitri Mendeleev formed the basis of the modern periodic law
- according to Mendeleev, the properties of any element are an average of the properties of its neighbours in the periodic table.

SELF-ASSESSMENT EXERCISE

Give the names of the scientist whose work contributed to the development of the periodic table?
6.0 TUTOR-MARKED ASSIGNMENT

1. What property did Mendeleev use to classify the element in his periodic table?
2. Enumerate four defects in the ‘Mendeleev’s periodic table.
3. Assuming that the element Ca had not been discovered, predict using the properties of the known element surrounding Ca its own properties such as its atomic weight and density.

7.0 REFERENCE/FURTHER READING

UNIT 2 MODERN PERIODIC LAW

CONTENTS

1.0 Introduction
2.0 Objectives
3.0 Main Content
   3.1 Modern Periodic Law
   3.2 The Long Form of the Periodic Table
   3.3 Nomenclature of Element having \( Z > 100 \)
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 Reference/Further Reading

1.0 INTRODUCTION

In Unit 1, you learnt about efforts made by several scientists to systematize the knowledge they gained through their observations and experiments. The effort of these scientists resulted in the formation of the periodic table and the periodic law. You learned about the works of such great pioneers as J. N Dobereiner, A de Chancourtois, John Newlands Lothar Meyer and Dmitri Mendeleev.

The periodic table of today has many similarities with that formed by Mendeleev, but differs from the Mendeleev table in some significant ways which we shall see in the next unit. Also in the past, element were named by their discoverers. In some cases such a practice has led to disputes between scientists who have discovered the same elements working independently in different parts of the world.

This has prompted the International Union of Pure and Applied Chemists (IUPAC) to device a method for naming newly discovered elements.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- state the modern periodic law
- explain the relative positions of K and Ar, Co and Ni and Te and I on the periodic table
- state the relationship between the atomic number and the periodic classification of elements
- apply IUPAC nomenclature rules in naming new elements having \( Z > 100 \).
3.0 MAIN CONTENT

3.1 Modern Periodic Law

In the previous section, you studied how D. Mendeleev classified elements and formed his periodic table. You must have noticed that there were anomalies in Mendeleev's original periodic table. There was for example, no place for lanthanides and actinides and in some instances; elements of higher atomic weight were placed before those of lower atomic weights example Co before Ni and Te before I. He could not predict the existence of noble gases, nor could he properly place hydrogen.

Between 1869 -1907, Mendeleev tried to improve his table. However, the most significant improvement of his periodic table came through the discovery of the concept of atomic number in 1913 by Henry Moseley, who suggested that the atomic number of an element is a more fundamental property than its atomic weight. Mendeleev's periodic law was therefore accordingly modified. This is now known as the Modern Periodic Law which states that: "the properties of elements are periodic functions of their atomic numbers".

Arrangement of the elements in order of their increasing atomic number removes most of the anomalies of Mendeleev's periodic table. The positions of K and Ar, Co and Ni, Te and I do not remain anomalous any longer since atomic number not atomic weight is used in arranging the elements.

As isotopes of an element have the same atomic number, they can all be placed at one and the same place in the periodic table. We know that the atomic number cannot be fractional. It increases by the integer from one element to the next. It has thus placed a limit on the number of elements. Today, 109 elements (from 1 to 109) have been discovered and any more elements that may be discovered in future will be beyond 109.
Fig. 2.1 The Modern Periodic Table in the Form Devised by Mendeleev

3.2 Long Form of the Periodic Table

You have now seen that in the modern form of Mendeleev periodic table, elements are arranged in seven horizontal rows and eight vertical columns. Normal and transition elements belonging to A and B subgroup of a group were placed in one and the same column of the table.

For example Sc and Ga both in group III A and B, Ti and Ge both in group IVa and b. In the long form of the periodic table (Figure 2.2) elements are arranged in eighteen vertical column by keeping the elements belonging to A and B subgroups in separate columns. Note that in the new arrangement, Sc is in group III B whereas Ga is now placed in group IIIA.
You would have also noticed that the group VIII B of Mendeleev's periodic table contains three triads Fe, Co, Ni, (4th period), Ru, Rh, Pd (5th period) and Os, Ir, Pr (15'th period). In the long form of the table, each element of the triad is kept in a separate column. So the group VIIIB occupies three columns of the table. You can see therefore that, the long form of periodic table is an extension of the modern periodic table.

Fig. 2.2 Long Form of the Periodic Table

Originally, Mendeleev gave A and B designation to the groups, containing normal and transition elements, respectively. However in his periodic table, this division into A and B group is often done arbitrarily. In different books for the elements of III to VIII groups, this designation of A and B groups is often reversed. To avoid this controversy, International Union of Pure and Applied Chemistry [IUPAC] has adopted Arabic numerals 1, 2, 3 ....18 as then newest group designation in the form of the periodic table. In this system therefore, the alkali and non alkaline earth metals constitute group 1 and 2, transition elements of Sc to Zn families become groups 3, 4, 5, 12 and finally the P block elements become groups 13, 14, 18 of the table.
3.3 Nomenclature of Elements Having Z > 100

It has been a historical practice to allow the discoverer of the elements to assign the element's name. In recent times, this has led to some controversy because elements with very high atomic number are so unstable that only minute quantities of them, sometimes only one or two atoms are prepared before scientists claim credit for their discovery. This has led to the questions of the reliability of the data and whether the said new element has in fact been discovered.

For example both American and Soviet scientists claimed credit for discovering element 104. The Americans named it rutherfordium and the Soviet scientist named it Kurchotovium. To avoid this problem, the IUPAC has made an official recommendation that until a new element discovery has been proved, a systematic nomenclature be applied according to the following IUPAC nomenclature rules:

1. The names be derived directly from the atomic number of the element using the following numerical root.

**Table 2.1: Numerical Root for Atomic Number of Element**

<table>
<thead>
<tr>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td>nil</td>
<td>un</td>
<td>bi</td>
<td>tri</td>
<td>quad</td>
<td>pent</td>
<td>hex</td>
<td>Sept</td>
<td>oct</td>
<td>enn</td>
</tr>
</tbody>
</table>

2. The root be put together in the order of the digit which make up the atomic number and be terminated by "ium and ending occurring in the names of the metallic elements as these are the final "n" of enn be dropped when it occurs before 'nil' and 1' of `bi' and `tri.' be dropped when it occurs before `ium'.

3. The symbol of the element be composed of the initial letters of the numerical roots which make up the names.
Table 2.2: The Systematic Names and Symbols of Elements having $Z = 101$ to 106 Derived by Application of IUPAC Nomenclature Rules

<table>
<thead>
<tr>
<th>Atomic Number</th>
<th>Systematic Names</th>
<th>Symbol</th>
<th>Trivial Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>101</td>
<td>Unnilinium</td>
<td>Unu</td>
<td>Mendelevium</td>
</tr>
<tr>
<td>102</td>
<td>Unnilbium</td>
<td>Unb</td>
<td>Nobelium</td>
</tr>
<tr>
<td>103</td>
<td>Unniltrium</td>
<td>Unt</td>
<td>Lawrencium</td>
</tr>
<tr>
<td>104</td>
<td>Unnilquadium</td>
<td>Unq</td>
<td>-</td>
</tr>
<tr>
<td>105</td>
<td>Unnilpentium</td>
<td>Unp</td>
<td>-</td>
</tr>
<tr>
<td>106</td>
<td>Unnilhexium</td>
<td>Unh</td>
<td>-</td>
</tr>
</tbody>
</table>

To further enhance our understanding of the rules, let us work out the name of the element 101.

*We start with the 1st 1 of the 101*
*You have un,*
*Then 0 you have nil*
*Then I again you have un*
*You end it with 'ium'*
*The name of element 101 is therefore un + nil + un +ium that is unnilinium.*

4.0 CONCLUSION

In conclusion, you can say that the periodic table in its current form is the result of years of painstaking research backed by consensus of opinions of leading scientists on the form it should take.

5.0 SUMMARY

In summary, you have studied the following in this unit:

- the periodic law first proposed by Dmitri Mendeleev had to be modified
- the modification came as a result of the recovery of the concept of the 'atomic number' in 1913 by Hendry Moseley
- the introduction of the concept of atomic number further clarified the arrangement of the elements in the periodic table and removal any ambiguities which were observed
- the atomic number rather than the atomic weight is the most important determinant of the properties of an element
- as a result of a consensus reached by IUPAC the recommendation of any newly discovered element must follow IUPAC nomenclature rules.
SELF-ASSESSMENT EXERCISE

Which group of elements appears in the modern periodic table but did not appear in Mendeleev's original table? Why?

6.0 TUTOR-MARKED ASSIGNMENT

1. What will be the name of an element whose atomic number is 110? Explain how you got your answer.
2. How would you use the modern periodic law to remove the anomalies with regards to the positions of ILK an Ar in Mendeleev’s periodic table?

7.0 REFERENCE/FURTHER READING

UNIT 3   ELECTRONIC CONFIGURATION

CONTENTS

1.0   Introduction
2.0   Objectives
3.0   Main Content
   3.1   Rules Governing the Filling of Electrons in Orbitals
   3.2   Electronic Configuration of all the Elements in the Periodic Table
   3.3   Electronic Configuration of Ions
   3.4   Electronic Configuration and Division of Element into Blocks
4.0   Conclusion
5.0   Summary
6.0   Tutor-Marked Assignment
7.0   References/Further Reading

1.0   INTRODUCTION

In the last unit, you have learnt about the periodic table. The periodic law gives rise to the periodic arrangement of the element according to their atomic numbers, which indeed means according to the number of electrons in their orbital. In the next unit you will be studying the distributions of the electrons within the atom (the electronic configuration), and how they govern the properties of the elements.

The electronic configurations of isolated atoms of elements are usually verified experimentally by a detailed analysis of atomic spectra. In this unit, we are not discussing these methods. Instead we are going to discuss the process of filling in of electrons in the various atomic orbital.

2.0   OBJECTIVES

At the end of this unit, you should be able to:

- state the principle involved in determining which electron goes into which atomic orbital
- fill out correctly electrons in a given atom once given the number
- list the four blocks of elements on the periodic table and determine to which block an element belongs if given the electronic configuration.
3.0 MAIN CONTENT

3.1 Electronic Configuration of Atoms

Rules governing the filling of electrons in orbital

The electronic configuration of atoms can be predicted with the help of Aufbau or the building up process. In the Aufbau process, it is assumed that there exist a set of empty hydrogen like orbital around the nucleus of an atom. The electronic configuration of the atom in the ground state is then derived by adding electrons one at a time to the orbitals of the lowest energy in the sequence shown by arrows in Table 3.1.

Table 3.1: Order or Filling of Atomic Orbitals in Polyelectronic Atoms

The order in which the orbitals are filled as shown in Table 3.1 is governed by the n +1 rule. According to this rule, in the building up of electronic configuration of the elements, the sub-shell with the value of n +1 fills first. This rule reminds us that the energy of sub shells of multi electron atoms depends upon the value of both the quantum numbers n and 1, but mainly on the value of n. For example, to determine which of the sub shells 5s or 4p fills first. We follow the rule thus: for the 5s sub shell the value n +1 = 5 + 0 = 5; for 4p sub shell also the value of n + 1
= 4 + 1 = 5, but the 4p sub shell has the lower value of the principal quantum number n and therefore it fill first. Filling of electrons in orbitals is also governed by Pauli’s Exclusion Principle and Hund’s rule.

According to the Pauli Exclusion Principle, no two electrons in the same atom can have the same value of n, l and m, they will differ in their ms values. In order words, an orbital can have at most, two electrons of opposite spin. Since there is only one s orbital for any given value of n, it can contain only two electrons. However, the three p orbitals for any given value of n can contain six electrons, the five d orbitals, for any given value of n can hold a total of ten electrons and the seven f orbitals can have fourteen electrons. Permitted combinations of all the four quantum numbers for the electrons in different orbitals are given below in Table 3.2.

**Table 3.2: Permitted combinations of Quantum Numbers for S, P, d and f Orbitals**

Hund's rule of maximum multiplicity states that, as far as possible in a given atom in the ground state, electrons in the same sub shell will occupy different orbitals and will have parallel spins. That means that when electrons are added to orbitals of the same energy such as three p orbitals or five d orbitals, one electron will enter each of the available orbital, two electrons in separate orbitals feel less repulsion than two
electrons paired in the same orbitals. For example, carbon in the ground state has the configuration $1S^2 \ 2S^2 \ 2Px^1 \ 2Py^1$ rather than $1S^2 \ 2S^2 \ 2Px^2$.

So far you have studied the rules governing the filling of electrons in the orbitals of atoms. We shall now consider the electrons configurations of all the elements in the periodic table; these are given in table 3.3.

**Table 3.3: Ground State Electronic Configuration of Gaseous Atoms**
3.2 Electronic Configuration of all the Elements in the Periodic Table

Period 1

This is the smallest of all the periods of the table. Hydrogen (Z = 1) and helium (Z = 2) are the two elements belonging to the period. The electronic configuration of hydrogen and helium are 1s\(^1\) and 1s\(^2\) respectively. Thus the 1s orbital which is only corresponding to n = 1 is completely filled. The 1s\(^2\) configuration of helium is usually represented by [He]. So anytime you see [He] electron configuration it represents 1s\(^2\).

Period 2

This period contain elements from lithium (Z = 3) to neon (Z F 10). In lithium and beryllium, the filling of 2S orbital takes place, then in the
next six elements from boron to neon, the 2p orbitals are filled. Neon thus has the electronic configuration of [He] 2S² 2P⁶ which as was done in the case of He, is represented by [Ne]. (An electronic configuration of [Ne] means 1S² 2S² 2P⁶. At this stage, the shell having n = 2 is complete.

**Period 3**

Similar to period 2, this period also consists of 8 elements from sodium (Z = 11) to argon (Z = 18) these elements 3s and 3p orbitals are successively filled in the sequence just as was done in period 2, thus argon has the electronic configuration [Ne] 2S² 2P⁶ represented as [Ar] although the third principal shell (n = 3) can accommodate 10 more electron in 3d orbitals filling of 4s orbital takes place first because of its lower energy.

**Period 4**

This period contains 18 elements from potassium (Z = 19) to krypton (Z = 36). In k and Ca, the first two elements of this period, the successive electrons go into the 4s orbitals giving them the configuration [Ar] 4S¹ and [Ar] 4S² respectively. Then in the following 10 elements (Sc, Ti, V, Cr, Mn, Fe, Co, Ni, Cu and Zr) filling of hitherto unoccupied 3d orbitals takes place. Thus the electronic configuration of zinc becomes [Ar] 3d¹⁰ 4S².

Occasionally, an electron from 4s orbitals is shifted out of turn to the 3d orbitals due to higher stability of half filled and completely filled orbitals, for example, Cr (Z = 24) and Cu (Z = 29) have the configuration [Ar] 3d⁵ 4S¹ and [Ar] 3d¹⁰ 4S¹ instead of the expected [Ar] 3d⁸ 4S² and [Ar] 3d⁹ 4S² respectively. After the 3d level is filled, in the next six elements of this period, that is Ga, Ge, As, Se, Br and Kr, the 4p orbitals are gradually filled and Kr has the electronic configuration [Ar] 3d¹⁰ 4S² 4P⁶ represented as [Kr].

**Period 5**

The next 18 elements from rubidium (Z = 37) to Xenon (Z = 54) belong to this periods In building up of the atoms of these elements, 5s4d and 5p orbital are successively filled just as the 4s 3d, Cd, 4p are filled in the elements of period 4. In Rb (Z = 37) and Sr (Z = 38), the 5s orbital is filled. After that in elements from Y (Z = 39) to Cd (Z = 48) filling of 4d orbitals takes place. You can see from table 3.1 that once again there are minor irregularities in the distribution of electron between 4d and 5s orbitals. For example Mo (Z = 42) and Ag (Z = 47) have respectively [Kr] 4d⁵ 5S¹ and [Kr] 4d¹⁰ 5S¹ configurations similar to those of Cr and
Cu respectively. Anomalous electronic configuration of Nb, Ru, Rh and Pd cannot be explained in simple terms. You have to therefore, remember them as exceptions. Now in the next six elements, that is I, Sn, Sb, Te, I and Xe filling of Sp orbitals take place and thus Xe ($Z = 54$) attains $[\text{Kr}] \, 4d^{10}S^25p^6$ configuration.

**Period 6**

This period contains 32 elements from caesium ($Z = 55$) to radon ($Z = 86$) in which the 6s, 4f, 5d and 6p orbitals are filled. The first two elements of this period have configurations analogous to those of corresponding member of the lower periods, thus caesium ($Z = 55$) and barium ($Z = 56$) have $[\text{Xe}] \, 6S^1$ and $[\text{Xe}] \, 6S^2$ configuration respectively. According to Aufbau principle in the next element La ($Z = 57$), the additional electron should enter 4f orbital. Instead it goes to the 5d orbitals and La has the configuration $[\text{Xe}] \, 5d^1 \, 6S^2$ but why? The extra electron in the building up of La atom goes to 5d orbital instead of 4f orbital because in La atom, the 5d and 4f orbitals have almost the same energy and hence, the electron is free to enter any of these two orbitals.

In the next 14 elements from cerium ($Z = 58$) to luteum ($Z = 71$), the 4f orbital is successively filled pertaining to $[\text{Xe}] \, 4f^1 \, 5d^1 \, 6S^2$ and $[\text{Xe}] \, 4f^15d^1 \, 6S^2$ configuration, respectively, but you should remember, it is only ce ($Z = 58$) Gd ($Z = 64$) and Lu ($Z = 71$) that 5d orbitals have one electron while in all the remaining Lanthanides the 5d orbitals remain vacant.

After Lutecium, successive electrons occupy 5d orbitals and the electronic configuration builds up from $[\text{Xe}] \, 4f^{14} \, 5d^26S^2$ to for hafnium to $[\text{Xe}] \, 4f^{14} \, 5d^{10} \, 6S^2$ for mercury the homologue of zinc and cadmium.

Again a minor departure from a steady increase in the number of d electrons occurs. For example, gold has $[\text{Xe}] \, 4f \, 145d^9 \, 6S^2$, and as you can see, has to do with the greater stability of half filled/fully filled orbitals. Finally the period is completed with successive occupation of the Gp orbitals from thallium, $[\text{Xe}] \, 4f^{14} \, 5d^{10} \, 6S^2 \, 6p^1$ to randon,$[\text{Xe}] \, 4f^{14} \, 5d^{10} \, 6S^2 \, 6P^6$.

**Period 7**

This period is still incomplete and contains 23 elements from francium ($Z = 87$) to unnitennium ($Z = 109$). In these elements, electrons are filled in 7s, 5f and 6d orbitals. Francium ($[\text{Ru}] \, 7S^1$), radium ($[\text{Ru}] \, 7S^2$) and antinium ($[\text{Ru} \, 36d^1 \, 7S^2$) have electronic configurations analogous to those of caesium, barium and lanthanum respectively. Thorium has the configuration $[\text{Ru}] \, 6d^37S^2$. Therefore in the 13 elements from
protactinium ($Z = 91$) to lawrencium ($Z = 103$) filling of 5f orbitals takes place successively. However, out of these only Pa ($Z = 91$), U ($Z = 92$), Np ($Z = 93$), Cm ($Z = 96$) and Lr ($Z = 103$) have an electron in 6d orbitals. In the rest of the elements, the 6d orbitals remain vacant, thus the electronic configuration of Lr ($Z = 103$) is [Ru] 5f$^{14}$ 6d$^{2}$ 7S$^{2}$. The next six known elements of this period are members of 6d transition series which have the configurations [Ru] 5f$^{14}$ 6d$^{2}$ 7S$^{2}$ to [Ru] 5f$^{14}$ 6d$^{7}$ 7S$^{2}$.

Having examined the electronic configuration of elements in the periodic table, you can see from Table 3.4 that the elements occupying the same group of the periodic table have the same valence-shell electronic configuration. In order words, the elements having the same valence shell electronic configuration recur periodically, that is after intervals of 2, 8, 8, 18, 18 and 32 in their atomic number. Therefore periodicity in the properties of elements can easily be understood.

### 3.3 Electronic Configuration of Ions

So far, you have studied the electronic configuration of neutral atoms of elements. I am sure you will be interested in knowing the electronic configuration of ions that are obtained by removal of electrons from the elements. When the gaseous iron atom having [Ar] 3d$^{6}$ 4S$^{2}$ ground state electronic configuration looses an electron, the Fe$^{+}$ ion is formed. This ions has its minimum energy in the configuration [Ar] 3d$^{7}$, although the iso electronic manganese atom has the configuration [Ar] 3d$^{5}$ 4S$^{2}$ in the ground State. Similarly, the ground state of the Fe$^{2+}$ and Fe$^{3+}$ ions are [Ar] 3d$^{6}$ and [Ar]3d$^{5}$ respectively rather than [Ar] 3d$^{5}$ 4S$^{1}$ and [Ar] 3d$^{3}$4S$^{2}$ which are ground states of iso-electronic atoms of chromium and vanadium respectively. Evidently the differences in nuclear charge between Fe$^{+}$ and Mn, Fe$^{2+}$ and Cr and Fe$^{3+}$ and V are important in determining the orbital to be occupied by the electrons.

However, along the series of ions carrying the same charge, the electronic configuration often changes much more regularly than the electronic configuration of the corresponding atoms. Thus for dispositive ions Sc$^{3+}$ to Zn$^{2+}$, the ground state electronic configuration changes regularly from [Ar]3d$^{1}$ to [Ar] 3d$^{10}$. For dispositive ions, there is a similar regular change from [Ar] for Sc$^{3+}$ to [Ar] 3d$^{9}$ for Zn$^{3+}$. For tripositive ions of lanthium elements, there is a regular change from [Xe] 4f$^{1}$ for Ce$^{3+}$ to [Xe] 4f$^{14}$ for Lu$^{3+}$. Since the chemistry of elements is essentially that of their for Lu ions, the regularities in configuration of ions are much more important than the irregularities in the electronic configuration of the neutral atoms.
3.4 Electronic Configuration and Division of Elements into Blocks

Elements of the periodic table have been divided into four blocks s, p, d, and f depending upon the nature of the atomic orbitals into which the differentiating or the last electron enters.

The s-block elements: In these elements the differentiating electron enters the `ns' orbital. Alkali and alkaline earth metals of groups (1A) and 2(11A) belong to this block. As you know the valence shell electronic configuration of these groups are ns1 and ns2 respectively. We also know that each period of the periodic table begins with alkali metals. All the elements in this block are metals.

The p-block elements: In the elements belonging to this block, the p-orbitals are successively filled. Thus the elements of the group 13 (111A), 14(IVA), 15(VA), 16(VIA), 17(VIIA) and 18(zero) are members of this block, since in the atoms of these elements, the differentiating electron enters the np orbitals. The ns orbital in the atoms of these elements are already completely filled so they have the valence shell electronic configuration ns2np1-6.

Note that the elements of s- and p-blocks are also known as normal representative or main group elements.

The d-Block Elements — The elements in which the differentiating electron enters the (n-1)d orbitals are called d-block elements. These elements are placed in the middle of the periodic table between the sand p-block elements. The electronic configuration of the atoms of the elements of this block can be represented by (n-1) d1-10 ns0-12. These elements which are also called transition elements are divided into four series corresponding to the filling of 3d - 4d — 5d — or 6d — orbitals. While the 3d, 4d, and 5d series consist of 10 elements each, the 6d series is incomplete and has only seven elements viz: Ac (Z = 89) and from Unq 9Z = 104) to Une (Z = 109). The element from Sc (Z = 21) to zn (Z = 30), Y (Z = 39) to Cd (Z = 48), La (Z = 57) and from Hf (Z = 72) to Hg (Z = 80) are the members of 3d, 4d, and 5d series respectively.

Note: d-Block elements are also known as transition elements.

The f-block elements — The elements in which the extra electron enters (n-2)f orbitals are called the f-block elements. The atoms of these elements have the general configuration (n-2) f1-14 (n-1) d0-1 ns2. These elements belong to two series depending upon the filling of 4f and 5f orbitals. Elements from Ce (Z = 58) to Lu (Z = 71) are the members of the 4f series, while those from th (Z =90) to Lr (Z = 103) belong to the...
5f series. Elements of 4f series which follow lanthanium in the periodic table are known as **Lanthanides** whereas those of 5f series following actinium are called **Actinides**. All these elements are collectively referred to as **Inner-transition** elements because of filling of electrons in an inner (n-2) f sub shell.

**Note that f-block elements are also known as inner transition elements.**

### 4.0 CONCLUSION

In conclusion, you have seen that the order in which electrons occupy atomic orbitals is governed by certain rules and principles. These rules determine how many and which electrons occupy valence shells. It is the valence shells that determine the kind of reaction an atom will be involved in. It is therefore very important that the order of filling of orbitals is properly understood.

### 5.0 SUMMARY

In summary, you have studied the following in this unit:

- that the filling in of electrons into their orbitals is governed by:
  - the Aufbau principle which assumes that there exist a set of empty hydrogen like orbitals into which electrons can be added
  - The n+1 rule, which states that in building up electronic configuration of the elements the sub shell with the lowest value of n =1 fills first
  - The Pauli Exclusion principles which states that no two electrons in the same atom can have the same value of all four quantum numbers
  - The Hund's rule which states that as far as possible in a given atom in the ground state, electrons in the same sub shell will occupy different orbitals and will have parallel spins
- that the electronic configuration of ions changed regularly
- that the elements in the periodic table are divided into four blocks viz: s-p- d- and f blocks.

### SELF-ASSESSMENT EXERCISE

i. What principles or rules are violated in the following electronic configuration? Write the names of the principle or rule in the space provided alongside each configuration.
(i) 1S²2S³
(ii) 1S²2S² 2Px² 2py¹
(iii) 1S²2Px²

ii. Write the electronic configuration of the atoms whose atomic numbers are:

(i) 21
(ii) 24
(iii) 29

6.0 TUTOR-MARKED ASSIGNMENT

1. Explain Pauli's exclusion principle.
2. State Hund's rule of maximum multiplicity.

7.0 REFERENCE/FURTHER READING

UNIT 4 ATOMIC RADII

CONTENTS

1.0 Introduction
2.0 Objectives
3.0 Main Content
   3.1 Atomic Radii
   3.2 Covalent Radius
   3.3 Van der Waal’s Radius
   3.4 Metallic Radius
   3.5 Ionic Radius
   3.6 Factors affecting Atomic Radii
   3.7 Periodicity in Atomic Radii
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 Reference/Further Reading

1.0 INTRODUCTION

In Units 1 — 3, you studied the development of the periodic law and the periodic table. We learned about the properties of elements being periodic function of their atomic numbers. We learned how electrons are arranged in their orbitals. Arrangements that give rise to similarities and differences in the properties of elements whose valence electrons appear in the same group and those whose valence electrons are in different groups respectively. These differences in the properties arise due to differences in atomic properties such as size of the atoms as measured in terms of radii.

In this unit, you will be studying about different types of atomic radii, factors affecting atomic radii and periodicity in atomic radii.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

• define accurately "Atomic radii"
• distinguished between various forms of atomic radii
• list and explain with 80% accuracy, the two factors that affects atomic radii
• explain using examples periodicity in atomic radii.
3.0 MAIN CONTENT

3.1 Measurement of Atomic Radii

Atomic radii are the measure of the size of the atom. Atomic radii are important because other atomic properties like ionisation energy, electron affinity and electronegativity are related to them. The wave mechanical picture of an atom depicts an atom as composed of a compact nucleus surrounded by an electron cloud. This electron cloud does not have a definite boundary surface like that of a ball. There is a definite but very small probability of finding an electron at an infinite distance from the nucleus of the atom. However, this does not mean that the atom is indefinitely large; therefore, we have to find a way to define the size of an atom. Accordingly, the radius of an atom can be defined as the distance from the centre of the nucleus to the point where the electron density is virtually zero.

Now that we have defined the size of an atom, we have to tackle the problem of measuring that size. We are immediately confronted with the problem of defining and accurately measuring which size we meant. Thus, if we are measuring the size of an atom when it is occupying a lattice site of the crystal, the value will be different from one when it is colliding with another atom in the gaseous state. Furthermore, the size of a neutral atom will be different from the one when it is present as a cation or anion. Consequently, we cannot have one set of atomic radii applicable under all conditions. It, therefore, becomes necessary to specify the bonding conditions under which the size is being measured. Pertaining to the four major types of bonding, the atomic radii to be measured are:

i. Covalent radius
ii. Crystal or metallic radius
iii. Van der Waals radius
iv. Ionic radius

3.2 Covalent Radius

Covalent radius can be defined as one half of the distance between the nuclei of two like atoms bonded together by a single covalent bond.

If in a homonuclear diatomic molecule of A2 type (e.g. F2, Cl2, Br2, I2) rA-A is bond length or inter nucleus distance and rA is the covalent radius of the atom A, then rA = 1/2 rA-A. The internuclear distance r c-c between two carbon atoms in diamond is 154pm, so the covalent radius
of carbon, $r_e$ is equal to 77pm. Similarly, the $r_{ci} - r_{cl}$ for solid is 198 pm. $r_{cl}$ is therefore 99pm.

In the heteronuclear, diatomic molecule of AB type, if the bonding is purely covalent, then the bond length $r_{A-B}$ is equal to the sum of covalent radii of A and B that is $r_{A-B} = r_{A} + r_{B}$. Thus covalent radii are additive. It is possible to calculate the radius of one of the atoms in a heteronuclear diatomic molecule of AB type. If we know the internuclear distance $r_{A-B}$ and radius of the other atom. For example, the Si—C bond length in carborundum is 193 pm and covalent radius of C is 77, so you can calculate the covalent radius of Si as follows:

$$r_{Si-C} = r_{Si} + r_{C}$$

or

$$r_{Si} = 193 - 77 = 116 \text{ pm}$$

As stated earlier, the above relation holds good only if the bond between the atoms A and B is purely covalent. If there is a difference in the electronegativities of the bonded atoms, it causes shortening of the bonds. Schoemaker and Stevenson have proposed the following relationship between the shortening of the bond and the electronegativity difference of the atoms:

$$r_{A-B} = r_{A} + r_{B} - 0.07 (X_{A} - X_{B})^2$$

Here $X_A$ and $X_B$ are the electronegativities of A and B respectively.

Multiplicity of the bond also causes a shortening of the bond. Usually a double bond is about 0.86 times and a triple bond about 0.78 times the single bond length for the second period elements. Covalent radii of the elements are listed in Table 4.1
Table 4.1: Covalent and Van der Waals Radii of Elements

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
<th>13</th>
<th>14</th>
<th>15</th>
<th>16</th>
<th>17</th>
<th>18</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>120</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>He</td>
<td></td>
<td>154</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Li</td>
<td>152</td>
<td>133</td>
<td>110</td>
<td>91</td>
<td>74</td>
<td>59</td>
<td>47</td>
<td>36</td>
<td>29</td>
<td>24</td>
<td>20</td>
<td>16</td>
<td>13</td>
<td>10</td>
<td>7</td>
<td>4.5</td>
<td>3.8</td>
<td>3.2</td>
</tr>
<tr>
<td>Be</td>
<td></td>
<td>133</td>
<td>110</td>
<td>91</td>
<td>74</td>
<td>59</td>
<td>47</td>
<td>36</td>
<td>29</td>
<td>24</td>
<td>20</td>
<td>16</td>
<td>13</td>
<td>10</td>
<td>7</td>
<td>4.5</td>
<td>3.8</td>
<td>3.2</td>
</tr>
<tr>
<td>B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ne</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Li</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Be</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>O</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ne</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

3.3 Van Der Waal's Radius

In the solid state, non-metallic elements usually exist as aggregates of molecules. The bonding within a non metal molecule is largely covalent. However, individual molecules are held together by weak forces known as Van der Waals forces. Half of the distance between the nuclei of two atoms belonging to two adjacent molecules in a crystal lattice is called Van der Waal's radius. Table 4.1 lists the values of Van der Waals radii of some elements. Figure 4.1 illustrate the difference between the covalent and van der Waals radii of chlorine.

Fig 4.1 Covalent and van der Waals Radii of Solid Chlorine

It is evident from the figure that half of the distance between the nuclei X and X' of the two non-bonded neighbouring chlorine atoms of adjacent molecule A and B is the van der Waal's radii of chlorine atom. On the other hand half of the distance between the two nuclei X and Y in the same molecule is the covalent radius of chlorine atom. Thus van der Waal's radii represent the distance of the closest approach of an atom to another atom it is in contact with, but not covalently bond to it. Values of van der Waals radii are larger than those of covalent radii.
because van der Waals forces are much weaker than the forces operating between atoms in a covalently bonded molecule.

### 3.4 Metallic or Crystal Radius

Metallic or crystal radius is used to describe the size of metal atoms which are usually assumed to be loosely packed spheres in the metallic crystal. The metal atoms are supposed to touch one another in the crystal. **Metallic radius is defined as one-half of the distance between the nuclei of two adjacent metal atoms in the close packed crystal lattice.** For example, the inter-nuclear distance between two adjacent Na atom in a crystal of sodium metal is 382 pm. Therefore, metallic radius of Na metal is 382 that is 191 pm.

The metallic radius depends to some extent on the crystal structure of the metal. Most metals adopt a close packed (hcp) or cubic close packed (ccp) lattice (see Figure 4.2)

![Fig. 4.2 Types of Metal Lattices: (a) Hexagonal;(b) Cubic close packed (c) Body-centred cubic](image)
Table 4.2: Metallic and Ionic Radii of Elements

In both these structure, a given metal atom has twelve nearest neighbours. However, a significant number of metals adopt a body centred cubic lattice (bcc) in which the number of nearest neighbours is eight. The number of nearest neighbours of a metal atom in a lattice is known as the coordination number of the metal. Experimental studies on a number of metals having more than one crystal lattice have shown that the radius of a metal in an eight coordinate lattice is about 0.97 of the radius of the same metal in a twelve coordinate environment. Table 4.2 gives a set of twelve coordinate radii for metal atoms. Compare these with the covalent radii or van der Waals radii in Table 4.1.

The metallic radii are generally larger than the corresponding covalent radii. Although both involve a sharing of electrons this is because the average bond order of an individual metal-metal bond is considerably less than one and therefore the individual bond is weaker and longer than the covalent. This does not mean that the overall bonding is weak as there are a large number of these bonds, eight or twelve per metal atom. On the other hand, the metallic crystal lattices are stronger than the van der Waals forces.

3.5 Ionic Radius

Ionic radius is defined as the distance between the nucleus of an ion and the point up to which the nucleus has influence on the electron cloud. In order words, it may also be defined as the distance of the closest approach from the centre of ion by another ion. Ionic radius is usually
evaluated from the distance determined experimentally between the centres of nearest neighbours. Thus, if we wish to estimate the ionic radius of Na\(^+\) we may measure the internuclear distance between Na\(^+\) and Cl\(^-\) ions in the NaCl crystal lattice. This distance is the sum of radii of Na\(^+\) and Cl\(^-\) ions. From the electron density maps obtained by x-ray analysis, it has become possible, in some cases, to apportion the internuclear distance into the radius of cation and anion. A small member of ionic crystals has thus been studied and the ionic radii of some of the elements have been determined. These radii have become the basis for assigning the ionic radii of most of the other elements.

Ionic radii are of two types, cation radii and anion radii. All common cations are smaller than all common anion except for rubidium and caesium cations (largest single atom cations). This is not too surprising since not only is there a loss of electron(s) from a partially filled outer shell on cation formations, but there is also an increase in the overall positive charge on the ion.

Conversely, in anion formation the addition of an electron to an atom increases the size due to increase in inter-electronic repulsion in the valence-shell and decrease in effective nuclear charge. In general, there is a decrease in size of anions to covalent radii of corresponding atoms to cations thus in the series of isoelectronic species (e.g. N\(^3-\), O\(^2-\), Ne, Na\(^+\), Mg\(^2+\) and Al\(^3+\)). The greater the effective nuclear charge, the smaller is the radius of the species. In Table 4.2; radii of some of the common ions have been listed.

### 3.6 Factors Affecting the Atomic Radii

So far, we have defined and explained types of atomic radii. We shall now turn our attention to two of the factors that affect them.

(a) **Principal Quantum Number (n):** As the principal quantum number increases, the outer electrons get farther away from the nucleus and hence the atomic radius generally increases.

(b) **Effective Nuclear Charge (Z\(^*\)):** The magnitude of the effective nuclear charge determines the magnitude of the force of attraction exerted by the nucleus on the outermost electrons. The greater the magnitude of effective nuclear charge, the greater is the force exerted by the nucleus on the outermost electron.

Hence, the electron cloud of the outermost shell is pulled inward nearer to the nucleus and consequently its distance from the nucleus. That is, atomic radius decreases. Effective nuclear charge Z\(^*\) is the amount of positive charge felt by the outer electrons in an atom. It is always less than the actual charge Z of the nucleus of the atom. This is because
electrons in inner shells partially shield the electrons in the outer shell from nuclear attraction. The effective nuclear charge felt by the outer electron depends upon the actual nuclear charge and the number and type of inner screening electrons. It can be calculated by subtracting the screening or shielding constant, S from the atomic number Z thus \( Z^* = Z - S \).

You can estimate the value of screening constant, S, with the help of Slater’s rules in the following manner:

i) Write out the electronic configuration of the element in the following order and groupings:

\[(1s) \ (2s, \, 2p) \ (3s, \, 3p) \ (3d) \ (4s, \, 4p) \ (4d) \ (5s, \, 5p) \ (5d) \ (5f) \ (6s, \, 6p) \text{ etc.} \]

ii) Electrons in any group higher in this sequence than the electron under consideration contribute nothing to \( S \). For example, in Ti atom (electronic configuration \( 1s^2 \ 2s^2 \ 2p^6 \ 3s^2 \ 3p^6 \ 3d^2 \ 4s^2 \)). The two electrons in 4s orbital will contribute nothing towards the screening for an electron in 3d orbital.

iii) Then for electrons in an ns or np orbitals:

(a) All other electrons in the (ns, np) group contribute \( S = 0.35 \) each except for the electron in is which contribute \( S = 0.30 \)
(b) All electrons in (n-1) shells contribute \( S = 0.85 \) each
(c) All electrons in (n-2) or lower shells contribute \( S = 1.00 \) each

iv) For an electron in an nd or nf orbital

(a) All electrons in the same group that is nd or nf contribute \( S = 0.35 \) each.
(b) Those in the groups lying lower in the sequence than the nd or nf group contribute \( S = 1.00 \) each.

In order to demonstrate the application of Slater’s rules, we shall now calculate the \( Z^* \) for an electron in N, K and Zn atoms.

A Electronic configuration of N = (Is\(^2\)) \((2s \ 2p^3)\)

Grouping (Is2) \((2s^2 \ 2p^3)\)

Value of screening constant for an electron in 2p orbital will be \( S = (4 \times 0.35) + (2 \times 0.85) = 3.10 \)

Hence:

\( Z^* = Z - S = 7 - 3.10 = 3.90 \)
B  Electronic configuration of K = Is\(^2\) 2s\(^2\) 2p\(^6\) 3s\(^2\) 3p\(^6\) 4s\(^1\)  Grouping of orbitals will be (Is\(^2\)) (2s\(^2\) 2p\(^6\)) (3s\(^2\) 3p\(^6\)) (3d\(^{10}\)) (4s\(^2\))  Value of screening constant for an electron in 4s orbitals will be:  
\[ S = 90.85 \times 8 \]  
\[ + (1 \times 10) = 16.80 \]  Hence effective nuclear charge is given as:  
\[ Z^* = Z - S = 19 - 16.80 = 2.20 \]

C  Electronic configuration of Zn = Is\(^2\) 2s\(^2\) 2p\(^6\) 3s\(^2\) 3p\(^6\) 4d\(^{10}\) 4s\(^2\)  Grouping of the orbitals gives (Is\(^2\)) (2s\(^2\)2p\(^6\)) (3s\(^2\)3p\(^6\))(3d\(^{10}\)) (4s\(^2\)). Value of screening constant for S for an electron in 4s orbital will be  
\[ S = (0.35 \times 1) + (0.85 \times 18) + (1 \times 10) = 25.65 \]  hence the effective nuclear charge felt by 4s electron will be  
\[ Z^* = Z - S = 30 - 25.65 = 4.35 \]

If we consider a 3d electron in Zn, the grouping is as above, but the effective nuclear charge felt by the 3d electron will be  
\[ Z^* = Z - S = 30 - [(9 \times 0.35) + (18 \times 1)] = 8.85 \]  Thus you can see an electron in 3d orbitals in Zn is more strongly held by the nucleus than that in 4s orbital.

Table 4.3 contains a list of values of nuclear charge for electron in valence shell in the first thirty elements calculated by Slater's rules. You can see from the table that there is a steady increase in Slater's Z* across rows of the periodic table. Effective nuclear charge felt by electrons also depends on the oxidation state of an atom in a compound.

The higher the oxidation state of the atom, the higher will be the effective nuclear charge felt by the electrons and therefore, smaller will be the atomic radius. Thus the ionic radius of Fe\(^{3+}\) ion will be smaller than that of the Fe\(^{2+}\) ion. Similarly, covalent radius of bromine in BrCl\(_3\) will be then that in BrCl.
3.7 Periodicity in Atomic Radii

Now that we know the various types of atomic radii and the factors that affect them, we will consider the periodicity in them. Before doing that however, we would like to emphasize that trends observed in one type of radii (example covalent radii) are generally found in the other type of radii also (example ionic and metallic radii). Two general periodic trends are found for all types of atomic radii. These are the atomic radii decreases along a period and generally increase down a group in the long form of the periodic table (Figure 4.3). These changes in the atomic radii can be related to the charges in effective nuclear charge and the principal quantum number in the periodic table.

If you examine Table 4.3 you will find out that there is a steady increase (by 0.65 units) in the value of Z* from alkali metals to halogens for the elements of period 2 and 3, but there is no change in the value of n because the electrons fill the same principal shell. As a result of this
there is a steady decrease in the covalent radius from 123 and 165 pm for Li and Na to 64 and 99 pm for F and Cl respectively.

In comparison to the above, the decrease in covalent radii across the transition series is much smaller. As you know, electrons are successfully filled in the (n-1)d orbitals across a transition series and hence screen the size determining ns electrons from the nuclear charge more effectively. Therefore across a transition series, there is only a small increase in effective nuclear charge (by 0.15 units), therefore only a small increase in effective nuclear charge decrease in atomic radius from one element to another takes place.

In 3d series, covalent radius decreases from 144 pm for Sc to 115 pm for Ni. Then in copper and zinc due to completion of 3d sub shell, the electronic charge density in this sub shell becomes very high which increases the inter electronic repulsion. As a result, covalent radii of Cu and Zn increase slightly to 117 and 125 pm respectively. Thus across the elements of the first transition series, there is an overall decrease in covalent radius by 19 pm which is much less than that across seven normal elements of Period 2 (59 pm) and Period 3 (57 pm). But due to this, the covalent radii of elements from Ga to Kr following Zn becomes much smaller than that expected by simple extrapolation of the values for elements of Period 2 and 3 for example, the covalent radii of Al and Ga are equal whereas the covalent radii of elements Ge, As, Se, Br are only slightly larger than those of corresponding elements (Si, P, and Cl) of Period 3. The rate of decrease in the size across the Lanthanide series is even less than that across the first transition series.

In the Lanthanide elements, filling of (n-2)f orbitals take place, while simultaneously the nuclear charge increases. The electrons in the (n-2)f orbital shield the ns electrons, (which largely determine the size, from the increase in nuclear charge) almost completely (S = 1.00) As a result of this, there is only a small decrease in the atomic radius from one element to another. But there are 14 elements in the series. There is a total of contraction of 13 pm across the series from Ca (Z = 57) to Lu (Z = 71). This is known as lanthanide contraction, because of which the atoms of elements (HF to Hg) following Lu are usually smaller than they would be if the lanthanide had not been built up before them. Lanthanide contraction almost exactly cancel out the effect of the last shell added in the sixth period and therefore, the transition elements of 4d and 5d series have almost the same atomic radii.

On descending any group of the periodic table, the number of electron in the valence shell remains constant but the number of shells around nucleus increases monotonically, so that the effective nuclear charge felt by valence electrons stays nearly the same. So with increase in principal
quantum number \((n)\) of the valence shell, an increase in atomic radii is generally observed down any group of the periodic table. For example as shown in Figure 4.3, there is an increase in atomic radii of alkali and alkaline earth metals as we proceed downward in the group.

However as pointed out earlier, with the inclusion of 3d transition elements in Period 4 increase in the radii of elements from Ga to Br is smaller than expected. Similarly, because of inclusion of Lanthanide elements in period 6, atoms of the transition elements of this period (Hf to Hg) are almost of the same size as atoms above than in period 5 (Zr to Cd). After that, only a small increase in size of elements of period 6 (tc to Al) as compared to the size of elements above them in period 5 (In to I) is observed.

Fig. 4.3 Increase in Atomic Radii of Alkali and Alkaline Earth Metals Downward the Group

4.0 CONCLUSION

Having gone through this unit, we can conclude that knowledge of the size of an atom is indeed very essential. It is through the knowledge of the atomic radii that we can predict accurately the reaction of the atom.

5.0 SUMMARY

In this unit, you have studied the following:

- the definition of atomic radii
- the various types of atomic radii viz: covalent, van der Waals metallic and ionic radii
- factors affecting radii that is the principal quantum number and the effective nuclear charge \(Z^*\)
- periodicity in atomic radii nuclear charge \(Z^*\)
- periodicity in atomic radii.
SELF-ASSESSMENT EXERCISE

i. Assuming that the atoms are touching each other, what would be the inter-nuclear distance between two fluorine atoms in F₂?

ii. Arrange the following iso-electronic species in order of decreasing atomic radius. Na⁺, Mg²⁺, Al³⁺, Si⁴⁺, N³⁻, O²⁻, F, Ne.

6.0 TUTOR-MARKED ASSIGNMENT

1. How does atomic size vary in a group and in a period? Give reasons for the variation.
2. Arrange H₂, H⁺ and H⁻ in order of increasing atomic radius.
3. What are isoelectronic ions? How does their size vary with the change of atomic number?

7.0 REFERENCE/FURTHER READING

UNIT 5    IONISATION ENERGY

CONTENTS

1.0     Introduction
2.0     Objectives
3.0     Main Content
   3.1    Factors affecting Ionisation Energy
   3.2    Periodicity in Ionisation Energy
   3.3    Trends in Ionisation Energy
   3.4    Trends in Successive Ionisation Energy
4.0     Conclusion
5.0     Summary
6.0     Tutor-Marked Assignment
7.0     References/Further Reading

1.0     INTRODUCTION

When elements react, they do so by gaining, losing or sharing of electrons. This process of gaining, losing or sharing of electrons is usually accompanied by energy changes. You know that the electrons to be shared, gained or lost are bound to the nucleus of its atom by an electrostatic force of attraction. In order to remove an electron from an atom, its force of attraction has to be overcome. This can be done by supplying energy. The energy required to remove the least strongly bond electron from an isolated gaseous atom in its ground state is known as the ionisation energy. This process can be represented by the following equation:

\[ M_{(g)} \rightarrow M^{+\text{g}} + e \]

Since more than one electron may be removed from an atom, the energy required for the above process is called the first ionisation energy. The second ionisation energy is the energy required to remove an electron from a univalent cation that process is represented by this reaction.

\[ M^{+\text{g}} \rightarrow M^{2+\text{g}} + e \]

The second ionisation energy is much larger than the first ionisation energy. This is because in this case an electron is being removed from a positively charged cation. Similarly you can define third, fourth and higher ionisation energies. The SI unit of ionisation energy which we will use throughout this course is Kilojoules per mole.
In this unit, you will be studying the factors that affect ionisation energy, periodicity in ionisation across the periods, trends in ionisation energy, down the groups and trends in successive ionisation energy.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- define ionisation energy
- differentiate between first, second, subsequent ionisation energies
- list the factors that affect ionisation energies
- explain trends in ionisation energies across period and down the groups.

3.0 MAIN CONTENT

3.1 Factors Affecting Ionisation Energies

The ionisation energy \( I \) of an outer valence electron is related to the effective nuclear charge felt by the electron and its average distance from the nucleus as stated in the equation below:

\[
I = \frac{Z^* e^2}{2} \times (I/r)_{av}
\]

Where \( Z^* \) is the effective nuclear charge, \( e \) is the charge on electron and \((I/r)_{av}\) is the average value of the electron from the nucleus, thus the higher effective charge felt by the electron, the higher will be the ionisation energy. Also the further the electron is from the nucleus, the lower will be the ionisation energy and vice versa.

In addition to the above, the ionisation energy also depends upon the relative stabilities of the sub shell from which the electron is removed.

As we have seen before, completely filled and half filled sub shells are comparatively more stable, so removal of an electron from them requires more energy. The valence shell electronic configurations of noble gases are exceptionally stable and therefore their ionisation energies are the highest in the irrespective period.

3.2 Periodicity in Ionisation Energy across Periods

In the previous sub-section we defined and identified factors which affect ionisation energies. In the next sub-sections, we shall examine the variation in ionisation energy across the periods and down the group in
the periodic table, values of ionisation energies of elements are given in Table 5.1

Table 5.1: Values of Ionisation Energies of Elements in KJ mol\(^{-1}\)

![Periodic Table](image)

The variation in ionisation energy in a particular group or period is best shown by a graph showing ionisation energies against atomic number. Figure 5.1 shows the plot of first ionisation energies of the elements of the first six periods against their atomic numbers. As is evident from the Figure, the first ionisation energy generally increases from alkali metals to noble gases across any row of the periodic table. But the increase is not perfectly regular.

![Graph](image)

Fig. 5.1 The Plot of First Ionisation Energies of the Elements of the First Six Periods against their Atomic Numbers
You have seen from an earlier section that across any row of the periodic table, the effective nuclear charge steadily increases and the atomic radii decreases. These two effects reinforce each other to increase the ionisation energies across a period. Thus, the ionisation energies of the alkali metals are the lowest and those of the noble gases are the highest in their respective periods. However as pointed out earlier, the increase is not smooth and some anomalies are observed. For example in the elements of Period 2, in spite of increase in Z* and decrease in r, the first ionisation energies of B and O are lower than those of Be and N respectively. However, these anomalies in the trend in ionisation energy can be explained by electronic structures of these elements.

In the case of beryllium, the electron is removed from the filled 2s sub shell whereas in boron, the electron is removed from the singly occupied 2p sub shell. The 2p sub shell is higher in energy than the 2s, so the 2p electron of boron is more easily removed than the 2s electron of beryllium. When we come to nitrogen, we will find out that we have a half filled 2p sub shell (electronic configuration Is²2s²3p³) while in oxygen the 2p sub shell is occupied by four electrons. The fourth electron in this 2p sub shell is in an orbital already occupied by another electron, so it experiences considerable repulsion. As a result, this electron is more easily removed than one of the electrons from a singly occupied orbital in nitrogen atom. Thus the ionisation energy of oxygen becomes less than that of nitrogen.

Similar anomalies are observed in elements of Period 3 where the first ionisation energies of magnesium and phosphorous are higher than those of aluminium and sulphur respectively. We have earlier seen that across the transition series, the increase in effective nuclear charge and consequent decrease in atomic radius is small. Therefore, increase in the first ionisation energies is also small. However, following the transition elements, the first ionisation energy drops abruptly in gallium, indium and thallium. This again is due to the removal of an electron from a singly occupy np orbitals which are of relatively higher energy than the ns orbital of Zn, Cd and Hg.

### 3.3 Trends in Ionisation Energy down the Groups

We have learnt from a previous sub section that on moving down a group of s- and p-block elements, effective nuclear charge remains almost steady. But there is a general increase in the atomic radius due to increase in the value of the principal quantum number n, thus the dominant factor in determining the ionisation energies of the elements on moving down the group is their atomic radius rather than the effective nuclear charge. Therefore, as expected, the first ionisation
energies decrease down the groups in the case of the main group elements in the periodic table. But in the case of transition elements opposite trends are observed. Thus, the first ionisation energies of the corresponding elements of 3d and 4d series are almost similar but these are smaller than the first ionisation energies of the elements of 5d series. Certainly the higher values of ionisation energies of the 5d transition elements are consistent with the relatively smaller size of their atoms.

3.4 Trends in Successive Ionisation Energies

We have already defined successive ionisation energies, that is second, third etc. Values of eight successive ionisation energies of the first twenty elements are listed in Table 5.2. It is evident from the values in the table that the successive ionisation energies of an element inevitably become larger because the removal of successive electron leaves a higher charge on the nucleus to hold the remaining electrons. It is also clear from the table that the difference between successive ionisation energies of the same element is not constant. Big jumps occur whenever an electron from a sub shell of lower principal quantum number is removed for the first time. For example, for alkali metals the second ionisation energies are much higher than the first.

For alkaline earth metal, the third ionisation energies are much larger than the second and for the halogen, the eighth ionisation energies are much greater than the seventh. These however cannot be explained on the basis of increase in nuclear charge alone. The stabilities of closed shell configuration similar to those of the noble gases are more important in these cases.

So far in our study, you have seen that ionisation energy generally increases across a period and decreases down the group in the periodic table. Accordingly, the tendencies to form cation that is metallic character decreases across a period and increases down the group. For example, in Period 3 metallic character decreases from Na to Cl whereas in the elements of Group 14, C is a non metal, Si and Ge are semi metals or metalloid whereas Sn and Pb are metals.

4.0 CONCLUSION

We can conclude this unit by observing that chemical reaction inevitably involves either a gain, a loss or sharing of electrons by atoms. It follows therefore that, whether a reaction takes place or not, the ease with which it can happen and how regularly it happens, depends very much on the factors which affect the formation of ions by the element.
5.0 SUMMARY

In this unit, you have learnt the following:

- the ability of an element to participate in a chemical reaction is governed by its ability to give up, gain or share its outermost electrons
- that this ability is measured in form of its ionisation energy
- that the ionisation energy can be defined as the energy required to remove the least strongly bond electron from an isolated gaseous atom in its ground state
- that the ionisation energy is affected by such factors as the effective nuclear charge of the atom and the reactive stabilities of the sub-shell from which the electron is removed.

SELF-ASSESSMENT EXERCISE

i. Which of the atoms having the following electronic configurations will have the highest first ionisation energy and why?
   (i) [Ne] 3s^2 3p^2
   (ii) [Ne] 3s^2 3p^3
   (iii) [Ne] 3s^2 3p^4
   (iv) [He] 2s^2 2p^3
   (v) [Ar] 3d^{10} 4s^2 4p^3

ii. In order of increasing ionisation energy for the atoms N, Ne, Na and P is:
   (i) Na, L,P, LN LNe
   (ii) N LNe C Na LP
   (iii) NL Na LNe LP
   (iv) Na LN LP LNe

6.0 TUTOR-MARKED ASSIGNMENT

Why is the second ionisation energy of sodium higher than that of magnesium?

7.0 REFERENCES/FURTHER READING


MODULE 2

Unit 1  Electron Affinity
Unit 2  Electronegativity
Unit 3  Hydrogen
Unit 4  Manufacture of Hydrogen
Unit 5  Ionic or Salt - Like Hydrides

UNIT 1  ELECTRON AFFINITY

CONTENTS

1.0  Introduction
2.0  Objectives
3.0  Main Content
   3.1  Electron Affinity
   3.2  Factors affecting Electron Affinity
   3.3  Periodicity in Electron Affinity
4.0  Conclusion
5.0  Summary
6.0  Tutor-Marked Assignment
7.0  References/Further Reading

1.0  INTRODUCTION

You will recall that in the last unit (Unit 5) of Module 1, you studied the process of removal of electron from a neutral atom. You learnt that the energy required to remove an electron from an isolated neutral atom in gaseous state is known as the ionisation energy. In this unit, you will study the process of addition of an electron to the gaseous atom in its ground state. In effect, what you are going to learn about is the reverse of the process of ionisation which we studied earlier on. We shall be talking about electron affinity.

2.0  OBJECTIVES

At the end of this unit, you should be able to:

- define electron affinity
- list the three factors affecting electron affinity
- discuss how electron affinity varies across periods and groups.
3.0 MAIN CONTENT

3.1 Electron Affinity

Electron affinity of an atom is a measure of its ability to accept an electron to form an anion. It is defined as the energy released or absorbed when an electron is added to the gaseous atom in its ground state. It can be represented by the following equation in which \( EA \) represents electron affinity of \( X \).

\[
X_{(g)} + e^- \rightarrow X^-_{(g)} + E_A
\]

When one mole of chlorine atom accepts one mole of electron, 348 kilojoules of energy is released. So the electron affinity of chlorine is \(-348 \text{ KJmol}^{-1}\).

In the reverse process of removal of an electron from the chloride ion to form a chloric atom, obviously an equal amount of energy has to be supplied. So, the electron affinity can also be expressed as the ionisation energy of the anion. Thus, the electron affinity of chlorine atom is clearly the ionisation energy of the chloride ion. Electron affinities are difficult to measure because the accurate values for all elements are not available. Values for some elements are given in Table 1.1.

You learnt in the preceding unit that energy is required to overcome the attractive force of the nucleus before an electron is removed from an isolated neutral atom in a gaseous state. It follows therefore that the reverse process of the addition of an electron to the neutral atom should release energy. Thus, the electron affinities of most elements are negative. However, a few elements are known to have positive values for electron affinity which means that the electron must be forced unto the neutral atom to form an anion. For example, nitrogen, alkaline earth metals and noble gases have positive values. All second and higher electron affinities also have large positive values.

This is not surprising, since the second and the subsequent electrons must be forced on against the negative charge of the anion. For example:
Note that as it is conventionally acknowledged, a negative value means an exothermic process. That is the reaction $X(g) + e \rightarrow X(g)$ is exothermic.

Also by high electron affinity, it is meant that the electron affinity is large and negative.

**Table 1.1: Electron Affinities of some Elements in KJmol⁻¹**

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>-73</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Li</td>
<td>-60</td>
<td></td>
<td>-9</td>
<td>-141</td>
<td>-328</td>
<td>+54</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Na</td>
<td>+30</td>
<td>-53</td>
<td>-72</td>
<td>-200</td>
<td>-348</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>-48</td>
<td>-53</td>
<td>-77</td>
<td>-195</td>
<td>-325</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca</td>
<td>-48</td>
<td>-48</td>
<td>-120</td>
<td>-77</td>
<td>-195</td>
<td>-325</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ba</td>
<td>-48</td>
<td>-48</td>
<td>-110</td>
<td>-181</td>
<td>-270</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### 3.2 Factors Affecting Electron Affinity

Factors affecting electron affinities are generally the same with those affecting ionisation energies. These factors are:

- **Atomic Radius:** When an electron adds on to any atom, the nucleus of the atom holds it by an electrostatic force of attraction which depends upon the effective nuclear charge and size of the atom. The smaller the size of the atom, the greater will be the force of attraction of the nucleus for the extra action added. Therefore the higher will be the electron affinity of the atom thus; more energy will be released in picking up an electron.

- **Effective Nuclear Charge:** The higher the effective nuclear charge, the greater the force of attraction exerted by the nucleus on the added electron and hence, higher will be the electron affinity of the atom.

- **Electronic Configuration:** Electronic configuration of the atom also plays an important role in determining the magnitude and sign of electron affinity. Halogens can achieve a stable noble gas configuration by accepting just one electron. Therefore, they have large negative (exothermic) electron affinities. On the other hand the noble gases with closed shell $ns^2 np^6$ configuration, beryllium and magnesium with $np^2$ (stable due to filled sub shell) and nitrogen having $np^2$ $np^3$ (stable due to half filled p sub shell)
configurations strongly resist the addition of any electron. Therefore, the electron affinities of these elements are either zero or have small positive values.

### 3.3 Periodicity in Electron Affinity

We have so far defined electron affinity and considered the factors that affect it. In this sub section, we will discuss how the electron affinity varies in the provided descriptions.

#### Trends across periods

On moving from left to right in a period, the size of an atom decreases and effective nuclear charge increases. Both these factors favour an increase in the force of attraction exerted by the nucleus on the extra electron. Consequently, the electron affinity generally increases across a period though irregularly. Thus electron affinities of alkaline metals have small negative values indicating their reluctance to form an anion. On the other hand, electron affinities of halogens in a period have the highest negative values which reflect their ability to form anions most readily. As explained earlier, the electron affinities of noble gases, beryllium, magnesium and nitrogen have small positive values.

#### Trend across groups

We know from the previous section that on moving down the group of s- and p- block elements in the periodic table, the effective nuclear charge remains almost steady, but there is a general increase in atomic radius due to increase in the value of the principal quantum number n. As a result, the electron affinity generally decreases down any group in the periodic table. This is evident from the values given listed in Table 1.1 values of electron affinities of second row non metals that is B, C, N, 0, F are however against the general trend, being smaller than those of corresponding elements that is Al, Si, P, S, Cl of Period 3. This is apparently an indirect result of the small size of the atoms of these elements that is B, C, N, 0, F. Thus, the crowding of electrons in the smaller outer shell of an atom of an element of Period 2 makes mutual repulsion of electrons substantially greater than that in the relatively larger outer shell of an atom of an element of Period 3. Therefore even though an electron added to an atom of an element of Period 2 is closer to the nucleus than one added to an atom of an element of Period 3, the greater inter electronic repulsion in a smaller shell leads to a lower electron affinity.
4.0 CONCLUSION

In conclusion, we can say the electron affinity of an element determines how easy or difficult it is for that element to participate in a chemical reaction.

5.0 SUMMARY

In this unit, you have studied the following:

- that the electron affinity of an element is the energy released or absorbed when an electron is added to a gaseous atom in its ground state
- that it is affected by three factors viz: atomic radius, effective nuclear charge and electronic configuration
- that the electron affinity varies across periods and groups in the periodic table.

SELF-ASSESSMENT EXERCISE

i. List three factors that affect electron affinity?
ii. Why does the electrons affinity generally deceased down a group.

6.0 TUTOR-MARKED ASSIGNMENT

1. Explain why the second electron affinity values are always positive.
2. Explain the difference between electron affinity and electro-negativity.

7.0 REFERENCES/FURTHER READING


UNIT 2 ELECTRONEGATIVITY

1.0 INTRODUCTION

Earlier, you studied ionisation energies and electron affinities of isolated gaseous atoms. These quantities are a measure of the tendency of isolated atoms to lose or gain electrons. In this unit, you are going to learn about electronegativity of an element. The electronegativity of an element is a measure of the power of an atom in a molecule to attract shared electrons to itself. Unlike ionisation and electron affinity, it is not a directly measurable physical quantity but rather, a theoretical concept for which several numerical scales have been developed.

In this unit, the three most important scales that have been developed for measuring electro-negativity are going to be discussed.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- define the concept of electronegativity
- calculate, given all the necessary impact, the electronegativity of an element, using Pauling, Mulliken-Taffe and Alfred-Rochow electro negativity scales
- discuss with at least eighty percent accuracy periodicity in electro negativity.
3.0 MAIN CONTENT

3.1 Pauling Electronegativity Scale

As you know in homonuclear diatomic molecule like A₂ (A-A) and B₂, (B-B) the electron pair is equally shared between the atoms bonded together. In a heteronuclear diatomic molecule of the AB type however, the situation is quite different. In the process of formation of a bond between A and B. Atom A slowly start stripping off its electrons thereby becoming a partially cationic species (it becomes partially positively charge). As the positively develops on A, its tendency to attract electron increases. Meanwhile, a similar process also takes place on atom B, thus both atoms A and B start attracting the electron pair towards themselves. They continue doing so until the tendencies of both the atoms in the bonded state to attract the electron pair towards themselves balance.

If one of the atoms say B, has higher tendency to attract the electron pair toward itself compared to that the other atom say A, the electron pair will spend more time on B than A. Partial charges will thus be generated on A and B. This tendency of attracting the shared electron pair toward itself by an atom a molecule has been termed the electro negativity of the element

The concept of electro negativity was first developed by Pauling. He defined it on the basis of the patterns desirable in the single bond energies of elements which were derived from the thermo chemical data. He realized that bond energy, $E_{A\text{-}B}$ between two unlike ion like atom A and B is greater than $\sqrt{E_{A\text{-}A}} \sqrt{E_{B\text{-}B}}$ where $E_{A\text{-}A}$ and $E_{B\text{-}B}$ are bond energies of A-A and B-B homonuclear bonds. He assigned the course of this excess bond energy $E_{A\text{-}B} - \sqrt{E_{A\text{-}A}} \sqrt{E_{B\text{-}B}}$ to the electrostatic attraction between partially charged atoms are separated due to difference in electron attracting tendencies of A & B that is the difference in the electro negativity of A and B. He was able to derive a relationship which can be stated as:

$$\Delta X_{A\text{-}B}^- = \sqrt{E_{A\text{-}B} - \sqrt{E_{A\text{-}A}} \sqrt{E_{B\text{-}B}}}$$

Where $X_A$ is the electro negativity of element A and $X_B$ is that of element B.
Knowing bond energies, it is possible to calculate the difference between the electro negativities of the two elements. This formula only gives the difference in the electro negativities of the two elements and not the absolute value assigned to a particular element. The largest electro negativity difference is that between fluorine the most electronegative element and calcium, the most electropositive element that came out to be 3.3. Pauling assigned arbitrarily a whole number value 4.0 for fluorine so that values of electro negativity of all elements remain positive. Table 2.1 shows values of electronegativity of different elements (bold faced) as calculated by Pauling using his Formulae.

### 3.2 Mulliken — Jaffe Electronegativity Scale

Mulliken defined electronegativity as the mean value of first ionisation energy and first electron affinity. Both quantities are given positive values if loss of electron involves absorption of energy and negative values if gain of electron involves release of energy. Thus electronegativity $X_A$ of atom A is given by the following relationship.

$$X_A = \frac{(I)_A + (E_A)_A}{2}$$

According to this relationship, a very electronegative element has very high ionisation energy. So it will be difficult to remove its electrons. It also has a very high electron affinity. Hence, a very stable species results when electrons are added. On the other hand, an element of low electro negativity will have a low ionisation energy and low electron affinity. So it loses an electrons readily and has little tendency to pick up electron.

It is very difficult to measure electron affinity for all elements (as we learned in Unit 1). Therefore this method is not universally applicable. The electro negativity values on Mulliken Scale are about 2.8 times those of Pauling's values. The trends in the variation of electronegativity are however the same.

### 3.3 Alfred—Rochow Electronegativity Scale

According to Alfred Rochow, electronegativity is equated to the force of attraction between an atom and the electron separated by a distance equal to the covalent radius of the atom. The force of attraction is expressed according to coulomb's law as:

$$F = \frac{Ze^2}{r^2}$$
Where Z* is Slater's effective nuclear charge e the electronic charge and r- the covalent radius.

**Table 2.1 Electronegativity Values of Element**

Electronegativity is a measure of the attraction that an atom has for electrons in a bond it has formed with another atom. The ability of an atom to attract electrons depends upon the charge on the atom and the hybridisation of the atom. An atom which has acquired a positive charge will tend to attract electrons to it more readily than will a neutral atom. In turn, a negatively charged atom will be less attractive to electron than a neutral atom.

Hybridisation also affects electronegativity because of lower energy and hence, greater electron attracting power of s-orbital. Thus hybrid orbitals having greater s-character possess higher electronegativity. An atom in sp hybridised state will be more electronegative than the same atom in Sp² hybridised state which will in turn be more electronegative than the same atom in SP³ hybridised state. Thus the carbon atom in CH₄, C₂H₄ and C₂H₂ has different values of electronegativity.
So you can say that electronegativity is not a constant quantity. All the electronegativity scales give only average values of electronegativities of element in different bonding environments.

All the electronegativity scales we studied give only relative values of electronegativities. These values are nevertheless useful in making quantitative comparisons between elements. Electronegativities can be used to predict the value of the bonding that a compound will have. The larger the difference between the electronegativities of the two elements, the more polar will be the bond between these elements. An electronegativity difference of about 107 corresponds to a partial ionic character of about 50%. So, a bond can be considered predominantly ionic, if the difference in the electronegativities of the bonded is more than 1.7. On the contrary, a difference in the range of 0.4 to 1.7 results in a covalent bond's partial ionic characters or polar covalent bond.

### 3.4 Periodicity in Electronegativity

Electronegativity values of elements show fairly discernible periodic trend throughout the periodic table. The trend is similar to that of ionisation energies. Thus, as expected, electronegativity of elements increases sharply across a row of S and P-block element. This is as a result of the sharp increase in effective nuclear charge of these elements example from lithium to fluorine. However, across a series of transition elements, the increase in electronegativity is much smaller. This is because the additional electron is being added to an inner shell which provides relatively good shielding for the outer electron from the nucleus. On moving down a group of representative elements, for example in the lithium group, there is a general decrease in electronegativity. The decrease is relatively small except between the first two elements.

The much greater electronegativity of lithium row elements correlates well with their small size. As expected, the elements of Period 4 from gallium onwards that is Ga, Ge, As, Se and Br have greater electronegativities than would be expected by extrapolation from values for the first two elements in the respective groups. This is due to the insertion of transition elements because of which the effective nuclear charge of these elements is greater than that, if the transition elements were not there. Similarly, the presence of the lanthanide elements is responsible for greater electronegativity of the elements of 5d series than would be expected by extrapolation from values of the elements of 3d and 4d series.
4.0 CONCLUSION

Let us conclude this unit by pointing out that, studies of ionisation energy and electron affinity show us the tendency of elements to loose or gain electron while going into a relationship. In this unit on the other hand, we studied the power of these elements while in a relationship with each other to attract to it the electrons in the bond between them.

5.0 SUMMARY

In summary, you have learnt the following from this unit:

- electronegativity is the tendency of an atom to attract toward itself the shared electron pair of a bond in which it is involved
- electronegativity can be measured by using any of the three scale developed by either, Pauling, Mulliken or Alfred Rochow
- the measurements vary according to the method used in measuring it
- the values of electronegativity vary across periods and groups of the periodic table.

SELF-ASSESSMENT EXERCISE

Name the least and most electronegativity elements in the periodic table.

6.0 TUTOR-MARKED ASSIGNMENT

Explain why the electro negativity values of noble gases are zero while those of halogens are the highest in each period.

7.0 REFERENCES/FURTHER READING


UNIT 3  HYDROGEN

CONTENTS

1.0  Introduction
2.0  Objectives
3.0  Main Content
   3.1  Position of Hydrogen in the Periodic Table
   3.2  Isotopes
   3.3  Deuterium compounds
   3.4  Tritium
   3.5  Ortho and Para Hydrogen
4.0  Conclusion
5.0  Summary
6.0  Tutor-Marked Assignment
7.0  References/Further Reading

1.0  INTRODUCTION

In the Units 1 and 2, you studied the development of the periodic table and periodicity in the properties of elements. I am sure that you noticed the fact that the very first element in the periodic table is hydrogen. The hydrogen atom consists of only one proton and one electron. In spite of that, hydrogen forms more compounds than any other element. In addition, it is the most abundant of all the elements in the universe (73.9% by weight). It is also the principal element in the solar atmosphere. However although hydrogen is very much in abundance (0.14% by weight) on earth, it exists only in the combined state.

Seeing how important hydrogen is and its peculiarity of being the first element in the periodic table, you will like to know more about hydrogen. In this unit, you will be studying some important aspects of the chemistry of hydrogen. You will also be studying its position in the periodic table, its isotopes and other forms.

2.0  OBJECTIVES

At the end of this unit, you should be able to:

- justify the position of hydrogen in the periodic table
- describe isotopes of hydrogen
- differentiate between Ortho and Para forms of hydrogen.
3.0 MAIN CONTENT

3.1 Position of Hydrogen in the Periodic Table

The position of hydrogen in the periodic table is of particular interest. Hydrogen is the first element of the periodic table, with an electronic configuration of \(1s^1\). This electronic configuration is similar to the outer electronic configuration of the alkali metals (\(ns^1\)). On the other hand, like halogens, it is one electron short of the corresponding inert gas helium (\(1s^2\)). Hydrogen, therefore, show some properties similar to alkali metals, while some others are similar to those of the halogens.

Like alkali metals, hydrogen forms halides, oxides and sulphides. We have seen in Unit 5 of Module 1 that the alkali metals have tight tendency of losing their outer most electron to form \(m^+\) ion. Hydrogen also forms \(H^+\) ion, but it does not do so, under normal conditions, because the ionisation energy of hydrogen (1312 kg mol\(^{-1}\)) is much higher than that of the alkali metals, e.g. Li-520; Na-495; k kg mol\(^{-1}\).

With high ionisation energy, hydrogen resembles halogens. (Their first ionisation energies are fluorine (1618 kg mol\(^{-1}\)) chlorine (1255 kg mol\(^{-1}\)) bromine (1142 kg mol\(^{-1}\)) and iodine (1007 kg mol\(^{-1}\))). Due to its high ionisation energy, hydrogen forms large numbers of covalent compounds by sharing a pair of electrons. Hydrogen like halogens, forms a diatomic molecule by sharing a pair of electrons between the two atoms. By picking up an electron, hydrogen forms the hydride ion (\(H^-\)), just like the halogens form the halide ion (\(X^-\)). From the previous discussion, it is clear that hydrogen resembles both the alkali metals as well as the halogens. So hydrogen can be placed with either of them in the period table. However conventionally, it is kept along with the alkali metals in Group 1 in the periodic table.

3.2 Isotopes of Hydrogen

Atoms of an element which have the same atomic number but different mass number are called Isotopes. Hydrogen has three different isotopes having mass numbers 1, 2, and 3 called ordinary hydrogen or Protium \(^1\)H, deuterium (D) or \(^2\)H and Tritium (T) or \(^3\)H respectively. These Isotopes differ from one another in respect of the presence of neutrons. Ordinary hydrogen has no neutrons, deuterium has one and tritium has two neutrons in the nucleus (Figure 3.1).
Fig. 3.1 Isotopes of Hydrogen

Deuterium is also called heavy hydrogen. These Isotopes have the same electronic configuration and therefore their chemical properties are almost the same. The only difference is in the rate of reactions. For example, protium has a lower energy of activation than deuterium in its reaction with halogens and therefore, reacts faster.

The physical properties of hydrogen, deuterium and tritium differ considerably due to their large mass differences. You can study more on the important physical properties of hydrogen, deuterium and tritium.

3.3 Deuterium Compounds

Naturally occurring hydrogen contains 0.0156% deuterium. Like water, (H₂O) which is the oxide of hydrogen, deuterium also forms an oxide, D₂O, which is known as Heavy Water. It can be obtained from ordinary water which contains 0.016% of deuterium oxide. This can be done either by fractional distillation or by electrolysis. Hydrogen is liberated more quickly than deuterium at the cathode and the residual liquid continuously gets richer in deuterium content on prolonged electrolysis of water. Deuterium oxide is used as a moderator in nuclear reactions.
since it slows down neutrons quickly. Physical properties of $\text{H}_2\text{O}$ and $\text{D}_2\text{O}$ and also differ from each other as in the case of $\text{H}_2\text{D}_2$. Hydrogen and deuterium are obtained by similar methods.

Many deuterium compounds, similar to those of hydrogen, are obtained from $\text{D}_2\text{O}$

We can also employ exchange reactions like those given below for the preparation of deuterium compounds:

3.4 Tritium

Tritium differs from the other two isotopes of hydrogen in being radioactive. Naturally occurring hydrogen contains nearly 10-15% tritium. The concentration of tritium increased by over a hundred fold when thermonuclear weapon testing began in 1954 but is now subsiding again as a result of the ban on atmospheric weapon testing.

Tritium was first obtained synthetically by the bombardment of deuterium compounds such as ($\text{ND}_4\text{SO}_4$) with fast deuterons.
It is now prepared on a large scale by irradiation of lithium-6 in the form of Li/Mg alloy of LiF with slow neutrons in a reactor.

The following reaction occurs in nature:

Tritium is radioactive and decays by emission of a beta — particle. Its half life period is 12.3 years

Tritium can be easily incorporated into biological molecules because it behaves chemically, just like ordinary hydrogen. The radiation tritium gives off within an organism as a result of its decay can cause many diseases including cancer.

3.5  Ortho and Para Hydrogen

Ortho and Para are two different forms of hydrogen molecule. These different forms arise as a result of differences in the direction of nuclear spin. When two hydrogen atoms combine to form a molecule, there are two possibilities. The two nuclei will either spin in the same direction (parallel spins) to give the form called Ortho Hydrogen, or they would spin in opposite directions to give Para Hydrogen. (Figure 3.2). This phenomenon is known as Spin Isomerism.
Due to spin isomerism, differences in the internal energy of the two forms of hydrogen arise, causing differences in the physical properties like boiling point, specific heat and thermal conductivity. Para hydrogen has a lower internal energy than that of ortho hydrogen. Hydrogen gas is an equilibrium mixture of Ortho and Para hydrogen. The ration of Ortho to Para hydrogen varies with temperature as shown in Figure 3.3.

Evidently, this ratio increases with the rise in temperature up to a temperature of about 300K (27°C) and remains constant thereafter. The percentage of hydrogen at 300K and above is 75%. This means it is not possible to get 100% ortho hydrogen at any temperature.

The equilibrium mixture of Para and Ortho hydrogen changes to almost 100% Para hydrogen when cooled to nearby absolute zero. Para hydrogen is stable for weeks in the absence of catalysts like activated charcoal, Fe, Ni Pt, O₂, NO₂ etc. these catalyse the conversion of Para to Ortho hydrogen, deuterium and tritium also exhibit spin isomerism and exist in Ortho and Para forms.
However the ratio of Ortho to Para forms in deuterium and tritium is different from that in hydrogen. The variation of Ortho/Para ratio at different temperatures is also different if you look carefully in Figure 3.3; you observe that tritium resembles hydrogen more closely than deuterium in this respect.

4.0 CONCLUSION

We can conclude this unit by observing that hydrogen holds a unique position in the periodic table. It is the first element in the periodic table and also exhibits properties of the alkali metals as well as that of the hydrogen. Hydrogen also exists in different forms thus explaining some of its properties.

5.0 SUMMARY

In this unit, you studied the following:

- hydrogen with an electronic configuration of Is' occupies a unique position in the periodic table
- hydrogen has Isotopes known as deuterium tritium and the normal hydrogen we know
- the weights of these Isotopes are H = 1 Deuterium (D)2 and Tritium (T) 3
- hydrogen has in addition to H, D and T two other forms, Ortho and Para Hydrogen
- these two forms arise as a result of differences in spins of the two molecules that make up the H2 molecule.

SELF-ASSESSMENT EXERCISE

Write ‘T’ for true and ‘F’ for false in the given books for the following statements about Ortho and Para forms of hydrogen:

i. Ortho and Para hydrogen are different due difference in their nuclear spins.
ii. Ortho and Para hydrogen are different due to difference in their electron spins.
iii. Physical properties Ortho and Para hydrogen are similar.
iv. Para hydrogen is more stable at lower temperatures.
v. At look percentage of Ortho hydrogen in the mixture is 70%.
6.0 TUTOR-MARKED ASSIGNMENT

1. Why does hydrogen resemble alkali metals?
2. Explain the formation of hydride ion.
3. Why are the chemical properties of Isotopes similar?

7.0 REFERENCES/FURTHER READING


UNIT 4 MANUFACTURE OF HYDROGEN

CONTENTS
1.0 Introduction
2.0 Objectives
3.0 Main Content
   3.1 Manufacture of Hydrogen
   3.2 Properties of Hydrogen
   3.3 Uses of Hydrogen
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 References/Further Reading

1.0 INTRODUCTION

In the last unit (Unit 3), you studied the unique position of hydrogen in the periodic table, you also studied the various isotopes of hydrogen and also the two forms of the hydrogen molecule.

In this unit, you shall be studying the ways hydrogen is manufactured, its properties and uses.

2.0 OBJECTIVES

At the end of this unit you should be able to:

- list at least three methods used in the manufacture of hydrogen
- explain using appropriate equations and examples, what takes place during the process of the manufacture of hydrogen
- list and explain at least five properties of hydrogen
- list and explain with eighty percent degree of accuracy at least four uses of hydrogen.

3.0 MAIN CONTENT

3.1 Manufacture of Hydrogen

Most if not all the methods used to manufacture hydrogen make use of water. Water is a natural abundant source for the manufacture of hydrogen. Water can be reduced to hydrogen either chemically or electrically.
(a) **Manufacture by chemical means**

The common reducing agents are Coke, (obtained from the destructive distillation of coal), carbon monoxide or hydrocarbon.

(i) The hydrogen is manufactured by allowing steam to react with red hot coke at about 1250k

\[
C + H_2O \xrightarrow{1250K} CO + H_2
\]

(ii) The mixture of CO and H₂, known as **water gas**, is also called **synthesis gas** because it is used for the synthesis of methanol and other hydrocarbons.

(iii) Hydrogen is also produced by the reaction of gas (chiefly methane) with steam in the presence of nickel catalyst.

\[
CH_4 + H_2O \xrightarrow{N_i \text{ catalyst}, \ 920-1170K} CO + 3H_2
\]

Similar reaction can occur with other hydrocarbons. In both cases above CO is converted to CO₂.

\[
[CO + H_2O \xrightarrow{\text{Catalyst}, \ 450-650K} CO_2 + H_2]
\]

Hydrogen and carbon dioxide can easily be separated from each other by bubbling the gas mixture through water in which CO₂ is fairly soluble and H₂ is virtually insoluble.

At higher temperatures, in the presence of catalysts (silica, alumina) hydrocarbon break up and rearrange in what are called cracking reactions. These reactions, which are used in refining of petroleum, produce hydrogen as a bi-product. One example of simple cracking reaction is cracking of propane.
(b) Manufacturing by electrolysis

i. Electrolysis of acidified water using platinum electrodes is a convenient source of hydrogen (and oxygen). On a large scale, very pure hydrogen (>99.95%) can be obtained from the electrolysis of aqueous solution of barium hydroxide between nickel electrodes. Hydrogen obtained by electrolysis of water is relatively expensive because of the cost of electrical energy.

ii. Hydrogen can however be obtained economically as a by-product in the electrolysis of brine during the manufacture of sodium hydroxide. During electrolysis, there is a competition at the anode between the oxidation of chloride ion and the oxidation of water.

When a concentrated salt solution (brine) is used, the first reaction preferentially takes place at the cathode, the reaction is the reduction of water because it is more easily reduced than Na⁺.

The anode and the cathode reactions are combined to give the reactions thus:
In the laboratory hydrogen can be prepared by the reaction of water or dilute acids on electropositive metals such as alkali metals alkaline earth metals, the metals of group 12 (e.g. Zn) and the lanthanides. The reaction can be explosively violent with alkali metals (e.g. Rb) convenient laboratory methods employ sodium amalgam or calcium with water or zinc and tin with hydrochloric acid.

### 3.2 Properties of Hydrogen

(i) Hydrogen is the lightest element known. It is colourless, odourless and tasteless gas. The hydrogen molecule is

(ii) Thermally stable and has little tendency to dissociate at normal temperatures, the reaction \( \text{H}_2 \rightarrow \text{H}(g) = +436 \text{kg mol}^{-1} \) being an endothermic one. However, at high temperature in an electric or under ultraviolet light, \( \text{H}_2 \) does dissociate. The atomic hydrogen produced, exists for less than half a second after which it recombines to give molecular hydrogen and liberates a large amount of energy (436kg mol\(^{-1}\)), in form of heat. Most of the transition metals catalyse the combination reaction of hydrogen.

(iii) Atomic hydrogen is a powerful reducing agent and reduces copper, silver and mercury salts to the metallic state of. e.g.

\[
\text{AgNO}_3 + \text{H} \rightarrow \text{Ag} + \text{HNO}_3
\]

(iv) It combines with alkali metals to form hydrides e.g \( \text{Na} + \text{H} \rightarrow \text{NaH} \)

(v) It reduces sulphur to hydrogen sulphide \( \text{S} + 2\text{H} \rightarrow \text{H}_2\text{S} \)

(vi) Carbon monoxide is reduced to formaldehyde.

\[
\text{CO} + 2\text{H} \rightarrow \text{HCHO}
\]

(vii) It also reacts with oxygen at room temperature to form hydrogen peroxide.

\[
\text{O}_2 + 2\text{H} \rightarrow \text{H}_2\text{O}_2
\]

(viii) \( \text{H} \) is used for wielding.

Atomic hydrogen is produced by passing ordinary hydrogen through electric arc maintained between two electrodes. The atoms set free are carried away by a stream of incoming hydrogen gas. These free atoms recombine at once on coming in contact with a metallic surface liberating a large amount of heat and thus arising temperature of the metal to say 4000 — 5000k. This
principle is utilised in the making of the 'atomic hydrogen welding torch' (Figure 4.1) It provides an opportunity of welding at a very high temperature but in a reducing atmosphere.

(ix) Despite the fairly high bond dissociation energy of the hydrogen molecules it is moderately reactive and forms strong bonds with many other elements. It reacts with almost all elements except the noble gases.

(x) Hydrogen reacts with alkali and alkali earth metals by accepting an electron to form conic hydrides; e.g. KH, CaH₂

(xi) With non-metals it forms covalent hydrides, e.g. NH₃, H₂O and HF.

![Fig. 4.1 Atomic Hydrogen Welding Torch](image)

(xii) Hydrogen is easily oxidised to water and; therefore it acts as a very good reducing agent in a variety of situations

(xiii) Hydrogen is used in metallurgy to reduce metal oxides to metals in cases where carbon cannot be used because the metal can form carbide. Such metals include Mo and W.

\[
\begin{align*}
WO + 3H₂ & \rightarrow W + 3H₂O \\
(NH₄)M₅O₄ + 3H₂ & \rightarrow M₅₂NH₃ + 4H₂O
\end{align*}
\]

(xiv) Hydrogen adds on the multiple bonds in organic compounds. In the presence of catalysts such as finely divided nickel, palladium
or mixtures of metal oxides, unsaturated organic compound are thus reduced to saturated compounds. For example:

\[
\begin{align*}
\text{CH}_3\text{CH} = \text{CH} \text{CH}_3 + \text{H}_2 & \xrightarrow{278\text{K}} \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_3 \\
2 - \text{butene} & \text{Pd, 1 atm} \\
\text{CH}_3\text{C} = \text{N} + \text{2H}_2 & \xrightarrow{673\text{K}} \text{CH}_3\text{CH}_2\text{NH}_2 \\
\text{Methylvicyanide} & \text{ZnO/\text{Cr}_2\text{O}_3} \\
\text{CH}_3\text{CH}_2\text{NH}_2 & \text{ethylamine}
\end{align*}
\]

(xv) Catalytic hydrogenation of unsaturated liquid vegetable oils to solid edible fats illustrates the industrial application of the reduction reactions; for example the reduction of an oleate (ester of oleic acid) to ester of stearic acid.

\[
\begin{align*}
\text{CH}_3\text{(CH)}_2\text{CH} = \text{CH}(\text{CH})_3\text{COOR} + \text{H}_2 & \xrightarrow{} \text{CH}_3\text{(CH)}_2\text{(CH)}_{16}\text{COOR} \\
\text{Olate} & \text{H}_2
\end{align*}
\]

(xvi) Hydrogen reacts with carbon monoxide in the presence of catalysts to form methanol.

\[
\begin{align*}
\text{CO} + 2\text{H}_2 & \xrightarrow{\text{Cobalt}} \text{CH}_3\text{OH}
\end{align*}
\]

This reaction is known as hydro-formulation reaction and is used in the industrial preparation of methanol.

### 3.3 Uses of Hydrogen

Some of the uses of hydrogen include the following:

1. The largest single use of hydrogen is in the syntheses of ammonia which is used in the manufacture of nitric acid and nitrogenous fertilizers.

2. Hydrogen is used in the hydrogenation of vegetable of oils and the manufacture of methanol.

3. In space crafts, hydrogen gas is used in fuel cells (fig 3.2) for generating electrical energy and for providing clean drinking water to the astronauts. In a fuel cell, electrical energy is generated by the reaction of hydrogen leaf. This is sometimes
called "cold combustion" A hydrogen oxygen fuel cell may be having an alkaline or acidic electrolyte.

The one in Figure 4.2 has porous carbon electrodes and KOH as electrolyte.

![Figure 4.2 A Hydrogen-Oxygen Fuel Cell with KOH Electrolyte and Porous Carbon Electrodes](image)

The half cell reactions are given below:

With acidic electrolyte, the half-cell reacts irons are:

\[
\begin{align*}
    
    \text{H}_2(\text{g}) & \rightarrow 2\text{H} + 2\text{e}^- \\
    \frac{1}{2} \text{O}_2(\text{g}) + 2\text{H}^+ + 2\text{e}^- & \rightarrow \text{H}_2\text{O}(	ext{l}) \\
    \text{H}_2(\text{g}) + \frac{1}{2} \text{O}_2(\text{g}) & \rightarrow \text{H}_2\text{O}(	ext{l})
\end{align*}
\]

With acidic electrolyte, the half-cell reacts irons are:

\[
\begin{align*}
    \text{H}_2(\text{g}) & \rightarrow 2\text{H} + 2\text{e}^- \\
    \frac{1}{2} \text{O}_2(\text{g}) + 2\text{H}^+ + 2\text{e}^- & \rightarrow \text{H}_2\text{O}(	ext{l}) \\
    \text{H}_2(\text{g}) + \frac{1}{2} \text{O}_2(\text{g}) & \rightarrow \text{H}_2\text{O}(	ext{l})
\end{align*}
\]

We can see from the equations above that the electromotive force of the cell remains the same whether we use alkaline or acidic electrolyte. This
is because we are using the same reactants at the electrodes in both cases.

Fuel cells have several advantages over other sources of energy. Firstly in a fuel cell unlike in the dry cell or storage battery (which requires recharging also), the cathode and anode reactants are continuously supplied so that energy can be indefinitely withdrawn from it. Secondly, in a fuel cell energy is extracted from the reactants under almost ideal conditions. Therefore, the thermodynamic efficiency of the fuel cells are higher than of most of the ordinary combustion processed. Fuel cells have efficiencies approaching 75% whereas power plants that burn fuels have efficiencies of only about 40%. Combustion of hydrogen is a highly exothermic reaction and produces no pollutants:

$$2 \text{H}_2(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{H}_2\text{O}(l), \quad \Delta H = -\text{572kJ}$$

Liquid hydrogen is therefore, used as a rocket fuel.

It is now clear that world reserves of fossil fuels like coal, oil and gas are finite, so they cannot last forever. Nuclear and hydroelectric power cannot meet the world's energy needs. Moreover, these resources post a danger to the world's environment. With these facts in mind there is now an active search for alternative source of energy. In addition to solar power, hydrogen is being considered a potential fuel for the future.

Hydrogen as a fuel has many advantages over the conventional fossil fuels and electric power. It is available in unlimited quantities in sea water. It is pollution free because the major product of its combustion is water with only traces of nitrogen oxides. It releases greater energy per limit weight of fuel in comparison to gasoline and other fuels. Hydrogen can be transported as a gas in high pressure pipelines, as a liquid in tankers and even as a solid in form of metal hydrides. Unlike electricity hydrogen can be stored and used when needed.

Hydrogen, however, has the following disadvantages viz:

Hydrogen like electricity is a secondary source of energy because it is produced using energy from a primary source such as coal, nuclear fission or sun.

Preparation of hydrogen through electrolysis is not economical at present in fact more energy has to be spent in electrolysis of water than what can be liberated by burning hydrogen as a fuel.
Decomposition of water by solar energy in presence of catalysis is known as photochemical decomposition of water. Using catalysts scientists in France have been able to achieve the efficient decomposition of water under visible and ultra violet light. If this process can be made industrial, a convenient method of converting solar energy directly to a useful form of store chemical energy will be available.

4.0 CONCLUSION

We can conclude this unit by observing that the major material for the manufacture of hydrogen is water. Since water is abundant, the production of hydrogen should be simple. However, it is not efficient to produce hydrogen by electrolysis. This is non economical because of the cost of electricity. The single most important use of hydrogen is in the manufacture of ammonia which is used in the production of nitric acid and nitrogenous fertilizers. Hydrogen is also used in space crafts as a source of fuel cell.

5.0 SUMMARY

In summary, you have studied the following in this unit:

- the major source for the manufacture of hydrogen is water
- there are two primary methods for the manufacture of hydrogen:
  a. manufacture by chemical means
  b. manufacture by electrolysis
- even though manufacturing by electrolysis would have been preferred it is too expensive, because of the cost of the energy needed for electrolysis
- the largest use of hydrogen is in the manufacture of ammonia, which is used to manufacture nitric acid and nitrogenous fertilizers
- hydrogen is made use of in the production of fuel cells for space crafts
- hydrogen is also used in extraction of metal and hydrogenation of vegetable oils.

SELF-ASSESSMENT EXERCISE

i. Complete the following chemical reactions which take place during the formation of hydrogen.
ii. While five important industrial uses of hydrogen

6.0 TUTOR-MARKED ASSIGNMENT

Write chemical equation for the following reactions:

1. Formation of methanol from coal.
2. Reduction of methyl cyanide.
3. Conversion of Oleic acid into stearic and
4. Reduction of ammonium molybdate to molybdenum.

7.0 REFERENCES/FURTHER READING


UNIT 5     IONIC OR SALT-LIKE HYDRIDES

CONTENTS

1.0      Introduction
2.0      Objectives
3.0      Main Content
        3.1      Ionic or Salt - Like Hydrides
        3.2      Covalent Hydrides
        3.3      Metallic Hydrides
4.0      Conclusion
5.0      Summary
6.0      Tutor-Marked Assignment
7.0      References/Further Reading

1.0      INTRODUCTION

In the previous units, you became aware of the fact that hydrogen forms more compounds than any other element. You also saw that despite the fairly high bond dissociation energy of the hydrogen molecule, it is moderately reactive and forms strong bonds with many other elements. You also learnt that it forms ionic and covalent hydrides with metals and non metals respectively.

In this unit you will be studying the formation of hydrides, their types and characteristics of the hydrides.

2.0      OBJECTIVES

At the end of this unit, you should be able to:

- list the three classes of hydrides
- discuss for each of the different types of hydrides and their properties.

3.0      MAIN CONTENT

As already pointed out at the beginning of this unit, hydrogen combines with a number of elements to form hydrides. As electronegativity of the element increases the stability of the hydrides also increases. Three types of hydride compounds are formed depending upon the electronegativity of the element. There are classified into:

i.     Ionic or salt like or salric hydrides
ii.    Covalent or molecule hydrides
iii.   Metallic or non-stoichiometric hydrides.
3.1 Ionic or Salt — Like Hydrides

These are formed by metals with low electronegativity values and are more electropositive with respect to hydrogen. These hydrides are formed by transfer of an electron from the metal to the hydrogen atom. Hydride ion is a peculiar chemical species and in contrast to proton which has small size, it is unusually large. It is larger than any of the negative ions except iodide.

The reason for this apparent paradox lies in the lack of control by a single nuclear proton over two naturally repelling electrons.

Alkali and Alkaline-earth metal of Groups 1 and 2 are sufficiently electropositive and force the hydrogen atom to accept an electron to form the hydride ion, if eg Lithium hydride Li H- and calcium hydride Ca2+ (l) 2.

Ionic hydrides are formed by heating metals in hydrogen at 973K. Ionic hydrides are white crystalline solids. They have high melting points and conduct electricity in liquid state, liberating hydrogen at the anode. Their density is higher than that of the metal.

They are powerful reducing agents especially at high temperatures e.g.

\[
\begin{align*}
2 \text{CO} + \text{NaH} & \rightarrow \text{HCOO Na} + \text{C} \\
\text{SiCl}_4 + 4 \text{NaH} & \rightarrow \text{SiH}_4 + 4 \text{NaCl} \\
\text{PbSO}_4 + 2 \text{CaCH}_2 & \rightarrow \text{PbS} + 2 \text{Ca(OH)}_2
\end{align*}
\]

Li+ H and Na+ H are used in making valuable reducing agents like lithium aluminium hydride (LiAlH4) and sodium-boro-hydride (NaBH4). The complex hydrides are frequently used in the reduction of aldehydes, ketones, acids and their derivatives to give alcohols.

\[
\begin{align*}
2 \text{COOH} & \overset{\text{LiAlH}_4/\text{NaBH}_4}{\rightarrow} \text{R} - \text{CH}_2 \text{OH} \\
>\text{C} = \text{O} & \overset{\text{LiAlH}_4/\text{NaBH}_4}{\rightarrow} >\text{CH} - \text{OH}
\end{align*}
\]
3.2 Covalent Hydride

These bonds are formed by elements of comparatively higher electronegativity such as the P-block elements and Be and Mg. The bonds formed in this class of hydrides are mostly covalent in character but in some cases, for example, in HF, the bond may be partially ionic.

The covalent hydrides can be prepared either by direct reaction of non-metals with hydrogen under suitable conditions or by the reaction of H2O or acids or nitrides, carbides, bonides, silicides, stanides of alkali and alkaline earth metals or by the reduction of halides. These are illustrated by the following reactions:

These hydrides have molecular lattice made up of individual saturated covalent molecules, with only weak van der Waals forces and in some cases along with hydrogen bonds. This accounts for their softness, low melting and boiling points, their volatility and lack of conductivity. Some covalent hydrides are unstable in the presence of air, e.g. stannane, SnH4.

Some covalent hydride hydrides of Groups 2 and 13 are electron deficient. These have structures between ionic and covalent hydrides. These are either dimeric, e.g. boron hydride (B₂H₆), or polymeric, e.g. beryllium hydride (BeH₂)n, magnesium hydride (MgH₂)n and aluminium hydride (AlH₃)n.

3.3 Metallic Hydrides

When heated, hydrogen reacts with many transition metals (lantharindes and actinides) to form metallic hydrides. Most of these have metallic appearance and are less dense than the parent metal. They all conduct heat and electricity though not as well as the parent metal.
They are almost always non-stoichiometric, being deficient in hydrogen. For example, Ti H$_{5.8}$, VH$_{0.56}$, CrH$_{17}$ NiH$_{0.6}$, TaH$_{22.76}$, LaH$_{28}$ YbH etc. Most of these hydrides are stable to water up to 375K but are quantitatively decomposed by acids and show some reducing properties.

Formerly, these hydrides were formed as interstitial compounds in which hydrogen was thought to be accommodated in the interstices in the metal lattice producing distortion but no change in its type. But recent studies have shown that except for hydrides of nickel palladium, cerium and actinium, other hydrides of this class have lattice of a type different from that of the parent metal. For example, the hexagonal close packed lattice of some lanthanides is transformed to a face-centred cubic lattice in their dihydrides. As pointed out earlier, these interstitial hydrides are poorer conductors of electricity, exhibit less Para magnetism and are more brittle than the parent metal. These characteristics suggest that hydrogen is present in the metal lattice as hydrogen atoms rather than as hydrogen molecules. The single electron of hydrogen is paired with an electron of the metal, thereby reducing the extent of metallic bonding. Breaking of the H-H bond is in agreement with the fact that there metals catalyse reactions of hydrogen.

4.0 CONCLUSION

You can conclude this unit by putting the fact that despite the fairly high bond dissociation energy of the hydrogen molecule, it reacts with certain elements to form three types of hydrides; ionic, covalent and metallic hydrides.

5.0 SUMMARY

In this unit, you learnt the following:

- hydrogen leads with certain metals to form hydrides
- these hydrides are classified into three viz ionic or salt —like hydrides, covalent hydrides and metallic hydrides
- ionic hydrides are powerful reducing agents, covalent hydrides are soft have low melting points and are poor conductors of electricity.

SELF-ASSESSMENT EXERCISE

i. Silt is an example of which of the following type of hydrides
   (a) Ionic
   (b) Interstitial
   (c) Metallic
   (d) Covalent.
ii. List three properties of ionic hydrides.

6.0 TUTOR-MARKED ASSIGNMENT

What types of bonding do you expect of?

1. Sodium hydride
2. Methane
3. Ammonia

7.0 REFERENCES/FURTHER READING


**UNIT 1 HYDROGEN BONDING**

**CONTENTS**

1.0 Introduction
2.0 Objectives
3.0 Main Content
   3.1 Hydrogen Bonding
   3.2 Intermolecular Hydrogen Bonding
   3.3 Intra-Molecular Hydrogen Bonding
   3.4 Effects of Hydrogen Bonding
      3.4.1 Boiling Point and Melting Point
      3.4.2 Water Solubility
   3.5 Polarising Power of H⁺
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 References/Further Reading

**1.0 INTRODUCTION**

A very important aspect of the chemistry of hydrogen is its formation of hydrogen bonds. When hydrogen is covalently bonded to a highly electronegative element like F, O, or N, the electronegative element attracts the electron pair towards itself giving rise to an induced positive charge (5+) on the hydrogen atom and negative charge (5-) on the electronegative atom for example:
Ho + Xo where X is the electronegative atom. When this happens the hydrogen, due to its positive character, will attract another electronegative atom of the neighbouring molecule forming a bond. This bond is known as hydrogen bond. This is illustrated below

```
--------------- H δ⁺ X δ -------------- H δ⁺ X δ ---------------- H δ⁺ X δ ----------------

--- denotes covalent bond.
- denotes hydrogen bond.
```
In this unit, you will be studying the types of bonding, the effect of hydrogen bonding on boiling and melting points. Also, the effects of hydrogen bonding on water solubility of the compound containing hydrogen bonded elements.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- define hydrogen bonding
- list and discuss the two types of hydrogen bonding
- list and discuss the effect of bonding on boiling and melting points
- discuss the effect of bonding on the solubility of a substance
- discuss giving examples, the polarising power of H+.

3.0 MAIN CONTENT

3.1 Hydrogen Bonding

Hydrogen bond can be defined as the attractive force which binds hydrogen atom of one molecule with electronegative atom of another molecule, generally of the same compound.

The hydrogen bond energy is only about $7 - 59 \text{ kg mol}^{-1}$ compared to the normal covalent bond energy of $389 - 665 \text{ kg mol}^{-1}$ for H-N, H-O and H—F bonds. This hydrogen bond is much weaker than a covalent bond. Obviously, its length is also much more than the covalent bond. There are two types of hydrogen bonding. These are:

(i) Intermolecular hydrogen bonding
(ii) Intra-molecular hydrogen bonding.

3.2 Intermolecular Hydrogen Bonding

In this case, two or more molecules of the same element are involved in hydrogen bonding. Some common examples of intermolecular hydrogen bonding occurring between the molecules of the same compound are HF, H$_2$O, alcohols etc.
Examples of the intermolecular hydrogen bonding between two different kinds of molecules are as following.

3.3 Intra-Molecular Hydrogen Bonding

Intramolecular hydrogen bond is formed between two atoms of the same molecule. As a consequence of this, generally a five or six membered ring called chelate ring is formed.
You have seen above, that intramolecular hydrogen bonding takes place in molecules like 6-nitrophenol and salicylaldehyde.

Intramolecular hydrogen bonding does not take place in p-nitrophenol because of the large distance between the two groups (NO$_2$ and QH) in p — nitrophenol

This type of hydrogen bonding is not possible. It does however show the usual intermolecular hydrogen bonding.

It is significant to note that the vast majority of intra-molecular hydrogen bonding occurs where a five or six membered ring can be formed because of the stability associated with such rings.
3.4 Effects of Hydrogen Bonding

Hydrogen bonding plays a very significant role in determining the properties of compounds. In this section we shall discuss its effect on the melting point, boiling point and solubility in water.

3.4.1 Boiling Point and Melting Point

If you examine the values for the melting and boiling points shown in Figure 1.1, you will see that the melting and boiling points of the hydrides of Group 14 elements i.e. CH₄, SiH₄ and GeH₄ and SnH₄ show a general increase with increase in molecular weight. However, NH₃ in Group 15, HF in Group 17 and water have abnormally high melting and boiling points as compared to other hydrides in their respective groups in the periodic table.

This anomaly is explained on the basis of hydrogen bond formation. In compounds where the molecules are linked by hydrogen bonds, some extra energy is required to break the intermolecular hydrogen bond and this is responsible for their higher boiling and melting points.

Intra-molecular hydrogen bond however has the opposite effects. For example, in ortho-nitrophenol the groups present in ortho position are involved in intramolecular hydrogen bonding thus preventing the intermolecular hydrogen bond formation, i.e. association of the molecule. Due to the intramolecular chelated structure, o-nitrophenol is seen to be volatile where as p-nitrophenol is not.

3.4.2 Water Solubility

Solubility of a substance increases markedly when hydrogen bonding is possible between the solvent and the solute molecules. For example, lower alcohols like, methanol, ethanol etc are highly miscible with water due to the hydrogen bonding with water molecules.
3.5 Polarising Power of H\(^+\)

We know that the polarising power of a cation, i.e. its ability to distort or polarise an anion is directly proportional to its positive charge and inversely proportional to its size. We can also say that the polarising power of a cation is proportional to the ratio of its charge to its size. This ratio is known as the ionic potential of the cation. As the hydrogen cation, i.e. proton is vanishingly small, it has a very high ionic potential and a vast polarising power.

As result of this polarising power for protons, H\(^+\) hardly exists freely. They are generally found associated with other molecules. For example, with ammonia and water, these form species like NH\(^+\), H\(_3\)O\(^+\), SO\(_2\)H, HgO\(^+\), etc. The aquated proton species are represented as H\(^+\). H Enthalpy of formation of these aquated proton species is large (-1075kg mol\(^{-1}\)). It is mainly because of this reason that many covalent hydrides (H-X) are acidic in aqueous solution, i.e. they release H\(^+\) ions even though H — X bonds are often very strong in them.

4.0 CONCLUSION

We can conclude this unit by stating that hydrogen bond formation is a very important phenomenon and it helps in explaining the behaviour of some compounds. Variations in boiling and melting points and solubility of substances are explainable by the existence of hydrogen bonds.
5.0 SUMMARY

In summary, you have studied the following in this unit:

- hydrogen bond is defined as the attractive force which binds hydrogen atom of one molecule with the electronegative atom of another molecule, generally of the same compound
- there are two types of hydrogen bonds:
  (i) Intermolecular hydrogen bond
  (ii) Intramolecular hydrogen bond
- the hydrogen bonding in substances affect their melting points, boiling points and their solubility in water
- because of the high polarising power of $\text{H}^+$, it hardly exists freely. It always exists in association with other molecules such as in $\text{NH}_4^+ \text{H}_2\text{O}^+ \text{H}_3\text{O}^+$ etc.

SELF-ASSESSMENT EXERCISE

i. What is the effect of hydrogen bonding on the properties of $\text{H}_2\text{O}$, HF and $\text{NH}_3$?

ii. Why is $\text{H}_2\text{O}$ a liquid and $\text{H}_2\text{S}$ a gas at room temperature?

iii. Why are Group 2 metals harder and have higher melting points as compared to the Group 1 metals?

6.0 TUTOR-MARKED ASSIGNMENT

1. Which hydrogen bond do you expect to be stronger and why?
   (a) $\text{S}---\text{H}-----\text{O}$ or (b) $\text{O}------\text{N}-----\text{S}$

2. Explain why the boiling points of hydrogen halides follow the trend

3. When a hydrogen bond is symbolised by $\text{X-H}.........\text{Y}$, what do the solid and dotted lines represent? Which distance is shorter?

7.0 REFERENCES/FURTHER READING


UNIT 2   GENERAL PHYSICAL AND CHEMICAL CHARACTERISTICS OF THE ALKALI METALS

CONTENTS

1.0 Introduction
2.0 Objectives
3.0 Main Content
3.1 Alkali Metals
3.2 Occurrence
3.3 Extraction of Alkali Metals
3.4 Uses of Alkali Metals
3.5 Physical Properties
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 References/Further Reading

1.0 INTRODUCTION

You will no doubt recall that at the beginning of our studies we learnt how the efforts of leading scientists resulted in the formation of the periodic table as we know it today. You recall that the end product of that effort is the arrangement of the elements according to their atomic numbers. You also recall how in the last few units, we discussed that the properties of the elements in a periodic table are indeed a periodic function of their atomic numbers as stated by the periodic law. We also saw that elements belonging to the same groups have the same properties.

In earlier units, you studied hydrogen and learnt about its unique position. Hydrogen, you recall gas properties which resemble the Group 1 elements in some respects and the Group 17 elements in others.

In this unit, you will be studying the elements of Group 1, their occurrence extraction and uses. You will also be studying their physical properties.

The elements of Group 1 and 2 are called the S-block elements because the outermost electron(s) in these elements occupy the S-orbital. Group 1 elements consist of Li, Na, K, Rb, C and Fr. They are called Alkali Metals because they foam hydroxides which are strong Alkalis.
2.0 OBJECTIVES

At the end of this unit, you should be able to:

- list for each member number of the alkali group where they can be land
- give for each member at least one use
- discuss, the atomic size, density, melting and boiling points, ionisation energy and electro negativity of members of the group.

3.0 MAIN CONTENT

3.1 Alkali Metals

Alkali metals are useful as metals as well as in the form of their compounds.

3.2 Occurrence

The alkali metals are highly reactive so they do not occur in the free state in nature. They occur in the combined form in the earth’s crust in the following relative abundance: Sodium 2.27%, potassium 1.84% lithium, rubidium and calcium in trace amounts $1.8 \times 10^{-3}$ %, $7.8 \times 10^{-3}$ % and $0.26 \times 10^{-3}$ % respectively.

Sodium as sodium chloride is the most abundant metal in sea water (M1.08%). Lithium occurs in alumino silicate rocks, e.g. spodumene, LiAl(SiO$_3$)$_2$ and Lepidolite (Li, Na, K)$_2$ (F,OH)$_2$. Sodium in rock salt, NaCl, in Chile Saltpetre NaNO$_3$, and in Cryolite, Na$_3$AlF$_6$. Potassium in carnallite, KCl, in MgCl$_2$. 6H$_2$O, in Salpetre KNO$_3$ and in Kainite, kCl, MgSO$_4$. 3H$_2$O.

Rubidium and calcium are rare elements and generally occur in small quantities along with other alkali metals. For example, carnallite contains up to 0.94% rubidium chloride, Lepidolite, and containing about 0.2 to 0.7 per cent of calcium expressed as calcium oxide. Francium being a radioactive element with a very short half life period (21.8 minutes) occurs in very minute’s traces in nature.

3.3 Extraction of Alkali Metals

Lithium and sodium are extracted by electrolysis of their fused (molten) chlorides.
Potassium is obtained by the reduction of its chloride with sodium vapour. This reduction by Na appears to be contrary to the normal order of reactivity, K>Na. However, at about 1150K the following equilibrium is set up:

\[
\text{Na}(g) + K^+_{(L)} \rightleftharpoons \text{Na}^+_{(l)} + K(g)
\]

Since potassium is more volatile, it distils off more readily displacing the equilibrium to the right and allowing the forward reaction to proceed. Rubidium and Caesium, can be prepared by the reduction of their chlorides with calcium metal at 1000K under calcium pressure. Rubidium and caesium salts are obtained during the recrystallisation of other naturally occurring alkali metal salts. Francium is produced as a result of a — emission (1%) during the radioactive decay of actinium (Figure 2.1)

![Figure 2.1 Production of Francium as a Result of Decay](image)

### 3.4 Uses of Alkali Metals

The alkali metals are very good conductors of heat and electricity. However, due to their highly reactive nature they cannot be used for this purpose.

Sodium in polyethylene enclosed cables is used in some underground high voltage transmission applications.

Because of the high specific heat and thermal conductivity, liquid sodium is used as a coolant in nuclear reactors. You must have seen bright yellow lights on the streets and particularly on the highways. These are sodium vapour lamps and the light from them can penetrate far well.
Caesium has the distinction of being the metal from which electrons are ejected most easily on exposure to light. This phenomenon is called photoelectric effect. Photocells, which are a device for converting light into electricity, are based on this phenomenon.

Other every day uses of alkali metal compounds include the following:

Lithium in the form of lithium stearate is used for the production of lubricating greases.

The hydrides of lithium and sodium viz \( \text{LiAlH}_4 \) and \( \text{NaBH}_4 \) are used as reducing agents in synthetic organic chemistry. Lithium and potassium compounds are used in picture tubes of colour televisions.

Can you imagine food without common salt! Apart from being an essential constituent of food, sodium chloride has many other important industrial uses like in the manufacture of NaOH, \( \text{Na}_2\text{CO}_3 \), \( \text{Cl}_2 \), and \( \text{H}_2 \) gases. Apart from sodium chloride, other compounds of sodium also have many uses. Caustic soda or sodium hydroxide is used in making soaps, sodium carbonate also known as washing soda, in laundering and the manufacture of glass, sodium bicarbonate as baking soda in baking powder in medicine and in fire extinguishers.

Potassium compounds also have many uses. Potassium hydroxide is used in liquid detergents. Potassium superoxide is used in breathing apparatus, potassium chlorate in matches and explosives and potassium bromide (KBr) in photography. Potassium nitrate is used along with charcoal and sulphur in gun powder. Potassium is a major component of plant fertilizers, where it is used in form of chloride and nitrate salts.

### 3.5 Physical Properties

Most of the physical properties of the alkali metals are directly relocated to atomic properties of elements. Variation of physical properties from one element to the other in a group is governed by the trends of the various atomic properties earlier discussed in Module 2. We shall now apply them to understand the group trends in the various physical properties of the Group 1 elements given in Table 2.1
Table 2.1: Properties of Group 1 Elements

<table>
<thead>
<tr>
<th>Properties</th>
<th>Lithium (Li)</th>
<th>Sodium (Na)</th>
<th>Potassium (K)</th>
<th>Rubidium (Rb)</th>
<th>Caesium (Cs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atomic number</td>
<td>3</td>
<td>11</td>
<td>19</td>
<td>37</td>
<td>55</td>
</tr>
<tr>
<td>Electronic configuration</td>
<td>[He]2s</td>
<td>[Ne]3s</td>
<td>[Ar]4s</td>
<td>[Kr]5s</td>
<td>[Xe]6s</td>
</tr>
<tr>
<td>Atomic weight</td>
<td>6.939</td>
<td>22.898</td>
<td>39.102</td>
<td>85.47</td>
<td>132.905</td>
</tr>
<tr>
<td>Covalent radius (pm)</td>
<td>123</td>
<td>156</td>
<td>203</td>
<td>216</td>
<td>235</td>
</tr>
<tr>
<td>Ionic radius (pm)</td>
<td>60</td>
<td>95</td>
<td>133</td>
<td>148</td>
<td>169</td>
</tr>
<tr>
<td>Boiling point (K)</td>
<td>1620</td>
<td>1154</td>
<td>1038</td>
<td>961</td>
<td>978</td>
</tr>
<tr>
<td>Melting point (K)</td>
<td>453</td>
<td>371</td>
<td>337</td>
<td>312</td>
<td>301.5</td>
</tr>
<tr>
<td>Density (10^3 x kg m^-3)</td>
<td>0.53</td>
<td>0.97</td>
<td>0.86</td>
<td>1.53</td>
<td>1.87</td>
</tr>
<tr>
<td>Electronegativity (Pauling)</td>
<td>1.0</td>
<td>0.9</td>
<td>0.8</td>
<td>0.8</td>
<td>0.7</td>
</tr>
<tr>
<td>Electronegativity (A/R)</td>
<td>1.155</td>
<td>1.0</td>
<td>0.9</td>
<td>0.9</td>
<td>0.85</td>
</tr>
<tr>
<td>Ionisation energy (kJ mol^-1)</td>
<td>520</td>
<td>495</td>
<td>1.0</td>
<td>403</td>
<td>374</td>
</tr>
</tbody>
</table>

**Atomic size**

From your studies in Unit 4 of Module 1, you will recall that the alkali metals are the largest in their corresponding periods in the periodic table. The size of the atom or its ion increases on descending the group (Table 2.1). This is due to the addition of an extra shell of electrons as we move down the group from one element to the next. The addition of the extra shell of electrons out-weighs the effects of increased nuclear charge and thus there is an increase in size from Li to Cs. This trend is shown in Figure 2.2.

![Fig. 2.2 Trend in Ionic Radii of Group 1 Elements](image)

**Density**

Related to atomic size is the density of the elements. Density can be defined as mass per unit volume. For solids the density is a function of
atomic weight, size of the atom and the structure of the solid (i.e. the closeness of the packing of the atoms).

There are two general trends observed in the densities of the elements in the periodic table. Along a period there is a general increase in density because of the increase in the size of the atom. Thus, in a particular period the alkali metals have the lowest density, considering the solid elements only. In a group also density increases on going down the group. Since the elemental structuring are often the same within any group, the factors which determine the density are atomic mass and volume. As you can see from Table 2.1 density increases as we move from Li to Cs. This means that the increase in atomic weight from one element to the next in passing down the group overweighs the effect of increase in the size of the atom. There are, however, some exceptions to this general trend and in this particular group of alkali metals you can see from Table 2.1 that the density of potassium is less than that of sodium. Thus potassium is an exception in this trend.

**Melting Points and Boiling Points**

These metals are soft and can be cut with a knife. It's a result of increase in size and repulsion of the non-bonded electrons, their cohesive energy and tendency for metallic bonding decreases down the group and thus softness increases as we go down from Li to Cs. These metals have low melting and boiling points which also reflect the low values of cohesive energy between the atoms. Their melting and boiling points decreases as we go down the group (Figure 2.3).

![Fig. 2.3 Trend in the Melting Point of Group 1 Elements](image)
Thermal and Electrical Conductivity

In alkali metals, electrons of the noble gas core efficiently shield the lone valence shell electron from the nuclear charge. Therefore the effective nuclear charge felt by the electron in the valence shell of an atom of an alkali metal is the least and their atoms are the largest in respective periods. As a consequence, the sole valence electron is very loosely held by the nucleus. It can move freely from one metal ion to the other in the lattice. This makes the alkali metals good conductors of heat and electricity. This loosely bond electron is also responsive for the silvery lustre of the alkali metals when freshly cut.

Ionisation energy

By losing the loosely bond solitary outermost electron, these elements can acquire the electronic configuration of the preceding noble gas elements. They have, therefore, a high tendency of giving up this electron to form univalent cations. Their first ionisation energies are the lowest in the respective periods and so they are the most reactive of all metals. As we go down the group, their atomic size increases, their ionisation energies decrease, resulting in an increase in their reactivity. The effective nuclear charge felt by the electrons increases after the removal of one electron from the atom of any element and hence, their second ionisation energies are always higher than the first. It is even more so in the case of the alkali metals, because their charged ions (Li\(^+\), Na\(^+\), K\(^+\) etc.) have the stable noble gas configuration of the proceeding group. Removal of an electron from a stable noble gas configuration is extremely difficult. These metals, therefore, form univalent cations only. Figure 2.4 shows the trend in the first conisation energies of the alkali metals down the group.

![Fig. 2.4 Trend in First Ionisation Energies of Alkali Metals](image)
On account of their low ionisation energies these elements have a high tendency to form cations. In other words, they have a high electropositive or metallic character which increases as we move down in the group from Li to Cs. In fact, Rb and Cs are so highly electropositive that they emit electrons even when exposed to light. That is they exhibit photo electric effect.

**Electronegativity**

Because alkali metals have a tendency to lose electron easily rather than to gain, values of electronegativity of these elements are very small. In fact, alkali metals are the least electronegative elements in the periodic table. As expected the electronegativity decreases on moving down the group.

**Ionic Character of Compounds**

Compounds formed by alkali metals with highly electronegative elements like halogens and oxygen, are largely ionic in nature because of the large electronegativity difference.

You can see that the trends in ionic character (Figure 2.5) show that the ionic character increases with increase in cation size and decreases with increase in anion size. Because of the small size of Li\(^+\), it has more polarising power and therefore favours covalent bonding.

**Solubility, Lattice Energy and Hydration Energy**

Alkali metal salts like halides, oxides, hydroxides, carbonates, sulphates etc exhibit some interesting trends in their solubility in water. First let us remind ourselves that lattice energy is the driving force for the formation of an ionic compound and its stability. Lattice energy is directly proportional to the charge on the ions and inversely proportional to the distance between the cation. This distance is taken as the sum of radii of cation and anion (rc + ra).

Let us again remind ourselves that lattice energy is the enthalpy change when one mole of crystal lattice is formed from the isolated gaseous ions and hydration energy is the enthalpy change when one mole of solute is dissolved in water. In a group, the charge on cations remains constant. Thus, lattice energy depends mainly on the size of the cation.

Similarly, hydration energy varies with the charge and size of the cation. The higher the charge and the smaller the size of the ion, the more is the hydration energy. In a group, lattice energy and hydration energy decrease as we move down. While the decrease in lattice energy favours
the solubility, the decrease in hydration energy makes the compound insoluble.

![Diagram of Ionic Character of Alkali Halides Metals](image)

**Fig. 2.5 Trend in Ionic Character of Alkali Halides Metals**

For the salts of small anions (like F, O²⁻, OH⁻ etc.) the lattice energy which is inversely proportional to \( r_c + r_a \), is very sensitive to the change in the size of the cation, anion being very small in size has little contribution in the total \( (r_c + r_a) \) and decreases sharply as we move down the group. Thus, in these salts, the decrease in lattice energy is greater than the decrease in hydration energy and therefore, the solubility of these salts increases as we go down the group. For example, in the case of alkali metal fluorides, the solubility increases in the order LiF < NaF < KF < RbF < CsF.

For the salts having large anions (SO₄²⁻, F, NO₃⁻, HCO₃⁻, CO₃²⁻, etc) as \( r_a >> r_c \), the radii of the cation has little contribution in the total \( (r_c + r_a) \) and decreases sharply as we move down the group. Thus, in these salts, the decrease in lattice energy is greater than the decrease in hydration energy and therefore, the solubility of these salts increases as we go down the group. For example, in the case of alkali metal carbonates, lithium carbonate is highly soluble while the solubility of calcium carbonate is very little.

Another important factor contributing to the solubility of the compound is the match in the size of the cation and anion. Whenever there is a mismatch, e.g. cation is small, anion is large or vice-versa, this will result in the increased solubility of the compound. Let us keep the cation constant, say calcium. If we, then change the anion from fluoride to iodide, then the solubility of the compound will vary as CsF > CsCl > CsBr > CsI. Thus, calcium fluoride will be most soluble and calcium iodide will be the least. Similarly, from lithium fluoride to lithium iodide: the solubility will increase in the order: LiF < LiCl < LiBr < LiI.
Table 2.2: The Main Trends in the Properties of Alkali Metals

<table>
<thead>
<tr>
<th>increasing:</th>
<th>Li</th>
<th>Na</th>
<th>K</th>
<th>Rb</th>
<th>Cs</th>
</tr>
</thead>
<tbody>
<tr>
<td>electropositivity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>density</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>atomic radii</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>atomic volume</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>reactivity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>reducing power</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>anion stabilization</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>solubility of salts having small anion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- The density of K is less than that of Na:

There can be many more examples which can be explained on the basis of above reasoning. Table 2.2 shows a summary of the trends discussed in this section.

4.0 CONCLUSION

We can conclude this unit by observing that the alkali metals as a group are very reactive, in fact so reactive that they can hardly be found in a free state. They occur in a combined form. Alkali metal compounds are very useful for everyday life. They are good conductors of heart as well as electricity.

5.0 SUMMARY

In summary, you have studied the following:

- the alkali metals can only be found in a combined state
- the alkali metals are very reactive
- the alkali metals are very useful, some of their uses include being used as electrical conductors.

They are also used at homes for example sodium chloride is used as table salt.

SELF-ASSESSMENT EXERCISE

i. Given below are some of the statements about the alkali metals. Write "T" if True and "F" if you think it is false.
- Sodium is the most abundant alkali metal in the earth
- Sodium is the most abundant metallic element in sea water
• Alkali metals occur in the free state in nature
• Lepidolite is an ore of lithium
• Atomic radii of the alkali metals decrease down the group
• Ionisation energy increases from lithium to francium
• Melting and boiling points of alkali metals decrease down the slope.
• Lithium is the highest of all the metallic elements
• Solubility of alkali metal fluorides in water increases down the group
• Ionic character of alkali metal halides decreases down the group.

ii. Explain in brief why the hydride bridge in (BeH$_2$)$_n$ considered to be electron deficient, but not the halide bridge in (BeCl$_2$)$_n$

6.0 TUTOR-MARKED ASSIGNMENT

1. Why are the elements of Group 1 called alkali metals?
2. Why do alkali metals not occur as free elements in nature?
3. Explain why:
   i. alkali metals are good conductors of electricity
   ii. lithium has the highest ionisation energy in the group
   iii. sodium forms +1 ion and not +2 ion
   iv. group 1 elements form ionic compounds
4. Explain why hydroxides become stronger alkalis on descending the group.
5. Why do peroxides and super oxides oxidize in aqueous solution?

7.0 REFERENCES/FURTHER READING


UNIT 3  COMPOUNDS OF ALKALI METALS

CONTENTS

1.0  Introduction
2.0  Objective
3.0  Main Content
   3.1  Compounds of Alkali Metals
   3.2  Oxides and Hydrogen
   3.3  Sulphides
   3.4  Hydrides
   3.5  Carbides
   3.6  Thermal Stability of Salts
4.0  Conclusion
5.0  Summary
6.0  Tutor-Marked Assignment
7.0  References/Further Reading

1.0  INTRODUCTION

In the last unit, you studied the elements of group 1 otherwise known as, Alkali metals. You studied the occurrences, extraction, uses and physical properties of the alkali metals.

In this unit, you will be studying the chemical properties of the alkali metals. You will also study the oxides, hydroxides sulphides, hydrides carbides and thermal stability of salts of the alkali metals.

2.0  OBJECTIVES

At the end of this unit, you should be able to:

- list the types of oxides formed by alkali metals
- explain using equations, the reactions involved in the formation of the oxides
- explain using equations the formation of sulphides, carbides and hydrides by alkali metals
- explain using examples the stability of salts formed by alkali metals.
3.0 MAIN CONTENT

3.1 Compounds of Alkali Metal

In Unit 2 you studied the important physical properties of the alkali metals. In accordance with their highly electropositive character these metals are very reactive and are powerful reducing agents reacting with water and most non-metals. They form crystalline ionic salts with high melting and boiling points. These salts are usually soluble in water giving conducting solution. Now we shall discuss some of the important classes of these salts.

3.2 Oxides and Hydroxides

As alkali metals are very reactive their lustre is lost in air due to the formation of oxide with atmospheric oxygen. Three types of oxides are formed by the alkali metals: normal oxides having $O^2-$ ion and the peroxides having $O_2^2-$ $[O-O]$ ion, both of which are dia-magnetic and colourless. The third one which is coloured and paramagnetic is super oxide containing $O^2^-$ ion. Controlled oxidations of these metals are as follows:

\[
\begin{align*}
2 \text{Li} & \xrightarrow{\frac{1}{2} O_2} \text{Li}_2\text{O} \xrightarrow{\frac{1}{2} O_2} \text{Li}_2\text{O}_2 \\
2 \text{Na} & \xrightarrow{\frac{1}{2} O_2} \text{Na}_2\text{O} \xrightarrow{\frac{1}{2} O_2} \text{Na}_2\text{O}_2 \\
2 \text{K} & \xrightarrow{\frac{1}{2} O_2} \text{K}_2\text{O} \xrightarrow{\frac{1}{2} O_2} \text{K}_2\text{O}_2 \xrightarrow{O_2} 2\text{KO}_2 \\
2 \text{Rb} & \xrightarrow{\frac{1}{2} O_2} \text{Rb}_2\text{O} \xrightarrow{\frac{1}{2} O_2} \text{Rb}_2\text{O}_2 \xrightarrow{O_2} 2\text{RbO}_2 \\
2 \text{Cs} & \xrightarrow{\frac{1}{2} O_2} \text{Cs}_2\text{O} \xrightarrow{\frac{1}{2} O_2} \text{Cs}_2\text{O}_2 \xrightarrow{O_2} 2\text{CsO}_2
\end{align*}
\]

In the scheme shown above the products underlined are the main products when the metals are burnt in a free supply of air. You may notice in the above scheme that Lithium forms normal oxide, sodium forms peroxide while potassium, rubidium and caesium form superoxide as the main product. All the Group 1 metal oxides are strongly basic and react vigorously to give hydroxide.
As shown, the peroxides and the super oxides on reaction with water give H$_2$O$_2$ which in turn is a powerful oxidising agent. Thus the peroxides and the super oxides are also oxidising in nature.

The basic strength of the hydroxides increase down the group. As the charge density (charge/size ratio) of the cation decreases between M$^+$ and 0H$^-$ also decreases. So, OH$^-$ can be liberated readily into the solution and O as we go down the group.

### 3.3 Sulphides

Alkali metals reacts with sulphur to form two types of sulphides: simple sulphides of Na$_2$S and polysulphides like Na$_2$S, where n — 2, 3, 4, or 6.

These polysulphides have a zig-zag chain structure as shown below.

![Zig-zag chain structure of polysulphides](image)

### 3.4 Hydrides

Alkali metals react with hydrogen and form ionic hydrides, MH. These hydrides on reaction with water liberate hydrogen. Thus they are a useful source of hydrogen:

\[
\text{MH} + \text{H}_2\text{O} \rightarrow \text{MOH} + \text{H}_2.
\]

Lithium hydride on reaction with AlCl$_3$ in other solution forms lithium aluminium hydride which is a useful reducing agent in organic chemistry.
Similarly, sodium hydride forms sodium borohydride which is also used as a reducing agent.

### 3.5 Carbides

Lithium reacts with carbon to form ionic carbides, whereas similar carbides of other metals are not formed on reacting with carbon. They can, however, be formed on heating the metal with acetylene or when acetylene is passed through a solution of the metal in liquid ammonia:

![Chemical reaction]

These carbides contain the carbide ion \((\text{C} = \text{C})^{2-}\) on hydrolysis they give acetylene. Hence, are termed as acetylides:

![Chemical reaction]

Alkali metals also form covalent compounds such as methylthium, LiCH₃ and ethylsodium NaC₂H₅. These come under the separate class of organometallic compounds. The main reactions of group 1 elements are summarised in Table 3.1

<table>
<thead>
<tr>
<th>Table 3.1: The reaction of the Group 1 elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>2Li + O₂ (excess) → Li₂O</td>
</tr>
<tr>
<td>2M + S → M₂S</td>
</tr>
<tr>
<td>M + H₂O → MOH + ½ H₂</td>
</tr>
<tr>
<td>M + ROH → MOR + ½ H₂</td>
</tr>
<tr>
<td>M + ½ H₂ → MH</td>
</tr>
<tr>
<td>M + ½ X₂ → MX</td>
</tr>
<tr>
<td>3Li + ½ N₂ → Li₃N</td>
</tr>
<tr>
<td>M + NH₃ (l) → [M(NH₃)₁]⁺ + c(NH₃) → M⁺NH₂ + ½ H₂</td>
</tr>
<tr>
<td>2M + C(orC₂H₂) → M₂C₂ (acetylides) + H₂O → C₂H₂</td>
</tr>
<tr>
<td>M + Hg → amalgams</td>
</tr>
</tbody>
</table>
3.6 Thermal Stability of Salts

The case with which a salt decomposes is related to the enthalpy of formation of the salt. (The standard enthalpy of formation of a compound is the enthalpy change when one mole of the compound in the standard state is formed from the elements in the standard state). The enthalpy of formation \( \Delta H_f \) of a salt \( MA \) (\( M \) is the metal, \( A \) the anion) is given by

\[
\Delta H_f = (H + I) + (H - E_A) - H_{\text{latt}}
\]

Where \( H \) is the enthalpy of atomisation, \( I \) is the ionisation energy and \( EA \) is the electron affinity. Since for any salt in a particular group the terms involving the anion alone remain constant the value of \( \Delta H^\circ F \) for such compounds is dependent upon the sum of the enthalpy terms of the particular metal (\( H \) atom + I metal) and the lattice energy \( \Delta H_{\text{latt}} \). The larger the lattice energy, the more negative the enthalpy of formation and so, the more stable is the compound. All these terms become smaller on descending the series from lithium to caesium. The relative stabilities of the salts are therefore decided by the parameter which decreases more rapidly — the lattice energy or the sum of the metal enthalpies.

In the salts having small anions of high charge density, eg \( F^- \), \( N_3^- \), \( O_2^- \) etc. the change in lattice energy is much dependent on the size of the cation and decreases rapidly on descending the group. Thus, as the size of the cation increases, lattice energy decreases more than the change in the sum of the metal enthalpies. Therefore as we go down the group the stability of these salts having small anions decreases. Thus in alkali metal fluorides, the stability decreases in the order \( \text{LiF} > \text{NaF} > \text{KF} > \text{RbF} > \text{CsF} \).

The opposite trend is observed in the stability of the salts containing large anions of low charge density, eg \( \text{Br}^-, \text{I}^-, \text{NO}_2^- \) etc. In such cases, the lattice energy is relatively insensitive to the change in cation size and there is more rapid decrease in ionisation energy and atomisation enthalpy on descending the group. The lower values of these favour the stability of the compounds. Thus, the stability of the compounds having large anions increases as we move down the group from lithium to caesium.

The stability of the compounds can also be explained by using the concept of polarising power. The simple idea is that, as the charge density of the metal ion increases, the thermal stability of the salts of
large polarising anions, relative to some decomposition product decreases. In general, the least polarising metal ion are those of the most electropositive metal ions and these form the most stable salts with large anions. In other words small cations form stable salts with small anions and large cations form stable with large anions.

Let us take for example carbonate of Group 1 metals. The carbonates of sodium potassium and caesium are resistant to the heat of a burner flame. However, lithium carbonate decomposes to its oxide and carbon dioxide under the same conditions.

\[ \text{Li}_2\text{CO}_3 \rightarrow \text{Li}_2\text{O} + \text{CO}_2 \]

The tendency of Li\(\text{CO}_3\) to undergo thermal decomposition may be explained in terms of the gain in electrostatic attraction that occurs when extremely small Li\(^+\) ion combines with the smaller oxide ion rather than the much larger carbonate ion. The other carbonates of group (Na — Cs) are more stable because the cations have a lower charge density and are considerably larger in size and so their decomposition is less favourable energetically.

All the metals except lithium form stable bicarbonates (Lithium bicarbonate is formed only in aqueous solution and has not been isolated). When we heat the alkali metal bicarbonates, they are decomposed to carbonates and simultaneously, carbon dioxide and water are liberated.

\[ 2\text{NaHCO}_3 \rightarrow \text{Na}_2\text{CO}_3 + \text{CO}_2 + \text{H}_2\text{O} \]

The thermal stability of group 1 hydroxides also follows a similar trend as that of carbonates. Thus with the exception of LiOH which on heating decomposes to Li\(_2\)O, all other group 1 hydroxides are stable.

Similarly, lithium nitrate also decomposes on heating to give Li\(_2\)O NO\(_2\) and O\(_2\) but all other alkali metal nitrates decompose on strong heating to nitrates liberating oxygen.

\[ 4\text{LiNO}_3 \rightarrow 2\text{Li}_2\text{O} + \text{O}_2 + 4\text{NO}_2 \]

\[ 2\text{NaNO}_3 \rightarrow 2\text{NaNO}_2 + \text{O}_2 \]
4.0 CONCLUSION

We can conclude this unit by observing that alkali metals being very reactive form oxides hydroxides and peroxides. Also, that the stability of alkali salts depends on the enthalpy of formation of the salt.

5.0 SUMMARY

In summary, you have studied the following:

- alkali metals form oxides, hydroxides and peroxides
- three types of oxides are formed viz normal oxides, peroxides and super oxides
- three normal oxides, peroxides and super oxides
- the alkali salt's stability is dependent upon the enthalpy of formation.

SELF-ASSESSMENT EXERCISE

i. Explain briefly why alkali metals act as strong reducing agents.

ii. The thermally least stable alkali metal fluoride is;
   (i)  LiF
   (ii) NaF
   (iii) KF
   (iv) RbF
   (v)  CsF

iii. Why is it that Be forms mostly complexes with tetrahedral structure?

6.0 TUTOR-MARKED ASSIGNMENT

1. Why do peroxides and superoxide oxidise in aqueous solution?
2. Explain why lithium forms more complexes?

7.0 REFERENCES/FURTHER READING


UNIT 4  SOLVATION OF ALKALI METAL IONS

CONTENTS

1.0  Introduction
2.0  Objectives
3.0  Main Content
   3.1  Solvation of the alkali metal ions
   3.2  Solutions of alkali metal in liquid ammonia
   3.3  Complexation behaviour of alkali metals
   3.4  Anomalous nature of Lithium
4.0  Conclusion
5.0  Summary
6.0  Tutor-Marked Assignment
7.0  References/Further Reading

1.0  INTRODUCTION

In Unit 2, you studied the formation of oxides, hydroxides, carbides and sulphides by the alkali metals. In this unit you will be studying the behaviour of the alkali metals in solutions. You will also study the complexation behaviour of the alkali metals. In your study of the chemical properties of the alkali metals, you must have noticed that lithium behaved differently from the rest of the alkali metals. In this unit, you will be studying the anomalous behaviour of lithium.

2.0  OBJECTIVES

At the end of this unit, you should be able to:

• discuss the solvation of alkali metals describing the behaviour of each member of the alkali metals when in solution of water
• describe using equations the behaviour of the alkali metals when dissolved in liquid ammonia
• explain what is meant by complexation (complex formation) by describing the role of the alkali metal in relation to the surrounding molecules
• list at least four anomalous behaviour of lithium.
3.0 MAIN CONTENT

3.1 Solvation of the Alkali Metal

When a metal is surrounded by solvent molecules, the phenomenon is called Solvation of the metal ion. When the solvent is water the phenomenon is now called Hydration.

Hydration is therefore solvation with water as the solvent. The alkali metal ions are highly hydrated. The smaller the size of the ion, the greater its degree of hydration. This is because the smaller the size, the more will be its charge density and the more will be its attraction for the polar solvent molecules (Unit 1). Thus Li$^+$ ion, which is the smallest, gets more hydrated than Na$^+$ and so on. The degree of hydration decreases on moving down the group. As a result of differences in their degree of hydration, the hydrated ionic radii of the alkali metal ions decrease as we go down the group from lithium to caesium. Lithium has the largest hydrated radius while Cs$^+$ has the smallest hydrated radius in the first group. You will agree that the smaller the size of the ion and the lighter it is, the more will be its mobility and thus conductance. In this regard we should expect the highest conductance of the alkali metals, but it is not so. This is because the hydrated Li$^+$ is the largest of all the alkali metal ions. In solution its mobility is less and so Li$^+$ ion is the least conducting in solution. The ionic conductance in solution actually decreases in the order Cs$^+ >$ Rb$^+ >$ Na$^+ >$ Li$^+ >$ K.

3.2 Solutions of Alkali Metals in Liquid Ammonia

All the alkali metals are highly soluble in liquid ammonia giving a deep blue colour. The solubilities of the metals for 100g of ammonia are Li, 10g; Na, 25g and K, 49g. The dissolution of the alkali metal is accompanied by its dissociation into the metal ions and the electrons. The metal ion and the electrons then get associated with ammonia solvent molecules. Electrons associated with the solvent are known as Solvated Electrons.

\[
\text{Na}_\text{(s)} + \text{NH}_3(\text{l}) \rightarrow \text{Na}^+(\text{NH}_3)_x^+ \text{e} (\text{NH}_3)_y
\]

The alkali metal solutions in liquid ammonia are highly conducting and behave almost as metals. Their specific conductivities are almost the same, because the anion i.e. solvated electron, in all the cases is the same. The small difference in the conductivity is due to the value of the metal itself. The solution of alkali metals in liquid ammonia is blue in
colour due to the presence of solvated electrons, and therefore the solutions are also paramagnetic. With increasing concentration there is a decrease in Para magnetising suggesting that the electron can get associated to form diamagnetic electron pairs although there may be other equilibria also.

\[
\text{Na}_s (\text{dispersed}) \rightleftharpoons \text{Na} \text{ (in solution)} \rightleftharpoons \text{Na}^+ e^-
\]

On increasing the concentration above 3M, the colour of the solution changes to copper bronze having metallic lustre, because the metal ions form clusters. Apart from lithium other alkali metals can be recovered unchanged from solution. Lithium, in ammonia solution form a complex of the type \([\text{Li} (\text{NH}_3)_4]^+\)

The blue solutions of alkali metals are moderately stable at temperatures where ammonia is still a liquid. The reaction that results in the formation of an amide:

\[
\text{Na} + \text{NH}_3(c) \rightleftharpoons \text{NaNH}_2 + \frac{1}{2}\text{H}_2
\]

Can occur photochemically and is catalysed by transition metal salts. The alkali metal solutions in liquid ammonia are powerful reducing agents and are used for this purpose in inorganic and organic reactions.

### 3.3 Complexation Behaviour of Alkali Metals

A complex compound can be defined as a compound with a central atom or ion surrounded by a group of ions of molecules called "ligands". These ligands are usually bond to the metal by a "coordinate bond" i.e. a bond formed by the donation of a lone pair of electrons from one atom (of the liquid) to the other (metal/ion). Although both metal and the legend are usually capable of independent existence as stable chemical species, yet when the complex is formed, it generally retains its identity in solution. For example, in solution, Fe ++ and CN can exist independently but once the complex \([\text{Fe(CN)}_6]^{-4}\) is formed it exists in solution as such. It does not dissociate into Fe+ and CN - , as a result it will not give any positive result when tested in Fe + and CN-. It is thus complex specie. The most stable complexes would be formed by the lightly polarising cations. This is because they have a strong tendency of
interacting with electron clouds of other anionic or neutral electron rich species (ligands).

According to the model above, a very weak coordinating ability is expected in the group 1 metals because of their large size and low charge of the cations $M^+$. According to this view, stability of the complexes of the alkali metals should decrease in the order Li $>$ Na $>$ K $>$ Rb $>$ Cs and this is the observed trend.

Alkali metals form few complexes mostly chelates with the ligands like B- ducetones, nitrophenils, nitroso naphthols etc. as shown in Figure 4.1. They have low stability.

![Fig. 4.1 Source Complex of Alkali Metal Ions](image)

Lithium, being the most polarising cation of all the alkali metals forms tetrahedral complexes with ligands like NH$_3$, L$_5$H$_5$N etc. With ammonia it forms the complex of the type $[\text{Li (NH}_3\text{)}_4]\text{I}$, whereas with pyridine a complex of the type $[\text{LiCl(C}_5\text{H}_5\text{N})_2(\text{H}_2\text{O})]$ formed.

### 3.4 Anomalous Natures of Lithium

On descending any group of S or P block elements of the periodic table, one notices that, there is a general decrease in electro negativity or increase in electro negativity. The difference in electro negativity between the first and second elements of each group is much greater than that between any two successive elements. This is reflected in the properties of the elements. Thus not only is the element more electronegative than the other elements of the group, but it are much more electronegative than expected by simple extrapolations.

This trend of bigger than expected differences in properties between the first member of a group and the rest of the elements is shown in group 1 where Li elements markedly different from the rest of the members of the group.
Let us take a look at these summaries again as they apply in Group 1. Due to the very small size of lithium, the metallic bonding between the atoms in the metallic lattice is very strong giving rise to strong cohesive forces. This is shown in its relatively higher melting point, and boiling point, hardness and homonuclear bond energy. The relatively higher attraction of lithium for its outer electron results in its relatively higher centre electronegativity ionisation energy, hydration energy electron affinity and of course smaller atomic radii relative to the other homologues.

Similar anomalies are also found in the chemical properties, but the different appear greater as we shall see in the following accounts.

(i) Lithium salts of large polarisable anions are thermally less stable than those of other alkali metals e.g. Lithium carbonate decomposes at 950K, whereas no decomposition of sodium carbonate takes place below 1050K.

(ii) Lithium does not form solid bicarbonate trioxide or superoxides, because these are unstable at room temperature. On the other hand those of other alkali metals require a higher temperature to effect their decomposition.

(iii) Lithium salts of anions of high charge density are less soluble than those of other alkali metals. The halides of lithium are more soluble in organic solvents.

(iv) Lithium forms stable salts with anions of high charge density owing to their high lattice energy. For examples, in air lithium forms the normal oxide, whereas the others form higher oxides lithium reacts with nitrogen to form nitride, Li$_3$N the others do not react. Lithium hydride is more stable than the other hydrides and lithium carbide is formed more easily with acetylene.

(v) Lithium reacts slowly with water.

(vi) Lithium forms more stable covalent bonds than other alkali metals and, therefore, forms more stable complex compounds as earlier seen in section 3.3 of this unit. For example, lithium cannot be recovered unchanged from its liquid ammonia solution. Owing to the formation of [Li(NH$_3$)$_4$]$_4$.

4.0 CONCLUSION

We can conclude this unit by stating that the alkali metal ions are very soluble in both water and liquid behaviour. Also that lithium shows the
anomalous ammonia observed for first members of any group of the elements in the periodic table.

5.0 SUMMARY

In summary, you have learnt the following from the foregoing unit:

- the alkali metals are highly soluble in water and in liquid ammonia
- Li+ ion is the least conducting in solution, when compared to the ions of the other members of the alkali metal ions in solution.
- the stability of alkali metal complexes is low
- the stability of the complexes decrease as you go down the group. ie Li > Na > K > Rb > Cs
- lithium shows anomalous behaviour in relation to the other members of group 1. That is that in each of the properties shared by members of the group, Li shows wide differences between it and the next member of the group immediately below it.

SELF-ASSESSMENT EXERCISE

i. Briefly explain the variation of conductivity in alkali metal salts giving reasons for the trend.
ii. Why are alkali metals poor complexing agents?
iii. Why does ammonia acts as a Lewis base
iv. Why is hydrazine unstable?

6.0 TUTOR-MARKED ASSIGNMENT

1. Explain why lithium forms more complexes than the other elements of group I.
2. Give the trend in the solubility of alkali metal iodides

7.0 REFERENCES/FURTHER READING


UNIT 5 ALKALINE EARTH METALS

CONTENTS

1.0 Introduction
2.0 Objectives
3.0 Main Content
   3.1 Alkaline Earth Metals
   3.2 Occurrence of Alkaline Earth Metals
   3.3 Extraction of Alkaline Earth Metals
   3.4 Uses of Alkaline Earth Metals
   3.5 Physical Properties of Alkaline Earth Metals
   3.6 Solubility, Lattice Energy and Hydration Energy
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 References/Further Reading

1.0 INTRODUCTION

In Units 2 — 4, you studied the general characteristics of group 1 elements, i.e the Alkali metals, and their compounds. Groups 1 and 2 elements belong to the s-block of the periodic table. Their electronic configurations show an outer shell of ns\(^1\) and ns\(^2\) for alkali and alkaline earth metals respectively. s-block elements are known to be very reactive metals and generally form ionic compounds. In this unit you will be studying the elements of Group 2 consisting of beryllium, magnesium, calcium, strontium, barium and radium. Elements Ca, Sr, Ba and Ra are called alkaline earth metals, because their earths (earth is the old name for a mineral oxide) are alkaline in nature. (Remember that the alkali metals are so called because they form hydroxides which are strong alkalis).

Beryllium is not counted as an alkaline earth metal since its oxides is not alkaline. We shall start our study of the alkaline earth metals by studying their occurrence, extraction uses and physical properties.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- list at least four places where the alkaline earth metals can be found
- describe at least two methods used in the extraction of the alkaline earth metals
- list at least three uses of members of the group
compare the physical properties of members of the group with each other.

3.0 MAIN CONTENT

3.1 Alkaline Earth Metals

The alkaline earth metals, like alkali metal are very reactive; therefore, do not occur free in nature. All of them are found in form of their salts. Let us now study their occurrences, extraction and uses. We shall also go on to study their physical properties.

3.2 Occurrence

Beryllium, the first member of the group is found in small quantities in the silicate mineral, phenacite, $\text{Be}_2\text{SiO}_4$, and beryll, $3\text{BeO}\cdot\text{Al}_2\text{O}_3\cdot0.6\text{SiO}_2$. Magnesium (2.76%) and calcium (4.66%) are among the eight most abundant elements in the earth's crust.

Magnesium (0.13%) is the second most abundant metallic element next only sodium (chloride) in sea water. It occurs as magnesite, $\text{MgCO}_3$; dolomite $\text{MgCa(CO}_3)_2$; kieserite, $\text{MgSO}_4\cdot\text{H}_2\text{O}$ and carmallite, $\text{KMgCl}_3\cdot0.6\text{H}_2\text{O}$ in the earth's crust.

Calcium occurs extensively as calcite and lime — stone ($\text{CaCO}_3$) in many mountain ranges. Calcium and magnesium are very important biologically too. Calcium is found in the bones of animals and human beings.

Magnesium is found in the green (chlorophyll) plants.

Strontium (0.038%) and barium (0.039%) are much less abundant and occur as carbonates and sulphates. These metals are well known because they occur as concentrated ores and are easy to extract. Radium is extremely scarce (10 -10 %) and it is a radioactive element.

3.3 Extraction of Alkaline Earth Metals

These metals are extracted by electrolysis of their fused chlorides, though magnesium has been manufactured by the carbon reduction of its oxide.

Beryllium is obtained by the electrolysis of molten beryllium chloride. Sodium chloride must be added to the melt as an electrolyte because $\text{BeCl}_2$ is covalent and, therefore is a very poor electrical conductor. During the electrolysis, the less active metal Be is produced at the...
cathode and C12 is evolved at the anode. Calcium is extracted from fused calcium chloride using a graphite anode and iron cathode.

Strontium chloride and Barium chloride are fused for the extraction of strontium and barium respectively.

### 3.4 Uses of Alkaline Earth Metals

Beryllium is used for making atomic fuel containers because it absorbs very few neutrons and does not become radioactive. Being transparent to X-rays it is used as a window material in X-ray apparatus.

It has a number of uses as alloys, e.g. when mixed with Cu, Beryllium increases the strength of Cu. Six fold. Beryllium alloys are non-sparking, therefore, they are used in making hand tools for use in the petroleum industry. Magnesium, because of its lightness, is used as a construction alloy material, e.g. in aircrafts. For this purpose it is alloyed with aluminium.

Magnesium is also used as a reducing agent in the extraction of some metals like titanium and uranium. It forms Gringard reagents RMgX, which are important organic reagents.

Calcium, strontium and barium as free metals do not find extensive uses because they are very reactive. Calcium oxide (quicklime) is a constituent of glass, mortar and portland cement.

### 3.5 Physical Properties

The alkaline earth metals are quick soft metals, but are harder than the corresponding Group 1 elements. This is because of their two valence electrons which participate in metallic bonding. They are good conductors of electricity. In pure form they are silver coloured, but on exposure to the atmosphere, the silvery lustre is lost, because of the formation of an oxide layer on the surface of the metal. Their physical properties are given in Table 5.1
The atoms of the alkaline earth metals are smaller than those of the corresponding Group 1 elements. This is because of the increase atomic number. Because of the resulting increase in effective nuclear charge, valence shell electrons are pulled in more firmly by the nucleus, thereby reducing the size of the atom.

Similarly, their ionic radii are also smaller than those of Group 1 elements, because the removal of two orbital electrons increases the effective nuclear charge even further.

The elements are denser than Group 1 metals because they have two valence electrons per atom for bonding the atoms into a metallic lattice and as a result more mass can be packed into a smaller volume.

The density decreases slightly on moving down the group from Be to Ca but increases considerably thereafter up to Ra.

The atomic/ionic radii increase from Be to Ra due to the effect of extra shells of electrons added. This outweighs the effect of increased nuclear charge. Group 2 metals have higher melting points when compared to group 1 metals. The reason being the +2 charge on the cations in the metallic lattice causing them to be more strongly attracted to the 'Sea of electrons and making it difficult to pull them apart.

The first ionisation energy (Table 5.1) of alkaline earth metals is more than that of corresponding alkali metals. This is because the alkaline earth metals have higher nuclear charge and are smaller in size. The electrons are therefore more tightly held to the nucleus. The second

---

**Table 5.1: Properties of the Group 2 Metals**

<table>
<thead>
<tr>
<th>Property</th>
<th>Beryllium Be</th>
<th>Magnesium Mg</th>
<th>Calcium Ca</th>
<th>Strontium Sr</th>
<th>Barium Ba</th>
<th>Radium Ra</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atomic number</td>
<td>4</td>
<td>12</td>
<td>20</td>
<td>38</td>
<td>5688</td>
<td>9096</td>
</tr>
<tr>
<td>Electronic configuration</td>
<td>[He]2s²</td>
<td>[He]3s²</td>
<td>[Ar]4s²</td>
<td>[Kr]4s²</td>
<td>[Xe]6e²</td>
<td>[Rn]7e²</td>
</tr>
<tr>
<td>Atomic weight</td>
<td>9.012</td>
<td>24.312</td>
<td>40.08</td>
<td>87.62</td>
<td>137.34</td>
<td>226.02</td>
</tr>
<tr>
<td>Ionic radius (pm)</td>
<td>31</td>
<td>65</td>
<td>99</td>
<td>113</td>
<td>135</td>
<td>135</td>
</tr>
<tr>
<td>Covalent radius</td>
<td>89</td>
<td>136</td>
<td>174</td>
<td>191</td>
<td>198</td>
<td>198</td>
</tr>
<tr>
<td>Boiling point (K)</td>
<td>3243</td>
<td>1380</td>
<td>1760</td>
<td>1607</td>
<td>1413</td>
<td>1700</td>
</tr>
<tr>
<td>Melting point (K)</td>
<td>1553</td>
<td>934</td>
<td>1118</td>
<td>1062</td>
<td>998</td>
<td>700</td>
</tr>
<tr>
<td>Enthalpy of hydration</td>
<td>-2455</td>
<td>-1900</td>
<td>-1565</td>
<td>-1415</td>
<td>-1275</td>
<td>-1275</td>
</tr>
</tbody>
</table>

Density (10³ g cm⁻³): 1.85, 1.74, 1.54, 2.6, 3.62, 5.5
Electropositivity: 1.5, 1.2, 1.0, 1.0, 0.9, 0.9
Ionisation energy (KJ mol⁻¹): 900, 758, 590, 549, 592, 599
ionization energy of these elements is almost twice their first ionization energy. This is because once one electron has been removed, the effective nuclear charge felt by the orbital electrons is increased, so that the remaining electrons are more lightly held and hence much more energy is needed to remove the second electron.

However, their second ionisation energy is less than that of the corresponding alkali metals because of stability of a closed shell configuration of the univalent cations that are formed in the cases of the alkali earth metals.

The ionization energy of alkaline earth metals decreases on moving down the group.

The metals of this group (beryllium is an exception) form ionic compounds. This is because the assembly of positive and negative ions into a symmetrical crystal lattice results in the liberation of large amounts of energy.

Electropositive character and the reducing property (tendency to lose elections) increase on moving down the group.

Since alkaline earth metals loose electrons easily, they form divalent anions which have noble gas structure with no unpaired electrons. Their compounds are diamagnetic and colourless, unless the anion is coloured. Ca, Sr and Ba compounds give characteristics flame colourations which are used to identify them - Ca(brick red flame), Sr (crimson red flame) and Ba (apple green flame).

3.6 Solubility, Lattice Energy and Hydration Energy

The solubility of alkaline earth metal compounds shows some interesting trends. The metal ions are easily hydrated, e.g. MgCl$_2$.6H$_2$O, CaCl$_2$.6H$_2$O, BaCl$_2$.2H$_2$O. The hydrogen energies of these metals ions are much greater than those of alkali metal ions (Table 5.2) because of their smaller size and increased cationic charge. The lattice energies of alkaline earth metal salts (Table 5.2) are also much higher than those of alkali metal salts.

Hydration and lattice energies decrease with increase in size of metal ions. Decreasing lattice energy favours increased solubility, whilst decreasing hydration energy favours decreased solubility. If on moving down the group the hydration energy decreases more rapidly than the lattice energy, the compound becomes less soluble. This occurs with most of the compounds except for fluorides and hydroxides for example, solubility of sulphates decreases from BeSO$_4$ to BaSO$_4$. Due to their
small ionic radii, Be\(^{2+}\) and Mg\(^{2+}\) have high hydration energies. Because of that BeSO\(_4\) and MgSO\(_4\) only slightly soluble in water, whereas SrSO\(_4\) and BaSO\(_4\) are almost insoluble in water. In the case of fluorides and hydroxides, the lattice energy decreases more rapidly than the hydration energy. This causes a reverse trend i.e., the fluorides and hydroxides increase in solubility on moving down the group.

Table 5.2: The Lattice Energies of Alkaline Earth Metal Salts

<table>
<thead>
<tr>
<th>Element</th>
<th>(\Delta H_{\text{latt}}) (kg mol(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Be(^{2+})</td>
<td>-2454</td>
</tr>
<tr>
<td>Mg(^{2+})</td>
<td>-1921</td>
</tr>
<tr>
<td>Ca(^{2+})</td>
<td>-1577</td>
</tr>
<tr>
<td>Sr(^{2+})</td>
<td>-1443</td>
</tr>
<tr>
<td>Ba(^{2+})</td>
<td>-1304</td>
</tr>
</tbody>
</table>

Enthalpies of hydration, OH\(^{-}\), alkaline of alkali earth metal ions M\(^{2+}\) and lattice energies \(\Delta H_{\text{latt}}\) of their oxides, carbonates fluorides and iodides in kg mol\(^{-1}\)

4.0 CONCLUSION

In conclusion we can say alkaline earth metals, like metals, are only found combined in form of their salts, because of their reactivity. Alkaline earth metals are usually extracted by electrolysis. They are used as alloys and some e.g. Beryllium are used as fuel containers.

5.0 SUMMARY

In this unit, you have learned the following:

- alkali earth metals are very reactive
- because of this reactivity, alkaline earth metals exist mainly in combined forms, as salts
- alkaline earth metals are extracted by electrolysis
- that beryllium is used as an alloys with other metals
- the atoms of the alkaline earth metals are smaller than those of the corresponding Group 1 elements.
SELF-ASSESSMENT EXERCISE

i. Explain briefly why of all nitrogen halides, only NF₃ is Stable

ii. The mechanism of the hydrolysis of phosphorous trichloride, involves the formation of an intermediate four coordinate species. Why is it impossible for NC₁₃ to hydrolyse using the same mechanism?

iii. Why are Group 2 elements smaller in size than their counter parts in Group 1?

6.0 TUTOR-MARKED ASSIGNMENT

1. Explain why the first ionisation energy of beryllium is greater than that of lithium, but the position is reversed in the case of the second ionisation energy.

2. Explain why beryllium forms covalent compounds.

7.0 REFERENCES/FURTHER READING


MODULE 4

Unit 1  Reactivity of Alkaline Earth Metals
Unit 2  Complexing Behaviour of Alkaline Earth Metals

UNIT 1  REACTIVITY OF ALKALINE EARTH METALS

CONTENTS

1.0  Introduction
2.0  Objectives
3.0  Main Content
   3.1  Reactivity of Alkaline Earth Metals
   3.2  Thermal Stability of Oxy Salts
4.0  Conclusion
5.0  Summary
6.0  Tutor-Marked Assignment
7.0  References/Further Reading

1.0  INTRODUCTION

In the Module 3, you studied some of the physical properties of the alkaline earth metals. In this unit, you will be studying their chemical properties. You will also study the stability of oxy salts of alkaline earth metals.

2.0  OBJECTIVES

At the end of this unit, you should be able to:

- write the equations showing the reactions of all the alkaline earth metals with oxygen
- describe the action of each member of the alkaline earth metals with acid
- describe the action of each member of the alkaline earth metals with water
- describe the structure of BeO
- describe using equations the formation of the metal halides
- describe the formation of hydrides by alkaline earth metals
- describe the thermal stability of oxy salts.
3.0 MAIN CONTENT

3.1 Reactivity of Alkaline Earth Metals

Alkaline earth metals are less reactive than alkali metals as they are less electropositive than they are.

Table 1.1 shows some chemical reactions of alkaline earth metals.

Reactivity of alkaline earth metals increases with atomic number down the group. Let us consider these re-activities at a time.

(a) All the metals liberate hydrogen form acids, although beryllium reacts slowly. Beryllium liberates hydrogen when treated with sodium hydroxide solution. The reaction can be represented thus:

$$\text{Be} + 2\text{H}_2\text{O} + 2\text{OH}^- \rightarrow [\text{Be(OH)}_4]^{2-} + \text{H}_2(g).$$

This shows the anomaly in the behaviours of beryllium, in having amphoteric properties

(b) All the alkaline earth metals burn in oxygen to form oxides, MO. Oxides can also be formed by thermal decomposition of MCO$_3^-$, M(OH)$_2$, M(NO$_3$)$_2$ and MNO$_4$. With the exception of Beryllium oxide which is covalent, all other oxides are ionic in nature.
BeO has a wurtzite (hexagonal ZnS) structure (Figure 1.1). In this structure, each ion has four nearest neighbours distributed tetrahedrally around it. Others have sodium chloride type of structure, i.e. each metal ion, $M^{2+}$, is surrounded by six $O^{2-}$ ions and each $O^{2-}$ is surrounded by six metal ions (Figure 1.2). CaO is prepared on a large scale by heating calcium carbonate in lime kilns and is used in the manufacture of Sodium Carbonate, Calcium carbide, bleaching powder, glass and cement.

(c) Barium peroxide, $\text{BaO}_2$ is formed by passing air over heated $\text{BaO}$, at 800K. Strontium peroxide $\text{SrO}_2$, is obtained in a similar way at high temperature and pressure. Calcium peroxide, $\text{CaO}_2$ is obtained as a hydrate by treating $\text{Ca(OH)}_2$ with hydrogen peroxide $\text{H}_2\text{O}_2$ and then dehydrating the product.

Magnesium peroxide, $\text{MgO}_2$ is obtained only in the crude form by using hydrogen peroxide but no peroxide of beryllium is known. These peroxides are ionic solids having $(O-O)_{2-}$ ion and can be considered as salts of very weak acids, $\text{H}_2\text{O}_2$. The peroxides on treatment with dilute acids form $\text{H}_2\text{O}_2$.

(d) Alkaline earth metals react less readily with water than alkali metals to give hydrogen and metal hydroxides. Beryllium does not react with water or steam even at red heat. Magnesium reacts with hot water and Ca, Sr and Ba react with cold water to form the corresponding hydroxides. Beryllium hydroxide, $\text{Be(OH)}_2$ is amphoteric; the basic strength increases in the order Mg to Ba. Aqueous solutions of calcium and barium hydroxides are known as lime water and baryta water respectively.

$\text{Ca(OH)}_2$ reacts with $\text{CO}_2$ to form first an insoluble $\text{CaCO}_3$ which further reacts with $\text{CO}_2$ to give soluble bicarbonate.

Calcium and barium bicarbonates are stable only in solution and decompose on removal of water to give carbonates. This property of bicarbonates is the reason for **STALACTITE** formation (The downward growth of CaCO$_3$ formed on the roof of a cave by the
tricking of water containing calcium compounds) and **STALAGMITE** (he upward growth from the floor of a cave; by the tricking of water containing calcium compounds) formation.

(e) Metal halides, are obtained by direct combination with halogens as well as by the action of halogen on metals. Beryllium halides are covalent and other metal halides are ionic. Beryllium halides are hygroscopic and fume in air due to hydrolysis. They sublime and do not conduct electricity. Anhydrous beryllium halides are polymeric. Beryllium chloride vapours contain BeCl$_2$ and (BeCl$_2$)$_2$ but the solid is polymerised and can be represented as (BeCl$_2$)$_2$ (Figure 3.3).

![Fig. 1.2 Wurtzite (ZnS) Structure](image)

![Fig. 1.3 Rock Salt (NaCl) Structure](image)

![Fig. 1.4 Beryllium Chloride (a) Monomer (b) Dimmer and (c) Polymer](image)

The halides are hygroscopic and form hydrates. CaCl$_2$ is a well known drying agent and anhydrous MgCl$_2$ is important in the electrolytic extraction of Mg.
(f) All the Group 2 elements except beryllium form hydrides, MH$_2$, by direct combination with hydrogen.

Beryllium hydride can be formed by reducing beryllium chloride with lithium aluminium hydride LiAlH$_4$. All these hydrides are reducing agents which react with water to liberate hydrogen. Calcium strontium and barium hydrides are ionic and contain the hydride ion H-. Beryllium and magnesium hydrides are covalent and polymeric (BeH$_2$)$_n$ has an interesting structure. The polymeric solid contains hydrogen bridges between beryllium atoms (Figure 1.4)

![Fig. 1.4: Beryllium Hydride Polymer](image)

Each beryllium atom is bonded to four hydrogen atoms and each hydrogen atom forms two bonds as it bridges two Be atoms. Since Be has two valence electrons and H only one, it means that there are no enough electrons to form the usual type of bonds in which two electrons are shared between two atoms. Instead of this, center bonds are formed in which a "banana — shaped" molecular orbital covers three atoms Be.........H......:Be containing the two electrons. The monomeric molecule BeH$_2$ if formed with normal bonds would have only four electrons in the outer shell of the beryllium atom and would be electron deficient. This would make the molecule very unstable. That is why BeH$_2$ exists as a cluster compound in which each atom shares its electrons with several neighbouring atoms and receives a share in their electrons in order to acquire a stable configuration.

(g) All the metals in the Mg - Ba series or their oxides react directly with carbon to give the carbides (acetylides), MC$_2$. These carbides are ionic in nature and have a NaCl type of structure (Figure 1.2) with M$^{2+}$ replacing Na$^+$ and C=--C 2- replacing Cl-. Beryllium forms methanide, Be$_2$C, with carbon, and acetylide BeC$_2$ with acetylene.

Magnesium on heating with carbon forms Mg$_2$C$_3$, which is an allyl ide since with water it liberates all ylene (methyl/acetylene). The naming of carbides depends on the hydrocarbon they liberate on reaction with water. If acetylene is liberated it is called acetylide. If methane is liberated it is methanide etc.:
Alkaline earth metals burn in nitrogen to form nitrides, $\text{M}_2\text{N}_2$ it requires a lot of energy to convert. The stable $\text{N}_2$ molecule into nitride ion, $\text{N}^{3-}$, and this is recovered from the very high lattice energies of the alkaline earth metal nitrides. Beryllium compound is rather volatile while others are not. They are all colourless crystalline solids which decompose on heating and react with water to liberate ammonia and form either the metal oxide or hydroxide, e.g.:

$$\text{CaC}_2 + 2\text{H}_2\text{O} \rightarrow \text{Ca(OH)}_2 + \text{C}_2\text{H}_2 \text{ (acetylene)}$$
$$\text{Be}_2\text{C} + 4\text{H}_2\text{O} \rightarrow 2\text{Be(OH)}_2 + \text{CH}_4 \text{ (methane)}$$
$$\text{MgC}_3 + 4\text{H}_2\text{O} \rightarrow 2\text{Mg(OH)}_2 + \text{CH} - \text{C} = \text{CH} \text{ (allene)}$$

### 3.2 Thermal Stability of Oxy Salts

All Group 2 elements form oxy salts. The thermal stability of the oxy salts increases with the increase in electro-positivity of the metal, it increases down the group. The salts of group 2 are thermally less stable than those of group 1. The carbonates of alkaline earth metals are stable at room temperature. On heating, they decompose into the corresponding oxides and carbon-dioxide.

$$\text{MCO}_3 \rightarrow \text{MO} + \text{CO}_2$$

The temperatures at which the carbonates decompose are:

$$\text{BeCO}_3 < 400\text{K}; \text{MgCO}_3 800\text{K}; \text{CaCO}_3 1200\text{K}$$
$$\text{SrCO}_3, 1550\text{K}, \text{BaCO}_3, 1650\text{K}.$$
The order of decomposition temperature of the sulphate is:

\[ \text{BeSO}_4: 850K; \text{MgSO}_4: 1150K; \text{CaSO}_4: 1400K; \text{SrSO}_4: 1650K. \]

Alkali metal nitrate decompose into nitrites on heating whereas alkaline earth metal nitrates decompose on heating to metal oxide, nitrogen dioxide and oxygen.

\[ \text{eg } 2\text{Ca(NO}_3)_2 \rightarrow 2\text{CaO} + 4\text{NO}_2 + \text{O}_2 \]

### 4.0 CONCLUSION

We can conclude this unit by stating that the alkaline earth metals are generally less reactive than the alkali metals. Some of the reactions of the alkali earth metals include their action on acid to liberate hydrogen and their reaction with oxygen to form oxides which with the exception of that of beryllium are ionic in nature.

### 5.0 SUMMARY

In summary, you have learnt the following in this unit:

- that alkaline earth metals are less reactive two than alkali metals
- that the alkaline earth metals react with acids to liberate hydrogen
- that all alkaline earth metals burn in oxygen to form oxides
- that the structure of BeO is the wurtzite (hexagonal ZnS) type, and that of other oxides is the NaCl of structure
- that the alkaline earth metals react less readily with water
- that the result of the reaction between alkaline earth metals and water is the formation of hydrogen and metal hydride oxides
- that alkaline earth metals combine directly with halogens to form metal halides which are hygroscopic in nature and some e.g. CaCl₂ are used as drying agents
- that all Group 2 elements with the exception of beryllium forms hydrides (MH₂) by direct combination with hydrogen
- that all the metals in the Mg — Ba series or their oxides react directly with carbon to give the carbides (acetylides) MC₂.
SELF-ASSESSMENT EXERCISE

i. Explain why noble metals like Au, Pt, Rh and Ir are not attached by nitric acid, but Au and Pt are dissolved by AQUA REGIA.

ii. What are the usual coordination number e.g. Be$^{2+}$ and Mg$^{2+}$? What is the reason for the difference?

6.0 TUTOR-MARKED ASSIGNMENT

1. Explain in brief why the hydride bridge is (BeH$_2$)$_n$ is considered to be electron deficient but not the halide bridge in (BeCl$_2$)$_n$.

2. Which is more stable to heat, beryllium carbonate or barium carbonate and why.

7.0 REFERENCES/FURTHER READING


UNIT 2 COMPLEXING BEHAVIOUR OF ALKALINE EARTH METALS

CONTENTS

1.0 Introduction
2.0 Objectives
3.0 Main Content
   3.1 Complexation Behaviour of Alkaline Earth Metals
   3.2 Anomalous Nature of Beryllium
4.0 Conclusion
5.0 Summary
6.0 Tutor-Marked Assignment
7.0 References/Further Reading

1.0 INTRODUCTION

In Unit 1, you studied the chemical properties of alkaline earth metals. You learnt about the reactions of the alkaline earth metals with oxygen, water and acids. You also studied the thermal stability of their oxy salts. In the course of our studies, you must have noticed that the first member of the alkaline earth metal always behave differently from the rest. In this unit, you will be studying the anomalous behaviour of beryllium. You will also be studying the complexation behaviour of the alkaline earth metals in this unit.

2.0 OBJECTIVES

At the end of this unit, you should be able to:

- list at least one complex that is formed by each member of the alkaline earth metals
- describe the structure of the complexes formed by members of the group
- list the properties in which beryllium differs from other members of the alkaline earth metals.

3.0 MAIN CONTENT

3.1 Complex-Forming Behaviour of Alkaline Earth Metals

An interesting property of the alkaline earth metals is their complex behaviour. As we have seen earlier, complex formation is favoured by small, highly charged cations with suitable empty orbitals of approximately the right energy with which the Ligand orbitals can
combine. Alkaline earth metal form more complexes as compared to alkaline metals. The tendency to form complex (mostly with 0 and N donors) decreases with increasing atomic number. Thus, of the heavier ions, only Ca\(^{2+}\) forms a complex with ethanol.

Beryllium having the smallest ion in the group tends to form complex most readily. It mostly forms complexes with tetrahedral arrangement because of the available orbitals as shown in Figure 2.1.

![Fig. 2.1 The Tetrahedral Structure of Tetrafluoroberyllates](image1)

Beryllium forms white crystalline molecular oxide — carboxylates of which basic **Beryllium Acetate** Viz \([\text{OBe}_4 (\text{CH}_3 \text{COO})_6]\) is typical. It is used for the purification of Be because of its volatility and solubility in organic solvents. Beryllium forms a number of chelates with legands like oxalates, \([\text{Be} (\text{C}_2\text{O}_4)_2]^{2-}\) (Figure 2.3) and B-diketonate anions.

![Fig. 2.2 TetrafluoroberyllaLe Complex Ion (BeF\(_4\))^{2-}\](image2)
In the hydrated salt, example BeCO$_3$.4H$_2$O and BeSO$_4$.4H$_2$O beryllium ions exist in the form [Be (H$_2$O)$_4$]$^{2+}$ where they show a coordination number of four. Magnesium is known to form a very important complex occurring in nature as in chlorophyll; a green pigment of plants which produces sugar in the presence of sunlight, carbon dioxide and water in a process known as photosynthesis.

Magnesium in chlorophyll is coordinated by four nitrogen atoms in the heterocyclic porphyrin ring system (Figure 2.4)

The rest of the alkaline earth elements from calcium to barium form complexes only with strong complexation agent such as acetylacetone,
ethylene diaminetetra acetic acid [EDTA] etc. In fact titrations are performed using EDTA in buffer solution to estimate the amount of Ca\(^{2+}\) and mg\(^{2+}\) present in water to determine the hardness of water.

### 3.2 Anomalous Nature of Beryllium

Beryllium, the first member of the alkaline earth metal group, differs from the members of the group just as lithium differs from three members of its group. In facts beryllium differs more from magnesium than lithium does from sodium. As we shall see in the later units, the anomalous nature of the first member of s- and p-block groups becomes more pronounced toward the middle of the table.

Beryllium also shows a diagonal resemblance to aluminium in the same way as lithium does to magnesium. The properties in which beryllium differ from magnesium it shares with aluminium in general. Let us now look at these properties one at a time.

A The cohesive properties of beryllium are much greater than those of magnesium and other elements in the group. As a result, it is much harder and has higher melting points.

B It has smaller atomic radii higher electron affinity and higher ionization energy.

C Its higher polarizing power leads to all its compounds being largely covalent with lower melting and boiling points, and enthalpies of formation (example BeF\(_2\), m.p 1073 K while for the rest of the group is about 1573 k). All the compound of Be are more soluble in organic solvents than the corresponding magnesium compounds. They hydrolyse in water and in this respect they rather resemble aluminium.

### 4.0 CONCLUSION

In conclusion, you have seen in this unit that alkaline earth metals form more complex than the alkali metals of Group l. We have also seen that complex formation favour smaller cations. The ability to form complexes decreases with increasing atomic number. Thus, Be forms more complexes than Mg, Ca, and Ba. Beryllium shows behaviours which differ from that expected from a member of Group 2.
5.0 SUMMARY

In this unit, you have studied the following:

- alkaline earth metals form complexes
- beryllium being smaller in size than the other member of the group forms more complexes
- magnesium forms important complexes that occur in nature. An example of such a complex is the one found in chlorophyll, the green pigment found in plants
- beryllium, the first member of the alkaline earth metal shows anomalous behaviour that is, it behaves differently than it is expected to do as a member of the group
- this anomalous behaviour is manifested in it having much higher melting and boiling points. It also has smaller atomic radii, higher electron affinity and higher ionisation energy.

SELF-ASSESSMENT EXERCISE

Explain why noble metals like Au, Pt, Rh and Ir are not attacked by nitric acid, but Au and Pt are dissolved by Aqua regia.

6.0 TUTOR-MARKED ASSIGNMENT

Why do alkaline earth metals form more complexes as compound to the alkaline metals? Give only two reasons.

7.0 REFERENCES/FURTHER READING
