Republic of Iraq
Ministry of Education
General Directorate of Curricula

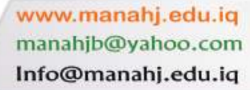


Fifth Class of Science Applied Branch

Revised A Specialized Committee From Ministry of Education

Scientific Supervisor : Dr. Huda Salah Kareem

Design Supervisor: Hiba Salah Mahdi





Translated from 9th Arabic Revised Edution

استناداً الى القانون يوزع مجاناً ويمنع بيعه وتداوله في الاسواق



Introduction

Chemistry is a practical, theoretical science. These key pillars have grown rapidly and the outcome has covered most aspects of our daily lives, mostly meeting the needs of society Environment and living conditions. The continuous coordination with the rest of the other biological sciences, especially physics, mathematics and life sciences.

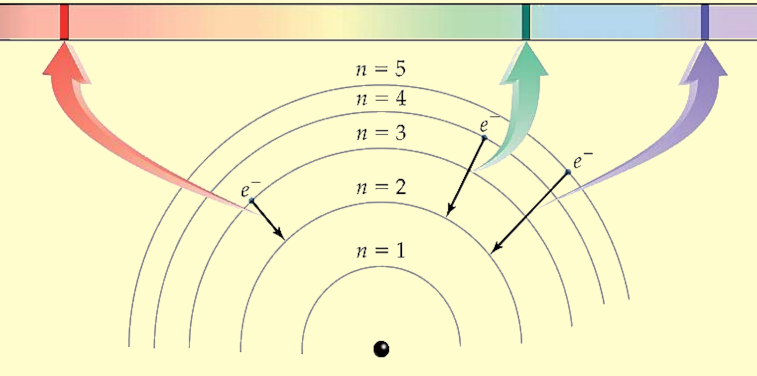
The Authoring Committee of the book of chemistry for the fifth grade of science; Interest in enriching the content of the vocabulary of the new curriculum in chemistry to encourage the student and interested to follow up on this vital science, taking into account the sensitive age for the student and taught in previous years of this science starting from the first secondary study. The committee emphasizes the need to support the theoretical material in the book with two main pillars: conducting practical experiments (including the practical needs of the teacher) and linking chemistry as much as possible with daily life and its impact is growing in industry, agriculture, nutrition and medicines cosmetics and others.

The chapters of the book included both horizontal and vertical expansion that these chapters were modern and simple at the same time. Subject to the use of the same symbols, formulas and references used in chemistry books for previous years to facilitate the student and teacher and to be the chemistry curriculum in the secondary stage a coherent gradient (with the addition of the update required by the various unit codes).

The committee assures of the chemistry teachers not to enter any symbols, formulas or terminology, which is waged by confusing the minds of students and adhering, as far as possible, by the quality of the questions and activities mentioned in end of each chapter - direct applied and deductive as well as mentioned exercises In the margins and the accompanying educational information illuminate the ideas of students and not be a stressful part In the exam questions and in order to complete the benefit. The committee considers that it organizes visits to qualitative and annual exhibitions and the establishment of trips - scientific and entertainment at the same time - to a factory near the school and informing students - on the ground - the successive stages of manufacturing raw materials, through to the final product, because it is a goal of great importance.

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Chapter One Development of the atomic concept

After completing this chapter, the student is expected to:

- Understand the importance of electrical discharge tubes experiments in electron and proton discovery
- Characterize the properties of cathode ray and channel ray.
- Know the value of a single electron's charge, its mass, and how it was found.
- Show the importance of nucleus discovery in the development of atomic structure.
- Know the phenomenon of electromagnetic radiation.
- Understand the meaning of quantum energy and the importance of quantum theory.
- Distinguish the difference between linear and continuous spectrum.
- Show the dual nature of the electron.
- Differentiate between the perception of the principal energy level according to Bohr's theory and mechanics waveform.
- Recognize the four quantum numbers and their importance.
- Determine the four quantum numbers of any electron in an atom.
- Know the principle of exception to Pauli.

Do you know

Zinc sulfide is a fluorinated substance light flashes when light falls on it.



Figure 1-1

Model for electric discharge tube.

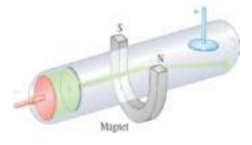


Figure 1-2

Cathode rays were affected by polarity magnetism.

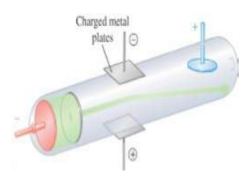


Figure 1-3

Cathode rays were affected by electrical plates.

1-1 Introduction

It came in Daltons atomic theory in one of its clauses stated that "atoms are not able to divide or fragment" but the electrical discharge experiments through gases and radioactivity phenomenon particales, meaning that atom is divisble.

1-2 Electron Discovery

When supplied with an electric discharge tube, which contains hydrogen gas under low pressure (Figure 1-1), with a zinc sulfate-covered reagent barrier as well as a metal plate near the cathode, with a narrow rectangular incision. When the electrical current passes, we see a luminous line across the barrier, and the explanation for that is beams or electric rays emanate from the cathode, and are moving to anode, the metal plate stops most of them, but a rectangular incision allows to this narrow beam runs through it, and hits the barrier and produces a luminous line. Because the beams appear to be coming from the cathode, they are known as cathode – ray. The most important properties of cathode rays are:

- 1 Moving in straight lines emitted from the negative pole towards the positive pole.
- 2 It consists of particles with a very tiny mass and moving very fast.
- 3 Affected by the electric field and attracted towards the positive electrode, which indicates it's a negative charge [Figure (1-3)].
 - 4 Affected by the magnetic field [Figure (1-2)].
 - 5 Ionize the medium in which it passes.

1-3 Proton detection

The first experiments conducted by Goldstein in 1886, to suggest that positive particles are also formed in the electric discharge tubes. In the Goldstein tube [Figure 1-4] there is an positive pole (Anode) towards the right and negative pole (cathode) on the left, which is formed from a piece of metal engraved by a hole, a detector barrier can be placed.

On the tube, to the left of the cathode. When passing an electric current a luminous line is formed, the formation of this line can be explanied as follows:electrons that are emitted from cathod are attracted towards anode and because there is hydrogen gas in the tube, electrons collide with Hydrogen neutral gas atoms, if the electrons have enough energy, they can flush away other electrons from the neutral atoms. positive particles remaining in its place are capture mostly by electrons and become neutral, a few numbers of these positive particles slip from the hole to the area behind the cathode "so called a channal ray". From the study of the properties of these rays found the following:

- 1- Attracted towards the negative pole, which confirms that it is positively charged.
- 2- It has mass and speed which indicates that it is matter particles and are usually heavier than electrons and its mass depends on the type of gas presents in the electric discharge tube.
 - 3- Affected by electric and magnetic fields.

1-4 Estimating the ratio of the charge of the electron to its mass

The first quantitative study on the deviation of electronic beams by electric and magnetic fields was carried out by the scientific Thomson in 1897 and Figure 1-5 illustrates a model, where he used a cathode-ray tube and, by using the properties of these rays. He found that these rays deviate from their straight path under the influence of the magnetic field but returns to its original path when the electric field is shed equal to the intensity of the magnetic field in a direction perpendicular to the direction of the magnetic field. From know the intensity of the electric and magnetic fields, it was possible to calculate the charge-to-mass ratio of particles. Thomson shows that these constant percentages do not change regardless of the metal which is used to make the cathode or the nature of the gas used inside the cathode ray tube. He has been found the ratio equal to 1.76 x 10¹¹ C/kg (coulomb / kilogram).

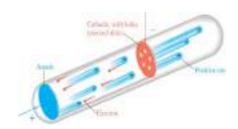


Figure 1-4
Goldstein tube

Exercise 1-1

What are the properties of cathode rays?

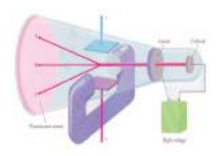


Figure 1-5 Model for Thomson tube.



Figure 1-6
A model of the Melican experiment.

Do you know

Rutherford's expression was when he saw the results "It was something great, almost unbelievable It was as if a shell was fired for range 15 inch on a thin translucent piece of paper then it bounced again and hit you.

1-5 Determination of electron charge

In 1909, Melican assigned an electron charge by placing small negative charge droplets between two electrodes as in Figure 1-6, the positive electrode above tries to attract oil droplets which is charged with negative charge and at the same time droplets tend to descend down by the power of gravity force. When the droplets stabilize in static state between the poles and from the known values of radius, density and intensity of the electric field enables Melican to calculate the carried charge on these droplets. The value is 1.6×10^{-19} C coulombs or one of their simple multiples because droplets can be charged with more than one negative charge. The charge of one electron must be equal the value of the smallest charge carried by the droplet 1.6×10^{-1} ¹⁹C and by using this value of the electron charge and the value of the ratio of the charge of the electron to its mass which is calculated by Thomson (paragraph 1 - 4 above) could calculate the mass of an electron from the electron charge found by Melican:

Electron mass=(The electron charge found by Melican)/(The ratio of the charge of the electron to its mass created by Thomson)

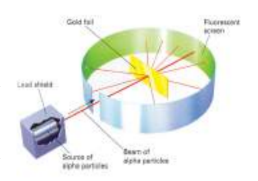
Electron mass =
$$\frac{1.6 \times 10^{-19} \text{ C}}{1.76 \times 10^{11} \text{ C/ kg}}$$

Electron mass = 9.1×10^{-31} kg

1-6 Detection of the nucleus

In 1910 Rutherford conducted experiment which tested, he was examining the dispersion of alpha particals by thin slices of metal. As expected, 99% of alpha particals penetrated the slice, some of them skewed sharply and a few of them were reflected along her trajectory [Figure 1-7]. That was not true at all for Rutherford because that the positive charge and mass in the metal slice are centralized in a very small area. According to Rutherford opinion, the atom has a nucleus or center, where its positive charge and mass are centralized.

The quantitative results of dispersion experiments illustrate (such as that carried out by Rutherford) that the nucleus of the atom has a diameter, which is equal to 10⁻¹³cm and atoms have diameters approximately 100,000 times the diameter nucleus (10⁻⁸cm).In other words, the nucleus occupies very little space from the total size of the atom and most of its size is a space occupied by electrons.



1-7 Detection of atomic number

If the nucleus is consist of protons, how many protons are that exist in a particular nucleus? This question remained until the answer came from scientific observations that are shown by Mossile in 1913 which led to the discovery of the atomic number.

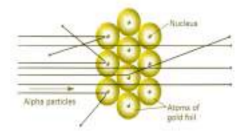


Figure 1-7

A model for the Rutherford expermant

Exercise 1-2

Who attributes the discovery of the nucleus of the atom?

Do you know

If the atom becomes bigger till the nucleus size as a point (.) then the whole atom would be bigger than a house.

Do you know ____

The atomic number (z) is significant as a special characteristic In chemistry, since it gives a number of protons the positive charge in the nucleus which is equal to the number of electrons out the nucleus which make the atom is neutral. And including (Z) represents the number of protons in a nucleus atom is a nuclear property that provides us with important information about the rest of the atom.

1-8 Detection of a neutron

In 1932 Chadwick threw a thin slice of beryllium in alpha particles, there were appeared rays resembling the rays of Kama which has very high energy from metal slice. Subsequent experiments showed that these rays represent the third model of the components of the atom, called by Shadwick as Neutron because its charge is neutral and its mass is roughly equal to that of the proton.

1-9 The configuration of electrons in an atom

The process of arranging the components in atom has gone through several theories. These theories were fading or evolving according to their ability to interpret physical and chemical phenomena. The model is considered to be an atom is consisting of a positive nucleus which the negative electrons are moving around it. This movement of electrons counteract the gravitational force generated by nucleus. Explanation of why these electrons do not radiate energy as they are observed in all other cases.

Moving electric charges under the influence of gravitational forces are losing energy. In the case of electrons, a loss of energy will result in a slow motion of the electron, which will gradually attract it to the nucleus in a spiral path approaching them to fall inside.

In this spiral path continuously gives energy as a continuous spectrum that resembles a semi - ray solar spectrum but this does not happen as the atoms are stable in their atomic construction. This atomic stability led scientists to present other theories about atomic construction.

1-10 Electromagnetic radiation

This expression includes different types of radiation, such as warmth it travels to us from a fireplace or light reflected on bright surfaces, or radiology used in hospitals. They are different but they are common in some intrinsic qualities, all these types of radiation are transmitted in the space in same speed equal to $3x10^8$ m/s, which is light speed. All of these types are wave-nature and their waves are similar to those which form above the surface of the water in which a stone is thrown [Figure 1-8]. We notice repetition in these waves, that is, the wave is repeated at regular intervals and the regular number of these waves is what is known as the frequency of the wave that occurs per second ,and when radiation passes at a specified point [Figure 1-9]. The waves are small and their frequency is high and the opposite is true and is related to length frequency wave in relation:

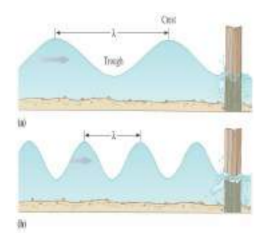


Figure 1-8

Model to form waves above the surface water.

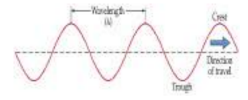


Figure 1-9

A model of the wave nature of light.

Where C: The speed of light in unit (m/s)

 \mathfrak{V} : Frequency in unit (1/s)

 λ : Wavelength in unit (m)

1.11 Quantum theory

 $v = \frac{C}{\lambda}$

Max Plank paved the way for the scientific revolution that emerged at the beginning of the twentieth century, by his hypothesis which is considered radiology electromagnetism is like a bundle of small energy groups which he called it Quantum. When he was studying the phenomenon of light emission from hot objects, Planck suggested that hot objects emit energy in small quantities is called (quantum). It is the minimum amount of energy that can be lost or gained by the body The process of losing or acquiring energy is not a continuous process, but rather batches or flows of energy so that the energy is lost or the acquired multiplier is a true multiple of that quantum or quantity. The amount of quantum depends upon the frequency of radiation,

and is associated with a constant (named Planck constant) and its value 6.63×10^{-34} J.s . Planck's energy is known as:

E=h v Where E: Energy in Joule (J) h: Planck constant in unit (J.s)

Einstein then came to expand Planck's theory of interpretation of the photoelectric effect phenomenon, a phenomenon of electrons emitting from a metal when the light is pointed to, [Figure 1-10]. The emission of electrons depends on the frequency of light, ie on its energy, if the frequency of atomic radiation hits the metal surface in low value, we are not observe the emission of electrons. When using increased frequencies, the frequency suddenly reaches at which it begins emitting electrons. Before that, however, the radiation intensity of the low frequency was increased the metal is not affected. If the radiation frequency exceeds that the minimum value of emission does not increase the number of electrons emitted but the energy carried by the electron increases. Einstein has used Planck's equation to calculate the quantum energy of light, which he called (photons) and photon is a particle of electromagnetic radiation has a mass of zero but holds a quantity of energy depends on the frequency of the electromagnetic wave. The photon energy has the lowest frequency needed to emit electrons by overcoming its binding energy that electrons relate to the atom. Einstein's interpretation of this phenomenon was the first application important for quantum theory. This interpretation has also given light a trait particle along with the wave characteristic.

1-12 Atomic Spectra

The scientific Newton passed ordinary sunlight through a glass prism, he found that the light decomposes into a range of colors firstly from violet and finally to red (Figure 1-11). Because there are no separate areas between one color to another, this spectrum is named with continuous spectrum. But it was observed that if the atoms of a pure element exposure into the gaseous state to heat or in the electric discharge tube in the case of low pressure emits radiation (spectrum) of a glowing element atoms is not connected or continuous but is consisted from a few lines of light separated by relatively large dark distances.

Do you know

Impose a car moving at a certain speed, if we want to increase its speed to reach higher speed the car must pass through all speeds that fall between lower speed and higher speed. It cannot be jump from speed to higher speed, as requires of the idea of quantum leaps.

Exercise 1-3

What is a photon?



Figure 1-10

Photoelectric effect phenomenon.

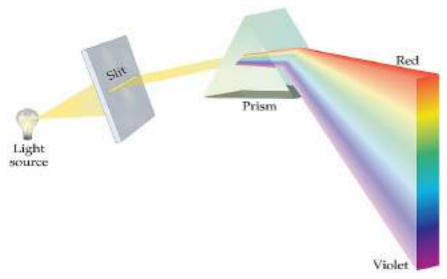


Figure 1-11

Continuous spectrum phenomenon.

Do you know

Each element has a special emission spectrum. Emission spectra are used to designate Identities of unknown samples and percentage determination components of stars.

call it (the line emission spectrum) because there are spaces between one color and another. It was later discovered that each element had a spectrum linearly distinguishes it from other elements [Figure 1-12]. linear emission spectrum shows that the emitted radiation from an atom emits a specific energy. In other words it is emitted in a quantum style rather than in a continuous manner.

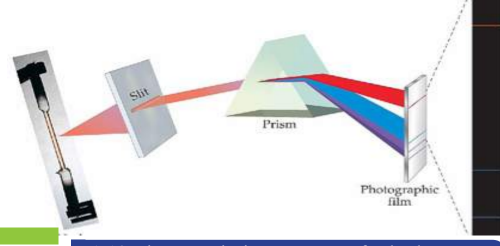


Figure 1-12

Linear spectrum phenomenon.

Exercise 1-4

That rainbow is a series of colors?

Discuss whether you consider this series of continuous or linear emission spectrum.

1.13 Linear emission spectrum for hydrogen

Classical theory claims that a hydrogen atom can be excited by any amount of energy that receive. So scientists predicted the emission of a series of continuous spectrum, but the hydrogen atom emits only limited frequencies of light, why, when the current passes in the hydrogen gas under low pressure. The potential energy of some of its atoms increases and the lowest atomic energy level is called (ground state). When the potential energy level of the atom is higher than the ground state led to atom to (excited state).

When the atom returns from excited level to the stable level, it lose the gained energy in form of electromagnetic rays (photons) and the energy of this photon is equal to the difference between first and second energy level [Figure 1-13]. Studies showed that hydrogen atoms release only specific frequencies of light. The energy difference between these energy levels of an atom is a specific difference and this means that the hydrogen electron in limited energy level.

1-13-1 Bohr's theory of hydrogen atom

After the discovery of the elements spectra, scientists tried to explain them based on the movement of the electron in the atom, but the attempt failed. Suppose the electron moves around the nucleus at a constant speed, in a curved path and because of laws of physics that require any partical charged as an electron undergo a process acceleration will lose energy on the image of electromagnetic radiation. This slows down the speed of the electron and therefore the electron will collide to nucleus and the atom collapses. Because atoms do not collapse, scientists had to challenge to explain how electrons rotate. The scientist Bohr used ideas of Planck and Einstein mentioned earlier in his theory that he managed of calculating the energy of an electron in hydrogen atom. Bohr was adopted in his theory on two facts: first, the atoms do not collapse and the second is that light emission from an atom is done at a certain frequency, which means that changes in the atom energy are certain and specific.

This shows that the electron exists in areas with specific energy and cannot exist between them because it is forced to specific energy levels in the atom.

In his theory, Bohr assumed that electrons have specific orbit with constant energy and volume [Figure 1-14]. Bohr theoretical hypotheses can be summarized as follows:-

- 1. The electron orbits in specific orbit (with a specified diameter) and a specific energy and do not radiate energy as a result of this rotation.
- 2. Energy is emitted from the atom in one case, which is the transmission of the electron from specific orbit to another orbit that has less energy than in first orbit.

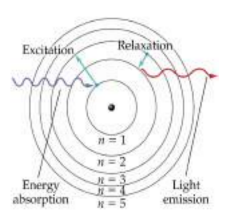
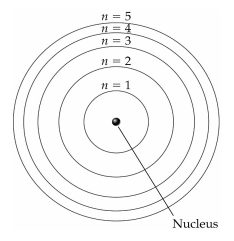


Figure 1-13

The transmission of an electron from a level of less energy to the level of the highest energy and the reverse.



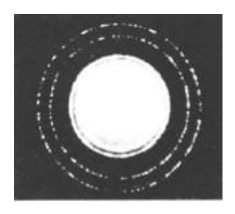


Figure 1-14

Major levels of energy with constant volume and constant energy.

Bohr's theory has proved useful in the structure of a hydrogen atom, but when scientists tried to apply the ideas in Bohr's theory to atoms of the other elements was failed because the rest of the atoms contained more electrons so the atomic spectra of these elements are more complex of the atomic spectrum of hydrogen. That means Energy levels is more complex and there are sub-levels of energy starting from the second level so he start thinking about it again and activate search for a more comprehensive theory.

1-14 The wave nature of the electron

We said that an electron is a particular with a specific mass and negative charge and it has the nature of light. De Broley (1924) suggested the possibility of duality nature (wavelength and particulate) in the case of the particles also depends on what Einstein and Blank achieved.

$E=mc^2$	(1)	Einstein equation
E=hv	(2)	Blank Equation
Since equation 1	and 2 are equal	
$hv = mc^2$		
Whereas $v = (c/\lambda)$	by substituting them	into equation (3).

Whereas $v=(c/\lambda)$, by substituting them into equation (3), we obtain:

$$mc^2 = h \frac{c}{\lambda} \qquad(4)$$

By omitting the value of c from either side in equation (4) it becomes:

$$mc = \frac{h}{\lambda} \qquad(5)$$

In order of equation (5) we get:

$$\lambda = \frac{h}{mc} \tag{6}$$

Momentum is known as the following relationship mc=p:

$$\lambda = \frac{h}{p} \qquad \dots (7)$$

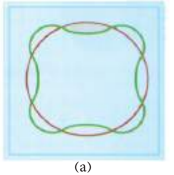
Based on this derivation can a torrent of moving electrons that leads to an interfering pattern, that is, a mobile electronic is similar to photons forming rays.

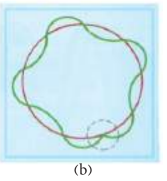
Exercise 1-5

Why Bohr theory failed?

1-15 Wave Mechanics

Despite the success of Bohr theory in the interpretation of the atomic spectrum of hydrogen as well in the interpretation of atomic spectra of systems in which one electron such as(He+ and Li2+) but did not produce good results when trying to explain the spectrum of atoms which has more than one electron. In 1936 the scientific Schrodenger used the mathematics to study the hydrogen atom. A science called wave mechanics or quantum mechanics has begun. Schrodenger solved a mathematical equation named after by his name and also called the wave equation. Schrödinger's idea is that instead of thinking about an electron moving circularly in a constant orbit, we have to assume a series of waves move within this stable orbit and that the perimeter of the orbit must be equal a simple multiple of the electron's wavelength. According to the assumption of Schrödinger electron wave behavior gives an estimate of the probability of existence electron within the wavelengths. This new concept is an alternative to the system of the fixed orbits assumed by Bohr in his theory rather than look of an electron as a particle orbiting in a specified radius orbit wave mechanics describes the motion of an electron in terms of the wave function (Figure 1-15), which we call orbital and it depends on the total energy and its potential energy and the coordinates of its location (z, y, x). Nature of this electron wave behavior makes our ability to determine the location of the electron and its momentum (the amount of movement) at the same time. Heisenberg's principle coined known by his name (the principle of inaccuracy) which states it is not possible to locate a particle and its momentum precisely at the same time is able to measure one with greater uncertainty in the accuracy of measuring the other.





Exercise 1-6

What does the excited state of the atom mean? and what happens to the electron during it?

Figure 1-15

The wave motion of an electron.

Since it is not possible to draw a path for electrons, the best thing to do is talking about the probability of finding an electron at a specific location within an atom. The probability of finding an electron at multiple points within an atom is mathmatical dilemma is very complicated. Quantum mechanics solves this dilemma as an electron as if it was a wave. In fact, the electrons wave properties appear under some conditions, so that the electron beam moving quickly, for example, appears diffraction, a characteristic of motion waveform. To identify any electron at its atom, solve the wave equation and know the values of the wave function, it requires three quantum numbers. These are real numbers relate to the potential energy and position of electron and electronic cloud shape. Fourth electronic quantum number must be set because the three theoretically quantum numbers were resulting from the solution of the Schrödinger equation are not sufficient to explain all observed properties of electrons in atoms. So the fourth quantum number was entered to fill this deficiency and it is called the electron spin quantum number because the electron can think of it as spinning around its axis as it moves around the nucleus. The fourth quantum number shows the direction of the spinning.

Exercise 1-7

How quantum mechanics was able to Explain the presence of an electron in a particular location in atom.

1-16 Quantum numbers

To know how electrons will be arranged we must examine the energy levels in the atom by studying the following quantum numbers:

1-16-1 The principal quantum number (n)

Energy levels in atom are determined by the principal quantum number and the larger the value of n the higher value of energy level and farther the electron from the nucleus. The value of n determines the size of the orbital and n take integers 1, 2, 3, etc.

1.16.2 The momentum quantum number (ℓ)

Determines the shape of orbital in which the electron is likely to be present and it is resulting from the movement of the electron around the nucleus. That each principal level n consists of one or more sub - (secondary) levels. The number of secondary levels in any orbital is equal to the number of principal quantum number, so n=1 consists of one

secondary level of energy s. The second principal level n =2 contains two secondary levels is s and p, and the third principal level, n=3, contains three secondary energy levels are s, p and d as well as the fourth principal level of energy n=4 contains four secondary levels of quantum numbers are s, p, d, and f. Each n value corresponds to a specific value for the quantum secondary number ℓ are integers starting with zero and ending with (n-1). If n=1, that ℓ will start with zero and end with (1 - 1) so there is one value for ℓ = zero but if n= 2 it is start with zero and end with (1 - 2) by1 so there are two values for ℓ is zero and one. So when n = 3 there are three values 0, 1, 2 and when n= 4 there are four values: 0, 1, 2, 3 and so on (Table 1-1).

Table (1-1) The values of ℓ and the symbols of the secondary				ndary		
levels indicated						
ℓ value	0	1	2	3	4	5
The letter signifying them	S	р	d	f	g	h

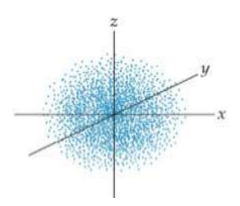
In our study we will only consider secondary levels s. p. d. f because they are only ones that are preoccupied with electrons in atoms in ground state. To determine the secondary level from any n principal level symbolically writes the value of n for the n principal level then the character assigned to the secondary level, for example the secondary level s from the second principal level has the symbol 2s and has values(n=2, $\ell=0$). The secondary level d of the third principal level is 3d and has quantum values (n = 3, $\ell=2$) and so on.

1-16-3 The magnetic quantities number (m ℓ)

Atomic orbitals can take the same shape around the nucleus but in different directions. Magnetic quantities number indicates the direction of the orbital around the nucleus where each secondary level consists of one or more orbitals. This number was used to explain the appearance of extra lines in the spectrum when atom placed in a magnetic field. A spatial distribution of an electron, spherical symmetric, ie, the probability of its existence is the same in all trends from the nucleus [Figure 1-16]. On the other hand the probability the presence of the electron p in some directions of the nucleus is more than in other. In fact, the probability of distribution for the electron p of two diffuse lobes to some extent, one on each side of nucleus [figure 1-17]. The p-level consists of three orbitals

Do you know

The letters s, p, d, f were selected from the letters the first sequences The main Sharp, principal, diffuse, and fundamental these words represent the forms of lines of linear spectra that are connected energy transitions.



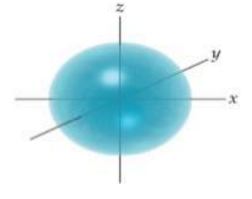


Figure 1-16
Asecondary level of S.

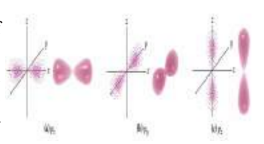
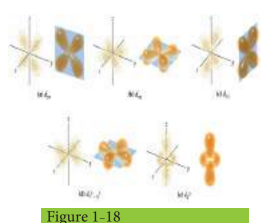


Figure 1-17
Secondary levels of p



Secondary levels of d

The secondary level d consists of five orbitals and the and secondary level f of seven orbitals and their spatial distribution is more complex to a large extent [Figure 1-18]. Each value of ℓ matched with the values of magnatic quantum numbers which are positive and negative integers. When the value of ℓ =0 there is value for m ℓ which is (+1, 0,-1). If ℓ = 1, the values of m ℓ are (+1, 0,-1) and if ℓ = 2 the values of m ℓ is (+2, +1, 0, -1, -2). When the value of ℓ =3 then the value of m ℓ are (+3, +2, +1, 0, -1, -2, -3). Table (1-2) summarizes quantum numbers of primary, secondary and magnetic.

Table (1-2) Values of quantum numbers of primary, secondary and mag-				
netic.				
principal	Secondary	Symbol	Magnetic quantum	Number of
quantum	quantum		number (mℓ)	orbitals at the
number	number			secondary level
(n)	(1)			
1	0	1s	0	1
2	0	2s	0	1
	1	2p	+1, 0,-1	3
3	0	3s	0	1
	1	3p	+1, 0,-1	3
	2	3d	+2,+1, 0,-1,-2	5
4	0	4s	0	1
	1	4p	+1, 0,-1	3
	2	4d	+2,+1, 0,-1,-2	5
	3	4f	+3,+2,+1, 0,-1,-2,-3	7

1-16-4 The electron spin quantum numbers (m_c)

We explained earlier that there is one orbital for the secondary level S, three orbitals for secondary level p, five for secondary level d and seven for the secondary level f and where these secondary levels can accommodates 2, 6, 10 and 14 electrons respectively. It follows that any orbital can accommodate two electrons but electrons differ in the same orbital by one important thing is that it has opposite spindle rotation. The reason for talking about the spin of the electron comes from sightings the magnetic behavior of the material can be obtained from the behavior of magnetic for single atoms, by Ottostron experiment.

In this experiment [Figure 1-19], a beam of silver atoms Neutralization (resulting from the evaporation of silver) passed between two magnetic electrodes. It was found that the beam splits into two separate beams, that is, half the atoms deflects in a certain direction and the rest deflects in the opposite direction.

To explain this behavior that each electron behaves like a fine magnet. This magnetism is produced by the spin of the negative charge because it is known that the spindle rotation of any charge is generated magnetic field. There are two opposite directions of the spin; we expect each electron to attract each other but that disability of the attraction between the electron orbital repulsion in their charge. Since the movement of the rotation of the two electrons is confined in only two directions so we have two values of the spindle quantum ms are $\pm 1/2$ and $\pm 1/2$.

It can summarize the characteristics of the electron in the atom As follows: -

- 1. The principal quantum number (n) and this characteristic indicates the order of the electrons configuration and increase the distance from the nucleus.
- 2. The momentum quantum number $(m\ell)$ describes the type of orbital that the electron operates from, where its spatial distribution is similar (for example, the electron s has spherical distributions

are symmetrical and the electrons p have symmetrical distributions along separated directions in the space).

- 3. The magnetic quantum number $(m\ell)$ and this characteristic defines any orbital from orbitals of the secondary level in which an electron is likely to be present.
- 4. The electron spin quantum number (m_s) and this characteristic determines either of the two possible directions of the spin of the electron.
- 5. When the four characteristics of the electron are determined in an certain atom, it will discover that it cannot exist in the same one atom another electron that has a set similar to those of its the four characteristics. This fundamental definition is known as (the Puli exclusion principle). This definition states that it cannot two electrons in the same atom have the same values for all quantum four number.

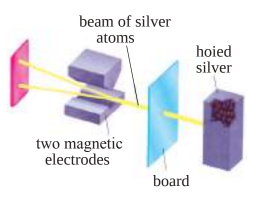


Figure 1-19
Ottostron experiment.

Do you know

To resemble the principal energy levels around an atom by drawer, that does not have a regular dimensional drawer. The distances are very high at first, then gradually diminishing whenever you move away these levels from nucleus.

1-17 How to write an electronic configuration

For the purpose of writing the electronic configuration of any atom, you must know the atomic number of that atom, whereas the number of electrons of an atom is equal to their atomic number (number of protons) if electrically equivalent. Well it should we know that the basic process in writing an electronic structure of an atom is that we start by filling the orbitals with electrons of the less energy and then the most energy, thus where they are arranged as follows: 1s 2s 2p 3s 3p 4s 3d 4p 5s 4d 5p 6s 4f 5d 6p 7s 5f 6d, In the case of secondary levels containing more than one orbital such as levels (p, d and f) electrons enter individually in these orbitals to avoid electric repulsion between charges according to the Hund's rule. This rule states that there is no duplication between two electrons in the secondary energy level only after orbitals occupies an individual firstly.

Exercise 1-8

Write the electronic configuration of following elements: $_{_{30}}Zn$, $_{_{20}}Ca$

Even if all orbitals of the secondary level now have an electron One in each of them as at the secondary level p³

1 1 1 1 1 then the fourth electron enters to double with one electron that preceded it to make full orbitals in the secondary level p⁴

1 1 1 1 and to overcome the repulsion between the two electron charges within a single orbital, the two electrons are represented by two opposite arrows.

Example 1-1

Write the electronic configurations for the following elements: $_{23}V \cdot _{19}K \cdot _{18}Ar$

Solution

If we note in the previous example of the elements that $_{23}$ V, $_{19}$ K That the electronic configuration of the first 18 electrons of them is (1s² 2s² 2p⁶ 3s² 3p⁶). This is the electronic configuration of [Ar] so we can simplify this configuration. So we will write the order of the first 18 electrons of potassium $_{19}$ K and vanadium $_{23}$ V as well as for the elements beyond which to [Kr]

As follows: $_{19}K$ [Ar] $4s^1$, $_{23}V$ [Ar] $4s^2$ $3d^3$

Example 1-2

Write the electronic configuration of the elements ₂₉Cu , ₂₄Cr Solution

$$_{24}$$
Cr [Ar] $4s^2$ $3d^4$

Note that level d needs an electron to become half saturated and atoms when their secondary levels are half full or full be more stable so we write the configuration as follows:

Cr. [Arl 4s¹ 3d⁵

 $3d^{10}$

follows:
$${}_{24}$$
Cr [Ar] $4s^1$ $3d^5$ As for ${}_{29}$ Cu [Ar] $4s^2$ $3d^9$ So we write it as follows:

₂₉Cu [Ar] 4s¹

Example 1-3

Write the electronic configuration for : $_{47}$ Ag , $_{36}$ Kr Solution

Exercise 1-9

Set the four quantum values of the last electron of each of the following atoms:

Example 1-4

Set the four quantum values of the last electron for each of the following element atoms: $_{54}$ Xe $_{23}$ V $_{17}$ Cl $_{3}$ Li Solution

$$_{3}Li$$
 $1s^{2}$ $2s^{1}$

- n=2 because the electronic configuration ended in the second level 2
- ℓ =0 because the electronic configuration ended with the secondary level s
- $m\ell = 0$ because the last electron is located in the secondary level s 1
- ms = +1/2 Because the last electron rotates clock wise (the first electron of the Orbital).

- n=3 Because the configuration ended with the third prin cipal level 3
- ℓ=1 Because the configuration ended with the secondary level p
- $m\ell = 0$ Because the last electron is located in the orbital which has a value of $\ell = 0$

ms = -1/2 Because the last electron is the second electron in the orbital.

$$_{23}V$$
 1s² 2s² 2p⁶ 3s² 3p⁶ 4s² 3d³

- n = 3 Because the highest main level contains electrons at the third
- ℓ=2 Because the electronic configuration ended with the secondary level d
- $m\ell = 0$ Because the last electron is located in the orbital whose value is $m \ell = 0$

ms = +1/2 Because the last electron is the first electron in the orbital.

$$_{54}$$
Xe $_{1}s^2$ $_{2}s^2$ $_{2}p^6$ $_{3}s^2$ $_{3}p^6$ $_{4}s^2$ $_{3}d^{10}$ $_{4}p^6$ $_{5}s^2$ $_{4}d^{10}$ $_{5}p^6$

- n = 5 Because the highest principal level contains elec trons is the fifth
- ℓ=1 Because the electronic configuration ended with the secondary level p
- $m\ell$ = -1 Because the last electron is located in the orbital whose value is m ℓ = -1

ms = -1/2 Because the last electron is the second electron in the orbital.

- Exercise 1-11

Exercise 1-10

atoms:

₅₅Cs, ₃₆Kr, ₂₆Fe

Write the four quantum num-

bers of the penultimate elec-

tron for each of the following

Write the electronic configuration of following ions ${}_{8}O = {}_{6} {}_{3}Li^{*1}$

Example 1-5

Write the four quantum numbers of the last electron for each of the following ions: $_{20}\text{Ca}^{2+}$, $_{9}\text{F}^{\text{-}}$ Solution

F fluoride ion is a fluorine atom that has gained an electron and its number of electrons becomes 10

$$_{9}F^{-}$$
 1s² 2s² 2p⁶

- n=2 Because the highest main level contains electrons is the second
- ℓ=1 Because the electronic configuration ended with the secondary level p

 $m\ell = -1$ Because the last electron is located in the orbital whose value is $m \ell = -1$

ms = -1/2 Because the last electron is the second electron in the orbital.

The Ca²⁺ calcium ion is a calcium atom lost two electrons and the number of electrons become 18

$$_{20}$$
Ca²⁺ 1s² 2s² 2p⁶ 3s² 3p⁶

n=3 Because the highest main level contains electrons is the second

ℓ=1 Because the configuration ended with the secondary level p

 $m\ell$ = -1 Because the last electron is located in the orbital which has a value of ℓ =-1

ms = -1/2 Because the last electron is the second electron in the orbital.

Example 1-6

Compare the four quantum numbers of the last electron to each of the two element atoms ₃Li and ₁₁Na Solution

$$_{3}\text{Li}$$
 1s^{2} 2s^{1}
 $n = 2$
 $\ell = 0$
 $m\ell = 0$
 $ms = +1/2$
 $_{11}\text{Na}$ 1s^{2} 2s^{2} 2p^{6} 3s^{1}
 $n = 3$
 $\ell = 0$
 $m\ell = 0$
 $ms = +1/2$

The difference is only in the principal level.

Example 1-7

If the last electron of an atom of an element has the following four quantum numbers

$$n = 3 \cdot \ell = 2 \cdot m \ell = +1 \cdot ms = -1/2$$

What is the atomic number of this element?

n=3 The principal level will be the third

 ℓ =2 The secondary level is the level and contains five orbitals

Exercise 1-12

Compare the four quantum numbers of an electron the last of both atomic elements.

Exercise 1-13

If the last electron had an element atom has the following four quantum numbers:

n=4.
$$\ell$$
=1. $m\ell$ =0. = ms = -1/2 write the electronic configuration of this atom and what its atomic number.

$$m\ell = +1$$

Located in the orbital indicator with the letter X

ms = -1/2 is the second electron

Exercise 1-14

Write the electronic configuration of ${}_5B$ atom Then write the quantum numbers for all the electrons in which .

So from previous information the last secondary level in the electronic configuration will end with a secondary 3d level which contains seven electrons then become an electronic configuration of secondary levels in this atom as follows:

$$1s^2$$
 $2s^2$ $2p^6$ $3s^2$ $3p^6$ $4s^2$ $3d^7$

adding the number of electrons above the secondary levels in the electronic configuration and the sum value represents the atomic number of this element.

So Atomic number = 27

Example 1-8

Write the electronic configuration of the 25Mn atom and write the quantum number of the electrons, the last secondary level.

Solution

$$_{25}Mn$$
 $1s^2$ $2s^2$ $2p^6$ $3s^2$ $3p^6$ $4s^2$ $3d^5$ $+2$ $+1$ 0 -1 -2

Electron	Quantum number			
	n	ℓ	mℓ	ms
First	3	2	+2	+1/2
Second	3	2	+1	+1/2
Three	3	2	0	+1/2
Four	3	2	-1	+1/2
Five	3	2	-2	+1/2

Chapter One Questions



- 1-1 Describe the model of electric discharge tube with drawing, explain the discovery of the electron?
- 1-2 What are the characteristics of channel rays.
- 1-3 What do the symbols (Z and A) mean?
- 1-4 Explain the effect photoelectric phenomenon. Inter-explanation Einstein to this phenomenon. And what gave this explanation on the nature of light.
- 1-5 What happens to an electron when it gains energy.
- 1-6 What is the importance of exposing the electric discharge tube for magnetic and electric fields when studying properties of electrons and protons.
- 1-7 Talk about the experience of Malikan and what is the important of that have find out
- 1-8 How Rutherford discovered the nucleus of an atom. What are the relationship between the size of an atom and its nuclei.
- 1-9 What are the difference between
- A- Rutherford's model and Thomson's model of atomic structure.
- B- Rutherford's model and Bohr model of atomic structure
- C- Forms of the principal levels of Bohr and quantum theory.
- D- Linear and continuous emission spectrum.
- 1-10 Derive the mathematical relationship λ =h/p based on Planck and Einstein equations.
- 1-11 What is the difference between the secondary level and the orbital? What are the numbers of orbitals in the first principal four levels.

- 1-12 What is the frequency wave? What are the units of frequency, then state the mathematical relationship between frequency and wavelength.
- 1-13 Explain the experience of Autostron, and explain its importance.
- 1-14 What quantum theory assumes.
- 1-15 Explain the following
- A- Non-repulsion of two electrons in the same orbital
- B- Millikan's experience is complementary to the experience Thomson.
- C- The secondary level s can accommodate only two electrons either the secondary level p accommodates just six electrons.
- D- The position and momentum of an electron in an atom cannot be set in high resolution simultaneously.
- E- Secondary level 3p is filled with electrons after secondary level 3s.
- F- The strongest attraction of the nucleus on the electron closest to it.
- 1-16 Define the four quantum numbers (n, ℓ , $m\ell$, ms) What do they benefit from each?
- 1-17 Draw the shape of the orbital when the value of $\ell = 0$ and when they are valuable $\ell = 1$.
- 1-18 What we mean by all of what comes.
- A- Photon.
- B- The wavelength.
- C- The double character of the electron.
- 1-19 What benefits from Hund's base in the of electronic configuration.

1-20 show

A- The Heisenberg principle.

B- Pauli's exception principle.

C- How an atom can emit a photon.

D- How Schrödinger was able to explain the spectrum of atoms which have a higher atomic number of hydrogen.

1-21 What are the values of the principal and secondary quantum numbers and magnetism in the following major levels (third and fourth).

1-22 If you know that the values of the four quantum numbers of the last electron for the atoms (A, B, and C), respectively.

A

$$n = 4$$
 $\ell = 2$ $m\ell = +1$ $ms = -1/2$

В

$$n = 2$$
 $\ell = 0$ $m\ell = 0$ $ms = +1/2$

 \mathbf{C}

$$n = 3 \ \ell = 1$$
 $m\ell = -1$ $ms = -1/2$

Write the electronic configuration of atoms (A, B, and C). What is the atomic number of each.

1-23 Mention at least two components ends up being distributed the last electronic levels are .s² d⁶

1-24 Write the electronic configuration of the following ions ₂₉Cu ⁺², ₂₉Cu ⁺¹

1-25 Write the electronic configuration of atoms and ions of the following ₇₉Au, ₃₄Se, ₁₆S⁻², ₃₇Rb, ₅₀Sn

1-26 Compare the four quantum numbers of the last electron for each of the atoms has the following elements ₃₈Sr, ₂₁Sc, ₂₀ Ca 1-27 Write the electronic configuration of the atom (9F) then write the quantum numbers of all the electrons. Show the principle that corresponds through your feedback to values quantum preparation for electrons.

1-28 Set the four quantum values for the electrons located in the last major level of each The following atoms 4Be, 15P

1-29 Set the four quantum values for the last electron only for each of the following atoms ₇₈Pt, ₃₅ Br, ₁₉K

1-30 choose the correct answer for the following:

1- When the North Pole finder is rounded to a magnet On cathode rays, the shining line is curved into:

A- Down

B- The highest

C- Not affected

2- Mass of one electron equal:

A- 1.76x1011 C/kg

B- 1.6x10⁻¹⁹ e

C- 9.1x10⁻³¹ kg

3- The discovery of neutron is attributed to the scientist:

A- Thompson

B- Henry Mosley

C- James Chadwick

4- The light wavelength is proportional to its frequency

A- directly

B- inversely

C- equal

5- In the photoelectric effect phenomenon if the radiation exceeds minimum emission value:

A- The number of emitted electrons increases

B- The number of electrons emitted decreases

C- The energy carried by the electron increases

6- Quantum number that determines the shape of the electronic cloud is:

A- Principal.

B- Secondary.

C- Magnetic.

7- The diameter of the atom is greater than the diameter of its nucleus

A-1000 times

B-10000 times

C-100000 times

8. If n = 2, there are:

A- one specific value for the secondary quantum number = zero.

B- two specific values for the number of secondary quantum = 0 and 1

C- three specific values for the number of secondary quantum = zero, 1 and 2.

9. The principle which states that two electrons cannot exist in the same atom would have the same values, of the four quantum is:

A- The Heisenberg inaccuracy principle.

B- The Pauli exclusion principle.

C- The principle of the rule of Hind.

10. If the electronic configuration ends at the secondary level 2p⁶ there is a possibility:

A- The atom should be only one element which is an element. Ne

B- That there are two element atoms, namely _oF and ₁₀Ne

C- That there is more than one atom element in addition to ₁₀Ne that these elements are lost or acquired electrons to be rearranged like an atom element ₁₀Ne

11- If the electronic configuration of an element's atom ends 4s² 3d⁵, so The four quantum numbers of the last electron be:

A- n = 3 $\ell = 2$ $m\ell = -2$ ms = +1/2

B- n = 4 $\ell = 2$ $m\ell = +2$ ms = +1/2

C- n = 4 $\ell = 2$ $m\ell = -2$ ms = +1/2

12- If the values of the quantum setting are four for the electron the last atom element is n = 3 $\ell = +1$ $m\ell = 0$ ms=+1/2 The atomic number of the element is:

A- 12

B- 13

C- 14

13- If the values of the four quantum numbers of the penultimate electron in element atom is

n = 2 $\ell = 1$ $m\ell = 0$ ms = -1/2

The atomic number of the element is: -

A-9

B- 10

C- 11

14- The electronic configuration of an atom of an element was

$$1s^2$$
 $2s^2$ $2p^6$ $3s^2$ $3p^3$

The four quantum numbers of electrons are insecondary level 3p³ will only differ

A- the secondary quantum number.

B- the magnetic quantum number.

C- the electron spin quantum number.

15- The four quantum numbers of electrons are in secondary level 4p² will only differ

A- the secondary quantum number.

B- the magnetic quantum number.

C- the electron spin quantum number.

16- The four quantum numbers of electrons are in secondary level 5p⁶ will only differ

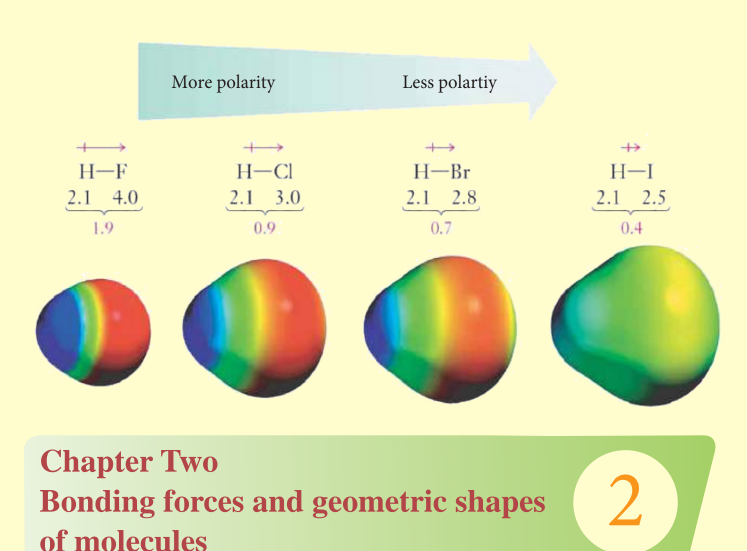
A- the secondary and electron spin quantum number.

B- the secondary and magnetic quantum number.

C- the magnetic and electron spin quantum number.

- 17. In his theory, Bohr assumed that electrons orbit at :
- A- fixed size and fixed energy orbitals.
- B- fixed size and variable energy orbitals.
- C- variable size and fixed energy orbitals.
- 18. In an electrostatic discharge tube postive particles that slides through the hole into an area behind the cathode are called
- A- cathode rays.
- B- channel ray.
- C- x ray.
- 19. If we assume that the atom includes only protons in its nucleus, it means:
- A- There is no difference in the atomic number of all atoms elements.
- B- There is no difference in the number of mass of all atoms elements.
- C- There is no difference in the mass number of all atoms one element.
- 20. Planck assumed when he was studying a phenomenon of light emission of hot objects:
- A- Hot objects release electromagnetic energy in the form of waves.
- B- Hot objects release electromagnetic energy as specific small amounts.
- C- Hot objects release electromagnetic energy as continuous small amounts.
- 21. The atom is excited:
- A- when the energy of the atom is at a stable level.
- B- when the potential energy level atom of becomes higher than the stable level.
- C- when energy is lost in the form of electromagnetic radiation (Photon).

- 22. Quantum number which indicates the direction of the orbital around the nucleus is:
- A- Secondary quantum number.
- B- Magnetic quantum number.
- C- Electron spin quantum number.
- 23. The principle quantum number is always equal for number:
- A-Secondary levels.
- B- Orbitals.
- C- Electrons.
- 24. The electron orbits in a fixed orbit with fixed diameter and energy Specific and specific energy (according Bhor assumption) and as a result of its rotation:
- A- Emit energy.
- B- Absorb energy.
- C- Does not emit energy.



After completing this chapter, the student is expected to:

- Understand why atoms combine together and the meaning of chemical bonds and their types.
- Understand the factors that determine the type of bonding and know how the atoms combine among them.
- Determine the properties of ionic compounds and distinguishes between their properties and those of covalent compounds
- Realize the absence of molecules in ionic compounds.
- Understand the concept of hybrid orbitals and their types.
- Recognize the principle of resonance and its effect on the effectiveness of some covalent compounds.
- Draw the geometric patterns of some molecules.
- Differentiate the bonds of sigma and the bonds of π .

Do you know

The number of elements already in nature is 94 elements. As for the elements prepared industrially are 26 elements due to chemists' efforts in world the number of elements is yet 120 elements. But the resulting compounds within the various branches of chemistry are in millions. These compounds are increasing with the days and the main and basic reason of that is nature of the atoms that are bonded together to form independent molecules or crystalline structures. Chemical bonds are the ladder, which if the chemist guided him in his laboratory to create new materials as we see it in our daily lives.

H	C	0
9	430	3
H ₂ O	CO_2	C_2H_6O

Hydrogen	Carbon	Oxygen	
atom (H)	atom (C)	atom (O)	
Water	Carbon	Ethyl	
molecule	dioxide	alcohol	
(H ₂ O)	molecule	molecule	
	(CO_2)	(C_2H_6O)	

Figure 2-1
Chemical bonding

2-1 Introduction

That all the atoms of the elements except those of noble gases possess chemical activity is uneven under normal conditions and can enter into chemical reactions to saturate the last energy level through loss or acquire or share electrons to access the electronic order to the nearest noble (inert) gas.

Chemical bonding is a bond atoms or ions together to form covalent molecules or ionic or other compounds are mainly used to understand and follow the chemical reaction process, which leads to the creation of new chemical compounds. This is increasing as days progress as a goal to improve the lives of human society [Figure 2-1]. For that and to the recall of the concepts that took place turning to it in advance in the fourth grade of science, let's review the most important continuing with our progress in the study of chemistry. Here we have to go through our imagination on some logical scientific questions we need to answer, in order to complete the general idea of how these bonds occur:

- * How do? and why? Atoms are bound together to form multiple molecules atoms were simple or complex?
- * Why different chemicals possess different physical properties from each other such as color, melting point, boiling point, acid and thermal conduction, electrical conductivity and solubility in different liquidsetc?
- * Why chemicals have different chemical qualities in terms of resistance to the effect of acids, bases or various chemical reagents and affected by temperature?

Many other questions need appropriate scientific answers to them to explain causes, and thus extrapolate other cases where extrapolation is an investigation to the desire of man and his curiosity to uncover the secrets of scientific knowledge. For the purpose of remind students by some of the concepts discussed in the previous year we will review some of them in the following items:

2-1-1 Chemical reaction

The chemical reaction between two elements is an interaction between atoms of two elements to form a new compound molecule has chemical and physical properties completely different from the characteristics of the elements involved in the reaction. Its purpose is to form orbitals saturated with electrons to achieve chemical stability. There are many scientific facts and notes must be in the thinking of students, when they are studying and learning of chemical reaction project as follows:

- 1. The nuclei of elements atoms do not enter as a part in the chemical reaction, the same is true for internal electronic orbital and their saturated and stable orbitals.
- 2. The chemical reaction is limited to the sharing of valence electrons located in the outer and unsaturated orbitals.
- 3. The atoms of the elements involved in the composition of the compound are strongly associated with abinding on the elements involved. Stability or weakness of bonds in the resulting compound depends on the nature of the bonding force between the atoms; the most important is the electronegativity of the involved atoms.
- 4. When fragmenting the compound into its initial elements, ie separating the two elements from each other, various chemical or physical methods are used to return them to two elements, that require the dismantling of the bonding force (ie, bond breaking). Bond breaking needs energy equal to that of emitted when it is configured.
- 5. When the atoms of the elements combine, the electronic configuration is done for unsaturated external atomic orbitals up to more stable state, where these atoms then have saturated molecular orbitals.

2.1.2 Lewis symbol and octet Rule

You knew from your previous study that the secondary third class is that electrons in the outer shell of the element's atom is configured in the Louis configuration in a mock manner so that the symbol of the chemical element is written surrounded by dots each point is representing one electron and every two adjacent points are an electronic pair. These points are distributed to the four sides surrounding the symbol so that do not exceed two points on each side.

Example 2-1

Write the Lewis symbol for two magnesium atoms ($_{12}$ Mg) and chlorine ($_{17}$ Cl)

Solution

Electronic configuration of magnesium atom:

 $_{12}$ Mg: $1s^2$ $2s^2$ $2p^6$ $3s^2$

So Louis' symbol for the magnesium atom is:

Electronic configuration of chlorine atom:

 $_{17}\text{Cl}: 1\text{s}^2 \quad 2\text{s}^2 \quad 2\text{p}^6 \quad 3\text{s}^2 \quad 3\text{p}^5$

So Louis' symbol for the chlorine atom is: CI

Exercise 1-2

Write Lewis symbol for atoms: 16S, 11Na, 10Ne

These electrons determine the nature of bonds between atoms and also determine chemical formulas for compounds resulting from the union of atoms with them. When atoms form covalent bonds, they share a number of electrons to be arranged similar to the noble gas closest to them which has eight electrons in its outer orbital (except helium who has only two electrons). This can be illustrated by writing Lewis to a number of molecules as follows:

Water molecule (H_2O) the electronic configuration of an atom (H) is $_1H$: $1s^1$

Lewis symbol of atom (H) is H.

And atom(O) is $1s^2 2s^2 2p^4$

Lewis symbol of atom (O) is .O:

Water molecule is H:O:

Exercise 2-2

Which one of these molecules apply octat rule to Its central atom the CH_4 or BeF_2 . Note that the atomic number of F = 9, Be = 4, H = 1, and C = 6

Ammonia molecule (NH₃) The electronic configuration of an atom (N) is N: $1s^2 2s^2 2p^3$ and the hydrogen atom as mentioned above.

Lewis symbol for the ammonia molecule is H:N:

You may notice that each of the atoms O and N in these compounds, it is surrounded by eight electrons and this rule applies to atoms in many compounds. This rule is called the (octat rule), where the central atom in the molecule is surrounded by eight electrons. This rule is not applicable for all the atoms in the molecules as in the phosphorus pentachloride PCl₅ and Boron trifluoride molecule BF₃



It notes that the central phosphorus atom was surrounded by ten electrons, as for the boron atom is surrounded by six electrons, so it does not agree with rule of eight. Authorized rule of eight is not general in all cases as there are many atoms that do not agree with this rule.

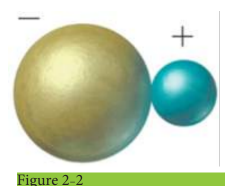
2-2 Types of chemical bonds

Chemical bonds are the phenomenon of the presence of atoms coherent together in molecule or crystal. The material atoms are bonded together by chemical bonds, the type and strength of the chemical bond depends on the electronic configuration of the constituent atoms that form bond. There are several types of chemical bonds can form between the atoms of different elements, which are:

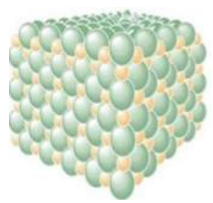
2-2-1 Ionic bond

Ionic bonds emerge between a metallic element and a non-metallic element through loss and gain electrons. It results from the interaction of two atoms, one of which have High electronegativity (such as atoms of halogen group elements) and others Have low electronegativity (e.g., alkaline earth metals group and metals group elements). In this case, the valence electron will move completely from the atom of the low electronegative element into the atom of the high electronegative element, and then we have two ions, the first is positive charge ions as a result of the loss of an valence electron, the second is negative ion as a result of the reception of this electron[Figure 2-2]. These different charged ions are then bound as a result of the electrostatic attraction to form the complex and neutral crystal lattice [Figure 2-3] like; sodium chloride NaCl, potassium chloride, KCl, magnesium chloride MgCl, and potassium fluoride KF and calcium chloride CaCl₂, as well as all hydrides of alkaline metals group elements and alkaline earth metals group elements, such as sodium hydride NaH. The most important properties of ionic compounds are:

- 1. It exists in form of a crystalline structure, a regular geometric arrangement of negative and positive ions.
- 2. It has very high melting and boiling point, to overcome the attraction forces between the negative and positive ions and to break the crystalline structure.
- 3. Inability of electrically conductive in solid state due to ions connection and their inability to move within the crystalline structure while becomes electrically conductive when melted or dissolved in water (then the ions will be free to move in the melted and aqueous solution).
- 4. Soluble in polar solvents such as water and do not dissolve in non-polar organic solvents such as gasoline or ether.



Ionic bond, of NaCl



1	Sodium ion
	(Na⁺)
1	Chloride ion
(march 1)	(Cl⁻)

Figure 3-2
Crystalline structure of sodium chloride.

2-2-2 Covalent bond

Covalent bond is formed when the valence electron is difficult to transmit a complete transition from one atom to another, in this case the pair is formed, from electronic contribution or participation of both atoms. Charges are not shown on atoms, covalent bonds often occur between nonmetals. The difference in the electronegativity value (see electronegativity values for some elements in Table 2.1) plays an important role in the formation of covalent bonds. It can lead to two types of covalent bonds:

A- Pure covalent bond:

This bonds that emerge between two atoms of a non-metallic element where they are similar in electronegativity, or between two elements atoms similar in electronegativity, where the difference in electronegativity is zero for the two case. The pair of electrons will spend equal time in the acquisition of both atoms [Figure 2-4].

An example of this type of covalent bond is the bond in the nitrogen molecule N_2 , the chlorine molecule Cl_2 and the oxygen molecule O_2 and in the fluorine molecule F_2 .

B- Polar covalent bond:

It is a family that arises between the atoms of two elements similar in electronegativity, the difference shall be greater than zero and less than 1.7 to participation with one or more electrons [Figure (2-5)]. Examples of this type of the covalent bonds are the bonds in the water molecules H_2O and ammonia NH_3 , hydrogen chloride HCl, hydrogen fluoride HF and aluminum chloride $AlCl_3$ and hydrogen bromide HBr. In this case carry one of the atoms is a negative partial charge (negative delta δ -) and the second atom positive partial charge (positive delta δ -).

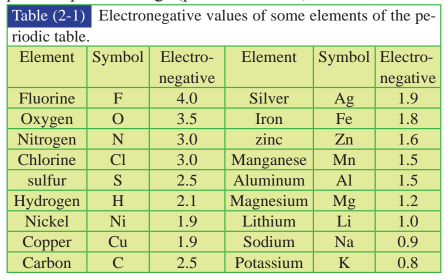




Figure 2-4
Pure covalent bond.

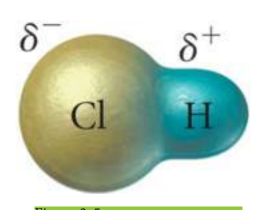


Figure 2-5
Polar covalent bond.

There are many types of covalent bonds that differ in number of electronic couplings bonding between atoms. That types of common covalent bond is a single bond, which share only one electronic pair, such as molecule F_2 , is involved, When participating in two electronic pairs, they are called double covalent bonds. In the case of participation in three electronic pairs makes it a triple covalent bond. An example of double bond is what we find in the oxygen molecule O_2 . An example of triple bond is what we find in the nitrogen molecule O_2 . Covalent bonds compounds are characterized by the following:

- 1. Low melting and boiling point, so do not need high heat energy, because the forces of attraction between their molecules are weak.
- 2. Do not conducted of the electric current because they are not form negative or positive ions in their melts or aqueous solutions.
- 3. Do not dissolve mostly in polar solvents as water while dissolve in organic solvents such as ether and benzene.

2-2-3 Coordinate bond

Coordinate bond is formed when one of the atoms provides a pair of electrons to another atom have the ability to receive this electronic pair to form bond. Then this pair will be shared between two atoms. The electron donor atom is the Lewis base and contains a pair of free electron such as an oxygen atom in a water molecule or a nitrogen atom in the ammonia molecule. The receiving atom is often a transitional metals (Lewis acid) because they have empty orbitals of the type d such as nickel or hydrogen atom ion.

We can say that coordinate bond Is a special type of covalent bond, except that the source of the electron pair is only from one atom, and the coordinate bond is longer and weaker than covalent bond.

Example 2-2

Graphically illustrate the emergence of the coordinate bond in the ammonium ion NH₄⁺

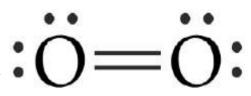
Solution

This ion consists of the binding of ammonia NH₃ to the hydrogen ion H⁺ in aqueous solution:

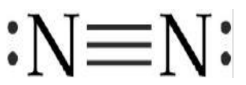
$$H: \overset{H}{\overset{}_{\stackrel{}{\overset{}}{\overset{}}}} \longrightarrow \begin{bmatrix} H: \overset{H}{\overset{}{\overset{}}{\overset{}}} \end{bmatrix} \xrightarrow{\stackrel{}{\overset{}{\overset{}}{\overset{}}}} \begin{bmatrix} H-\overset{H}{\overset{}} \rightarrow H \end{bmatrix}$$



Single covalent bond.



Double covalent bond.



Triple covalent bond.

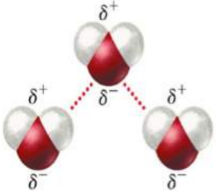
Exercise 2-3

Use the Lewis symbol for atoms F. O and N to illustrate the formation of covalent bonds single, double and triple.

Exercise 2-4

Represent the two ions H_3O^+ (BF $_4$ using Lewis symbol and interpreted the formation of coordinate bond between them.

H H H



Hydrogen bond in water molecule.

Note from the previous example that the nitrogen atom has a pair of electrons not involved in the NH₃ molecule can be involved, The hydrogen ion has an empty orbital that can receive this pair. When the ammonia molecule is close enough to hydrogen, it occurs, this pair will be shared between atoms as in covalent bond exactly and form NH₄⁺ ion. This type of bond is called coordinate bond and is referred to in the order of the Lewis symbol with an small arrow rather than a line that represents a covalent bond.

2-2-4 Metallic bond

A chemical bond that happens between the atoms of an element of metals, this bond is due to the metal atoms possessing electrons in their outer shells contribute to the formation of a crystal and these atoms has free electrons movement in this crystal. When metal atoms are binding together, they do not reach the electronic configuration of noble gases. Atoms of metals such as sodium and potassium, they are easy to lose their equivalent electrons, and become positive ions because their electrolysis is low. The strength of the metal bonds is affected by several factors, the most important of which is the charge density is equal to the ion charge / ion size (proportional to the number of orbits), where the ion charge is the charge that the metal gains after losing electrons in the last orbit (+1,+2,+3).

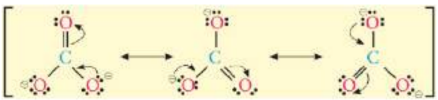
Therefore, the force of the metallic bond depend on the number of the valence electrons beam in the metal atoms, the more electrons of the valence beam, the greater the coherence of the metal to be more solid and higher in boiling point. Higher charge density on the ion increased bond strength and as a result higher melting point obtained. Many of the properties of natural metals are due to the nature of this bond, electrical conductivity and thermal conductivity of metals caused by the movement of free electrons between atoms.

2-2-5 Hydrogen bonding

Hydrogen bonds arise due to the attraction that occurs between the positive terminal (hydrogen atom) and the negative terminal atom has electronic pair or more. These three properties are limited to three elements only are; oxygen, fluorine and nitrogen atoms. Therefore we find water, ammonia, hydrogen fluoride molecules and others are agglomerated by effect of hydrogen bonds. The hydrogen bond is the cause of the high boiling point of water, the melting point of ice and the expansion of the volume of frozen water that led to ice floats on the water. Hydrogen bond is a weak physical bonding force between molecules and not a precise chemical bond: therefore its strength is much lower than other.

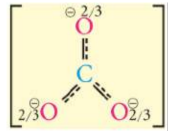
2-3 Resonance

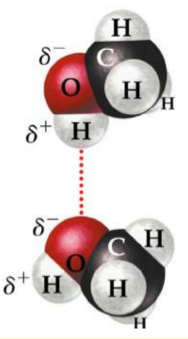
Some types of bonds can have more than a dot shape such as ozone (composition I, II) in which the central atom has a single bond with one atom and a double bond with another. The dots shape cannot tell us which of the atoms has a double bond. Both atoms have the same chance of a double bond. These two structures are likely to be called resonance structure or resonance. In fact, the structure of ozone is a hybrid resonance structure between its resonance structure (composition III). Rather than having a double bond and a single bond, in fact, the two bonds are equal, which is a middle state between the single bond and double bond, where three electrons are at each all time. Carbonate ion (CO₃-2) can also be drawn by any of the following formulas:



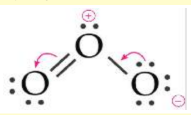
It is clear from the structures above that the central atom (C atom) is connected with the other atoms (O atoms) with two mono bonds and one double bond alternating their position on the three atoms and these potential structures are called resonance. In fact, the ion structure (CO_3^{-2}) is a hybrid resonance structure between the above three struc-

tures as following

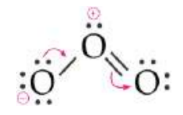




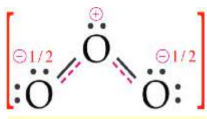
Hydrogen bond.



Composition I



Composition II



Composition III

Hybrid resonance structure between the two structures I.II

Resonance of ozone

Exercise 2-5

Draw the resonant shapes of the sulfate $ion(SO_4^{-2})$ and phosphate $ion(PO_4^{-3})$.

2-4 Geometric shape of the molecules

The molecules of chemical compounds take certain geometric forms that control in its composition a number of factors, the same that control the type of chemical bond (whether ionic, covalent or metallic), which are:

- 1. Number and type of atoms associated in the molecule.
- 2. Electronic configuration of element atoms involved in the formation of the molecule.
- 3. The ability of atoms to acquire, lose or share valence electrons.
- 4. The presence or absence of an empty secondary outer shell in the atom.

The atom reaches a more stable state and a minimum energy state so that the repulsion between its electrons in while atoms are attracting as much as possible. Theories explain the formation of the bonds and the forms of molecules and recall the most important now, these theories:

2-4-1 Valence Shell Electron-pair Repulsion Theory (VSEPR) This theory explains the configuration of atoms around a central atom based on the repulsion between pairs of electrons involved or not involved in the valence shell of the central atom. The repulsion between these pairs in minimum, when they are as far as possible from each other so as to achieve this greater gravitational attraction of the atoms makes them more stable and less energy. To illustrate, the geometry of some molecules can be studied:

BeF, molecule:

When Lewis symbol is written for both the ₄Be beryllium atom and the ₉F fluorine atom as follows: Be F:

The Lewis symbol shows that the beryllium atom shares the two fluorine atom by pair of electrons atoms to have a two pairs of electrons shared around beryllium atom. In order for the electrons to be in the lowest state of repulsion, they are distributed on both sides of the beryllium atom and the fluorine atoms according to the following:

So you notice that the three atoms are arranged on a straight line so that beryllium falls in the middle and the spatial shape of the molecule is linear and the angle between the two bonds is 180° [Figure 2-6].

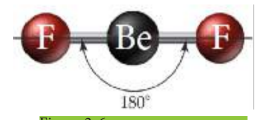


Figure 2-6
Linear form of beryllium fluoride molecule.

BCl₃ molecule:

The Lewis symbol for the boron chloride molecule BCl₃ is written as follows:

It is clear from this symbol that there are three pairs of electrons surrounded borons atom. In order to be less repulsive between pairs of electrons as possible, the chlorine atoms are distributed around the boron atom in the form of a triangle planar. The angles between its bonds are 120° (Figure 2-7).



The Lewis symbol for CCl₄ is as follows: Cl:C:Cl

It notes that there are four sets of electron pairs surrounding carbon atom, the repulsion between them at its lowest when chlorine atoms are distributed towards the heads of a regular tetrahedral so that the angle between each pair is 109.5 ° (Figure 2-8).

H₂O molecule O:H

It is clear from Lewis symbol of the water molecule that there are four pairs of electrons surround the oxygen atom, further note the composition of the oxygen atom in water molecule is similar to the cpmposition of a carbon atom in CCl₄ molecule. The pairs of electrons are distributed toward a angular shape. The presence of non-participating electronic pairs pushes two hydrogen atoms because they are spread over two vertical sides in a curved shape at an angle of magnitude (Approx.104.5°) [Figure 2-9].

NH₃ molecule H:N:H

It is clear from the Lewis symbol that there are four sets of electron pairs surrounded nitrogen atom and it is expected to be distributed as in the case of the CCl₄ molecule toward the form a trigonal pyramidal, but the presence of a non-participating electron pair push the three hydrogen atoms to be distributed on the heads of the three-base pyramid. The angle between the three bonds is about (107.3 °) [Figure 10-2].

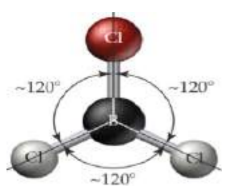


Figure 2-7
Triangle planar.

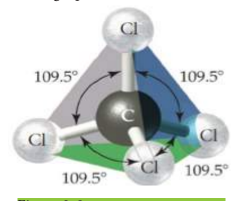
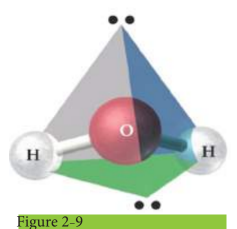
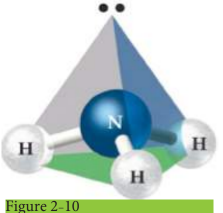


Figure 2-8 Regular tetrahedral.



Geometric shape of water angular.



Geometric shape of ammonia trigonal pyramidal.

2-4-2 Valence bond theory

The theory of repulsion of electrons pairs of valence shell was able to explain different shapes of molecules, but they could not explain how the electrons were distributed between the valence shells of the two constituent atoms of bond. One of the theories that contributed to this clarification theory of the valence bond, which was adopted in its interpretation of the formation of the covalent bond on the interference of atomic orbitals of the valence of the atoms where the electrons move around the nuclei and increase the electronic density in the area of interference between the two atoms. This leads to the two nuclei approaching each other, reducing their energy and increasing their attraction towards the overlap area of the orbitals, thus forming a covalent bond between them. This can be illustrated in the formation of covalent bonds of some simple molecules such as H₂, HF,Cl₂, O₂, N₂. Atomic orbitals overlap as follows:

A- Orbital overlap S of two hydrogen atoms in the hydrogen molecule

When writing the electronic configuration of the hydrogen atom, we notice that it has one electron at the secondary level (1s): $_1H:1s^1$

Bond forms in the hydrogen molecule consists of the overlap of the orbital 1s in the first atom with the orbital 1s of the second atom as shown in Figure 2-11:

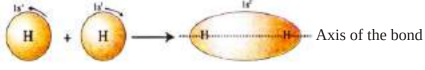


Figure 2–11

Vertical overlap of Orbitals S.

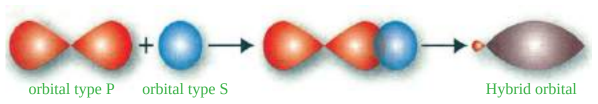
It is noted from the figure that the electrons of the bond are concentrated between the nuclei of the two atoms, where the electronic density increases in the area of interference around the axis connecting the two nuclei. This bond is called sigma (σ)

B- Orbital overlap s with orbital p from other atom as in HF molecule

When writing the electronic configuration of hydrogen atom, we note that it has one electron at the secondary level 1s $H: 1s^1$

When writing the electronic arrangement of the fluorescence atom is as follows: ${}_{9}F:1s^2 2s^2 2p^5$

It notice that there is a single electron in orbital in the fluorine atom, so a sigma bond is created, when the orbital p from the hydrogen atom overlaps with the orbital p (located on the vertical axis) where the electronic density is distributed around the axis connecting the two nuclei. This bond is also named sigma bond [Figure 2-12].



C- Orbital overlap p of the first atom with orbital p of the second atom

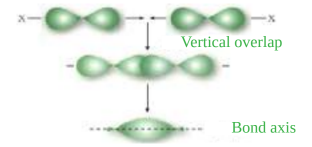
Orbitals p overlap in two ways as follows:

1- Orbital p overlaps with orbital p on the same axis

When orbital type p of atom is vertical overlap with orbital type p of another atom (at the same axis), then electronic density is distributed between two nuclei of the two atoms symmetrically along the axis connecting them. This bond is called sigma bond as in Cl, molecule.

$$_{17}CI$$
 : $1s^2$ $2s^2$ $2p^6$ $3s^2$ $3p^5$ 1 1 1 1 $_{3p^5}$

Atomic orbital approaches 3p, which contains one electron in both of the two atoms and it overlap between them and arises cloud of electronic density. It represents the sigma bond between the chlorine atoms [Figure 2-13].



2. Orbital p overlap with orbital p on the side axis

When an orbital p of an atom overlaps with an orbital p of another atom, the lateral interference occurs vertically, the electronic cloud will be distributed over the bond axis that bonds of the two atoms. These overlap will form a covalent bond type pi (π) , as oxygen molecule.

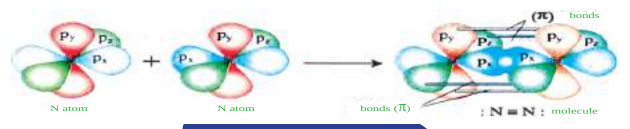
Figure 2–12

Two orbitals p.s vertical overlap.

Figure 2-13

Two orbitals p vertical overlap.

When lateral interference occurs horizontally, the electronic cloud distributes on the right and left axis of the bond axis that connected the two atoms, that gives bond type pi. Three cases can be found from overlap (vertical, lateral vertical, lateral horizontal) as shown between the two atoms of nitrogen molecule in [Figure 2-14].

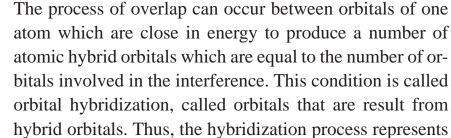


2-5 Orbital hybridization

and its products:

Figure 2-14

Formation of π bond and sigma bond in nitrogen molecule.



interference atomic orbitals have to produce hybrid orbitals similar in shape and size and equal in energy and more stable and lead to get stronger covalent bonds between atoms involved in molecule formation. Regarding following are some observations regarding the hybridization process

- 1. The hybridization process occurs in the same atom orbital after excitation the electrons that fall within the main outer plane and their task reducing the repulsion between electrons of resulted molecules.
- 2. Hybridization is done between atomic orbitals converging in energy together like (2s,2p), (3s,3p), (4s,4p).
- 3. The holographic geometry of hybrid atomic orbital differs from Stereoscopic geometry of atomic orbital before hybridization. The hybrid atomic orbital form consists of two lobes, one of which is large; where the cloud is relatively concentrated and the other is small and often neglects while drawing [Figure 2-15].
- 4. The name of the hybrid orbital is derived from the names and number of pure orbitals, as example the hybrid orbital (sp³) which is involved in the hybridization process. It means the participation of three atomic orbitals of type (p) with one atomic orbital type within the same principal electronic shell.

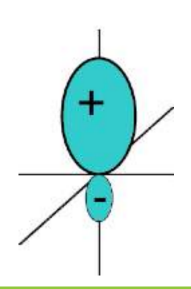


Figure 2-15
Hybrid orbital sp.

- 5. The number of hybrid atomic orbitals formed is equal to the number atomic orbitals involved in the hybridization process. Based on this is when the hybridization of three atomic orbitals of the type (p) with atomic orbital one of type (s) to form four orbitals of type (sp³).
- 6. The energy of atomic hybrid orbitals formed is equivalent or equal.
- 7. The ability of hybrid orbitals to interfere with atomic orbitals of another atom is greater than the susceptibility of not hybridized atomic orbitals that participate in the hybridization process, because atomic orbitals hybrids orbitals are more extensible in vacuum than the participated atomic orbitals not hybridized.

2-5-1 Types of hybrid orbitals

It can get types of hybrid atomic orbitals according to number and type of atomic orbitals involved. The most important will be addressed at the current school level.

A-Hybrid orbitals type (sp): According to the above observations, it know that this type of hybrid atomic orbital may be as a result of the participation of two atomic orbitals one of type (s) and the other of type (p).

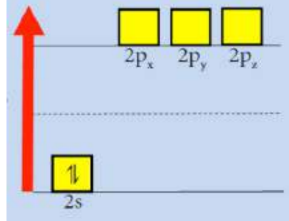
The overlap of these two atomic orbitals will lead to formation two hybrid orbitals type sp located on a straight line and at an angle of °180 in order to have the least repulsion between the two orbitals like the beryllium hydride molecule BeH₂. For the purpose of clarifying how that is happen, the following steps are followed:

1. The electronic configuration of beryllium atom:

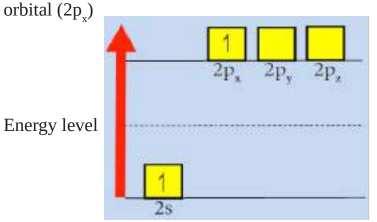
__Be 1s^2 2s^2

The orbitals of the casings in the stable state are expressed

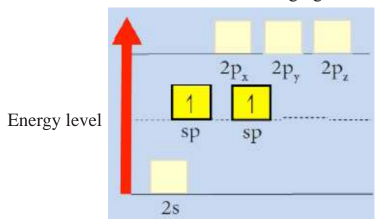
as follows:



2. One of the electrons will rise from the orbital 2s to the



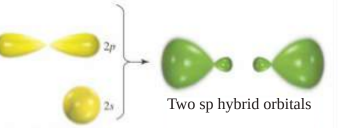
3. Then the interference occurs to generate two hybrid orbitals type sp in equivalent energy less than the energy atomic orbitals $2p_x$, $2p_y$, $2p_z$ and higher than the energy of atomic orbital $2s^2$ and on the following figure.

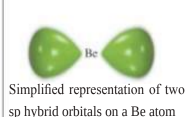


4. These two orbitals have the ability to form molecular orbitals with two hydrogen atoms orbitals (1s¹) to form two covalent bonds between Be and H and form the beryllium hydride molecule. The four steps above can be illustrated by the representation of stereotypes geometric shapes of atomic orbitals before and after hybridization. In this case we have two hydrogen atoms each one has an orbital (1s¹), They interfere with the two hybrid orbitals sp of beryllium atom. This overlap is done on one of the coordinates to give the linear shape of the resulting molecule as shown in Figure (2-16).

Figure 2-16

Composition of hybrid orbitals type sp from atomic orbital interference type s with two orbitals type p_x , p_y of the s-type.



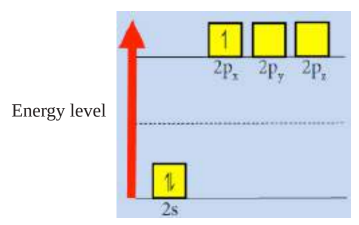


B- Hybrid orbitals of type (sp²): This type of atomic orbitals hybridized by the participation of one atomic orbital type (s) with two atomic orbitals type (p) within the same main orbital. The overlap of these three atomic orbitals will lead to formation three hybrid atomic orbitals type sp² fall at the same level between them is an angle centered at the center of the central atom of 120° . This will be led to less repulsion between the orbitals as it occurs in a trifluoride boron molecule BF₃. Below is an explanation of how this hybridization was achieved the following steps:

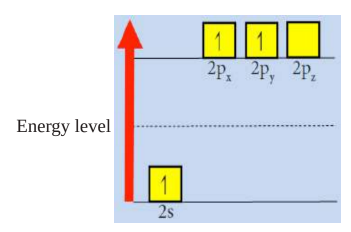
1. The electronic configuration of boron is:

$$_{5}B 1s^{2} 2s^{2} 2p^{1}$$

Orbital (1s²) is considered internally and will not participate in hybridization, so the representation of the electronic configuration of the outer shell of this atom is represented as follows:

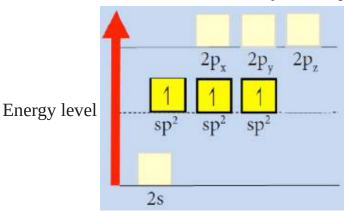


2. One of the two electrons in the orbital $(2s^2)$ goes up to the orbital p_v with the approximate energy:



3. Then the interference occurs to generate three hybrid atomic orbitals of type sp² has less equivalent energy than non-hybrid atomic orbitals

 $2p_x$, $2p_y$, $2p_z$ and higher than non-hybrid atomic orbital $2s^2$. These orbitals are distributed around the nucleus in a plane that shares two coordinates, and they have equal angles.



4. These three orbitals have the ability to form three molecular orbitals with three orbitals for three fluoride atoms to form a triple boron fluoride molecule.

The four steps above can be illustrated by the representation of stereotypes geometric shapes of atomic orbitals before and after hybridization. In this case we have three fluoride atoms interfering with sp² hybrid orbitals of boron atom. This overlap is done at one level as shown in Figure (2-17).

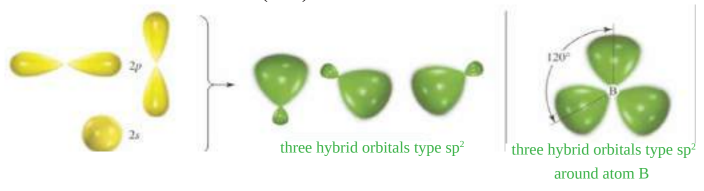


Figure 2-17

Composition of hybrid orbitals type sp^2 of atomic orbital overlap of type 2s with two orbitals p_x and p_y .

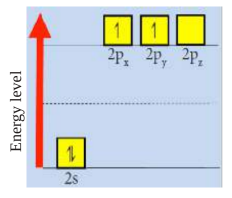
C- Hybrid orbitals type (sp³): This type of atomic orbitals hybridized by one atomic orbital interference of type (s) with three atomic orbitals of type (p).

The overlap of these the four atomic orbitals will produce four hybrid atomic orbitals type (sp³), these orbitals form a regular tetrahedral shape around the nucleus of the central atom with vertical angles of 109.5°,

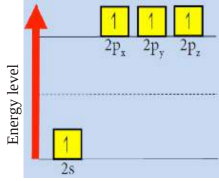
In order to have less repulsion between the orbitals as in the methane molecule CH₄. Below is an explanation of how the hybridization takes place according to the following steps:

1. The electronic configuration of carbon atom is: ${}_{6}C \ 1s^{2} \ 2s^{2} \ 2p^{2}$

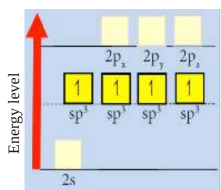
Orbital (1s²) is considered internally and will not participate in hybridization. So the representation of the electronic configuration of this atom is as follows:



2. One electron from orbital $(2s^2)$ rises to orbital $(2p_2)$



3. The interference is happened to generate four hybrid atomic orbitals from $\rm sp^3$ with equivalent energy less than energy of non-hybrid atomic orbitals $\rm 2p_x$, $\rm 2p_y$, $\rm 2p_z$ and higher atomic orbital energy of 2s. These hybrid orbitals are distributed in a regular tetrahedral shape around the central carbon atom with equal angles.



4. These four orbitals have the ability to form four molecule orbitals with four atomic orbitals for four hydrogen atoms (1s¹) to form a methane molecule. The four steps above can be illustrated by the representation of stereotypes geometric shapes of atomic orbitals before and after hybridization [Figure 2-18]. In this case we have four hydrogen atoms each one have orbital (1s¹) has an overlap with the hybrid orbital sp3 of carbon atom as shown in Figure (2-18).

Exercise 6-2

Explain how hybridization get in molecule for $SiCl_4$ according to the four steps, that we followed in the hybridization process (atomic numbers Cl = 17 and Si = 14).

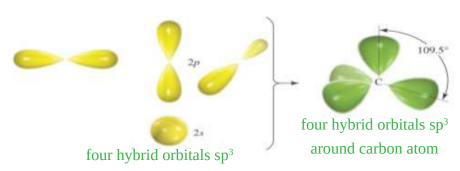


Figure 2-18

Composition of hybrid orbitals sp^3 of orbital overlap from 2s with three orbitals p_x , p_y , p_z .

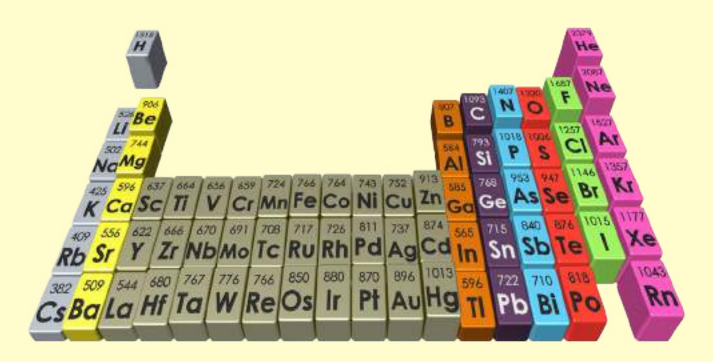
- 2-1 Reasons for each of the following:
- 1. The boiling point of water (100° C) is higher than the boiling point of hydrogen sulfide gas $(-60 ^{\circ} \text{ C})$.
- 2. The NH₄Cl ammonium chloride molecule contains three polar covalent bond, coordinate bond and ionic bonds.
- 3. Ionic compounds do not conduct electrical in solid state but for melting or solutions of ionic compound in water a good ability of electrical conductivity.
- 4. When you put a piece of ice in the water, float however when you put a frozen piece of gasoline in gasoline immerse.
- 2-2 Compare between each of the following:
- 1. Covalent bond and coordinate bond.
- 2. The sigma (σ) bond and pi(π) bond.
- 3. Hybrid orbital and non-hybrid orbital.
- 2-3 Draw a figure showing a raised carbon atom and an hybrid Carbon atom sp³
- 2-4 What is the hybridization situation in each of the following. Mentioning forms of the resulting compounds:
- 1. Oxygen in water.
- 2. Nitrogen in HN = NH
- 2-5 Draw resonance formulas for: CH₃COO⁻, O₃, CO₃⁻²
- 2-6 Why are the atoms of elements combined together? are the combinations of atoms of elements always produce compounds? Discuss your answer scientifically with at least two examples.

2-7

- A- What is chemical bond? Identify it accurately.
- B- Numbering only types of bonds you know.

2-8

- A- What is the ionic bond? What are the conditions of its composition?
- B- Why molecules are not formed in ionic compounds?
- C- What are the most important characteristics of ionic compounds?
- 2-9 What do we mean by hydrogen bonding? Explain this in example, draw hydrogen bonds between Methanol (CH₂OH) molecules.
- 2-10 Covalent bonds may be polarized, when is that?
- 2-11 What are the factors that determine the bonding between two atoms covalent or covalent polarized? or Ionic?
- 2-12 What are the metal bonds? what effect it has on the properties pure metals?
- 2-13 Rewrite the following statements as corrected (may) you find it from scientific mistakes:
- A- All compounds with covalent bonds didn't dissolve in water.
- B- Protons and neutrons are involved in the composition ionic bond.
- C- (π) bond is less energy than (σ) bond of the same molecule himself.



Chapter Three Periodic Table and Chemistry of Transition Elements

3

After completing this chapter, the student is expected to:

- Recognize the historical sequence of the emergence of the periodic table and attempts made to divide the elements.
- Arrange elements in a modern periodic table based on atomic numbers instead of their atomic masses.
- Able to determine the number of periods included in the periodic table and the group numbers.
- Recognize the periodic table parts.
- Understand the periodic properties of elements in the periodic table.
- Understand when an element has an atomic spectrum and that the atomic spectrum is a characteristic for the elements.
- Distinguish between a transitional element and a non-transitional element as recognize the internal transition elements.
- Distinguishe between the elements that are attracted towards the magnetic field and those that are not attracted.

3-1 Modern Periodic Table of Elements

After the discovery of the electron and the appearance of the concept of atomic number by Moseley in 1914, the elements were arranged in ascending order by increasing their atomic numbers rather than by increasing the atomic mass as arranged by Mandeleeff . That is, each element in the modern periodic table exceeds the element preceded by a single electron known as a distinct electron. This arrangement according to the increase in atomic numbers corresponds with the order of elements by increase energy levels from least to most energy. Thus, the image of the modern atomic table, [Figure 3-1] in the form of 7 horizontal periods and includes arranged groups in the form of 18 columns:

1. The first short period comprises hydrogen and helium.





2. The second and third short period, each consisting of 8 elements. They are arranged in the periodic table as follows:

	-3-0	-4-
-	1.4	Be
	11	12
	Na	Mg

5 B	e C	7 N	0	P P	10 Ne
za	14	15	16	17	18
Al	Si	P	5	CI	Ar

3. The fourth long period consists of 18 elements and they are arranged in the periodic table as follows:

Do you know

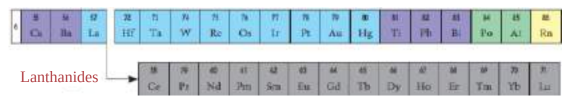
Due to the difficulty of putting lanthanides with lanthanum in the third group, is placed, in a separate row below the periodic table. The same is the case with actinides which is relative to the actinium element, they are also placed in another row below periodic table.



4. The fifth long period consists of 18 elements and they are arranged in the periodic table as follows:

	——Transitional elements ———																	
	E	3	P	R	41	4	6	#	-6	4	9	4	0	-	3	ī	2	N
,	15	St.	Y	Zı	Nb	Mo	Te	Rn	3th	Pi	Āg	Cd	In:	50	Sb	Te	1	Xe

5. The six long period consists of 32 elements and they are arranged in the periodic table as follows:



6. The seventh long period consists of 24 elements and they are arranged in the periodic table as follows:



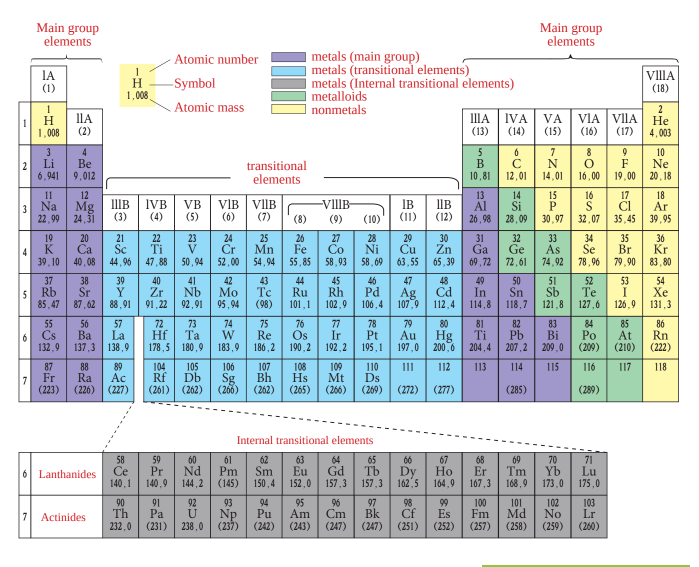


Figure 3-1

Modern periodic table.

The groups in the periodic table and its number (18 groups) divided into Group A (8 groups) and Group B (10 groups) as shown in Figure (3-1). These groups are arranged in vertical columns based on equal number of electrons in the outer shell (external valence electrons). Thus, the elements of a single group are similar in their chemical properties, despite the great difference in their atomic masses so as to equal the number of electrons that atom can lose or gain, or contribute during the entry into the chemical reaction. While in a single period the elements where arranged on the base of increase the atomic number, the elements are convergent in their atomic masses, yet they differ in their physical properties. For example, Nitrogen is near carbon and oxygen in the second period, despite their convergence in the atomic mass, they have different physical properties and it is worth noting that each of the energy level in elements atoms contain secondary energy level (sub), its number equal to the number of principal level. These principal level is filled with electrons as it increases in the energy where fills the lowest energy level(s) and then the higher energy level. This configuration in filling level is similar to the configuration of elements in periodic table by increasing atomic number.

3-1-1 Periodic Table Parts

The periodic table includes four parts arranged as follows [Figure (3-2)]:

First part: includes the first group IA and the second group IIA and both are ends with (ns) level known as alkali metals and alkaline earth metals respectively

Second part: include the elements in the groups:

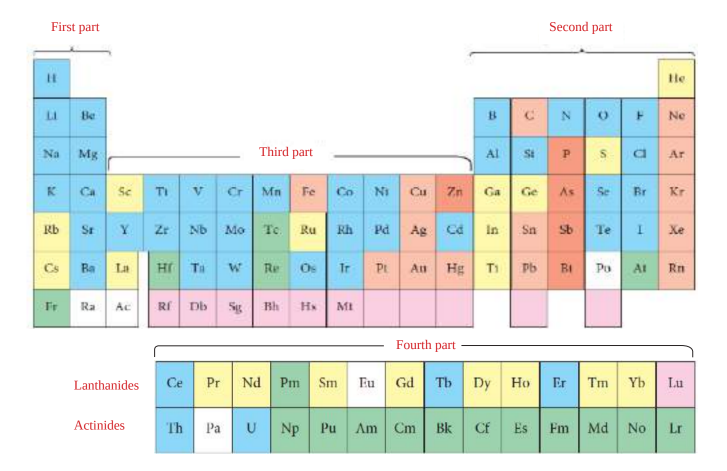
IllA, IVA, VA, VIA, VIIA, VIIIA (the last group is called 0). The elements of this area (Part I and Part II) are full the electrons in the sub-energy levels (s and p). The first and the second part are called the represented elements.

Third part: includes all the elements in subgroups B, namely:

IB, IIB, IIIB, IVB, VB, VIIB, VIIIB, (Group 0) which include (8B, 9B, 10B) and the outer shell of these elements is type s, d and d is not full of electrons is called transitional elements because they are transite by properties

between elements with the level s (group, IA, IIA) and elements with level p group (IIIA, IVA, VA, VIA, VIIA, VIIIA). They mediate the periodic table.

Fourth part: comprises two series of internal transitional elements:



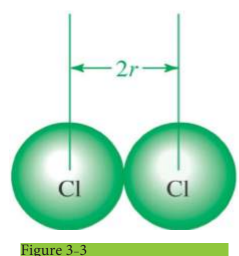
Many of the physical properties of the elements change periodically accordingly to position of these elements in the periodic table in terms of group and period, some of these features are discussed below.

A-Atomic size

Atomic size is a difficult characteristic for several reasons one of the most important reasons is that the probability of electronic distribution is influenced by atoms adjacent in the chemical compound and thus the size of the atom changes somewhat when moving from one case to another. Therefore, when checking any table of atomic radiuses, it should be remembered that tabulated values may be a meaning only when considering a relative comparison of volumes, Figure (3-3) shows

Figure 3-2

Parts of periodic table.



Method of measurement of atomic radii of elements

Method of measurement of atomic radii of elements derived from distances measured from the centers of adjacent atoms in pure elements and on this basis atomic size is defined as half distance between the centers of two identical atoms in a crystal and it is measured using X-rays.

In general, atomic radii are reduced per one period when moving from left to right (ie, atomic number increases) in the periodic table. This behavior can be explained by a table showing the change of atomic radii along the second period.

In the case of the group, the radius increases in one group from top to bottom as the atomic number increases.

The reason for this is the addition of electronic shells with increasing quantum numbers farther from the nucleus as shown in the table of atomic radii[Figure(3-4)].

In the transitional elements

1. The size decreases gradually in each transition series until the fifth element, ie to half of the period then gradually increases until the end of the series, where the size decreases due to the increase of attraction force, as the electron, which is added by increasing the number of atomic number form element to another enters the secondary shell orbitals (d).

2.Observations have shown that adding half of this number, ie 5 electrons, is accompanied by a state of stability and this electronic system block the effect of the nucleus so the attracted force will be less to the electrons that are added, which it explains the increase in size slightly after

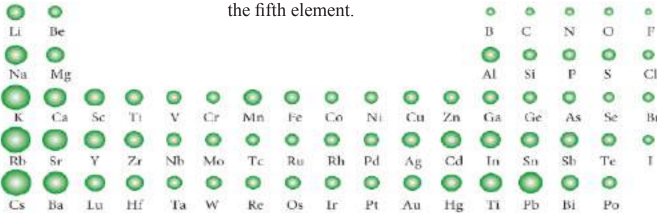


Figure 3-4

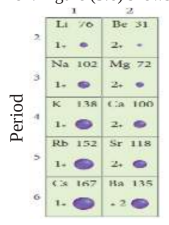
Changing atomic radii (ie. the sizes of atoms) of some elements in the period and group.

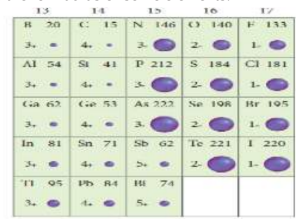
In the internal transitional elements, the size also decreases gradually by increasing the atomic number up to the seventh element (ie to half of the series) and then gradually increasing again and on the same basis of interpretation in point 2

For the radius ions of atoms it is known that atoms can lose or gain one or more electrons to form ions. Electrons have negative charge so atoms gain an extra charge when gain or lose electrons. So the ion is an atom or an atomic group has positive or negative charge.

When an atom loses electrons and is a positive ion is formed, the atom size becomes smaller. This is due to two factors: firstly, the electron that the atom loses often it's an equivalent electron, and the loss of electron led to form an empty external orbit, causing radius decreases. Secondly, electrostatic repulsion decreases between the remaining electrons and increases the attraction between them. The nucleus has a positive charge, allowing the electrons to get closer of the nucleus and [Figure (3.5a)] shows the decrease in the ion radius of sodium atom when it formed a positive ion.

Conversely, when an atom acquires electrons and form negative ions, their size increases because the addition of an electron to the atom generates electrostatic repulsion bigger with the outer-level electrons, and pushes them strongly outward. An increase in the distance between external electrons results in an increase in the amount of radius. [Figure (3.5b)] shows how the radius of the chlorine atom increases, when it forms negative ion. Figure (3.6) shows the ion radius of some elements.

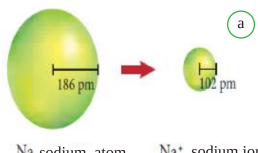


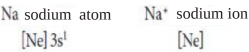


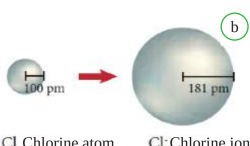
B- Ionization energy

It is the energy needed to extract an electron from a neutral atom in its gaseous state and to form positive ion as in the following equation: $M + \text{Energy} \longrightarrow M^+ + e^-$

This energy is consumed in extracting an electron from external valance electrons. The ionization energy is measured in units of electron volts (ev) and electron volts is a small energy equal 1.6×10^{-19} J. Energy is needed to remove the first electron is called first ionization energy. The result of that ion with one positive charge.







Cl Chlorine atom [Ne] 3s²3p⁵

Cl⁻Chlorine ion [Ne] 3s²3p⁶ or [Ar]

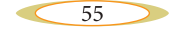
Figure 3-5

- a- positive ions smaller than their neutral atoms.
- b- negative ions are larger than their neutral atoms.

Figure 3-6

Ion radius of ideal elements measured in pico meters.

 $(1pm=10^{-12}m)$



The energy needed to extract a second electron is also called a second energy ionization. Thus, the second ionization energy is always greater than the first ionization energy because the positive nucleus charge attracts the second electrons strongly larger.

If the relationship between ionization energy and the atomic number of elements is plotted, it is noted that change occurs periodically Figure (3-7).

The noble elements are located at the great ends of it in order to stabilize their system. The alkaline elements also located on the lower ends due to the large size of the atomic size and that the quantum layer penultimate contains (8) electrons are characterized by a high degree of stability and work as a barrier obscures the effect of the nucleus charge on the valence electron is easy to extract.

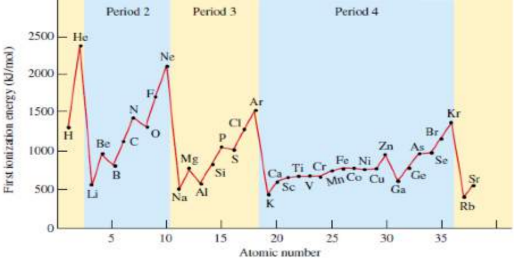


Figure 3-7

The relationship of ionization energy with the atomic number.

It is evident that the ionization energy increases in one period with increasing atomic number because of their small atomic sizes (radii) except atoms whose last shell is saturated or halfsaturated, where ionization energy is greater than ionization energy of the next atom. For example, that The ionization energy of ₇N is greater than the ionization energy of ₈O and the reason for this is due to that the last nitrogen shell is half-saturated in which three electrons and to be more stabile than oxygen despite being the largest atomic number. As well in the case of manganese 25Mn and iron 26Fe the Mn ionization energy is greater than Fe ionization energy for the same previous reason. In one group, ionization energy decreases as atomic number increases due to large atomic size which facilitates extraction the outer electrons of the atom, for example, the ionization energy reduce from lithium to Cesium and from beryllium to radium as shown in Figure (3-8).

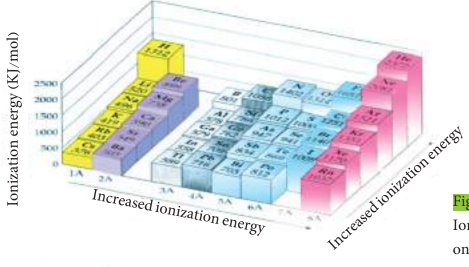


Figure 3–8

Ionization energy changes in one period and group.

C-Electron affinity

An electronic affinity is known as the amount of energy released from an neutral atom in the gaseous state when it acquires an electron forming negative ions, according to the following equation:

$$F_g + e^- \longrightarrow F_g + 328kJ/mol$$

Electronic affinity increases in periods by increasing the atomic number, the small atomic sizes make it easier for the nucleus to attract the electron. Electronic affinity of elements in a single group decreases as atomic number increases because of increasing atomic size. Just as there is first and second ionization energy, so is there first and second electronic attraction whilst some energy is released upon acquisition the first electron often absorbs some energy when second electron is acquired because there is a repulsive force between the negative ion and the acquired electron.

The emission of energy when an electron is added to an atom leads to the transition to lower energy case, ie to a more stable state and this explains the tendency of some atoms to acquire electrons as elements in the VII group in chemical reactions to reach a more stable and minimal state of energy.

The low values in the electronic affinity of noble gases and the elements to the far left of the periodic table are attributable to their inability on the formation of negative ions compared to fluorine (F), chlorine (Cl) and bromine which form easily negative ions as shown in Figure (3-9).

Do you know

The electronic affinity measures the severity of additional electron binding to the atom.

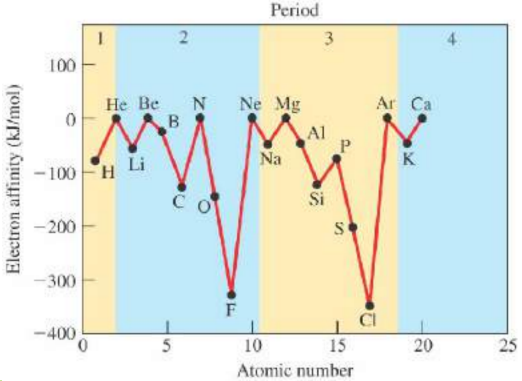


Figure 3-9
The relationship of electronic affinity with the atomic number.

Do you know

And now we ask the benefit of the value of electronegativity the answer to this question that one of these benefits is to predict which bonds are ionic and which covalent. It can also benefit of electronegativity in the polar prediction where two elements apart for electronegativity values whenever necessary, the bonds are more polar and the bond between H and Cl is more polar than those between (Cl and Br).

D-Electronegativity

Is the ability of atoms in molecules to attract electrons toward them of other atoms associated with them by chemical bond. There are values numerical elements as shown in Figure (3-10) these numbers describe the relative ability of an atom to form a bond, ie turns negative state and shared electron can then attract to it. Fluorine element has high electronegativity relative to any element in the periodic table because it located at the end of its period and on the top of its group. The elements of noble gases are not formed chemical bonds and did not agreed on their values yet. In general electronegativity increases in a single period from left to right as the atomic number increases. Existing elements at the far left of the periodic table (group IA, IIA) have low electronegativity.

The elements on the far right except the group (0) has high electronegativity and it is attributed for the elements group (VII) the following electronegativity to:

F=4.0, Cl=3.0, Br=2.8, I=2.5

The order of decrement in the electronegativity shall be orderd, other than the order of affinity in one group, the electronegativity decreases with increasing atomic number as we move from the top to down of the group.

ΙΛ (1)																
H 2,1	ILA (2)			Holove 1.								III.A. (13)	1VA (14)	VA (15)	VIA. (16)	VIIA (17)
Li 1.0	Be 1.5	-	<u> </u>	Above 1	*							B 2,0	C 2,5	N 8.0	3,5	F 4.9
No 9.9	Mg 1.2	111B	IVB (4)	VB (5)	VIB (6)	VIIB (7)	(8)	VIEB (9)	(10)	1B (II)	HB (12)	Al 1,5	5i 1.8	р 2.1	S 2,5	3.0
1C 0.8	Ca t,0	Sc 1,3	Ti t.5	V 1.0	Cr 1,t	Mn 1,5	Fe 1,8	Co 1,9	Ni 1,0	Cu 1,9	Zn 1,0	Ga 1,6	Ge 1.8	As 2.0	5e 2,4	Вт 2,8
R5 0.3	.Sε 1.0	Y 1,2	Zr 1.4	Nb 1.0	Mo 1,5	Tc 1,9	Ru 2,2	Rh 3.2	Pd 2.2	Ag 1,9	Cd 1.7	In 1.7	Sn 1,8	Sb 1.9	Te 2,1	2,5
Cs 0.7	23 ay 0 . 9	La 1,1	Hf	Ta 1,5	W 1,2	Re 1,9	Os 2,2	Ir 2,2	Pt 2,2	Λu 2,4	Hg 1.9	Ti	Pb 1,9	Bi 1.9	Po 2.0	At 2.2
Fc 0.7	FER 9.4	Ac 1.1														

E- Metallic and non-metallic properties:

Metals have a number of properties including metallic luster and electrical and thermal conductivity and high melting and boiling points such as iron, copper and zinc, while nonmetals, are characterized by no luster and gloss. They are often fragile, with low melting and boiling points such as sulfur, carbon, phosphorus and gases. The metalloid are elements their properties between metals and nonmetals such as boron and silicon. Their properties are graded in the periodic table are as follows:

In one period the metallic properties decrease and the nonmetal properties increase as they increase atomic number. We find that the elements of the beginning of the period are all metals and then less this property and a non-metallic property starts to appear as we go to the right of the period, that is, an increase atomic number for example in the second period lithium (₃Li) and beryllium (₄Be) metallic properties, while boron (5B) shows the properties of semi-metals. The rest comes second period elements such as carbon, nitrogen, oxygen and fluorine to appear properties of nonmetals where the atomic number increases. In one group metallic properties increase and Non - metallic properties decrease with increasing atomic number. All elements of groups (IA and IIA) are metals while elements of groups (VII and 0) are nonmetals. The rest of the groups all elements are not in one class, for example, in the fifth group shows (N) metallic properties, while (S) shows the behavior of semimetals and Bismuth comes the last element in the fifth group with metallic properties.

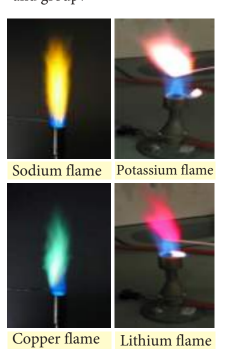
Figure 3-10

Electronegative values for some elements periodic table.

In periods, the elements of the first period (H and He) for metals are either in the next four periods there is a gradual transition from metallic properties to nonmetallic properties of properties. In the sixth period all its elements are metals except the last two elements are non-metallic elements either elements of the seventh period are all metals and show transitional elements and elements Lanthanides and actinides have metallic properties, as shown in Figure (3-11).

	ties, as shown in Figure (5-11).																		
	Reduction of metal properties																		
ties		1A			metals (transitional elements)										VlllA				
roper	1	1 H	llA		metals (Internal transitional elements metalloids nonmetals										lVA	VA	VlA	VllA	He
ents p	2	3 Li	4 Be		transitional									5 B	6 C	7 N	8 O	9 F	10 Ne
eleme	3	11 Na	12 Mg	lllB	lVB	VB	VlB	VllB	nents (8)	-VllB-	(10)	lB	llB	13 Al	14 Si	15 P	16 S	17 Cl	18 Ar
sition	4	19 K	20 Ca	21 Sc	22 Ti	23 V	24 Cr	25 Mn	26 Fe	27 Co	28 Ni	29 Cu	30 Zn	31 Ga	32 Ge	33 As	34 Se	35 Br	36 Kr
c tran	5	37 Rb	38 Sr	39 Y	40 Zr	41 Nb	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 Cd	49 In	50 Sn	51 Sb	52 Te	53 I	54 Xe
netalli	6	55 Cs	56 Ba	57 La	72 Hf	73 Ta	74 W	75 Re	76 Os	77 Ir	78 Pt	79 Au	80 Hg	81 Ti	82 Pb	83 Bi	84 Po	85 At	86 Rn
Increase of metallic transition elements properties	7	87 Fr	88 Ra	89 Ac	104 Rf	105 Db	106 Sg	107 Bh	108 Hs	109 Mt	110 Ds	111	112	113	114	115	116	117	118
reas					Intern	nal tra	nsitio	nal ele	ments										
Inc	6	Lantha	anides	58 Ce	59 Pr	60 Nd	61 Pm	62 Sm	63 Eu	64 Gd	65 Tb	66 Dy	67 Ho	68 Er	69 Tm	70 Yb	71 Lu		
	7	Actin	ides	90 Th	91 Pa	92 U	93 Np	94 Pu	95 Am	% Cm	97 Bk	98 Cf	99 Es	100 Fm	101 Md	102 No	103 Lr		

Figure 3-11 Change of metallic and nonmetallic properties in one period and group.



F-Flame Spectrum:

The spectrum of an element can be obtained by placing the element or its vapor in an electric discharge tube under low pressure and high voltage that irritates its atoms. It emits linear spectra and each element has a distinct linear spectrum, there is a relationship between the linear spectrum and the atomic structure of the element, that's called the atomic spectrum. For example, the hydrogen atom is in a stable state if the electron is found at the first energy level and when increased energy of an electron moves to a higher energy level and is said to be an atom is excited or irritated when an electron falls from an higher energy level to a lower energy level it loses energy equal to the difference between energy levels.

This energy appears as an electromagnetic radiation (Spectrum) accompanied by a color and has a specific wavelength and frequency for each element linear spectrum is characteristic of it when heating a calcium metal on a flame, it flames brick red and strontium is crimson barium is yellowish green, sodium yellow and potassium violet, cesium blue and rubidium dark red.

3-2 Transitional elements

3-2-1 Introduction

Transitional elements appear in the fourth, fifth, sixth and seventh periods of the periodic table and has an electronic configuration, orbitals electrons d and f play an important role.

These elements can be traditionally divided into two groups:

The set of elements d and the set of elements f

The set of d elements consists of three complete series and a fourth series incomplete. Each series of the third series have ten these are:

- 1.The first transition series: starting from the element scandium ($_{21}$ Sc) to zinc element ($_{30}$ Zn).
- 2. The second transition series: starting from the element yttrium ($_{30}$ Y) to cadmium element ($_{48}$ Cd).
- 3.The third transition series: starting from the element lanthanum ($_{57}$ La) to gold element ($_{30}$ Au).

The fourth transition chain starts with actinium element ₈₉Ac and ends with element daramstadtium ₁₁₀Ds, which is composed of (8) elements can be observed these chains of looking at the periodic table Figure (3-11). The F group consists of two chains and in each series 14 elements are called internal transition elements, the first series is called lanthanides and the second series is the actinides, and these two series are:

- 1. The lanthanide series starts from the cerium Ce atomic number 58 to the lutetium Lu atomic number 71.
- 2. The chain of actinides begins with the element thorium Th atomic number 90 to the element lorticium Lr atomic number 103.

In each series, the electronic configuration of the full energy levels remains the same the higher main quantum number is constant while internal energy levels are gradually filled d(n-1) and F(n-2) with increasing atomic number Table (3-1). In the chains of transition elements, orbitals 3d, 4d, and 5d are gradually filled, while the internal transitions are gradually filled with orbitals 4f and 5f. The transition element can thus be defined as the element, which has an electronic distribution in which orbitals d or f is partially full in the case of neutral or chemically unified atoms in their compounds.



Ions of salts of transition elements and their solutions from left to right.

 Mn^{2+} , Fe^{2+} , Co^{2+} , Ni^{2+} , Cu^{2+} , Zn^{2+}

Table 3-1 Electronic configurations of transitional elements						
The	e first	transition	series			
Atomic number	Symbol	Name	Electronic configuration			
21	Sc	Scandium	$[_{18} Ar] 3d^1 4s^2$			
22	Ti	Titanium	$[_{18} Ar] 3d^2 4s^2$			
23	V	Vanadium	$[_{18} Ar] 3d^3 4s^2$			
24	Cr	Chromium	[₁₈ Ar] 3d ⁵ 4s ¹			
25	Mn	Manganese	$[_{18} Ar] 3d^5 4s^2$			
26	Fe	Iron	$[_{18} Ar] 3d^6 4s^2$			
27	Co	Cobalt	$[_{18} Ar] 3d^7 4s^2$			
28	Ni	Nickel	$[_{18} Ar] 3d^8 4s^2$			
29	Cu	Copper	$[_{18} Ar] 3d^{10} 4s^{1}$			
30	Zn	Zinc	$[_{18} \text{Ar}] 3d^{10} 4s^2$			
The	e seco	nd transiti	on series			
39	Y	Yttrium	[₃₆ Kr] 4d ¹ 5s ²			
40	Zr	Zirconium	$[_{36} \text{Kr}] 4d^2 5s^2$			
41	Nb	Niobium	$[_{36} \text{Kr}] 4d^3 5s^2$			
42	Mo	Molybdenum	$[_{36} \text{Kr}] 4d^5 5s^1$			
43	Тс	Technetium	$[_{36} \text{Kr}] 4d^5 5s^2$			
44	Ru	Ruthenium	$[_{36} \text{Kr}] 4d^7 5s^1$			
45	Rh	Rhodium	[₃₆ Kr] 4d ⁸ 5s ¹			
46	Pd	Palladium	[₃₆ Kr] 4d ¹⁰			
47	Ag	Sllvar	[₃₆ Kr] 4d ¹⁰ 5s ¹			
48	Cd	Cedmium	$[_{36} \text{Kr}] 4d^{10} 5s^2$			
The	e third	d transition	n series			
57	La	Lanthanum	[₅₄ Xe]5d¹ 6s²			
72	Hf	Hafnium	$[_{54}$ Xe]4 f^{14} 5 d^2 6 s^2			
73	Ta	Tantaium	$[_{54}$ Xe]4 f^{14} 5 d^3 6 s^2			
74	W	Tungsten	$[_{54}$ Xe]4 f^{14} 5 d^4 6 s^2			
75	Re	Rhenium	$[_{54}$ Xe]4f ¹⁴ 5d ⁵ 6s ²			
76	Os	Osmium	[₅₄ Xe]4f ¹⁴ 5d ⁶ 6s ²			
77	Ir	Iridium	[₅₄ Xe]4f ¹⁴ 5d ⁷ 6s ²			
78	Pt	Platinum	[₅₄ Xe]4f ¹⁴ 5d ⁹ 6s ¹			
79	Au	Gold	[₅₄ Xe]4f ¹⁴ 5d ¹⁰ 6s ¹			
80	Hg	Mercury	[₅₄ Xe]4f ¹⁴ 5d ¹⁰ 6s ²			

The definition of the transitional element by the electronic configuration of neutral atom excludes copper, silver and gold whose electronic configurations are in stabilization ns¹ (n-1)d¹⁰ state as well as zinc, cadmium and mercury, ns² (n-1)d¹⁰ on the other hand if the presence of electrons in chemically united atoms is the only property excluding the elements Y, La and Ac.

3-2-2 General characteristics

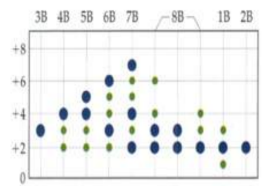
1 - Physical properties

All transition elements d are generally highly dense metals and low atomic volumes and high melting and boiling points. It is fused and boils the last elements of the group at distinctly lower temperatures as compared to other elements of the group. Mercury is the last element of the transition chain represents the observed exception of metals for being fluid under normal condition. Elements of the lanthanide chain possesses metallic properties which show a metallic luster and good conductivity of electrical and heat, as well as that the degrees of melting and boiling points as expected.

2. Chemical properties

A - Elements of a group d:

In general, group d elements are relatively ineffective with oxygen, halogens, sulfur, nitrogen, hydrogen and water vapor at normal conditions but at high temperatures are reacted with these reagents more easily. The group of transition elements reacts with hydrogen under certain conditions to form substances with structures called internal hydrides. This name originally implied that the arrangement of the metal atoms is roughly the same as that of the metal crystallization while the hydrogen atoms enter the distances. Although this is not the reality, the term remains traded. These hydrides occupy a volume larger than the metal volume, where it was formed from it and has a metallic appearance. It is prepared by direct union of metals and hydrogen at high temperatures. For example, platinum, palladium and iron are permeable metals to hydrogen at high temperatures. Tantalum absorbs hydrogen, creating an easily broken product. The elements of group d react directly when the carbon-element mixture is heated at temperatures above about 2200°C produced



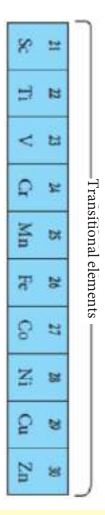
There are no zero oxidants in the first series of transition elements as shown In the larger circles in the graph.

carbides. The carbides have high melting degrees and are very hardened, which is on two groups with a general formula MC and M₂C such as titanium carbides, zirconium, hafnium, vanadium and others. These carbides are characterized by chemical inertness, such as titanium carbide TiC, it is not affected by water or aqueous solutions of hydrochloric acid even when 600°C. B. Elements of group f lanthanides:

Lanthanide metals are soft and more effective with known reagents of group d elements, so lanthanides react slowly with halogens forming MX₃ compounds and with oxygen forming-M₂O₃ at room temperature but easily ignite with these reagents at temperatures above 200°C and react with sulfur at boiling point to form M₂S₃ and nitrogen at a temperature higher than 1000°C to form MN. At temperatures above 300°C, lanthanides react rapidly with Hydrogen to form a kind of hydride and at high temperatures this reaction with boron and carbon gives borides and carbides respectively.

3-2-3 The first transition series

The elements of this series (from scandium to zinc) (Zinc) in the fourth period of the periodic table between calcium Ca in group (IIA), and Galuim Ga in group (IIIA). Table (3-2) shows the electronic configuration of these elements. These elements are called transition elements or group of d elements because they possess internal energy level 3d. It is proposed that the transitional elements of the fourth period or the first transition series should end with nickel element because the 3d energy level of the next two elements (copper and zinc) is full, and fact that copper shows multiple characteristics of the transition elements, while zinc shows intermediate characteristics between the transition elements and the main group elements. For this, it is appropriate to include both copper and zinc in the first series of transition elements. That the electronic configuration differences that distinguish transitional elements of other elements, lead to the emergence of special physical and chemical properties of the transition elements. These properties are not necessary unique behavior to the transitional elements. but combined they give transitional elements have a behavior that is distinct from that of any other type of element. These characteristics can be limited to the elements of the transition chain first:



series of the first transitional elements.

A- Metallic Characteristics:

One of the most prominent properties of the elements of the first transition series is that they are all of metals with high melting and boiling point and good conductors for heat and electrical. They are generally hard materials and are strong alloys with them and the possession of these qualities gives them a unique technological importance of its kind. Although the transition elements are denser and stiffer and have higher boiling points than the main groups, there is no systematic increase in the degree of these properties as the atomic numbers increase. That elements metal of this series are divided into two groups; first groups from Sc to Mn and the second from Mn to Zn with peaks at Ti and V and at Co, and Ni.

The division of the series into two groups has to do with the fullness of orbitals d. The energy level 3d of manganese is half full and then it becomes occupied by individual electrons fills by electrons pairs to become complete fullness at copper and zinc.

That electronic configuration 3d⁵ in chrome and 3d¹⁰ in copper gets by removal of an electron from the 4s level for the purpose of obtaining the most stable ranking. These elements are intended to release the largest number of external valence electrons for metal bonding, that it gives the highest binding energies and is reached in the middle of the series.

B - Oxidation cases:

There are more than one oxidation state in all transition elements, for example, iron has two oxidation states, +2 and +3, and an element atom Cobalt has two oxidation states of +2 and +3 of Cr also numerous oxidation states are +2, +3, +4, +5, and +6 in their ionic and covalent compounds. The oxidation state of the transition elements reaches +7, in the case of manganese as in the compound potassium permanganate KMnO₄ Table(3-2). The cause of the oxidation state is multiple for atoms of the transition elements are due to the number of electrons in the outer shell ns and (n-1)d to the atom of that element, where the loss of electrons from ns begins first and then in terms of the loss of these electrons one after the other that the number of electrons in d does not exceed five electrons. Each electron loses an oxidation state appear, so that manganese has an oxidation state of +1, +2, + 3, +4, +5,+ 6 and +7 due to the difficulty of losing all electrons in (n-1) d because of its need for high ionization energy so prefer to compose coordinate. The resulting ion is a strong oxidizing agent that draws electrons from neighboring atoms. The highest oxidation state of the atoms of the transition elements in the first series depends on:

- 1. Power of oxidizing agent.
- 2. Nature of the resulting compound

Table	Table 3-2 idation states reported by the first transition series									
Element symbol	Electronic configuration of the two orbitals (n- 1) 3d . ns	Multiple oxidative states	Highest oxidation case	The most stable oxidation state						
Sc	$4s^2 \ 3d^1$	+3	+3	+3						
Ti	$4s^2 3d^2$	+3 ، +4	+4	+4						
V	$4s^2 3d^3$	+2 , +3 , +4 , +5	+5	+4						
Cr	$4s^2 3d^5$	+2 , +3 , +6	+6	+3						
Mn	$4s^2 3d^5$	+2 , +3 , +4 , +5 , +6 , +7	+7	+2						
Fe	$4s^2 \ 3d^6$	+2 ، +3	+6	+3						
Со	$4s^2 3d^7$	+2 ، +3	+4	+2						
Ni	$4s^2 3d^8$	+2 , +3	+4	+2						
Cu	4s¹ 3d¹0	+1 ، +2	+3	+2						
Zn	$4s^2 3d^{10}$	+2	+2	+2						

From the Table (3-2) you can see the following:

- 1-The presence of familiar oxidative state (+2) when electronic loss $4s^2$
- 2- The increase in the number of oxidation state of scandium (Sc) to (Mn) and in the latter element the oxidation state is consistent with the loss of electrons 3d⁵, 4s²
- 3- The sharp decrease in the number of oxidants after manganese due to the difficulty of removing electrons after doubling.

C-Acid and base properties

The acidic and basic properties of the transition elements are based on Lewis' concept of oxidation state, so the higher the oxidation number of element, the lower the basic and the more acidity as shown in Table 3-3 for manganese in its oxides.

Table 3-3 Acid and base properties for magnesium transition element in its oxides								
Oxidation number	property	formula	Oxide name					
+2	basal	MnO	Manganese oxide					
+3	Weak basic	Mn_2O_3	Trioxide dimagnesium					
+4	Amphoteric	MnO_2	Dioxide magnesium					
+6	Acidity	MnO ₃	Trioxide magnesium					
+7	Rich acidity	Mn_2O_7	Heptoxide dimagnesium					

D-Formation of coordinate Complexes

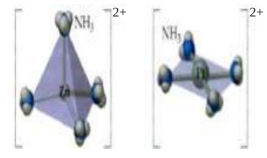
The positive ions of the transition elements shed strong electrostatic attraction on molecules or ions that contain one or more pairs of non-bonding electrons are known as such molecules or ions by (ligands). The result of this attraction is the so-called complex coordinate. Thus, we can say that the coordinate complex is an centralized atom is often a transitional element surrounded by a group of atoms, molecules, or ions are called ligands. The central atom often, they are transition metals, but ligandsmay be a negative ion of signal atom, such as a halide ion, or may be a polyatomic molecule, or an ion containing a donated atom belonging to oxygen or nitrogen group such as H₂O or NH₃ or CN or NO₂ and others. The term coordinate number uses to refer to the number of atoms that donated electrons related to the central atom. The coordinate number of the iron atom in the coordinate ion [Fe(CN)₆]-4 equals 6, and the oxidation state of iron equals +2. The number -4 represents the complex ion charge, which is equal to (algebraic sum of the coordinate charge number and the central ion charge), where the sum of coordinate charge number equal (-6) and central ion charge equal (+2) so, the difference between them is (-4) representing the complex ion charge. The coordinate number varies from one element to another but in the complex compounds of the transition elements in oxidation states +2 and +3 is usually 4 or 6. Examples of such complexities are [Co(NH₃)⁶] ⁺³ and $[Ni\ (CO)_{a}]^{+2}$.Common forms of such complexes are octahedral of the coordinate number (6), and tetrahedral of the coordinate numbers 4. There are other forms such as linear, binary, triangular, and others. These complex shapes will the student is informed about it in his university studies.

H- Color

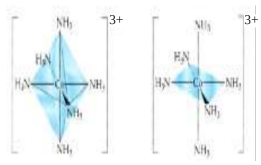
All complexes of the transition elements are characterized by their distinctive bright colors, for example $CuSO_4.5H_2O$ copper sulphate has a light blue color, while $Ni (OH)_2$ light green color and copper fourth ammonia (II) $[Cu (NH_3)_4]^{+2}$ dark blue and others.

I- Magnetic properties

Electrons have a charge and their motion creates two magnetic effects one is associated with the quantum number L and the other is accompanied by the quantum number s and these two effects give each electron the characteristics of a small magnetic pole.

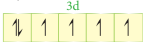


A octahedral complex.

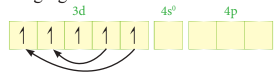


B tetrahedral complex.

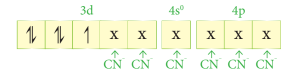
A magnetic moment in electron-filled shell is equivalent to the magnetic moment of each other and the atom as a whole will possess the resultant magnetic moment only if there are individual electrons in the valence orbit. When this happens it will make the outer magnetic field projected to direct the magnetic moment of the atoms toward that this field and this such behavior is called (paramagnetism). When all electrons are double in external orbitals, The case is induced by magnetic fields that reflect the magnetic field that generated it. This makes material repel with the external magnetic field, and this behavior is known as (diamagnetism). For example, we find iron as an element is attracted towards the magnet field because of its existence individual electrons in its outer shell where the outer shell is for the iron atom (3d).



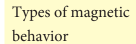
If the iron is within the coordinate ion as in $[Fe(CN)_6]^{-4}$ (Hexaciano ferrate(II)) it is not attracted to the magnetic field because the orbitals the outer shell of the iron ion (II) has been saturated with electrons granted by ligands. For the component $[Fe(CN)_6]^{-3}$ (Hexaciano ferrate(III)), it is where the triple iron ion gravitates toward the magnetic field (paramagnetism) due to the presence of a single electron in the outer orbital shell 3d of triple iron ion as shown in the following figure:

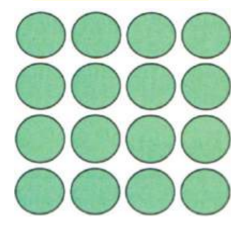


At the approach of the ligand(coordinate group), such as (CN⁻¹) has a high rang of orbits 3d of triple iron ion works on duplex electrons in 3d orbitals for iron ion as shown in following figure:

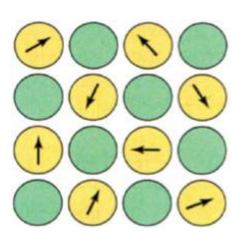


The electrons given by the coordinate groups are placed in the form of (X) differentiated from electrons not granted by coordinate groups (1)





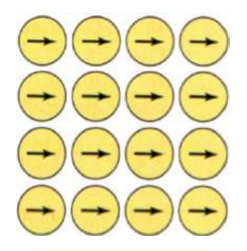
Diamagnetism



Paramagnetism

Information

The green color represents the electron pairs And the yellow color represents single electrons Single.



ferromagnetism

A complex ions that have a paramagnetic property is [NiCl₄]⁻², where the outer shell (3d) of the Ni⁺² ion contains two single electrons, so it shows a single paramagnetic property

$${}_{28}Ni^{2+}\ 1s^2\ 2s^2\ 2p^6\ 3s^2\ 3p^6\ 4s^0\ 3d^8$$

in orbitals



If we take the zinc component ($_{30}$ Zn) we observe from its electronic configuration that all the electrons in its outer shell are in pairs and so it is not attracted to the magnetic field.



Permanent magnets made of materials ferromagnetism

A third type of magnetic behavior is ferromagnetism, which is actually very rare but it is extremely importance. a special case of paramagnetism occurs in compounds that contain a large part of atoms or ions containing single electrons.

For this type of components and in appropriate conditions, the single electrons of each atom interact and regulate themselves with electrons. This process is repeated during all atoms in the component. Therefore, the effect leads to the construction of permanent magnets. Ferromagnetic behavior can be observed mainly among metals ,alloys and transition elements oxides and less impact in internal transition elements oxides.

Gouy balans

The material with dia or paramagnetic properties can be identified using a sensor called (Gouy balance). This device has a very sensitive balance attached in one hand the material whose magnetic properties are to be defined and the equivalent weights are placed the material in the second cuff as shown in Figure (3-12).

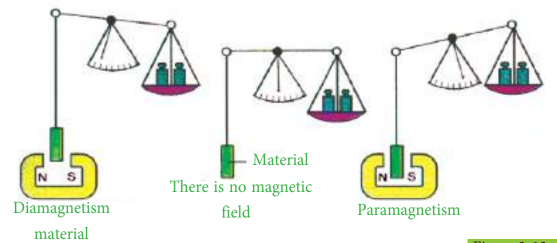


Figure 3-12

Gouy balans.

I-Effectiveness as catalysts:

The catalysts derived from the transition elements have great importance in many biological processes and systems and are indispensable in chemical industries too. Almost all of the transition elements ability to behave as catalysts either in their free or in component forms. This ability is likely to come from the use of d orbital or from the formation of compounds that can absorb and activate reactants. The ability of these compounds to behave as catalysts by ability to create low-energy pathways for reactions by either oxidation-induced changes or by the formation of suitable intermediate compounds.

3-2-4 Lanthanides and Actinides

1.Lanthanides

Lanthanum (La) (atomic number 57) shows the first element in the third series of transition elements which has electronic configuration [$_{36}$ Kr] $4d^{10}$ $5s^2$ $5p^6$ $5d^1$ $6s^2$.

The next element is cerium (58) and its atomic number is 58 and electronic configuration [36]Kr] 4d¹⁰ 4f¹ 5s² 5p⁶ 5d¹ 6s². Where electrons begin to full the energy level (peak level) 4F until we get to the element lutetium (Lu) has an atomic number (71) and then the 4d level returns to fullness. Elements from lanthanum to lutetium are called lanthanides the first internal transition series consists of 14 elements.

 58
 59
 60
 61
 62
 63
 64
 65
 66
 67
 68
 69
 70
 71

 Ce
 Pr
 Nd
 Pm
 Sm
 Eu
 Gd
 Tb
 Dy
 Ho
 Er
 Tm
 Yb
 LL

Lanthanides series

These elements are called a rare earth because they exist in uncommon mixtures as they were thought to be elements of soil or oxides.

The elements of lanthanides are very similar to each other, and their separation is a major problem because all their compounds are very similar, where the oxidation state (+3) appears and this state shows the predominant ionic properties are similar to the alkaline earth metal ions except that they are positive trilogy and not positive dualism. There is no aerobium (Eu) (one of the elements in this series) free in nature and can only be discovered when obtained within an nuclear fission output.

2. Actinides:

Similarly to the lanthanide series, the actinides begin with an element actinium (Ac) has an atomic number of 89 and has an electronic configuration: [54Xe] 4f¹⁴ 5d¹⁰ 6s² 6p⁶ 6d¹ 7s² Before we know about the possibility of the presence of elements after uranium has developed heavier natural elements such as thorium, erotectinium and uranium in the sixth period of periodical classification in tandem elements of hafnium, tatalium and tungsten. It was inferred that these elements were the beginning of a new series of four ten elements in which the sixth quantum level is filled in exactly the same way that the fifth quantum level is full by hafnium, tantalum and tungsten elements. The discovery of several post-uranium elements and the study of their qualities in fact demonstrate the emergence of a new internal transitional series that begins after actinium. So postactinium elements are now called actinides. Whatever the extent of oxidants possessed by oxidants, they always have an oxidative state within oxidants.

In the actinides, the element curium is one of the elements, its inner subshell casing is probably half-filled, and the vast majority, for its compounds curium (Cm) is a positive trilogy, while the elements are above americium have many oxidative states behave like +2, +3, +4, +5, +6 and an Berkelium element (Bk) after curium exhibits oxidative states +3, +4. Polymorphism in oxidative states of actinides elements to extent amersium elements makes the chemistry of the elements of this series very complex.



actinides series.

In addition, in cadolinium (Gd) the subshell level is filled in half with electrons. This is known particularly represents electronic configuration and ionization. Cadolinium forms Gd⁺³ ions only, (with the loss of three external electrons) and does not show inclination to add or lose electrons in a half-filled inner level. This behavior can be compared with the element that comes before cadolinium is an aerobium (Eu), and this element exhibits an oxidation state of +2 and also +3,the next element, terbium (Tb), exhibits oxidative states +3 and +4. We will take iron as an example of transition elements.

3-3- Iron

3-3-1 Introduction

Iron is the second metal after aluminum and the fourth element after oxygen, silicon and aluminum in proportion to the crust of the ground, believed to be the center of the earth, consists mainly of iron and nickel. Iron is spread in the earth's crust, united with other elements forming multiple ores; hematite which contains Fe₂O₃ and magnetite contains Fe₃O₄ and limonite contains FeOOH and siderite contains FeCO₃.

Iron is also the most important transitional element, it is one of the elements involved in the synthesis of blood hemoglobin and is also found with molybdenum in the synthesis of the nitrogen fixing enzyme. It is one of the necessary elements that are Chlorophyll for the food industry in the plant.

3.3.2 The location of iron in the periodic table

Iron falls in the periodic table and in the fourth period the eighth group B, and has the following electronic configuration: $_{26}$ Fe : $1s^2 \ 2s^2 \ 2p^6 \ 3s^2 \ 3p^6 \ 4s^2 \ 3d^6$

The fact that the iron element of the transitional valence fall within the two outer shell [ns, (n-1) d] but it does not show an oxidation state equal to this number (8).

The most common oxidation states in iron are +2 and +3, the process of losing two electrons from the iron atom to form an iron ion (II) or the loss of three electrons to form iron ion (III) depends on the reactants nature. The relationship between the oxidation states can be represented the following equation below: $Fe^{2+} \longrightarrow Fe^{3+} + e^{-}$



magnetite



limonite



hematite



The equation above shows that the iron ion(II) can oxidize (lose an electron) to become an iron ion(III), or the iron ion(III) reduces (acquires an electron) to the iron ion(II).

3-3-3 Properties of iron

Iron is a shiny white metal that is pure and is characterized by all the propertied of metals such as hardness, good thermal and electrical conductivity, and ability of hammering and, extrusion and other properties. It is a magnetized metals with melting point 1528°C, boiling point 2861°C and density 7.86 g/cm³ at 25°C

3-3-4 Iron reactions

1. Iron does not react in normal temperatures with dry air and does not react with water free from dissolved air, but reacts with oxygen in moist air to form a brownish reddish layer called rust.

This layer of iron is separated in the form of crusts, which is aqueous iron oxide and does not stop rust at the outer surface of the iron piece (due to permeability and incoherence of the oxygen layer formed)

This layer of iron is separated in the form of crusts, which is aqueous iron oxide and does not stop rust at the outer surface of the iron piece (due to permeability and incoherence of the oxygen layer formed)

$$4\text{Fe} + 3\text{O}_2 + \text{nH}_2\text{O} \longrightarrow 2\text{Fe}_2\text{O}_4.\text{nH}_2\text{O}$$

2. Iron is oxidized when heated to high temperatures (degree of redness) and in the presence of air forming iron oxide magnetic Fe_3O_4 as in the following equation:

$$3\text{Fe} + 2\text{O}_2 \xrightarrow{\Delta} \text{Fe}_3\text{O}_4$$

3.Iron reacts with water vapor, forming magnetic iron oxide and realizing hydrogen gas as in the following equa-

tion:
$$3\text{Fe} + 4\text{H}_2\text{O} \xrightarrow{\Delta} \text{Fe}_3\text{O}_4 + 4\text{H}_2$$

4. Iron reacts easily with dilute acids such as hydrochloric acid and sulfuric acid forming iron salt (II) and realizing hydrogen gas as in the following equations:

It reacts with concentrated acids such as hot concentrated sulfuric acid and nitric acid, forming iron salts and water, and rising SO₂ gas as in the following equation.

$$3Fe + 8H_2SO_4 \longrightarrow FeSO_4 + Fe_2(SO_4)_3 + 4SO_2 + 8H_2O$$
Iron sulfate (II) Iron sulfate (II)

that the most famous iron sulphate is iron pyrite, which is known as gold false because of the color similar to the color of gold, which is used in cheating gold without easily detected 5. When heating a mixture of iron filings and sulfur powder form Iron sulfide (II) as in the following equation:

$$Fe + S \longrightarrow FeS$$

6.Iron reacts with halogens (fluorine, chlorine and bromine) to produce iron halides (III) according to the equation:

$$2Fe + 3X_2 \longrightarrow 2FeX_3$$
; $X = F \cdot Cl \cdot Br$

For example, iron chloride (III) is produced from the passage of chlorine gas on the heated iron filings to the redness temperature as in the following equation:

$$2\text{Fe} + 3\text{Cl}_2 \longrightarrow 2\text{FeCl}_3$$

3-3-5 Iron extraction

Iron is produced artificially in the kiln blower by reducing iron ores, especially hematite and magnetite with carbon at 2000°C.

The method involves mixing iron ore with coke and limestone(mostly calcium carbonate) and add the mixture from the top hole of the bloating oven [Figure (3-13)].

Then the hot air from the nozzles is blown in a result of the high temperature of the mixture, many reactions happen within the furnace can be summarized. Firstly coal combustion to heat the furnace and form a carbon monoxide:

Carbon monoxide reduces iron ore to produce fusible iron and carbon dioxide:

$$Fe_2O_3 + 3CO \xrightarrow{\Delta} 2Fe + 3CO_2$$

The furnace heat also caused the decomposition of calcium carbonate into calcium oxide and carbon dioxide according to the equation:

$$CaCO_3 \xrightarrow{\Delta} CaO + CO_2$$

Then calcium oxide combines with sand to be molten calcium silicate according to the equation:

$$CaO + SiO_2 \longrightarrow CaSiO_3$$

The calcium silicate molten floats above the iron molten, because its density less than the density of iron, forming a layer called slag that prevents mixing of iron fusion with the material above it. The molten iron is drawn from time to time from the bottom of the oven, where it is poured into special molds and then called cast iron, which is used as iron ore in the following stages of iron manufacturing.

Do you know

That iron enters into a lot of the most famous cyanide compounds Fe₄ [Fe(CN)₆] is called a blue tincture Prussia which is used to remove yellowing water from sediment salts iron. called locally (Jweet).

The slag formed is also pulled from special holes below furnace, which is used in cement, concrete or concrete roads and other construction. Due to environmental concerns about the use of coke, alternative methods of iron processing have emerged, one of which is the reduction of iron using natural gas. Iron is produced in the form of a powder called spongy iron, which used in steel industry. The process consists of two main interactions namely: oxidation of natural gas with the help of auxiliary agent and

heat.
$$2CH_4 + O_2 \longrightarrow 2CO + 4H_2$$

the resulting H₂ and CO gases are then at 800-900°C reduction of iron ore to produce iron spongy.

$$Fe_2O_3 + CO + 2H_2 \xrightarrow{\Delta} 2Fe + CO_2 + 2H_2O$$

The sand is then removed by adding calcium carbonate in the next step to form slag as in the previous method.

This method of iron production is characterized by:

- 1. Does not require expensive coke and is not available.
- 2. The cost of its creation is much lower than the cost of setting up the blower oven.
- 3. Technology is simple and easy to use.
- 4. The iron produced is carbon-free while the cast iron is produced from the lower furnace contains approximate ly 4% carbon.

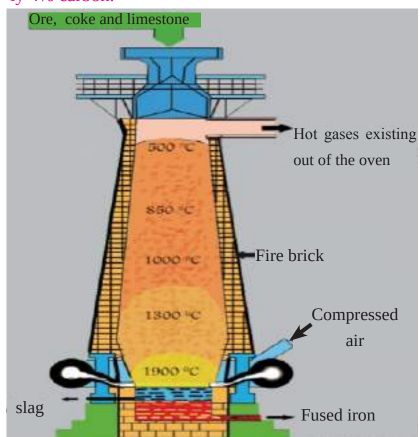


Figure 3-13
Blower furnace.

3-3-6 Iron types

We learned that the iron produced from the blower furnace is called cast iron contains impurities ranging from (6% to 8%) mostly carbon and silicon. Percentage change these impurities can get different types of iron ones:

A- Casting Iron

When re-melting cast iron and pouring into molds to make the required tools then it is called cast iron and it has two types depending on casting and cooling method. If you use metal molds for casting be the casting process is quick and the output is called white cast iron. If sand molds are used then the solidification process is slow and the product is called gray cast iron. In general, cast iron, known locally as 'Aheen', is characterized very rigid but brittle and does not withstand strong shocks. It is often used in making some parts for fireplaces, pipes and sewage covers.

B) casting (steel):

Iron produces steel from cast iron after oxidation of existing impurities in a special furnace, add the required amount of carbon (0.2% to 1.5%) and other elements according to the desired qualities of the desired steel quality. The properties of iron and steel depend on the ratio of carbon and

Chapter Three Questions



- 3-1 How the atomic size of the elements of the single period and single group change by increases the atomic number and why.
- 3-2 Which atoms in each of the following pairs are larger atomic size ($_3$ Li, $_4$ Be) ($_{16}$ S, $_8$ O), ($_{17}$ Cl, $_{35}$ Br).
- 3-3 Arrange the following according to the ionaization energy increasing, mentioning reason. Al, Al⁺, Al⁺², Al⁺³.
- 3-4 If you have the following elements (17Cl, 15P, 11Na) answer the following questions:
 - A- Any of the elements have a larger atomic size and which the smallest atomic size.
 - B- Arrange these elements as increasing electronegativity, mentioning the reason for this arrangement.
 - C- Arrange them with elements according to the increasing attraction electronic affiliation, mentioning the reason
 - D- Any of these elements in which the metallic properties are expected.
- 3-5 You have atoms ₃₄Se, ₁₆S, ₈O arranged them according to the following with the reason.
 - A- Electronic affinity.
 - B- Increasing electronegativity
 - C- Atomic radius increasing
 - D- Increasing ionization energy
- 3-6 Explain
- A- High ionization energy ₁₇Cl compared to ionization voltage ₁₂Mg
- B- Energy is released when the first electron is acquired, but many often absorb some energy when gaining the second electron.
- C- Fluorine ionization energy is greater

- than oxygen ionization energy.
- 3-7 How to obtain a spectrum for the element. Does the spectrum of the element appear when it absorbs energy or when it emits energy.
- 3-8 What is the coordinate number of the central atom and complex Ion charge of each of the following:
- A- Hexaciano ferrate [Fe(CN)₆](III)
- B- Quaternary ammonia copper [Cu(NH₂)₄](II)
- C- Quaternary chloro Nickel [NiCl₄](II)
- 3-9 Mention methods of industrial extraction of iron? Which is better and why?
- 3-10 Define each of the following
- 1- Transitional element.
- 2- Delements group.
- 3- Coordinated number.
- 4- Ligand.



Chapter Four Chemical Kinetics

4

After completing this chapter, the student is expected to:

- Calculate the reaction speed in terms of the change in the concentration of materiel over time.
- Determine the order of reaction and deduces the law of reaction speed.
- Explaine the chemical reactions occur collision theory and transition state theory.
- Determine the factors influencing reaction speed and how to control it to increase reaction speed.
- Distinguish between exothermic and endothermic reactions.
- Propose an acceptable reaction mechanism that conforms to the specified conditions

4-1 Introduction

We have learned from our previous study of chemical changes and simplified that they occur between the reactants to form the products, and expressed by a balanced chemical equation, which is useful in calculating the number of moles and masses of materials and the volumes of reactive and resulting gases. But the chemical equation did not tell us other details regarding the speed at which the reactants are converted into products as well as the number of steps that the reaction pass to create the products. All these changes are studied by chemical kinetics and concern by the following points:

- 1. The speed of the chemical reaction and how to measure it and the factors affecting it.
- 2. The reaction mechanism, ie the initial reactions that the reactants pass through to reach the final product and how they are expressed in balanced chemical equations.

Chemical kinetics is an important science because knowledge speed of the reaction and the factors that affect it, such as concentration, pressure, temperature, and the nature of the reactants, allow students to predict how quickly the reaction will reach at equilibrium. Also and that study of the reaction mechanism (the initial steps that the reaction passes through to the reaction of the reactants into the products), enables them, when they know, to control the flow of the reaction to obtain the products in the required quantities and in appropriate economic ways.

4-2 Chemical reaction speed

The definition of speed is familiar in our daily lives. If a car drives 70 kilometers in one hour, we say it has a speed of 70 Kilometer / hour, or student reads twenty pages of a book in 30 minute, or the people of Baghdad consume one million cubic meters of potable water per day. These expressions of speed share a common explanation that a change occurs over a certain period of time. The speed of the car expresses the change in place measured in kilometers per hour. For a student reading a book, the number of pages read increases twenty pages every 30 minutes. As for the people of Baghdad who consume one million cubic meters of potable water per day, water tanks will decrease by one million cubic meters during this time period.

Similar to chemical reactions, they occure at varying rate according to the change in the quantities of substances entering or leaving the reaction per unit of time. Chemical reactions vary in rate. Some of them are very fast when mixing the reactants in a fraction of a second, such as neutralization reactions between acids and bases and combustion reactions. Others have moderate rate that take several minutes to several months, such as iron rust and reactions that lead to the ripening of fruits and vegetables. There are very slow. reactions that take several years or millions of years to occur, such as reactions that lead to human growth and age and turn dead plants into coal. Note the Figure (4-1).

4-2-1 Measurement of chemical reaction speed

The rate of a chemical reaction is measured by knowing the concentration change in reactive or product material in unit of time (unit of time may be second (s), minute (min), hour (hr), day (day) or any other units of time). In this chapter we will use the molar concentration exclusively, which is defined as the number of moles of solute in one liter of solution (mole /L) expressed by a square arc [], where any symbol inside the arc means molar concentration. For example [X] means the molar concentration of material X. So we will express the rate of reaction speed in the mathematical relationship as the following:

Rate =
$$\frac{\Delta []}{\Delta t}$$

rate of reaction=(Change in concentration of one of the reactants/change in time)

$$R \longrightarrow P$$

Where Δ means change, and [] means molar concentration in unit (mol/ L). Therefore have the chemical reaction rate unit:

mole / L.t ((or).) mole L^{-1} t^{-1}

For the following general reaction:

Where R represents any (reactant) and P represents any product substance. The rate of this general reaction is calculated by



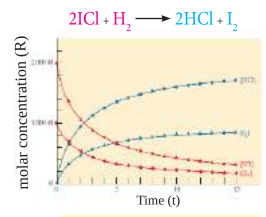






Figure 4-1

Differential reaction rate, from top to bottom: combustion is very fast reaction, ripening fruits and vegetables Moderate reaction rate, repel a iron reaction moderate rate and lower rate of fruit ripening, human growth reaction is very slow.



The concentration of reactants ICl and H_2 decreases, while the concentration of the resulting substances HCl and I_2 increases over time.

Figure 4-2
The concentration of reactant R decreases over time.

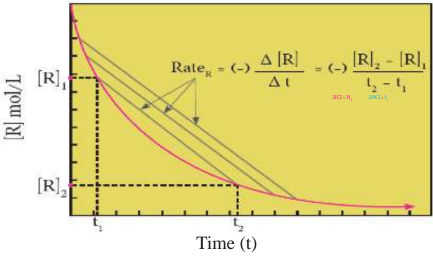
The relationship above and in terms of reactants R and as follows:

Rate of reaction in term R = (-)concentration of R/change in time

$$Rate_{R} = (-) \frac{\Delta [R]}{\Delta t}$$

If the reaction starts with a certain concentration of the reactant, let $[R]_1$ be in the time of (t_1) and as the reaction progresses, the concentration of the reactant material decreases, while concentration of product material will be increased. In time (t_2) concentration of the reactant material $[R]_2$, Note Figure (4-2), rate of reaction in term of R as follows:

Rate_R = (-)
$$\frac{\Delta [R]}{\Delta t}$$
 = (-) $\frac{[R]_2 - [R]_1}{t_2 - t_1}$



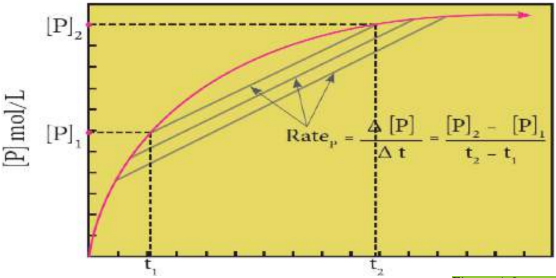
The negative signal is added in the law when expressing the reaction speed in terms of the change in the concentration of a reactant, because the reactant is consumed during the reaction and its concentration is lowered, so that the second concentration is less than the first concentration and the change in concentration is negative Then the negative signal is added to make the rate is positive .

When the reaction rate is measured in terms of the change in the concentration of the substance produced by the reaction P, the relationship is as follows:

Rate of reaction in term P = change in P concentration/ change in time

Rate_p -
$$\frac{\Delta [P]}{\Delta t}$$
 - $\frac{[P]_2 - [P]_1}{t_2 - t_1}$

The change in the concentration of the resulting substance Δ [P] is positive because its concentration increases over time and the second concentration is [P]₂ greater than the first concentration [P]₁, so no negative sign is added in the law. As shown in Figure (4-3).



Time (t)

Figure 4–3

The concentration of the resulting substance P increases over time.

Example 4-1

The concentration of material R change from 1.2 mole / L to 0.75 mole / L over 125 s depending on the following reaction:

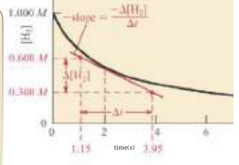
$$R \longrightarrow P$$

Calculate the reaction rate as a function R.

Solution

Rate_R = (-)
$$\frac{\Delta [R]}{\Delta t}$$
 = (-) $\frac{[R]_2 - [R]_1}{t_2 - t_1}$

Rate_R =
$$\frac{\text{(-) (0.75 - 1.20) mol/L}}{\text{(125 - 0) s}} = 0.0036 \text{ mol/L.s}$$



Exercise 4-1

Assume the following reaction:

$$3H_2 + CO \longrightarrow CH_4 + H_2O$$

The concentration of H_2 varies according to the above diagram. Find the speed of this reaction and the time needed to decrease the concentration of H_2 to 0.300 mol / L.

4-2-2 The relationship of reaction rate with the number of moles

The reaction rate varies in terms of the changing in the concentration of the substance depending on the number of moles of the substance in the balanced reaction equation. For example, the reaction between hydrogen and iodine to form hydrogen iodide as in the following equation:

$$H_{2(g)} + I_{2(g)} \longrightarrow 2HI_{(g)}$$

When one molecule of hydrogen is consumed (one mole), one molecule of iodine is also consumed (one mole), while two molecules (two moles) of hydrogen iodide is formed. ie rate of consumed of hydrogen or I_2 equal to half the rate of HI formation.

Rate =
$$-\frac{\Delta [H_2]}{\Delta t} = -\frac{\Delta [I_2]}{\Delta t} = \frac{1}{2} \frac{\Delta [HI]}{\Delta t}$$

or the formation rate of HI is twice rate of consumption H or I

$$\frac{\Delta [HI]}{\Delta t} = -2 \frac{\Delta [H_2]}{\Delta t} = -2 \frac{\Delta [I_2]}{\Delta t}$$

the equation can be write as follows

$$Rate_{(HI)} = 2Rate_{(H_2)} = 2Rate_{(I_2)}$$

Rate(HI) means the rate in term product HI concentration of, and so on for the rat in terms of the reactants H_2 and I_2 , we note that we are not put signals when rate is used because rate is always positive, while change in concentration is negative for reactants and positive for products substances.

For general reaction
$$aA + bB \longrightarrow gG + hH$$

Where (a, b, g and h) are the number of moles of substances in the balanced reaction equation, and the rate of this reaction is expressed in terms of the reactants produced as follows:

$$Rate = -\frac{1}{a} \times \frac{\Delta \left[A\right]}{\Delta t} = -\frac{1}{b} \times \frac{\Delta \left[B\right]}{\Delta t} = \frac{1}{g} \times \frac{\Delta \left[G\right]}{\Delta t} = \frac{1}{h} \times \frac{\Delta \left[H\right]}{\Delta t}$$

$$\frac{1}{a} Rate_{(A)} = \frac{1}{b} Rate_{(B)} = \frac{1}{g} Rate_{(G)} = \frac{1}{h} Rate_{(H)}$$

Exercise 4-2

The rate of formation of NH_3 in the following reaction is 0.15mol / L.min.

$$N_2 + 3H_2 \longrightarrow 2NH_3$$

So the rate in terms of N_2 consumption is equal n unit mol/L. min

A- 0.150

B- 0.075

C- -0.175

D- 0.200

E- 0.300

or

When the rate is expressed in terms of one mole of any substance involved, the reaction ,(general reaction speed) is used. It is the overall rate of reaction in terms of one mole of reactants or products, and as follow:

Rate =
$$\frac{1}{n_1} \frac{\Delta[J]}{\Delta t}$$

Where n_J represents the number of moles of J with its negative signal of the reactants and the positive of the products, it was –a or –b or g or h in the general reaction above.

Example 4-2

For the following reaction

$$2H_{2(g)}^+ O_{2(g)} \longrightarrow 2H_2O_{(g)}$$

- 1.Express the rate in terms of the change in concentration of H_2 , O_2 and H_2O .
- 2. Calculate the $\rm H_2O$ formation rate if the $\rm O_2$ consumption rate is equal 0.023 mol / L.s
- 3. Calculate the amount of H₂O formed after 18s Solution

$$1- Rate_{(H_2)} = (-) \frac{\Delta [H_2]}{\Delta t}$$

$$Rate_{(O_2)} = (-) \frac{\Delta [O_2]}{\Delta t}$$

$$Rate_{(H_2O)} = \frac{\Delta [H_2O]}{\Delta t}$$

$$\begin{array}{lll} 1 \; \Delta \; [H_2O] & & \Delta \; [O_2] \\ 2 \; \; \Delta \; t & \; \Delta \; t \end{array}$$

$$Rate_{(H_2O)} = 2 Rate_{(O_2)}$$

 $Rate_{(H,O)} = 2 \times 0.023 \text{ mol/L.s} = 0.046 \text{ mol/L.s}$

3- Rate_(H₂O) =
$$\frac{\Delta [H_2O]}{\Delta t}$$
 = $\frac{[H_2O]_2 - [H_2O]_1}{t_2 - t_1}$ = $\frac{([H_2O]_2 - 0) \text{ mol/L}}{(18 - 0)\text{ s}}$

 $[H_2O]_2$ = Rate_(H,O) mol/L.s × 18 s = 0.046 mol/L.s × 18 s

 $[H_2O]_2 = 0.828 \text{ mol/L}$

Exercise 4-3

Which one of the expressions below do not represent the suitable expression for the general rate of reaction

$$2A + 3B \longrightarrow F + 2G$$

$$A - \frac{-\Delta [A]}{\Delta t}$$

B-
$$\frac{-\Delta [B]}{3\Delta t}$$

$$C- \quad \frac{-\Delta \ [F]}{\Delta t}$$

D-
$$\frac{-\Delta [G]}{2\Delta t}$$

$$H_{-} \frac{-\Delta [A]}{2\Delta t}$$

Example 4-3

The reaction between ethylene and ozone gas is expressed by:

$$C_2H_{4(g)} + O_{3(g)} \longrightarrow C_2H_4O_{(g)} + O_{2(g)}$$

It was found that the concentration of O₃ changed as the reaction progressed in the table below

t/(s)	0	10	20	30	40	50	60
[O ₃] / 10 ⁻⁵ /(mol/L)	3.20	2.42	1.95	1,63	1.40	1.23	1.10

- A- Calculate the reaction rate during the first 10s of the reaction.
- B- Calculate the reaction rate during the last 10s of the reaction.
- C- What is the reason for the difference between the rate of reaction in both cases?

Exercise 4-4

For the following reaction:

$$4\mathrm{NO}_{(g)} + 3\mathrm{O}_{2(g)} \longrightarrow 2\mathrm{N}_2\mathrm{O}_{5(g)}$$

- A- Express the reaction rate in terms of change in the concentration of each substance over time.
- B- Calculate the rate of O_2 consumption if consumption rate of NO was equal $1.60 \times 10^{-4} \text{mol} / \text{L. s}$

Solution

A- Reaction rate during the first 10s of the reaction.

Rate_(O₃) =
$$-\frac{\Delta [O_3]}{\Delta t} = -\frac{[O_3]_2 - [O_3]_1}{t_2 - t_1}$$

= $-\frac{(2.42 \times 10^{-5} - 3.20 \times 10^{-5}) \text{ mol/L}}{(10 - 0) \text{ s}} = 7.8 \times 10^{-7} \text{ mol/L.s}$

B-The reaction rate during the last 10s of the reaction.

Rate_(O₃) =
$$-\frac{\Delta [O_3]}{\Delta t}$$
 = $-\frac{[O_3]_2 - [O_3]_1}{t_2 - t_1}$
= $-\frac{(1.10 \times 10^{-5} - 1.23 \times 10^{-5}) \text{ mol/L}}{(60 - 50) \text{ s}}$ = 1.3 × 10⁻⁷ mol/L.s

C- The reaction rate during the first 10s is equal to six times as reaction rate during the last 10 s, indicating that the reaction rate is not constant. It decreases over time in terms of the concentration of the reactant as a result of its consumption.

4-3 The reaction rate law

In general, the reaction rate is directly proportional multiplying the products the concentrations of the reactants and each concentration is raised to a given exponent.

The following general reaction:

$$aA + gG + hH \longrightarrow pP$$

The reaction rate is proportional to the concentrations of the reactants as follows:

$$Rate \propto [A]^{\alpha} [G]^{\beta} [H]^{\gamma}$$

Or in the form of equality

Rate =
$$K[A]^{\alpha}[G]^{\beta}[H]^{\gamma}$$

The student should note that the number of moles of the reactants a, g and h is not related to the ranks of the reactants α , β , and γ , but are derived from practical experiments only. The general rate of reaction is represented by Rate for any of the reactants on the prodacts as the following:

Rate =
$$-\frac{1}{a} \frac{\Delta[A]}{\Delta t}$$

Rate = $-\frac{1}{g} \frac{\Delta[G]}{\Delta t}$

Rate = $-\frac{1}{h} \frac{\Delta[H]}{\Delta t}$

Rate = $\frac{1}{h} \frac{\Delta[P]}{\Delta t}$

Where [A], [G] and [H] are the molar concentrations of A, G and H respectively.

 α is called the reactant order A and β is the order of the reactant G and γ is the order of the reactant H. The general order of the reaction n is defined as being equal to the sum of the orders of the reactants as follows:

$$n = \alpha + \beta + \gamma$$

n values are zero, 1, 2, or 3 and may be fractional. K is called the reaction rate constant and the relationship above framed by the law of reaction rate.

Exercise 4-5

For the following gaseous reaction:

$$A + B \longrightarrow C$$

Experimentally found that the law of reaction rate

Rate =
$$K[A]^2[B]$$

So the order of this reaction

- A- First order
- B- Second order
- C- Third order
- D- zero order
- E- half order

×The student notes that the rank of the reactant in the law of reaction rate may not correspond to number of moles of that reactant in the balanced equation, so it is necessary to create the term molecular reaction and compare it with the order of reaction.

The reaction molecule is defined as the number of molecules or ions of reactants in the balanced equation that can theoretically be found from the number of moles. Either reaction order is a value that can only be known by experience, and there is another difference that the order of reaction may be zero while the reaction molecule cannot be like this.

If the reaction is zero order, then the values of α , β and γ are equal to zero, meaning that the reactant concentration do not affect the reaction rate and therefore the law of rate for zero-order reactions is as follows:

If the reaction is a first order, the value $\alpha = 1$ the value of β and γ is zero, the law of rate is written as a

Rate=
$$K[A]$$

For example, the following reaction is of the first order *

$$2N_2O_5 \longrightarrow 2N_2O_4 + O_2$$

The law of reaction rate is:

Rate=
$$K[N_2O_5]$$

The general rank of the reaction is first (n = 1)If the second order reaction, the law of rate can be written as following forms:

Rate=
$$K[A][G]$$

or

Rate=
$$K[A]^2$$

So for other orders. The following examples illustrate the law of rate and orders of some reaction.

The law of reaction rate:

$$2NO_{(g)} + 2H_{2(g)} \longrightarrow N_{2(g)} + 2H_2O_{(g)}$$

Is

Rate=
$$K [NO]^2 [H_2]$$

* The reaction is of the first order for the $\rm H_2$ reactant and the second for the NO reactant. So the overall reaction rate is $\rm n{=}3$

For the following reaction:

$$(CH_3)_3CBr + H_2O \longrightarrow (CH_3)_3COH + HBr$$

The law of reaction rate is:

Rate=
$$K[(CH_3)_3CBr]$$

Reaction rate is first order for the reactant $(CH_3)_3CBr$ and it is zero order for the H_2O reactor and the overall order of the reaction is firstly, ie n = 1

The reaction between acetone and iodine has the presence of MnO₂ as a catalyst.

$$CH_3COCH_3 + I_2 \xrightarrow{MnO_2} CH_3COCH_2I + HI$$

It was observed that its rate does not depend on the concentrations of reactants and the reaction rate law for this reaction is

Rate=K

Thus, this reaction is zero order From the above example, it is clear that the law of reaction rate and its order cannot to be concluded by just considering the balanced chemical equation of the reaction, indeed this should be determined by experiment only.

The values of α , β and γ are not determined by the number of moles of the reactants a, g and h as in the general reaction: $aA + gG + hH \longrightarrow pP$

4-3-1 Determination of reaction orders

To illustrate how to set the values of α , β , γ experimentally. It is used several experimental methods, we will look at one here. This method involves making a number of experiments in which the concentration of a reactant is altered and maintained the concentrations of the other reactants are constant, so we find the order of the variable reactant, and so on for the other reactants. This method can be explained by the following examples:

Example 4-4

The first reaction at a given temperature:

$$2N_2O_5 \longrightarrow 2N_2O_4 + O_2$$

The reaction rate was measured at different concentrations of the reactant N_2O_5 . Data were obtained in the table below.

Exp. No.	$[N_2O_5]$ / mole/ L	Rate / mole / L.s
1	0.0113	6.7×10^{-6}
2	0.0084	5.0×10^{-6}
3	0.0042	2.5×10^{-6}

Determine the order of the reactant and deduce the law of reaction rate and its general reaction rate order.

Exercise 4-6

Determine the orders of the reactants and the general Reactions orders for each of the following reactions:

$$CH_3CHO_{(g)} \longrightarrow CH_{4(g)} + CO_{(g)}$$

Rate = $K[CH_3CHO]^{3/2}$

C-

$$H_2O_{2(aq)} + 3I_{(aq)} + 2H_{(aq)}$$

Rate =
$$K[H_2O_2][I]$$

$$2NO_{(g)} + Cl_{2(g)} \longrightarrow 2NOCl$$
Rate = $K[NO]^2 [Cl_2]$

Exercise 4-7

For the following gaseous reaction:

$$A + B \longrightarrow C$$

Experimentally found that the law of his rate

Rate=
$$K [A]^2 [B]$$

If the concentration of A is three times greater and the concentration of B is twice. The reaction rate is increased by:

A- 6

B- 9 C- 12

D- 18

E-36

Exercise 4-8

For the following gaseous reaction:

 $2NO + 2H_2 \longrightarrow N_2 + 2H_2O$ It has the following rate law:

Rate =
$$K[NO]^2[H_2]$$

If the NO concentration is halved and the H₂ concentration is tripled, the change in reaction rate relative to the first rate will be:

- A- Decreases by 3/4
- B- Increases by 3/4
- C- Increases by 2/3
- D- Decreases by 2/3
- E- Remains the same

Solution

1 - We write the reaction rate law

Rate =
$$K[N_2O_5]^{\alpha}$$

2- To determine the value (α) we choose any two experiments such as 1 and 2 and compensate the results of each experiment in the law of reaction rate and divide one equation by the other and simplify of fraction we get (α) value

$$\frac{\text{Rate}_{1}}{\text{Rate}_{2}} = \frac{\text{K} \left[\text{N}_{2}\text{O}_{5}\right]_{1}^{\alpha}}{\text{K} \left[\text{N}_{2}\text{O}_{5}\right]_{2}^{\alpha}}$$

$$\frac{6.7 \times 10^{-6}}{5.0 \times 10^{-6}} = \frac{K (0.0113)}{K (0.0084)}$$

$$(1.34)^1 = (1.34)^{\alpha} \implies \alpha = 1$$

Therefore the value of α is equal to the integer one, so the reaction is of the first order and its rate law is written as follows:

Rate = $K[N_2O_{\epsilon}]$

4-3-2 Calculating the value of the reaction rate constant

The reaction rate constant is a constant amount that does not change unless the temperature changes. Its value is calculated from the compensation of the results of an experiment in the law of reaction rate, as in the following example:

Example 4-5

Calculate the value of the rate constant of the reaction in Example 4.4.

Solution

To find the value of rate constant we compensate the results of one experiment in the law of reaction rate, that we got in the previous example as follows:

Rate =
$$K[N_2O_5]$$

Suppose we have chosen experiment number 1 and offset its values in the relationship above

$$6.7 \times 10^{-6} \text{ mol/L.s} = K (0.0113) \text{ mol/L}$$

$$K = \frac{6.7 \times 10^{-6} \text{ mol/L.s}}{(0.0113) \text{ mol/L}} = 5.95 \times 10^{-4} \text{ s}^{-1}$$

4-3-3 Rate constant units

The units of rate constant vary depending on the general order of the reaction and the unit of time used to express the reaction rate. It can be derived mathematically from compensation in the law of reaction rate, or from the application of the following law:

Unit of rate constant = $\frac{\left[\frac{\text{mol}}{L}\right]^{1-1}}{t} = \frac{\left[M\right]^{1-1}}{t}$

Where n represents the general reaction order and M molarity. Table (4-1) shows units of rate constant for different orders if time is expressed in (s)

Table 4-1	Uni	ts of rate constant corresponding to the general reaction order
Order (1	n)	Units of rate constant corresponding to the general reaction rate
0		$M. s^{-1}$ or mol $L^{-1} s^{-1}$ or mol/L. s
1		s ⁻¹ or 1/s
2		M^{-1} . s^{-1} or $L.mol^{-1}s^{-1}$ or $L/mol.$ s
3		M^{-2} . s^{-1} or L^2 mol 2 s $^{-1}$ or L^2 /mol 2 .s

Exercise 4-9

Use the constant rate value calculated in example 4–5 to calculate the consumption rate of N_2O_5 . When its concentration is equal to 0.1 mol/L at the same set temperature.

Example 4-6

For the following reaction

$$O_{2(g)} + 2NO_{(g)} \longrightarrow 2NO_{2(g)}$$

Determine the orders of the reactants and deduce the law of reaction rate, general reaction rate, and rate constant of experiments results in the table below:

Exp.No.	[O ₂] / mole/ L	[NO] / mole/ L	Rate / mole / L.s
1	1.10×10 ⁻²	1.30×10 ⁻²	3.20×10 ⁻³
2	2.20×10 ⁻²	1.30×10 ⁻²	6.40×10 ⁻³
3	1.10×10 ⁻²	2.60×10 ⁻²	12.8×10 ⁻³
4	3.30×10 ⁻²	1.30×10 ⁻²	9.60×10 ⁻³
5	1.10×10 ⁻²	3.90×10 ⁻²	28.8×10 ⁻³

Solution

Write reaction rate law Rate= $K [O_2]^{\alpha} [NO]^{\beta}$

To determine the value of (α) , select two experiments such as 1 and 2, where [NO] is constant in the two experiments while $[O_2]$ variable and compensate for the results of each experiment in the law of rate. Divided one equation over the other and simplify the fraction we get (α) value

Exercise 4-10

For the following reaction

$$X + 2Y \longrightarrow P$$

It is found first order for X and second order to Y.

The units of rate constant of this reaction if time in seconds is:

 $A - M \cdot s^{-1}$

 $B-M^{-2}$. s^{-1}

C- M-3. s

 $D-M^2$. s^{-1}

 $E-M^{-1}. S^{-1}$

Exercise 4-11

For the following reaction:

From the results of the three experiments in the table below:

$\boxed{ [C_3H_6O_2] }$	[OH]	Rate/ mol/L. s
0.040	0.040	0.000225
0.040	0.080	0.00045
0.080	0.080	0.00090

A- Determine the orders of the reactants.

B- Deduce the law of reaction rate.

C- Find the value of the constant rate.

$$\frac{\text{Rate}_{2}}{\text{Rate}_{1}} = \frac{K \left[O_{2}\right]_{2}^{\alpha} \left[\text{NO}\right]_{2}^{\beta}}{K \left[O_{2}\right]_{1}^{\alpha} \left[\text{NO}\right]_{1}^{\beta}}$$

$$\frac{6.40 \times 10^{-3}}{3.20 \times 10^{-3}} = \frac{K \left(2.20 \times 10^{-2}\right)^{\alpha} \left(1.30 \times 10^{-2}\right)^{\beta}}{K \left(1.10 \times 10^{-2}\right)^{\alpha} \left(1.30 \times 10^{-2}\right)^{\beta}}$$

$$(2)^{1} = (2)^{\alpha} \implies \alpha = 1$$

So the reaction is first order for O₂

To determine the value (β), we select experiments 1 and 3 where $[O_2]$ is constant in both experiments, while [NO] is a variable and we substitute the results of each experiment in the rate law and divided one equation over the other and simplify the fraction, we get a (β) value.

$$\frac{\text{Rate}_{3}}{\text{Rate}_{1}} = \frac{K \left[O_{2}\right]_{3}^{\alpha} \left[\text{NO}\right]_{3}^{\beta}}{K \left[O_{2}\right]_{1}^{\alpha} \left[\text{NO}\right]_{1}^{\beta}}$$

$$\frac{12.80 \times 10^{-3}}{3.20 \times 10^{-3}} = \frac{K \left(1.10 \times 10^{-2}\right)^{\alpha} (2.60 \times 10^{-2})^{\beta}}{K \left(1.10 \times 10^{-2}\right)^{\alpha} (1.30 \times 10^{-2})^{\beta}}$$

$$(4) = (2)^{\beta} \implies (2)^{2} = (2)^{\beta} \implies \beta = 2$$

So the reaction is second order for NO.

The general order of the reaction is equal

$$n = \alpha + \beta = 1 + 2 = 3$$

so the reaction rate law is Rate= $K[O_2]$ [NO]²

We use the information in one of the experiments in the table above and replace it in the law of reaction rate to get the value of the rate constant. We used information from experiment 2

Rate =
$$K [O_2] [NO]^2$$

6.4 × 10⁻³ mol/L.s = $K(2.20 \times 10^{-2})$ mol /L $(1.30 \times 10^{-2})^2$ mol² /L²

$$K = \frac{6.4 \times 10^{-3} \ mol/L.s}{(2.20 \times 10^{-2}) \ mol/L \ (1.30 \times 10^{-2})^2 mol^2/L^2} = 1721.4 \ L^2 \ /mol^2.s$$

Note that the unit of rate constant for third order reactions is $L^2 / mol^2.s$

4-4 Theories of reaction rate

A number of theories have been developed to explain how the reaction occurs. The most important of these theories is the theory of collision and the theory of the transition state or called the theory of activated complex. The first explained how the reaction occurs on the basis of the visible, while to the second explained the reaction in a way that happens inside molecules subject to interaction and we will briefly explain both theories.

4-4-1 Collision Theory

The rate of chemical reactions varies depending on the properties of the reactants particles (atoms, molecules, or ions) with varying reaction conditions.

To illustrate how chemical reactions occure and why their rates are differ. Collision theory had been submitted, the hypotheses of which are:

- 1. The chemical reaction occurs as a result of the collision of the reactants' particles, this theory assumed that the shape of the colliding particle was spherical.
- 2. The reaction rate is directly proportional to the number of collisions between the reactants particles per unit of time. Where you can say it whenever the number of collisions between the reactants particles increased the probability of the reaction occurs increased.
- 3. Not all collisions are effective and lead to the formation of products, in that the number of collisions between particles are enormous and about 1×10^{27} collisions per second between gas particles of 1 liter volume under normal conditions. Therefore, if such a large number of collisions lead to the formation of products so all the reactions will be complete at the moment of mixing of reactants. So most collisions are ineffective and does not lead to products.

To be effective, the collision must meet the following two conditions:

- 1. Collision molecules have a minimum of potential energy to overcome the power of repulsion among themselves when colliding and breaks the bonds of materials and turn them into products. This minimum energy necessary to reaction is called (activation energy).
- 2. The direction of the colliding particles is appropriate, that the particles are in an appropriate geometric space and in the right direction at colliding. This it creates the desired products. To illustrate this, consider Figure (4-4).

Exercise 4-12

What the texts below are erroneous with regard to hypotheses collision theory:

- A- Occurrence of collision between the reactant molecules is a basic condition of the reaction.
- B- All intermolecular collisions lead to the formation of products.
- C- In order for the products to be formed, the collision molecules must be in a appropriate suitable space.

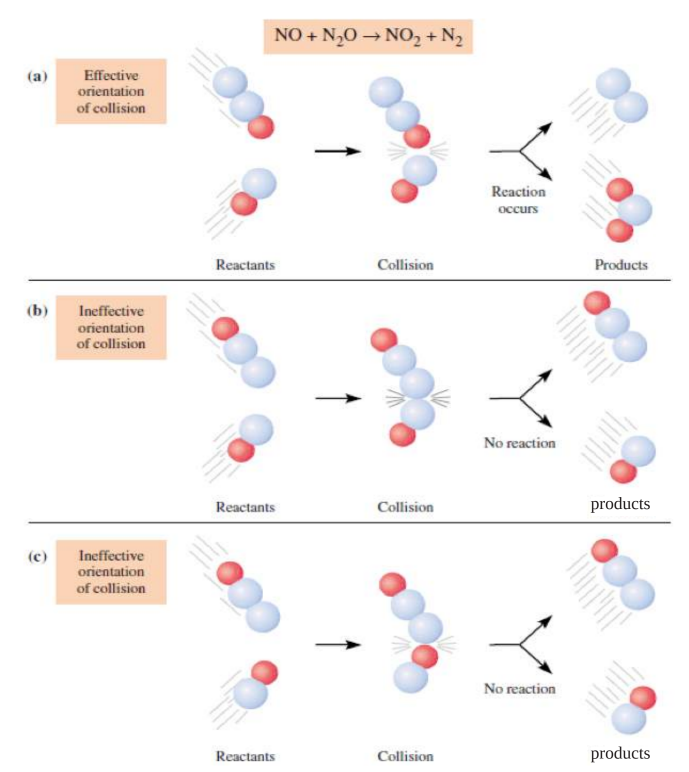


Figure 4-4

Some possible collisions between N₂O and NO molecules in the gas phase.

(a) collision that could be effective in producing the products.

(b. c) Collisions that would be ineffective orientation.

From the above we find that in order for reaction to occur according to this theory, the molecules must possess the minimum activation energy and that their collision is spatially oriented in the right direction. That's known that the internal (total) energy of any moving molecules are equal to the sum of the kinetic energy and the potential energy it possesses, and the molecules moving very quickly have high kinetic energy and low potential energy. When two moving molecules approach too fast, part of the Kinetic energy of each one will convert to potential energy, Because

of the repulsion between them is caused by the presence of electrons in their outer shells. At the moment of the collision, the two molecules stop moving and the kinetic energy of each is converted to the potential energy for collision, if that energy is less than the activation energy of the reaction will bounce off each other without an reaction (Non effective Collision) [Figure (4-4b and c)].

If they have enough kinetic energy, they will rush with great power and high speed, they can overcome the power of repulsion and entry the reaction(Effective Collision) [Figure (4-4a)].

In fact, the collision theory failed to give the true values of reactions rate when tested, the reason for the failure of this theory is to assume that the molecules are all spherical. This, in fact, applies only to a few number of molecules that are monatomic. Therefore, other scientists have proposed a new theory to explain how the reaction occurs, namely the theory of the transition state or called the theory of activated complex.

4-4-2 Transition State Theory

This theory showed that in all chemical reactions the products can not be formed directly, but must pass through the so-called state active transition, which is where formed the so-called (activated complex). It is a non-static active compound suggested that its synthesis is a compromise between the reactants and the products, so these theory are called (activated complex theory) and it is also in equilibrium with the reactants [Figure(4-5)]. The energy of this compound is always greater than that of the reactants and the products. It should be noted that the activated complex is often inseparable but can disintegrate either to produce products or to give reactants according to reaction conditions.

An important example to illustrate the idea of collision theory and transition state theory is the reaction between the I⁻ ion and methyl chloride CH₃Cl.

$$I^- + CH_3Cl \longrightarrow CH_3I + Cl^-$$

This reaction occurs according to the method shown in [Figure (4-6)]. The reaction begins with the collision of the iodide I ion with the methyl chloride molecule CH₃Cl.

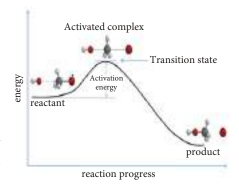
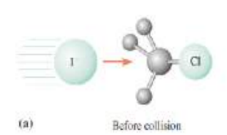


Figure 4-5

Reaction progress according to the theory of transition state.

From behind the C-Cl bond through the center of the three hydrogen atoms, where the bond begins between carbon and iodine while elongating and weakening between carbon and chlorine and the transition state of the reaction (the activated complex) is formed. Activated complex has three standard C-H bonds and two weak bonds (partial bonds) between carbon and Iodine C-I, carbon and chlorine C-Cl. As the link strength increases between carbon and iodine, the bond is established between them, while the bond breaks down between carbon and chlorine and reaction is done [Figure (4-6)a].

If the collision between the iodide ion and the methyl chloride molecule is in the wrong direction, the reactants remain unchanged. The reaction doesn't occurs, note the [Figure (4-6)b].



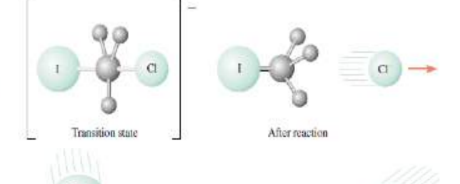


Figure 4-6

Some types of molecular collisions in the gaseous state.

- A- Effective collision leading to the formation of products.
- B- Collisions are ineffective because the direction of the colliding molecules is incorrect in addition to lack of energy Sufficient for an effective collision.

4-5 Heat of reaction

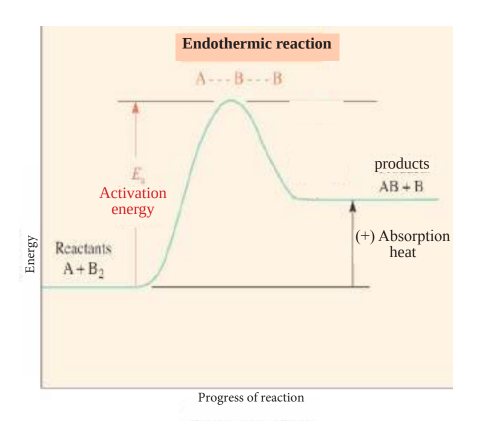
(b)

The rate of the chemical reaction is related to the activation energy, wherever the activation energy value is large, the reaction is slow, and the reaction is fast as the activation energy value is small.

The chemical reaction is accompanied by a change in energy due to the absorption or emission of a quantity of heat. This quantity represents the difference between the energy of the products and the energy of the reactants. They are called reaction temperatures and are calculated as follows:

Heat of reaction = Energy of products - Energy of reactants

The reaction is endothermic when the reaction temperature is a positive value, ie the energy of the products is greater than that of the reactants. As shown in Figure (4-7).



If the amount of reaction temperature is negative value, that is energy of product < energy of the reactants, so the reaction is exothermic. As shown in Figure (4-8).

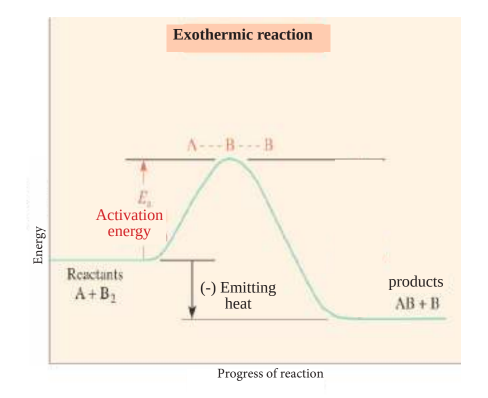


Figure 4-7

The following reaction scheme

$$A + B_2 \longrightarrow AB + B$$

When the energy of the products is greater than the energy of the reactants the reaction will be endothermic so signal heat value is (+)

Figure 4-8

The following reaction scheme

$$A + B_2 \longrightarrow AB + B$$

When the energy of the products is smaller than the energy of the reactants the reaction will be exothermic, so signal heat value is (-)





Figure 4-9

Increase in the speed of ignition of the splinter by increasing the concentration of oxygen.

(right) ignition of the splinter in the air (Left) burning flame glow inside a bottle filled with oxygen gas.



Figure 4-10

Phosphorus has two forms. White Phosphorus (above) ignites and burns quickly when exposed to oxygen air therefore, it should be stored under water.

Red phosphorus (under) reacts very slowly severe with air so it can be stored with open bottles.

4-6 Activation Energy

Activation energy is defined as the minimum energy required that the reactants must possess them in order for the collision to be effective.

4-7 Factors affecting reaction rate

Reactions are vary in rate, some slow and others fast under the same conditions, the reaction rate can be changed by controlling the factors affecting the reaction rate, namely:

5-7-1 Concentration

We observed from our study of the reaction speed law that increasing the concentration of a reactant often increases the reaction speed. Figure (4-9) shows effect of increasing oxygen concentration on reaction rate where the burning fragment lights in the air, containing oxygen gas 20%, but it glow in the flame when inserted into a filled bottle with Oxygen gas is caused by an increase in the concentration of Oxygen, which leads increased ignition rate.

It also the concentration has the effect of increasing the rate of reactions involving on reactants in the liquid state, the pressure has a similar effect on reactions involving reactants in the gaseous state. That increased pressure reduces the volume of gas and thus increases its concentration which increases of the chemical reaction rate.

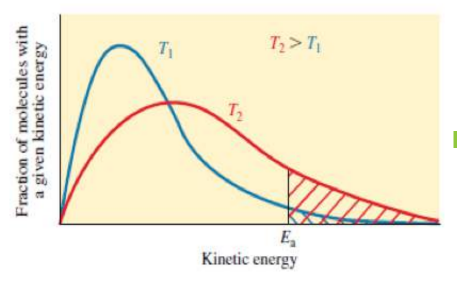
4-7-2 Physical nature and reactants nature

The rate of chemical reactions depends on the physical state of the reactants. If, for example, we had three pieces of metals, sodium, zinc and tin have the same size and we put each piece in a hydrochloric acid solution has the same concentration. We find that the sodium piece reacts strongly, the zinc piece will react at a lower rate, for the tin, it will react very slowly. So the a physical state of the reactants are very important to determine their effectiveness and rate of their reaction, as well as the effectiveness of white and red phosphorus [Figure (4-10)]

The surface area of a given mass of material increases whenever the size of the constituent particles is small, when the surface area increases, the area exposed to the reaction will increases, and the number of collisions increases, and reaction rate increases, [Figure (4-11)]. The surface area of the solid can be increased, either by dissolve it in a solvent, the solvent separates the particles from each other, or grinds it and turn it into a fine powder.

4-7-3 Temperature

The reaction rate increases as the temperature rises and it Figure 4-11 decreases when temperature decreasing. This is because high temperature increases the speed of the reacted molecules, then collision of molecules increases. So the number of molecules that has kinetic energy equal to or greater than the activation energy of the reaction increased, which can react to form the product, as shown in Figure 4-12.



Most reactions rate are doubled by 10 degrees (10°C) because of the increasing percentage of molecules involved in reaction as shown in Table 4-2.

The reaction rate doubles as the reaction temperature increases (10 °C)

Number of molecules involved in the reaction	Temperature
25 °C (298 K)	1.70 ×10 ⁻⁹
35 °C (308 K)	3.29 ×10 ⁻⁹
45 °C (318 K)	6.12 ×10 ⁻⁹



The chalk powder (pure carbonate calcium CaCO₃) reacts quickly with diluted hydrochloric acid because it has a large surface area, while the chalk finger with a small surface area reacts much slower.

Figure 4-12

Temperature increasing leads to a marked increase in the number of molecules with high energies equal to or greater than activation energy which will increase the number of collisions and the rate of reaction.

Do you know

The mixture of coal dust and air is a mixture capable of explosion at high temperature and may cause the explosion of coal mines, as well as the mixture of flour dust and air may cause an explosion for the same reason.

W

4-7-4 Catalyst

Enzymes are vital catalysts that increase the rate of reactions in the human body. When eating a meal containing protein, the digestive enzymes act on breaking down their molecules within a few hours, but without enzymes, the process takes several years.

Catalyst is a substance that increases the rate of reaction without consume through it. In most reactions the effect of adding catalyst is stronger than the effect of high temperature on reaction rate. Since the catalyst reduces the activation energy level of the reaction, so the number of colliding particles with kinetic energy is equal to or greater than the energy of activation increased, so that it can react to form the products, as shown in Figure (4-13). Note from the figure that reducing the activation energy for the reaction, leads to the opening of a new line that differs from the first line before adding the catalyst by creating a new activated complex with low energy. The effect of the catalyst on increasing the reaction rate of hydrogen with oxygen can be illustrated:

$$2H_2 + O_2 \longrightarrow 2H_2O$$

The reaction is hardly happening at the normal temperature, but is taking place quickly when adding a little platinum powder as a catalyst.

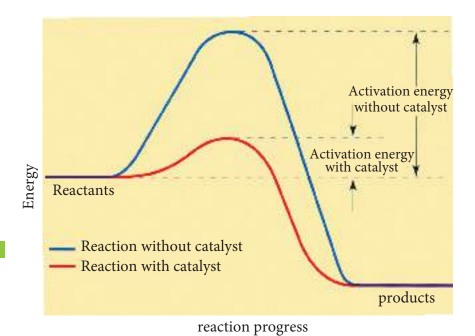


Figure 4-13

The catalyst reduces the activation energy and increases the reaction rate.

Chapter Four Questions



4-1 for the following reaction:

$$N_2O_5 \longrightarrow 2NO_2 + \frac{1}{2}O_2$$

The results in the following table:

t / hr	$[N_2O_5]$ / mol/L
0.00	0.849
0.50	0.733
1.00	0.633
2.00	0.472

Calculate the avarge of reaction rate in hour unit of the following .

A- From 0.00 to 0.50

B-From 0.50 to 1.00

C-From 1.00 to 2.00

4-2 Explain how the following factors affect chemical reaction rate

A-Temperature.

B- Surface area of reactants.

C- The nature of the reactants.

D- Concentration of reactants.

4-3 The following reaction is a first order for Br₂ reactant and it is second order for NO.

$$2NO_{(g)} + Br_{2(g)} \longrightarrow 2NOBr_{(g)}$$

A- Write the reaction rate law

B- How the reaction rate changes when the concentration of Br_2 is to be three times the initial concentration.

C- How the reaction rate changes when the NO concentration is doubled only.

D- How the reaction rate changes when the concentration of both Br₂ and NO is doubled.

4-1 for the following reaction:

 $Pt(NH_3)_2Cl_2+H_2O \longrightarrow pt(NH_3)_2(H_2O)Cl^++Cl^-$ The results in the following table:

Reaction rate law

Rate = $K [Pt(NH_3)_2Cl_2]$

Its value $K = 0.090 \text{ hr}^{-1}$

A-Calculate reaction rate when concentration is

pt(NH₃)₂Cl₂=(0.040M,0.020M,0.010M) B-How the consumption rate changes of Pt(NH₃)₂Cl₂ with its concentration changes.

C-What is the effect of changing the concentration change of Pt(NH₃)₂Cl₂ on formation rate of Cl⁻.

4-5 What effect does the catalyst on the following:

A- Heat of reaction.

B- Activation energy.

C- Reactants energy and product energy.

4-6 The reaction below is found that the formation rate at a given time is 0.036M.s⁻¹.

Rate in terms of change in A,B and D and general reaction rate (all in unit M.s⁻¹) is

$$2A+3B \longrightarrow 4C+2D$$

A- 0.018,0.027, 0.018, 0.009

B- 0.018, -0.027, 0.018, 0.009

C- 0.072, -0.048, 0.072, 0.144

D- -0.036, -0.036, 0.036, 0.009

E- -0.018, -0.012, -0.018, -0.018

4-7 Methyl acetate decomposes in a basic media to acetate ion and methyl alcohol, according to the following equation:

$$\begin{array}{c} \text{CH}_{3}\text{COOCH}_{3(\text{aq})}^{+}\text{OH}^{-}_{\ (\text{aq})} & \longrightarrow \\ \text{CH}_{3}\text{COO}^{-}_{\ (\text{aq})}^{+}\text{CH}_{3}\text{OH}_{(\text{aq})} \end{array}$$

Reaction rate law

Rate = $K [CH_3COOCH_3] [OH^3]$

K=0.14L/mol.s

A-What is the dissociation rate of methyl acetate when concentration of both CH₃COOH₃ and OH⁻ are 0.025M.

B-What rate of appearance CH₃OH in solution.

4-8 The following reaction

$$2NO_{(g)} + 2H_{2(g)} \longrightarrow N_{2(g)} + 2H_2O_{(g)}$$

Exp. No.	$[H_2]$	[NO]	Rate mol/L.s
1	2.5×10 ⁻³	5.0×10 ⁻³	3.0×10 ⁻³
2	2.5×10 ⁻³	15.0×10 ⁻³	9.0×10 ⁻³
3	10.0×10 ⁻³	15.0×10 ⁻³	36.0×10 ⁻³

From the results of the experiments in the following table:

- A- Conclude the law of reaction rate.
- B- Calculate the value of the rate constant.
- C- Calculate the reaction rate when the concentration NO, H₂ equal to 8.0x10⁻³M
- 4-9 For the following interaction and the results of experiments in the table below:

$$C_2H_{4(g)} + O_{3(g)} \longrightarrow 2CH_2O_{(g)} + \frac{1}{2}O_{2(g)}$$

Exp. No.	$[C_2H_4]$	[O ₃]	Rate mol/L.s
1	1.0×10 ⁻⁸	0.5×10 ⁻⁷	1.0×10 ⁻¹²
2	1.0×10 ⁻⁸	1.5×10 ⁻⁷	3.0×10 ⁻¹²
3	2.0×10 ⁻⁸	1.0×10 ⁻⁷	4.0×10 ⁻¹²

A - Deduce the law of reaction rate, then calculate K value.

- B Calculate the reaction rate when the concentration of each O_3 and C_2H_4 are equal to $2.0 \times 10^{-7} M$
- 4-10 Write the law of reaction rate for the following primary reactions:

$$A- Cl_{(g)} + H_{2(g)} \longrightarrow HCl_{(g)} + H_{(g)}$$

$$B- 2NO_{2(g)} \longrightarrow N_2O_{4(g)}$$

4-11 The following reaction

$$2NO_{2(g)} + F_{2(g)} \longrightarrow 2NO_2F_{(g)}$$

From the results of the experiments in the following table:

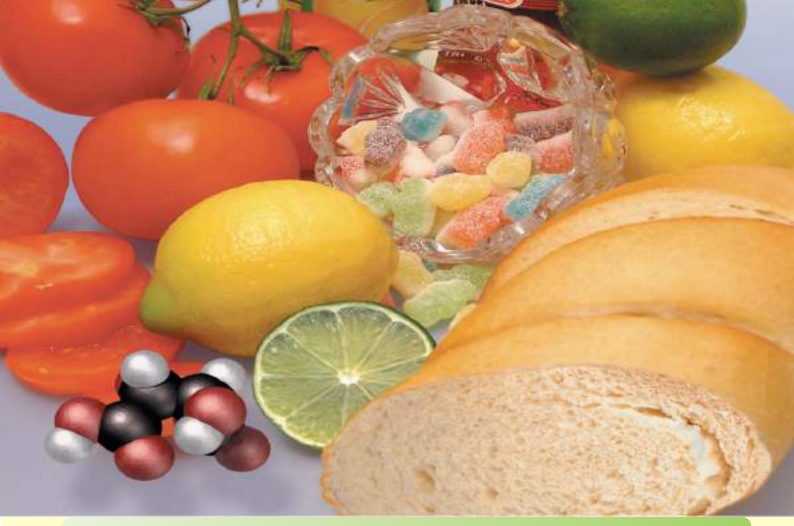
Exp. No.	[NO ₂]	$[F_2]$	Rate mol/L.s
1	0.001	0.005	2.0×10 ⁻⁴
2	0.002	0.005	4.0×10 ⁻⁴
3	0.002	0.002	1.6×10 ⁻⁴

- A- Conclude the law of reaction rate.
- B- What is the order of the reactants for each substance in the law of rate?
- C- Calculate the reaction rate constant.
- 4-12 The following reaction:

$$NO_{2(g)} + CO_{(g)} \longrightarrow NO_{(g)} + CO_{2(g)}$$

From the results of the experiments in the table below, specify the reactants orders, deduced the law of reaction rate and general reaction rate.

Exp.	[NO ₂]	[CO]	Rate mol/L.s
1	0.10	0.10	0.005
2	0.40	0.10	0.080
3	0.10	0.20	0.005



Chapter Five Acids, Bases and Salts

5

After completing this chapter, the student is expected to:

- · Recognize the properties of aqueous solutions of acids and bases.
- Identify the different molecular concepts of acids and bases according to Arrenius and Bronshted-Lauri and Lewis theories.
- Characterize the auto-ionization of water and the amphoteric properties of some substances.
- Recognize the reactions of acids and bases in aqueous solutions.
- Distinguish between the types of salts, the method of forming each and the properties of thier aqueous solutions.
- Recognize the indicator that used in neutralization reaction.
- Recognize the titration process and understand its importance.

4-1 Introduction

In nature, there are many acids, bases and salts that are used in different fields, for example, digestive juices in the human body contains a solution of hydrochloric acid at a concentration of about 0.1 molar and in the blood of human and also water components in body cells have a moderate acid action. The solution is in the car battery is composed of 40% by mass of sulfuric acid solution. On the other hand, sodium hydroxide is used in the manufacture of soap, paper industry and a number of other chemical industries. The salt also has extensive uses, baking soda is one of the salts of carbonic acid. Table salt (sodium chloride) uses to give a salty taste to food on the one hand and on the other hand it is used to preserve many types of food. Another example of the use of these substances is the application of calcium chloride salt to dissolve snow accumulated on public roads. This salt is also used in the treatment of people who has heart attacks. Ammonium salts are also used as nitrogen fertilizers to increase soil fertility. Most organic acids and their derivatives are found naturally, the vinegar we use have about 4% of acetic acid in its contents. The pain that you have feel when ants bite you is caused by the formic acid that these insects secrete into the body. Amino acids form protein modules in living organisms. Many other examples illustrate the importance and uses of these substances in everyday life.



5–2 Properties of aqueous solutions of acids and bases

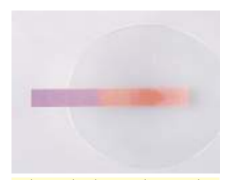
The aqueous solutions of most acids have distinctive characteristics attributed to the presence of positive hydrogen ion (H⁺) (proton) or crosses about hydronium ion (H₃O⁺) in its aqueous solution. The properties are:

- 1. Has an acidic taste.
- 2. Change the color of a number of indictors (pigments), for example change the pigment of a sunflower tint from blue color to red color
- 3. Acids react with most metals and release hydrogen gas (H₂).

- 4. Reaction with metal oxides and metal hydroxides to form salts and water.
- 5. Its aqueous solutions have the ability to conduct electrical current because of their full or partial ionization. The Table (5-1) shows some common acids.

The aqueous solutions of most bases also have distinctive characteristics, attributed to the presence of hydroxide ion (OH⁻) in its aqueous solution and qualities. The properties are:

- 1. Has a pungent taste.
- 2. It has a viscous soapy texture as in the case of aqueous solution of sodium hydroxide.
- 3. Change the color of a number of indictors, for example change the pigment of the sun flower tint from red color to blue color.
- 4. Reacts with acids (neutralize) to form salts and water.
- 5. Its aqueous solutions have the ability to conduct electrical current to their ability to be ions.



The acid solution changes the color of the pigment sunflower to red color.

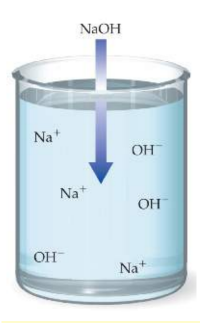


The basic solution changes the color of the pigment sunflower to blue color.

Table (5-1) Some of t	able (5-1) Some of the common acids and their uses.		
Acid name	Acid name Uses		
hydrochloric acid HCl	Purification of minerals and purification of mineral ores and preparation of some food. It is the main component of stomach acids.		
Sulfuric acid H ₂ SO ₄	Manufacture of fertilizers, explosives, glues and liquid used in the cars batteries.		
Nitric acid HNO ₃	Manufacture of fertilizers, explosives and adhesives.		
Acetic acid CH ₃ COOH	Plastic and rubber industry and in food preservation is the main component for vinegar.		
Carbonic acid H ₂ CO ₃	It is present in all carbonated drinks due to the reaction of carbon dioxide with water.		
Hydrofluoric acid HF	Clean the metal and polish the glass and engraving it.		

HCI CI CIT

Ionization of hydrochloric acid in water.



Ionization of sodium hydroxide in water.

5-3 Molecular concepts of acids and bases

In this section we will look at the molecular concepts of acids and bases according to the main theories are:

6-3-1 The Arrhenius Theory

In 1884 Arrhenius presented his theory of electrolysis dissociation, which later resulted in a theory for naming acids and bases which states on the following:

Acid is the substance that contains hydrogen which is ionized to given hydrogen ions (H⁺) in the aqueous solution.

By the definition of Arrhenius, HCl is considered to be Arrhenius acid because it produces H⁺ ions in aqueous solution:

$$HCl_{(aq)} \longrightarrow H^{+}_{(aq)} + Cl^{-}_{(ac)}$$

 $HCl_{(aq)} \longrightarrow H^{_+}_{(aq)} + Cl^{_-}_{(aq)}$ The base is the substance that contains the hydroxide group (OH-) which ionizes the data of the negative hydroxide ions in the aqueous solution.

According to Arrenius' definition of the base, NaOH is the Arenius base because it produces hydroxide ions OH in the aqueous solution:

$$NaOH_{(aq)} \longrightarrow Na^{+}_{(aq)} + OH^{-}_{(aq)}$$

 $NaOH_{(aq)} \longrightarrow Na^{_+}_{(aq)} + OH^{_-}_{(aq)}$ The process of neutralization according to this theory is defined as a union H⁺ ions and OH⁻ ions to form water molecules:

$$H_{(ag)}^+ + OH_{(ag)}^- \longrightarrow H_2O_{(l)}$$

Arrhenius theory successfully explained the acids reactions that have protons with metal hydroxides (hydroxide bases), although the application of this theory is limited, it has led to the development of more comprehensive theories to describe the properties of acids and bases.

Hydronium ion (hydrated hydrogen ion)

Arrhenius described hydrogen ions in water as H⁺ (H₂O)_n, (where n is an integer) this is because the attraction between H⁺ ions and oxygen atom carrying a negative partial charge, delta δ in the polarized water molecule. Although we do not know exactly how hydrogen ions hydrolyze in most aqueous solutions, we usually represent hydrogen ion as hydronium H₃O⁺. Thus, it is argued that the hydrated ion of hydrogen is to give aqueous acid solution its specific acidity properties. The hydrogen ion can be expressed as H⁺(aq) or hydrolyzed H₃O⁺ as follows:

$$H^+ + : \ddot{O} - H \longrightarrow H - \ddot{O} - H^+$$
 H
Hydrogen ion Water ion hydronium

5-3-2 The Bronsted – Lowry Theory

In 1923, scientists Bronsted and Lowry introduced the individually developed the theory of Arrhenius and thus emerged new theory known as Bronsted-Lowry theory.

According to this theory, acid is defined as the donor of the proton, and the base is the proton receptor.

These definitions of acid and base are highly general as any molecule or ion contains hydrogen and has the capability to release a proton is acid, while any molecule or ion can receive a proton is a base. So an acid reaction with a base can be defined as that. The reaction involves the transfer of a proton from the acid to the base. For this reason, the ionization process of hydrochloric acid (strong acid) in water, It is an acid-base reaction in which the water molecule behaves as a base because it acquires a proton. This can be expressed as follows:

First step (Arrhenius description)

$$HCl_{(aq)} \longrightarrow H^{\scriptscriptstyle +}_{(aq)} + Cl^{\scriptscriptstyle -}_{(aq)}$$

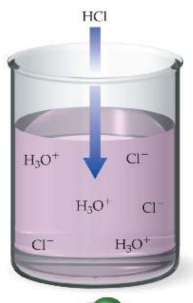
Second step

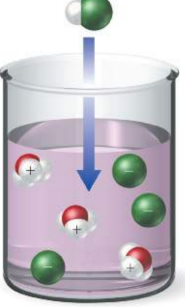
$$H_2O_{(l)} + H^+_{(aq)} \longrightarrow H_3O^+$$

Overall reaction (Bronsted-Lowry description)

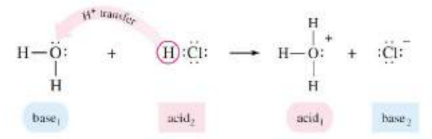
$$HCl_{(aq)} + H_2O_{(l)} \longrightarrow H_3O^+ + Cl_{(aq)}^-$$

The reactions of acids and bases can be described by the Bronsted-Lowry concept in terms of pairs (conjugate acidbase) and each pair is two different classes in a proton. In the previous equation, HCl (acid) with Cl (conjugate base) is in the sense of a pair (conjugate acid-base), while H_2O (base) with H_3O^+ (conjugate acid) ie is another pair of (conjugate base-acid) is as in the following figure:





Ionization of hydrochloric acid in water.



On the other hand, the ionization of hydrogen fluoride (weak acid). It occurs similarly but to a lesser extent and therefore represents the following:

In this equation HF is acid with F conjugate base, while H₂O represents a base with H₃O+ conjugate acid. In the forward reaction (->) HF and H₂O exhibit acid and base behavior respectively, while in the reverse reaction (\leftarrow) H₃O⁺ exhibits acid behavior (ie, a proton donor) and fluoride ion F exhibits base behavior (ie, a proton receptor). We note from the examples above that when a weak acid (HF) dissolves in water, HF molecules will produce a small amount of H⁺ ions. It can be acquired by one of the two basic classes in the solution F or H₂O. In fact, HF dissociates slightly, indicating that F is a stronger base than H₂O. When a strong acid (HCl) is dissolved in water, the HCl molecules will produce H⁺ ions that can be gained by one of the two basic classes in the solution Cl or H₂O. Since HCl dissociates completely in diluted aqueous solution, this means that Cl is lower base than H₂O. That all means that the weak acid has strong conjugate base and the strong acid has weak conjugate base. It can be generalized by saying: the greater the strength of acid, the weaker the conjugated base strength of his context and vice versa. These concepts must be used correctly; a strong acid or a weak acid (as well as a base) are qualities that you will use to describe a relative situation. When we say (in the previous example) that F is a strong base, it does not mean that it is so by comparing it with OH for example,

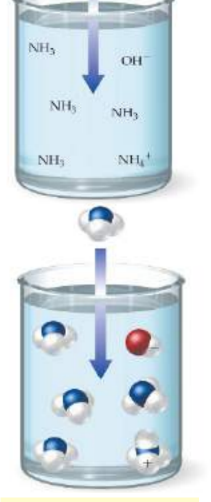
But we mean it is a stronger base relative to the conjugate base of the strong acid in the example (H₂O). In the aqueous solution of ammonia, the ammonia molecules behave as a weak Bronsted base behavior while the water molecules behave as acidic behavior and can be expressed as follows

$$NH_{3(aq)}$$
 + $H_2O_{(\ell)}$ \rightleftharpoons $NH_{4(aq)}$ + $OH_{(aq)}$ base₁ conjugated acid₁ conjugated base₂

As is evident in the reverse reaction, the ion NH₄⁺ behaves as conjugate acid of ammonia while the OH exhibits the behavior of a conjugate base of water. It is noted from the above that water exhibits acidic (proton-donor) behavior in its reaction with NH₃ while its behavior is basic (proton-receptor) at its reaction with HCl and HF, therefore, the behavior of water as acid or as a base depends on the other class contained in the solution, and this can be described that water has an amphoteric behavior and will explain this later.

5-3-3 Lewis Theory

In 1939 Lewis proposed the most comprehensive theory among other theories for the definition of acid and base. The base, according to Lewis' concept is any substance that can donate a non-bonded pair of electrons in thier chemical reactions, while the acid is any substance has a blank orbital and can accept the electronic pair of another class. This theory does not stipulate that the electronic pair must move completely from one atom to another but instead assume that the electronic pair which one atom has, becomes shared between the two atoms. So the neutralization process (acid-base reaction) identify as it is to form coordinate bond, and the reaction between trichloride boron with ammonia is an typical example of the reaction between Lewis acid-base.

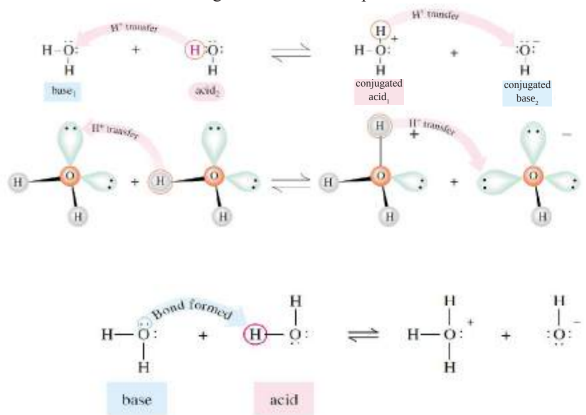


NH₂

The dissolution of ammonia in its aqueous solution.

Self-ionization of water

Careful measurements showed that pure water ionizes ever so slightly to produce equal numbers of hydronium ions and hydroxide ions. Where one water molecule donates a proton to another water molecule, and can to express this according to Bronsted concept:



The transition of the proton to the base involves the formation of a coordinate bond. It is clear that in the process of self-ionization, one of the molecules of water behaves acid and other molecule behaves as base behavior so it is said that water behavior is amphoteric behavior.

5-4 Amphoteric substances

As we already know, a certain substance can behave as acid or base behavior depending on the medium it is in. Amphoteric is general term describing the reactivity of a substance either as an acid or as a base. The amphoteric behavior describes the state in which the substance has an amphoteric property by gaining or losing a proton (H⁺). For a number of the metallic hydroxides that low soluble in water are an amphoteric property as they react with acids to form salts dissolved in water but at the same time they can dissolve by reacting with an excess of strong base. For example, aluminum hydroxide is a typical example of metallic hydroxide amphoteric as it exhibits a basic behavior by reacting with nitric acid to form salt as in the following equation:

$$Al(OH)_{3 (s)} + 3HNO_{3 (aq)} \longrightarrow Al(NO_{3})_{3 (aq)} + 3H_{2}O$$

When a solution of any strong base (such as NaOH) is added to the solid aluminum hydroxide powder, Al (OH)₃ exhibits acidic behavior and begins to dissolve, forming dissolved sodium aluminate as in the following equation:

$$Al(OH)_{3(s)} + NaOH_{(aq)} \longrightarrow NaAl(OH)_{4(aq)}$$

A number of other metal hydroxides have the same behavior, as shown in Table (5-2).

Table (5-2) Some	amphote	ric hydroxides							
metallic ion metallo	oids ion	Insoluble amphoteric	The complex ion is formed						
		hydroxide	with an excess of a strong base						
Beryllium ion	Be ²⁺	Be(OH) ₂	$[Be(OH)_4]^{2-}$						
Aluminum ion	Al^{3+}	Al(OH) ₃	$[Al(OH)_4]^-$						
Chrome ion	Cr^{3+}	Cr(OH) ₃	[Cr(OH) ₄]-						
Zinc ion	Zn^{2+}	$Zn(OH)_2$	$[Zn(OH)_4]^{-2}$						
Tin (II) ion	Sn^{2+}	Sn(OH) ₂	[Sn(OH) ₃] ⁻						
Tin (IV) ion	Sn^{4+}	Sn(OH) ₄	[Sn(OH) ₆] ²⁻						
Lead ion	Pb ²⁺	Pb(OH) ₂	[Pb(OH) ₄] ²⁻						
Arsenic(III) ion	As^{3+}	As(OH) ₃	[As(OH) ₄] ⁻						
Antimony(III) ion	Sb^{3+}	Sb(OH) ₃	[Sb(OH) ₄] ⁻						
Silicon ion	Si ⁴⁺	Si(OH) ₄	SiO_4^{4-} SiO_3^{2-}						
Cobalt ion	Co^{2+}	Co(OH) ₂	[Co(OH) ₄] ²⁻						
Copper ion	Cu^{2+}	Cu(OH) ₂	[Cu(OH) ₄] ²⁻						

5-5 Acid and base reactions in aqueous solutions

The reaction between acid and base which results salt and water is called neutralization reaction, most salts are ionic compounds. Table (5-3) shows common strong acids and bases

Table (5-3) Common strong	acids and bases							
Base	Acid							
Lithium hydroxide LiOH	Hydrochloric acid HCl							
Sodium hydroxide NaOH	Hydrobromic acid HBr							
Potassium hydroxide KOH	Hydroiodic acid HI							
Rubidium hydroxide RbOH	Perchloric acid HClO ₄							
Cesium hydroxide CsOH	Chloric acid HClO ₃							
Calcium hydroxide Ca(OH) ₂	Nitric acid HNO ₃							
Strontium hydroxide Sr(OH) ₂	Sulfuric acid H ₂ SO ₄							
Barium hydroxide Ba(OH) ₂	Chromic acid H ₂ Cr ₂ O ₇							

When any acid reacts with a base in stoichiometric will result a normal salt (equivalent). It does not contain hydrogen atoms or ionized hydroxide groups.

For example, the complete neutralization of phosphoric acid (H₃PO₄) with sodium hydroxide (NaOH) is produced a normal salt sodium phosphate (Na₃PO₄) according to the following equation:

$$H_3PO_{4(aq)} + 3NaOH_{(aq)} \longrightarrow Na_3PO_{4(aq)} + 3H_2O_{(l)}$$

If base is added less than the equivalent amount required to completely neutralize the acid, it will result acidic salts due to the ability of these salts to react with the base, as shown in the following equations:

$$H_3PO_{4(aq)} + NaOH_{(aq)} \longrightarrow NaH_2PO_{4(aq)} + H_2O_{(l)}$$

$$H_3PO_{4(aq)} + 2NaOH_{(aq)} \longrightarrow Na_2HPO_{4(aq)} + 2H_2O_{(l)}$$

There are many examples of this type of salts, including hydrogenated sodium carbonate (acidic), also called sodium bicarbonate (NaHCO₃).

When a polyhydroxide base (a base containing more than one hydroxide group in its chemical formula) is reacted with an equivalent amount of acid, a normal salt is formed as described in the following equation:

$$Al(OH)_{3(s)} + 3HCl_{(aq)} \longrightarrow AlCl_{3(aq)} + 3H_2O_{(l)}$$

When this type of base reacts with less amount of acid than is necessary to equivalent it, then strong base salts are formed (i.e. salts contains non-reactive OH groups (as shown in the following equations:

$$\begin{aligned} &\text{Al(OH)}_{3 \text{ (s)}} + \text{HCl}_{\text{(aq)}} & \longrightarrow & \text{Al(OH)}_{2}\text{Cl}_{\text{ (s)}} + \text{H}_{2}\text{O}_{\text{(l)}} \\ &\text{Al(OH)}_{3 \text{ (s)}} + 2\text{HCl}_{\text{(aq)}} & \longrightarrow & \text{Al(OH)Cl}_{2 \text{ (s)}} + 2\text{H}_{2}\text{O}_{\text{(l)}} \end{aligned}$$

It should be noted that the basic salts is not necessary to have a basic character, but they can react and neutralize acids as described in the following equation:

$$Al(OH)_2Cl + 2HCl \longrightarrow AlCl_3 + 2H_2O$$

5-6 Types of salts

As we learned earlier, when a solution of acid is neutralize with the base completely and accurately, the resulting solution is a salt derived from a pair acid-base. That such a situation often occurs in chemical reactions used in chemical analysis processes which will be necessary to calculate the acidic function (pH) or in other words the concentration of hydrogen ion in the solution. Salts are is a strong electrolyte, therefore salt will completely dissociated in the solution. If the acid and base are strong, the acidic value of aqueous salt solution remains constant does not change. If one or both pairs (the acid and the base that formed the salt) is weak, then the calculation of the acidic function of solution becomes more complecated, so it is appropriate to address this issue by dividing the salt into four main sections:

5-6-1 Salts derived from strong acids and strong bases reactions

Example of reaction of hydrochloric acid HCl with sodium hydroxide NaOH

The solubility of this group of salts does not affect the equilibrium process between hydrogen and hydroxide ions in water:

$$H_3O^+ + OH^- \rightleftharpoons 2H_2O$$

Therefore, the solution remains neutral.

5-6-2 Salts derived from weak acids (HA) and strong bases (MOH)

The weak acid HA reacts with the strong base of MOH as follows:

$$HA + MOH \longrightarrow MA + H_2O$$

The salt (MA) of this type completely dissociated in its aqueous solution:

$$MA \longrightarrow M^+ + A^-$$

As is known, the aqueous solution also contains small amounts of hydrogen and hydroxide ions resulting from dissociation of water molecules. Because acid HA is weak acid, it is partially ionized, so the amount of A⁻ ions that can be present in the solution with H⁺ ions is also small, and to maintain the equilibrium state in the solution, A⁻ ions combine with H⁺ ions to form the weak acid HA

$$H^+ + A^- \rightleftharpoons HA$$

The only source of hydrogen ions in the solution is the dissociation of more water molecules. That the continued dissociation of water molecules will produce quantities of hydroxide ion and hydrogen ion consumed from the solution to form the weak acid HA. All that leads finally to increase in the concentration of hydroxide ions in the solution at the expense of the concentration of hydrogen ions, and the solution becomes basic. This condition in which an ion (or ions) of salt reacts with water ions is called hydrolysis process. The hydrolysis process of a salt derived from a weak acid and a strong base can be expressed by the following equation:

$$MA + H_2O \implies HA + MOH$$

The property of the solution depends on the relative strength of the acid and the base resulting from the hydrolysis process, an example of this type of salts is sodium acetate (CH₂COONa) and format potassium (HCOOK).

5-6-3 Salts derived from strong acids (HA) and weak bases (B)

This type of salts is formed according to the following equation:

$$HA + B \longrightarrow (BH)A$$

This salt is fully ionized in the aqueous solution:

$$(BH)A \longrightarrow BH^+ + A^-$$

In the aqueous solution of salt (BH) A, the concentration of BH⁺ ion (formed by the complete dissolution of the salt) decreases, due to its association with the hydroxide ion (resulting from the dissolution of water molecules) to form the weak base B and reach to the equilibrium state, as in the following equation:

$$BH^+ + OH^- \rightleftharpoons B + H_2O$$

 $BH^+ + H_2O \rightleftharpoons B + H_3O^+$

Therefore the concentration of hydrogen ion in the solution will increase at the expense of the hydroxide ion (consumed to form the weak base), so the solution becomes acidic.

An example of this type of salts is ammonium chloride salt (NH₄)Cl and ammonium sulfate salt (NH₄)SO₄.

5-6-4 Salts derived from weak acids (HA) and weak bases(B)

In the aqueous solution of this type of salt, the following two instantaneous reactions occur:

$$A^{-}+ H_2O \rightleftharpoons HA + OH^{-}$$

 $BH^{+}+ H_2O \rightleftharpoons B + H_3O^{+}$

The property of the acidic or basic solution depends mainly on this case on the relative strength of acid and base (whichever is stronger than the second). If they are the same strength (acid dissociation constant = base dissociation constant) the aqueous solution of salt is neutral. If the acid dissociation constant is larger than the base dissociation constant. The aqueous solution of the salt is acidic and vice versa.

Information

Dissociation constant of weak acid and base means (ability of dissociation of weak acid and weak base to produce their ions in the solution).

5-7 Acid and bases indicators

Acid and base indicators are organic dyes (organic compounds), the color of which depends on the concentration of H₃O⁺ ions in the solution. The color of the indicators shows the amount of acidity or basic of the solution. The first indicators that used for this purpose was plant dyes such as sunflower (Litmus), but most of the indicators used now is laboratory-made materials. Most of the indicators for acid-base reactions is weak organic acids (symbolized as HIn), which have a color for its non-dissociated formula differ than the color of dissociated formula In⁻. For example, the non-dissociated blue bromophenol dye indicator (is a very weak organic acid) have yellow color, while the dissociated dye have a blue color as shown in the following equation for the indicator HIn:

$$HIn + H_2O \longrightarrow H_3O^+ + In^-$$

Color 1 yellow Color 2 blue

The color of the solution is determined by the ratio of the quantities of both dissociated In and non- dissociated HIn available in the solution. A type of indicator used in neutralization reactions is called general or comprehensive indicators (consisting of a combination of a number of acid-base indicators). These indicators show a gradient in color for a wide range of acidic function.

5-8 Acid and base solutions

Aqueous solutions of acids and bases are considered to be the most important solutions. These solutions are vitally important, as the biological reactions that occur within our bodies take place in aqueous acidic or basic media, and any difference, even if a slight concentration of these solutions may alter these reactions, it may result in impaired physiology. Acids and bases prepare in the laboratory in different concentrations according to what we have already learned about the preparation of solutions previously. For example, a solution of hydrochloric acid (HCl) can be prepared by dissolving hydrogen chloride gas HCl in water, and solutions of sodium hydroxide can also be prepared by dissolving solid NaOH in water. The preparation process each time is done by dissolving a known quantity (mass or number of moles) of the solute, and precisely, in a specific amount (mass or volume) of the solvent according to calculations made for this purpose.







Figure 5-1

The colors of the three solutions are common indicators at values for the acidic function of the solution (pH) ranging from 3-11 (A) red methyl. (b) blue bromothymol. (c) phenolphthalein.

Example 5-1

Calculate the molarity of a solution containing 3.65 g HCl in 2 L of solution (molar mass 36.5g/mol).

Solution

According to the law of the molar

$$M = \frac{n \text{ (mol)}}{V \text{ (L)}}$$

Calculate the number of moles of HCl from the following relationship:

relationship:

$$n_{HCl} = \frac{m(g)}{M(g/mol)} = \frac{3.65(g)}{36.5(g/mol)} = 0.1 \text{ mol}$$

So molarity is equal to:

$$M = \frac{n \text{ (mol)}}{V \text{ (L)}} = \frac{0.1 \text{ (mol)}}{2.0 \text{ (L)}} = 0.05 \text{ mole/L} = 0.05 \text{ M}$$

Example 5-2

Calculate the mass of Ba(OH)₂ (molar mass = 171 g / mol) needed to prepare 2.5 L of barium hydroxide solution at a concentration of 0.06 M.

Solution

Based on the law of molarity:

$$M (mol/L) = \frac{n (mol)}{V (L)}$$

From the definition of moles numbers

$$n_{HCl} = \frac{m(g)}{M(g/mol)}$$

Combining the two relationships above we get:

$$M \text{ (mol/L)} = \frac{\frac{m \text{ (g)}}{M \text{ (g/mol)}}}{V \text{ (L)}}$$

and then we get the mass of Ba(OH)₂

$$m(g) = M(mol/L) \times V(L) \times M (g/mol)$$

$$m(g) = 0.06 \text{ (mol/L)} \times 2.5(\text{L}) \times 171.3(\text{g/mol})$$

$$m(g) = 25.695 g$$

Exercise 5-1

Calculate the molarity of concentrated sulfuric acid solution in one liter (molar mass = 98 g/mol) if you know the mass percentage of the acid in the solution is 96.4% and the mass of the solution is equal 1.96 kg / L.

5-9 Titration

Laboratory (acid-base) reactions are usually used to measure the volume of a solution (known concentration) to react with a certain volume of another solution with unknown concentration and then the measured volume is used to calculate the concentration of the first solution, this process is called titration. Titration is the process in which the solution of one of the two reactants is added gradually from a cylindrical glass instrument inserted in such a way that it can accurately measure the volume of the solution and it is called the burette, to a solution of the other reactant typically found in a conical flask called a (Erlenmeyer flask) until the reaction between the two substances is complete. Then measured the volume of the solution that should added to complete the reaction.

To determine the point at which the titration process is stopped (the point at which the reaction between the acid and the base ends) is called the (end point) or the actual equivalence point where a substance color-changing is added that its color-changing at this point called the indicator.

For example, an acid solution with unknown concentration in the conical flask can be titrate by gradual addition of a standard sodium hydroxide solution (with a known concentration) from the burette [Figure (5-2)] and using the phenolphthalein indicator.

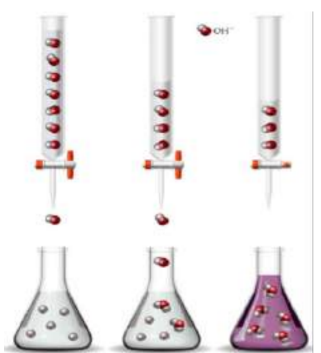






Figure 5-2

Titration of a solution of unknown concentrated acid against a standard solution of the base where the reaction endpoint is determined when the color of the added indicator to the titrated solution changes.

Example 5-3

What is the molar concentration of an acid solution for hydrochloric acid if know that 36.5 mL of it was necessary to react with 43.2 mL of 0.236M of sodium hydroxide solution?

Solution

The reaction equation shows that one mole of acid is equivalent to one mole of the base, or one mlimole of acid is equivalent to one mlimole of the base, so it calculates the number of mlimoles of sodium hydroxide involved in the reaction firstly. Then it can be calculated the molar of hydrochloric acid concentration because the volume of the acid solution is known.

Information

$$V_{HCl} = 36.5 \text{ mL}$$

$$M_{HCI} = ?$$

$$V_{NaOH} = 43.2 \text{ mL}$$

$$M_{NaOH} = 0.236 M$$

Using the dilution law of the solution

$$M_{HCl} \times V_{HCl} = M_{NaOH} \times V_{NaOH}$$

$$M_{HCl} \times 36.5 \text{ mL} = 0.236 \text{ M} \times 43.2 \text{ mL}$$

$$M_{HCl} = \frac{0.236 \text{ M} \times 43.2 \text{ mL}}{36.5 \text{ mL}} = 0.278 \text{ M}$$

Chapter Five Questions



- 5-1 How did Arrenius defind the acid, base and neutralization process? Explain this by example.
- 5-2 Using the Bronshted and Laurie theory, explain what is meant by each of the following with example:

A- acid B- conjugate base C- base D- conjugate acid

E- pairs of conjugate acid-base

5-3 Distinguish between acids and bases in each of the following using the theory of Lewis, indicating the donor and acquired of electron pair:

 $H_2O + H_2O \longrightarrow H_3O^+ + OH$ $HCl + H_2O \longrightarrow H_3O^+ + Cl^ NH_3 + HCl \longrightarrow NH_4^+ + Cl^-$

5-4 What is acid salt? Write balanced chemical equations to show how to get the following acidic salts from appropriate acids and bases:

A- NaHSO₃ B- KHCO₃ C- NaH₂PO₄ D- NaHS

5-5 Explain why

A- Salts solutions derived from strong acids and strong bases will be neutral.

B- Salts solutions derived from strong acids and weak bases are acidic.

C- Salts solutions derived from weak acids and strong bases are basic.

5-6 Calculate the molarity of the nitric acid solution if you know that 35.7 mL from it neutralized, in titration process, 0.302g from Na₂CO₃.

5-7 How much the volume of concentrated HCl acid(Its mass percentage = 36% and its density = 1.18 g/mL) is required to prepare 500 mL of dilute solution of the same acid at concentration of 2 M?

- 5-8 Calculate the mass of HNO_3 contained in 5 mL of concentrated acid solution (Its mass percentage = 69.8% and its density = 1.42 g / mL).
- 5-9 How much ammonium sulfate salt $(NH_4)_2SO_4$ (molar mass = 132.8 g/mol) required to prepare 400 mL of solution in a concentration of 0.25 M?
- 5-10 How much the volume of concentrated sulfuric acid(Its mass percentage = 98% and its density =1.84 g / mL) is required to prepare 100 mL of diluted acid solution(Its mass percentage 20% and its density = 1.14 g / mL).
- 5-11 Put (\checkmark) in front of the correct statement and a sign (\times) in front of incorrect statement as follows:
- 1. All Bronshted-Laurie acids and bases are considered Arrhenius acids and bases.
- 2. Ammonia reacts with water as Bronshted base.
- 3. The aluminum chloride (AlCl₃) is an acid Louis.
- 4. The conjugate base of strong acid behave as strong base behavior.
- 5. Boron trifluoride is considered Lewis acid.
- 6. Which of the following statements relates to Arrhenius theory are correct?
- 7. Acid-base reaction according to Bronshted-Lauri theory includes protons transmission.
- 8. One of the essential requirements for the Bronshted-Lauri statement is the presence of hydroxide ion in its chemical formula.



Chapter Six Polymer Chemistry

6

After completing this chapter, the student is expected to:

- Recognize on molecules (polymers) as large molecules built from small units called monomers.
- Understand polymerization of monomers process for the formation of polymers.
- Recognize some types of natural and synthetic polymers.
- Explain the effect of different general structural composition of polymers on their properties.
- Specify the types of polymers (polymers of addition and condensation polymers) and how they are prepared in laboratory.
- Describe the formation of polymers by addition polymerization as well as by condensation polymerization .
- Recognize rubber (as a type of polymer) and distinguish between natural rubber
- and synthetic polymer.
- Know some of common uses of synthetic polymers like plastics synthetic fibers.

6–1 Polymers (plastics)

Plastics, synthetic fibers, rubber, cellulose and proteins, the clothes we wear all these materials are polymers, starch and fatty are other examples of polymers. The word polymer consists of two segments, poly the Greek word means multi and the word mer is a molecule, and so on, polymer means polymorphic(multi-molecules). Polymers are huge molecules composed of a large number of small molecules called monomers. These monomers were connected by mostly covalent bonds in the form of long chains according to organic chemical reactions. One of the polymers contains approximately 1000 to 200,000 atoms, mostly carbon and hydrogen. The monomers are small molecules that may be organic or inorganic compounds that combine with each other by chemical reaction to form huge chains, which is polymer. A polymer consisting of units of these duplicate molecules is called duplicated unit, and the reaction process of these monomers of the formation of polymers is called polymerization process. So polymerization is a bonding process of small molecules called monomers come together to form huge molecules called polymers. If the monomer is denoted by the letter M and the number of monomers is n (n values are between 5 and 50,000), the polymerization process can be represented to form polymer (M)n as in the following equation:

$$\begin{array}{c} \text{Polymerization} \\ nM \xrightarrow{process} & \longleftarrow M \xrightarrow{n} \\ \text{n monomers} & \text{polymer with n duplicate units} \end{array}$$

6–2 Polymers types

Polymers were initially classified into two types:

6-2-1 Natural Polymers

They are found in nature such as starch, cellulose, proteins and natural rubber.

6–2–2 Synthetic Polymers

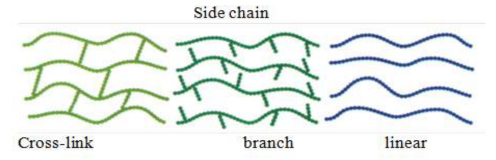
Which was prepared laboratorial or synthetically by human such as polyethylene, polyamide (nylon), polyvinyl chloride and other thousands of polymers are known nowadays. Polymers may be classified according to the effect of temperature on:

A-Thermoplastic polymers

This type of polymer is softened when it is heated where it can be reformed several times and then it hardens at a low temperature. These polymers are named plastics. The most famous examples are: polyethylene, polypropylene, polyvinyl chloride and polystyrene.

B-Thermosetting polymers

Do not soften when heated, but maintain its original shape. One of the famous example melamine, synthetic rubber. Polymers may also be classified according to their structural composition and effect of heating on, it is either linear or branched or crosses - linked as shown in Figure 6-1. When linear polymer is heating the molecules are free to move and slide easily forward and backward on top of each other as they are thermally unstable. As for heating the branching polymer and its molecules contain side chains that prevent the particles slide off each other easily but are likely still not thermally stable. The cross-link polymer has molecules adjacent are interrelated so it cannot slide over each other by heating. It remains in this state to maintain its shape and to be fixed thermally.



6–3 Synthetic polymers

synthetic polymers (which are made by Human) can be made by two types of polymerization, namely:

6–3–1 Addition Polymerization

A chemical reaction in which too many small unsaturated molecules (monomer containing double bond) are added to form one huge molecule, which is a polymer without any by-product (those polymers may be made of one type monomer). Examples of addition polymers are polyethylene (PE), polyvinyl chloride (PVC), polystyrene (PS) and polypropylene (PP), and we will look at the methods of preparation and compositions in a brief summary.

Figure 6-1

Types of polymer according to their structural composition.

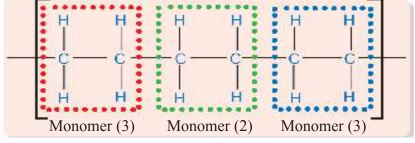


1. polyethylene (PE)

Polyethylene is prepared from the addition of ethylene molecules(CH2=CH2) for some of them to be a polyethylene molecule. As shown in the following equation:

The letter (n) represents the number of polymerized ethylene molecules which may be a number repeated units of approximately 50,000 or more. The addition reaction is repeated many times to form a polymer with length n of monomers (very long molecular chain). This reaction can be repeated hundreds or even thousands of times, the ethylene unit in the polymer chain is called the repeated unit, where one of the two bonds breaks between the two carbon atoms in the ethylene molecule (\square bond). Each atom is bound to a carbon atom of another molecule, and this process is repeated until thousands of ethylene molecules are bonded to form polyethylene as follows:

This equation can be written briefly as in the previous equation and figure 6-2 shows how the number of units of a molecule is known polymer (polyethylene), consisting of three repeated units.



Forms of ethylene

Polyethylene is usually prepared in various forms according to the conditions of the polymerization reaction. There are three forms of polyethylene according to temperature, pressure and catalysts used in the preparation:

- -

Figure 6-2

Number of units of polymer molecule (polyethylene).

A-High-Density polyethylene (HDPE)

It is formed when ethylene is heated to 100 ° C in a hydrocarbon solvent at normal atmospheric pressure a high density linear polymer is formed. Because linear molecules are very closely aligned, their size is small this makes the density is high, so the polymer remains strong and solid, making it plastic containers such as milk containers and juices.

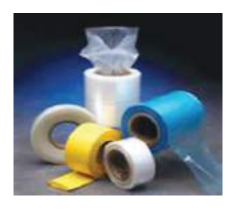
B-Low-Density polyethylene (LDPE)

It is formed when ethylene is heated to 200 ° C at high pressures (approximately 1000 atm) with little oxygen as the initiator of the reaction. Branched polymer is formed when removal of hydrogen atoms from the molecule and the addition of ethylene molecules instead in these sites, unlike the linear molecules because their molecules cannot convergence. So the density of branching chain is less than the density of the linear chain. So the polymer is less rigid than high density polyethylene and therefore used in making ordinary plastic bags.

C- Cross-linked polyethylene (CPE)

This type of polyethylene can be obtained when removed hydrogen atoms of polyethylene molecules, where two adjacent molecules in the chain are bonded and form an cross-link bonding between two molecules. The cross-type is known to be stiffer and stronger than high-density polyethylene, making things that require extreme stiffness. Figure (6-3) shows the different forms of polyethylene with their different uses.





Do you know __

The polymer chain is in adhesive tapes. low density polyethylene with a value of n about 10,000 units each series has an average molar mass of about 300,000.

Figure 6-3

Properties of various forms of polyethylene and some special uses to them.

Figure 6.4 shows some of the different uses of polyethylene.



Figure 6-4

Different uses of polyethylene.

2. polypropylene (pp)

It forms of the union of a large number of propylene molecules(CH₂=CH-CH₃) with catalysts in the following equation written briefly as follows:

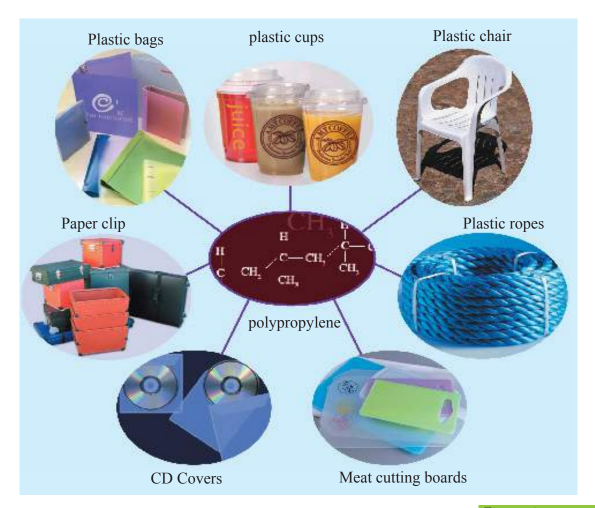
n
$$\begin{pmatrix} CH_3 & H \\ C = C \\ H \end{pmatrix}$$
 $\xrightarrow{\text{Catalysts}}$ $\begin{pmatrix} CH_3 & H \\ C - C \\ H & H \end{pmatrix}$ Propylene Propylene

This process can be repeated until thousands of propylene molecules bind to the polypropylene as follows:

Polypropylene has the following characteristics:

- 1. Easily formed, poured and resistant to heat and chemicals.
- 2. Unbreakable
- 3. Transparent and odorless
- 4. Used in the manufacture of medical instruments, toys, tubes and Figure (6-5)

Shows some different uses of polypropylene.



3. Polyvinyl chloride (PVC)

It consists of the bonding of a large number of vinyl chloride molecules (CH2 = CH-Cl) with catalysts, as the following simplified reaction:

n
$$\begin{pmatrix} H & CI \\ H & H \end{pmatrix}$$
 Catalysts $\begin{pmatrix} H & CI \\ C & C \end{pmatrix}$ Heat $\begin{pmatrix} H & CI \\ C & C \end{pmatrix}$ Polyvinyl chloride

PVC has the following qualities:

- 1. More durable, heat resistant and chemicals than polyethylene and propylene.
- 2. Cheaper
- 3. It is used instead of metal pipes in the manufacture of water pipes.
- 4. Water resistant and insulator so it enters the car brushless and raincoats industry.

Figure (6-6) show some uses of polyvinyl chloride.

Figure 6-5

Some different uses of polypropylene.



Molecular structure of vinyl chloride



Figure 6-6

Various uses of polyvinyl chloride.

4. Polystyrene (PS)

Polystyrene consists of the polymerization of a polymer molecule called styrene with the presence of suitable catalysts and the structural formula of styrene is as follows:

Polymer styrene to form polystyrene according to the following formula:

Polystyrene is a white solid that is easy to form and resist for acids and alkalis, used in the manufacture of synthetic sponge (cork), insulators, pipes, some household utensils and vegetable containers. Figure 6-7 illustrates some uses of polystyrene.

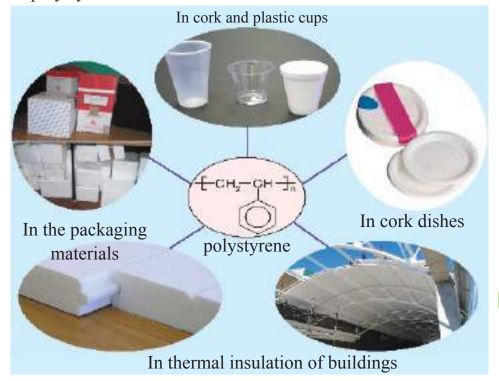


Figure 6-7
Different uses of polystyrene.

Cork is made of polystyrene, which is small balls as shown in Figure (6-8). It size varies depending on the type of cork and there is a monolith together. When it exposed to flame in an atmosphere saturated with oxygen (at least 30%), it ignites and because of this characteristic cork is used in the manufacture of high-temperature resistant materials, note Figure (6-9), which made polystyrene to compete with many other polymers prepared.

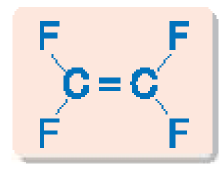




Figure 6-8

Cork is a small balls.

Figure 6-9 Cork resistance to heat in the normal atmosphere.



Structural formula of tetrafluoroethylene.



Figure 6-10
Frying pan coated with Teflon.

5. Poly tetrafloroethylene (Teflon)

This polymer is prepared from the tetrafluoroethylene monomer ($CF_2 = CF_2$), note the structural formula adjacent to the polymer.

Where molecules bonded to this monomer in the form of a long chain to form a substance known as Teflon. This substance contains a non-reactive and stable carbon fluorine bond at temperature is 325 °C. It has a very low friction coefficient and this it means that the material slides easily off the surface, making it an important material in the manufacture of parts of heat-resistant machines in addition to use in the manufacture of kitchen utensils that do not stick food (Tefal) Figure (6-10).

Teflon has the following characteristics:

- 1. Its extreme resistance to heat and chemicals
- 2.More fixed than any polymer material, whether natural or manufactured.
- 3. It does not burn and is not corroded by weathering. The most important uses are:
- 1. In the manufacture of tools exposed to heat.
- 2. In coating cooking utensils to prevent food sticking to them.
- 3. In the manufacture of heat resistant clothing.

Note from the examples above that added polymers are all from hydrocarbons, which they are very similar in form and chain the method of composition, differ in terms of groups that are related to the carbon atom; for example, the hydrogen atom in ethylene is replaced by the methyl group -CH₃ as in propylene, chlorine atom -Cl as in vinyl chloride, and the phenyl ring -C₆H₅ as in styrene. The four atom of hydrogen may be replaced by fluorine atoms -F as in tetrafluoroethylene. This causes a difference in the qualities of plastics produced from them.







Table 6-1 Structures of monomers and refined units of addition polymers and their names												
Monomer name	The repeated unit form in the polymer	Polymer name	Monomer form	uses								
Ethylene	$\left(\begin{array}{c} H \\ C = C \\ H \end{array}\right)$	Polyethylene	H H - C - H H	plastic bags «slices» strips .etc.								
Propylene	$(H)^{3} = C + H$	Polypropylene	CH ₃ H C - C - H H H	Glasses، Plastic								
Chloro ethylene (vinyl chloride)	$\begin{pmatrix} \mathbf{CI} \\ \mathbf{H} \end{pmatrix} = \mathbf{C} \begin{pmatrix} \mathbf{II} \\ \mathbf{H} \end{pmatrix}$	Poly chloride vinyl (PVC)	CI H C-C-T H H	Waterproof materials isolators، cylinders								
Styrene	$\begin{pmatrix} \mathbf{C}_{6}\mathbf{H}_{5} \\ \mathbf{C} = \mathbf{C}_{H} \end{pmatrix}$	Polystyrene	C ₆ H ₅ H 	Canning, roofing of buildings								
Tetrafluoroethylene	F F F	Teflon	F F C - C - F F	Non - stickutensils								

6-3-2 Condensation Polymerization

Monomers are bonded together with the removal of a simple molecule (by-product) such as water. Each polymer may contain two types of monomer molecules. It contains two functional groups which make each monomer associate with another and withdrawn a molecule as a by-product. One of the most important examples of this polymers type are polyamides (nylon) and polyester polymers (Terylene) and will show the preparation and properties of these polymers in brief.

1.Polyamide

Nylone (nylone) is the first common manufactured condensation polymer derived from New York and London. It was made of two monomers, one of which has the amine group(- NH₂) at both ends, The other monomer has a carboxyl group(-COOH) at both ends. These two monomers are easily linked together with removal molecule of water and they are formed polyamide and the bond roup between the two monomers for the nylon chain OH is called amide group. The following equation shows the nylon industry:

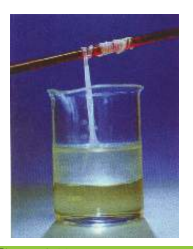


Figure 6-11

Nylon when prepared from hexagonal methylene Diamine and adipic acid and pull it in the form of filaments.

Do you know

One type of polyamide is called fiber (Kevlar) is used in armor bulletproof worn by military personnel.



Hexamethyl diamine

HO-C-(CH₂)₄-C-OH

Hexamethyl diamine

Adipic acid

$$(-N-(CH_2)_6-N-C-(CH_2)_4-C-O)_n+nH_2O$$

6.6 nylon

water

Figure 6-11 shows nylon 6,6 wrapping around the glass leg. We conclude from the equation that the reaction comprises two types of monomers: adipic acid monomer (HOOC(CH₂)₄COOH). The second monomer is hexamethylene diamine (H₂N(CH₂)₆NH₂) and the resulting polymer is called (6,6 nylon), which is to contain each carbon monomer (6) carbon atoms. This is one of the most manufactured polymers frequently used. Nylon is characterized by its low water absorption, which makes it limited to use in body contact clothes, but is also used in the manufacture of moisture resistant fabrics such as coats and umbrellas. Figure (6-12) shows the most important uses of nylon.



2. Proteins

When we mention polyamides we have to look at proteins. They are polymers that contain thousands of (H₂N-X-COOH) molecules that bind together and have the same bonding groups in Nylon, it is an amide group. It removes water from amino acids when it polymerized, so it is one of the condensation polymerization reactions. The polymerization of these acids can be represented as follows:

3. Polyesters

An example of polyesters is polyethyleneteraphatlate (PET). It is polyster is one of the most famous condensate polymers where it is prepared from the reaction of ethylene glycol (HOCH2CH2OH) with with terephthalic acid (COOH-C6H4-COOH) in the presence of catalysts according to the following reaction:

A water molecule is removed during the polymerization process and the bonding group between the repeated units in the polysters is the ester group. Polyethylene terephthalate is used in most bottles of carbonated beverages and water because it is a synthetic polymer that does not pollute these beverages.

Do you know

that the cartoons that withdraw cash and those used to mobilize phones Mobile charge and credit cards in banks and others are all made of the plastic is light, long-lasting, strong, cheap and easy to print and save magnetic information (CD) on them.

Table 6-2	Structures of condensate polymers, their repe	ructures of condensate polymers, their repeated units and some of their uses												
Polymer name	Repeated polymer unit form	Uses												
Polyurethane	$ \begin{cases} C-NH-R-NH-C-O-K-O \\ 0 & 0 \end{cases} $ As example $R_{\iota}K = -CH_{2} - CH_{2} - $	Foam filler for bedding, seats and com- position, spray insulators, spare parts for cars and industry footwear and packaging materials waterproof												
Polyethylene Terephthalate (Polyester)	$\begin{bmatrix} O - CH_2 - CH_2 - O - C - C - C \\ O \end{bmatrix}_n$	Tire: tapes: clothes and bottles and soft drinks												
6.6 Nylon (Polyamide)	$\begin{bmatrix} \text{NH} + \text{CH}_2 \\ \text{O} \end{bmatrix}_n \text{NH-C-} (\text{CH}_2)_4 - \text{C} \\ \text{O} \end{bmatrix}_n$	Home furniture, clothes, fiber carpets and fishing nets Fish												

Extraction of natural rubber from the rubber tree



Figure 6-13
Some uses of synthetic rubber.

Do you know

That the material Bakelite, the first plastic material, its was discoverer by Bakeland 1908, so attributed to him and uses bakelite in making hard tools such as Radio, television and telephone instruments.

4. Natural rubber and synthetic rubber

Natural rubber is extracted from the rubber tree (available in Malaysia and Brazil). Natural rubber is composed of a repeated unit called isoprene in its cis form (cis-isoprene): a 2-methyl 1,3-butadiene, which is polymerized in the following form:

Rubber produced from other monomers, in addition to isoprene, such as butadiene and styrene.

Note that the applications of pure natural rubber and syntheses relatively few because when it heated, individual molecules slide easily into front and back and on each other, so that the rubber become soft and viscous. It cannot be used to make needs, so sulfur is added to the rubber. It cannot be used to make needs, so sulfur is added to the rubber in its manufacturing processes turn it into a solid and strong material. Adding sulfur to rubber is called vulcanization. it is a process that leads to the entanglement of adjacent rubber molecules through sulfur atoms. Vulcanization makes the rubber usable in a wide field as rubber pipe industry, automotive tires ...etc. Note Figure (6-13). Another example is the rubber made from polymerization of Neoprene, which is 2-Chlorobutadiene with the following structural structure similar to isoprene. Note that 2-chlorobutadiene is similar to natural rubber isoprene monomer, except that the chlorine atom has replaced the methyl group in carbon atom number (2). As describe below

Chapter six Questions



- 6-1 Circle the correct t answer symbol in all of the following:
- 1. Which of the following materials is used for non-stick surfaces?
- A- Dichloro difluoromethane
- B- Polychlorofluoroethylene (Teflon).
- C- Tetrachloromethane
- D- Polyvinyl chloride
- 2. During the condensation polymerization reaction.
- A-The output becomes saturated
- B- Often forms of water
- C- Didn't form by- product
- 3. Which of the following molecules is heated with sulfur atoms during the vulcanization process.
- A- Isoprene B -Polyisoprene
- C-2-methyl 1,3butayin D-Butadiene
- 4. In the vulcanization process, what happens to the contiguous molecules when heated with sulfur atoms.
- A- branching
- B- interrelated cross-link
- C- slip over each other
- D- collide
- 5. Any polymers of the following is added polymer.
- A- Nylon B- Polystyrene
- C- protein D- terlene
- 6. Any of the following hydrocarbon molecules can polymerize it.
- $A-CH_4$ $B-C_2H_6$
- $C C_3 H_6$ $D C_4 H_{10}$
- 7. Biodegradable plastic.
- A- It burns easily and produces toxic fumes.
- B- can be absorbed through the roots of
- C-break down by bacterial decomposition.

- 8. Plastic has pollution problem because many plastics are plastic.
- A- Very flammable
- B- burns to create toxic fumes
- C- decomposes into toxic products
- 9. The polymer that does not melt when heated but it retains its original form.
- A-Thermally unstable B-Thermally stable
- C- linear D- branching
- 10. Large molecules consist of a number of units the small ones are related in organic reactions are.
- A- Monomers
- **B-** Polymers
- C- Functional groups
- D- carboxylic acids
- 11. Which of the following are small units associated to each other in a polymer during organic chemical reactions.
- A- Monomers B- Micr
 - **B-** Micropolymers
- C- branched polymers D- linear polymers
- 12. Small units that are associated during organic reactions to be large molecules.
- A- Must be similar
- B- must be different
- C- can be similar or different
- 13. Linear polymers
- A- They have freely moving molecules
- B- Have molecules that slide easily when heated
- C- (a) and (b) together
- 14. The polymers are associated in cross-link shape
- A- Thermally unstable
- B- retains its shape when it is heated
- C- have side chains.
- 15 . In an cross-link polymer, contiguous molecules.
- A- linked to by chains
- B- slide forward and back on each other

when heated

C- regular in layers that can slip apart when heated

16. Either of the following forms the two main types for polymers?

- A- Thermally unstable and thermally fixed plastic
- B- Linear and branching polymers
- C- Adding polymers and condensation polymers
- 17. A polymer is formed during a chain addition reaction among the monomers that have a bilateral bond?
- A polymer addition
- B- condensation polymer
- C- Branched polymer
- D- Cross-link polymer
- 18. In the addition reaction, monomers associated through reaction involved
- A- Identical functional groups
- B- Different functional groups
- C- Bilateral links
- 19. A polymer usually consists of two alternating monomers
- A polymers added
- **B-** Condensation polymers
- C- Branched polymers
- 20. Linear molecules are stacked together
- A- Cross-link polyethylene
- B high density polyethylene
- C low density polyethylene
- D Neoprene
- 6-2 Explain why:
- 1. Natural rubber is thermally unstable polymer.
- 2. Alkane is not suitable as a polymer for the added polymer.
- 3. Cooking handles are made of a fixed thermally polymer.
- 4. No single functional group molecule

- for a condensation reaction to form a polymer.
- 5. Branched polymer needs to be fused to quantity of heat is greater than the quantity it needs linear polymer.
- 6. Vulcanized rubber withstand higher temperatures of synthetic rubber.
- 6-3 Classify the following polymers into polymers natural or synthetic.
- 1. cellulose
- 2. nylon
- 3. proteins
- 4. Polyisoprene
- 5. Polypropylene
- 6-4 Draw the structural composition of vinyl chloride.
- 6-5 Write the polymerization reaction of vinyl chloride to form polyvinyl chloride.
- 6-6 Teflon is proven from any other polymeric material, whether synthetic or natural.
- 6-7 Define the following: vulcanization, plastics, polymers, monomers, polymers, addition polymer.
- 6-8 In a chemical reaction, two small molecules were bound, a water molecule produced what type of reaction occurred.
- 6-9 Does the reaction of addition increase the saturation of a molecule or reduces it.
- 6-10 What are the two reactions can be produced polymers.
- 6-11 What are the difference in Structural compositions of the three types of polyethylene HDPE, LDPE, CPE
- 6-12 What does the molecular structure of neoprene differs structure of natural rubber.
- 6-13 Can ethanol acid be used as a monomer in a condensation polymer. Justify your answer.



After completing this chapter, the student is expected to:

- Understand what aromatic compounds and their characteristics and identify benzene and its composition. Draw its shape and accommodates its own properties.
- Recognize the meaning of resonance in chemical compounds and identifie the resonant forms of some aromatic compounds.
- Named aromatic derivatives of benzene.
- Distinguish between additive and substitution reactions in aromatic compounds.
- Recognize electrophilic substitution reactions in benzene and some of its derivatives.
- Recognize phenols and their properties and some uses.
- Understand what are nonhomogeneous heterocyclic compounds and some examples.

Do you know

that the name of aromatic compounds is derived from the Latin word (aroma), which means fragrance or smell because these compounds have distinctive scents. Though that is a lot of these compounds are toxic and some of them are considered carcinogens as some compounds produced from tobacco smoking. Some of these compounds too of great importance in the pharmaceutical industry such as aspirin and tetracycline.

7–1 Introduction

A group of organic compounds have been discovered with a great deal of unsaturation, but they are unique in their stability and since the derivatives of these compounds have aromatic odors called aromatic compounds or aromatics. This label has lost its actual meaning and gained a theoretical significance to describe the properties of that type of unsaturated compounds.

The first compound in this group is benzene that enables the scientist Faraday to obtain it of the first time in 1825 from lighting gas. Coal tar is an important source of gasoline and large quantities is produced from petroleum hydrocarbons. This type includes organic compounds are benzene and its derivatives and similar compounds in terms of chemical composition and effectiveness, such as naphthalene and anthracene.



7–2 Benzene composition

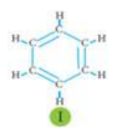
Benzene is the first aromatic compounds and its molecule consists of six carbon atoms linked together in the form of a regular hexagonal ring and each carbon atom is connected to a hydrogen atom and double bonds and single-carbon atoms alternate between carbon atoms (composition I). Or each angle in the hexagon symbolizes a carbon atom with a sphere combines with hydrogen and rotates the double and single bonds (II), and can compensate for the rotation of the double and a single by a circle inside the loop installation(III).



Schematic representation of a benzene molecule.

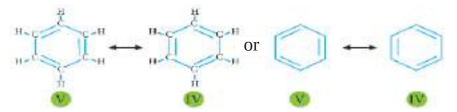






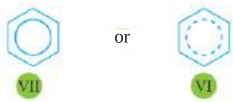
7-2-1 Resonance in benzene

In 1865, the scientific Kekule proposed the two formulas (IV) or (V) for benzene as follows

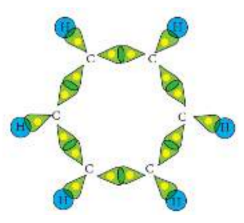


It is clear that the carbon atoms in the formulas (IV and V) occupy similar places but differ in the places assigned to the double bonds. The exchange of these links sites is called resonance (Ringing). That benzene is actually not a reciprocal state between the two states of resonance IV and V are not part of the molecules in the form of resonance and another part of the molecules in the form of the other resonance. But the real form of benzene is modified for the two shapes is called the resonance hybrid. Where we observe it's in place there is a double bond in one of the shapes, it will be single in other shape. When the average is taken, we get a hybrid form of resonance containing of six carbon -carbon bonds, which are identical and of equal length as the average case between the length of the single bond and the length of the double bond.

Therefore, the shape of benzene is usually painted in the form of a hexagonal ring, it has an inner loop (VI and VII composition) rather than in the form of successive double and double aces (IV and V composition) because it describes the situation is more accurate. However, drawing in the form of double bonds and individual successive is better in the case of monitoring the movement of electrons as in reaction mechanism state.



In brief, benzene is a molecular formula C_6H_6 containing six identical carbon atoms and six hydrogen atoms. The C-C bonds have equal lengths and it is in a median case between the length of the single bond and the length of the double bond.

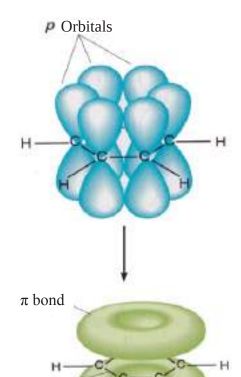


A diagram of how carbon atoms bonded in a benzene molecule.

Do you know

The C_6H_6 benzene, we are studying is completely different from gasoline used in internal combustion engines. The latter is a mixture of 5 to 10 carbon hydrocarbons.

7-2-2 Special characteristics of benzene



An electronic cloud for a π bond in a benzene molecule.

σ bond

1. Stability

It is a stable compound as indicated by its reactive properties and relative resistance to chemical changes. That most unsaturated compounds tend to engage in addition reactions in which binary or triple bonds are saturated into individual bonds. Note that cyclohexane reacts easily with the bromine dissolved in carbon tetrachloride, forming 1,2-Dibromocyclohexane, whereas benzene does not fully react under the same conditions.

$$+ Br_2/CCl_4 \longrightarrow Br_{Br}$$
 $+ Br_2/CCl_4 \longrightarrow No Reaction$

In order for benzene to react with bromine, a catalyst (FeBr3) must be used to enter a substitution reaction (substitution) and not an addition interaction where the loop remains preserved in its shape and this is evidence

on the stability of benzene.

2. Resonance Energy

To understand the energy of the resonance, we explain the following:

When a chemical reaction occurs, a change in thermal energy. For example, cyclohexane hydrogenation is a heat-emitting reaction, releasing 120 kJ energy per mole of cyclohexane.

At the hydrogenation of 1,3 cyclohexane (cyclohexane containing two π -bonds) we expect the amount of energy to be released 240 kJ per mole, twice the liberated energy at the hydrogenation of cyclohexene because there is twice the number of double bonds in the cyclohexene. The real energy measured for this reaction was 232 kJ Per mole, it is very close to expected.



According to the same logic, we expect that the energy released by hydrogenation of gasoline is three times the liberated energy at hydrogenation of cyclohexane, ie 360 kJ per mole. The actual energy of this has been found. The reaction is 208 kJ per mole and is very different to our expectation.

The difference between the expected energy value and the real energy value is 152 kJ per mole and this amount of energy is called a resonance energy. This means that benzene contains 152 kJ per mole of energy is less than expected. So that gasoline is more stable than expected by 152 kJ per mole.

3. Carbon-Carbon Bonds Length

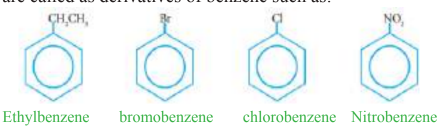
Physical measurements showed that the lengths of carbon-carbon bonds in benzene are equal and are of average length between the length of a single C-C and double C = C.

C-C	C=C	C-C
Single	double	for benzene
1.54A°	1.34A°	$1.40A^{\circ}$

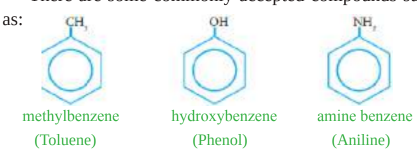
7-2-3 Benzene Derivatives Names

There are some systems in the label as follows:

1.1 Monocompensation Components: These compounds are called as derivatives of benzene such as:



There are some commonly accepted compounds such



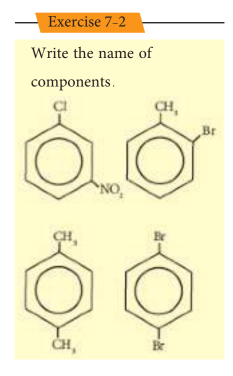
Exercise 7-1

Type the following compounds names as derivatives of benzene.



2. Dual-compensation Compounds:

When there are two groups on the benzene ring, it is not enough to mention the names of the two groups only, but they must be located on the benzene ring. Where the carbon atoms are numbered for the benzene ring and the two groups take the smallest figures, for example, 1,2 binary bromobenzene is:





The terms ortho may be used to denote the site 2 and (meta) to denote site 3 or (para) to denote on site 4 for the main compensator)

ortho X ortho meta meta

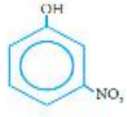
for single-compensation benzene.

If the two groups are different, they are mentioned in the nomenclature according to the alphabets one after the other and followed by the word benzene with its location at the beginning of the name or called the compound as a derivative

> 1,4 chloronitrobenzene para nitrochlorobenzene para chloronitobenzene

Where we notice in the second label we considered that the molecule of chlorobenzene is a benzene monocompensation by chlorine. In the third label it was considered that the benzene mono-compensation in nitro group, so it is the basis in the designation. Dual-compensation benzene compounds are named in this way and as in the following examples:

1,3 Nitro hydroxybenzene Meta-nitrophenol Meta-hydroxy nitrobenzene



1,2 Bromo iodide benzene Ortho iodide bromo benzene Ortho bromo iodide benzene



3. Multiple Compensation Compounds:

When there are more than two groups on the benzene ring in which case it should be numbered the locations of these groups, where the name Ortho, Barra and Meta are unacceptable. These groups are named as a derivative of benzene or as a derivative of common names if any. If the associated group are large, the compound is named in the general designation of hydrocarbons. It takes the longest hydrocarbon chain as the base of the name and benzene is named as a compensating group, where it is called Phenyl. If benzene is replaced by a group, it is called Aryl.

We will be satisfied with this stage by naming mono and dual-compensation compounds. We postpone the designation of multi-compensation compounds to advanced levels.

7-2-4 Preparation of benzene

A-In laboratory

Benzene is prepared in laboratory by heating sodium benzoate with sodium hydroxide (NaOH) in a glass distillation device where, benzene is obtained from the distillate at 80°C.

B- Industrially:

Benzene industrially is prepared in several ways, the most important of which are:

1. From phenol: Phenol is heated with Zn-dust in a distillation apparatus and then benzene is obtained from an 80°C distillate.

2. From benzene sulfonic acid: benzene sulfonic acid is heated with dilute hydrochloric acid or dilute sulfuric acid to boiling point and under high pressure according to the following equation:

7-2-5 Properties of benzene

1. Physical properties

A- Colorless liquid, flammable, has a specific aromatic smell and it is toxic.

B - boiling point 80 °C and freezing point 5 °C.

C - Its density is less than the density of water and didn't mixed with it.

D - a good solvent for non-polar organic materials such as grease, oils, resins and others.

2. Chemical properties

Benzene is a relatively stable chemical compound compared to unsaturated compounds for the presence of resonance phenomenon.

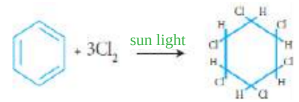
It is not affected by concentrated bases or concentrated hydrochloric acid nor by strong oxidizing agents such as potassium permanganate but suffers from a number of reactions, such as combustion and addition and substitution.

A- Combustion: Benzene is burned with a bright and flame due to the ratio of its high carbon content 92.3% and gives carbon dioxide and water with heat liberation.

B- Additive reactions: Chlorine is added to benzene in the presence of light and there is a reaction accompanied by popping the reactants leading to the formation Hexachlorocyclohexane as described in equation A:

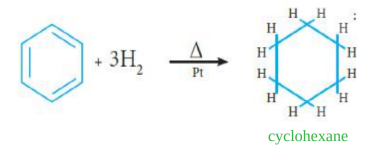


Burning of benzene with a bright and flame.



Hexachlorocyclohexane

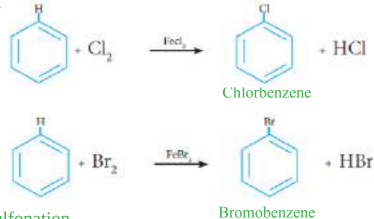
Benzene is also reduced to hydrogen at high temperatures and under high pressure in the presence of a catalyst such as platinum to cyclohexane, as in the following equation:



C-Substitution reactions (compensation): One of the most important reactions of benzene and its derivatives is the possibility of replacing (compensation) one of the hydrogen atoms by atom or different group (e.g. alkyl group R, nitro group NO₂, sulfonic group SO₃H, halide X or acetyl group $^{\text{O}}_{\text{-C-CH}_3}$ in the presence of an appropriate catalyst, which is help for reaction. Examples of substitution reactions is:

A-Halogenation

It is the process of replacing one of the hydrogen atoms with a halogen atom (such as chlorine Cl_2 or Br_2) with a catalyst such as ferric $FeBr_3$ chloride $FeCl_3$ or ferric bromide.



B- Sulfonation

It is the process of replacing one of the hydrogen atoms with the sulfonic group SO₃H, for example benzene reacts with the dark concentrated sulfuric acid at room temperature to compose benzene sulfonic acid.

C- Nitration

Replace a hydrogen atom with a nitro group NO_2 where benzene reacts with a mixture of concentrated nitric and sulfuric acid at a temperature of 45 °C to compose of nitrobenzene.

D- Friedle- Crafts reactions

1. Friedel-crafts Al-kylation: It is the process of replacing one hydrogen atom in the alkyl group (R) with an appropriate catalyst. When benzene reacts with alkyl halide (R-X) in presence of catalyst like dry aluminum chloride (AlCl₃) lead to compose of alkyl benzene

$$+ R-X$$
 $\xrightarrow{AIX_3}$
 $+ HX$
 $= alkyl benzene$

Example 7-1

Type the Friedel-crafts Al-kylation for benzene using methyl chloride (CH₃Cl) in the presence of dry aluminum chloride as a catalyst and named the produced compound.

Solution Benzene reacts with CH₃Cl in the presence of dry AlCl₃ according to the following reaction:

2. Friedel-Kraft acetylation: is the process of replacing an atom hydrogen in the acetyl group (-R-C). Benzene reacts with acetylcholine (R-C-CI) in the presence of dry aluminum chloride to compose of acetyl benzene.

Example 7-2

Type the Friedel-kraft acetylation of benzene and name the produced compound.

Solution Benzene reacts with in the presence of dry AlCl₃ according to the following reaction:

Chapter seven Questions



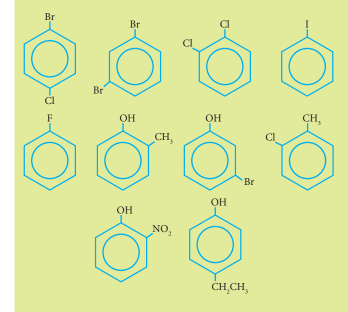
- 7-1 How did the concept of aromatic changed?
- 7-2 What is the source of aromatic hydrocarbons?
- 7-3 Why is the composition of benzene a resonance hybrid?
- 7-4 When benzene behaves as a non-saturated hydrocarbon?
- 7-5 How can you distinguish of benzene in its reactions from alkanes and alkenes? What caused the relative stability of its molecule?
- 7-6 What are the possible isomers of nitrochlorobenzene?
- 7-7 How do you distinguish between cyclohexane and benzene?
- 7-8 Write the chemical equations for the reaction of benzene with :

A- Cl₂/FeCl₃

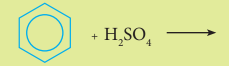
B- CH₃CH₂Cl/AlCl₃

C- CH₃COCl/AlCl₃

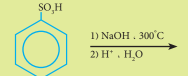
7-9 Named the following components

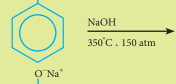


- 7-10 Draw the formulas for the following compounds:
- 1. Para-dichlorobenzene
- 2. 1, 3 bis fluorobenzene
- 3. 1, 2 iodochlorobenzene
- 4. Para bromochlorobenzene
- 5. Meta-bromo sulfonic acid
- 7-11 Complete the following equations?











- 7-12 What are the synthetic methods for preparing benzene? Write them in detail?
- 7-13 Write the composition formulas of the following compounds
- 1. 1, 3, 5 Tri-Bromo Benzene.
- 2. Meta Chloro Toluene.
- 3. Ethyl-benzene.
- 4. Para dichlorobenzene.
- 7-14 Expressed in composition formulas the reaction of benzene with ethyl chloride?
- 7-15 By chemical equation and composition formulas, write the product reduction process of benzene by hydrogen and with the presence of platinum?
- 7-16 Write the reaction of chlorine addition to benzene in presence of light?
- 7-17 Draw of the following compounds:
- A- Meta-fluoro-chlorobenzene
- B-2-Bromo-4-Chloropridine
- 7-18 Place a (\(\sqrt{} \) in front of the correct statement and (\(\sqrt{} \)) in front of the wrong statement:
- 1. Aromatic compounds are organic compounds with high nonsaturation.
- 2. The compensation reactions introduced by benzene are evidence on its high stability.

- 3. The terms Ortho, Meta and Para are used in the designation of binary and multiple benzene derivatives.
- 4. Benzene burns with a bright and smoky flame due high carbon content.
- 5. Benzene is easily reduced by hydrogen to cyclohexane.
- 6. Halogenation is the process of replacing an Hydrogen atom in benzene by sulfonic group .SO₃H
- 7. Reagents rich in electrons are called reagents looking for electrons.

			Е	lem	ent I	Eiect	roni	ic Co	nfig	urat	tions	5							
Elements									sul	olev	rels								
Elements	1s	2s	2p	3s	3p	3d	4s	4p	4d	4f	5s	5p	5d	5f	6s	6р	6d	7s	7p
1-Hydrogen	1																		
2-Helium	2																		
3-Lithium	2	1																	
4-Berylium	2	2																	
5-Boron	2	2	1																
6-Carbon	2	2	2																
7-Nitrogen	2	2	3																
8-Oxygen	2	2	4																
9-Fluorine	2	2	5																
10-Neon	2	2	6																
11-Sodium	2	2	6	1															
12-Magnesium	2	2	6	2															
13-Aluminum	2	2	6	2	1														
14-Silicon	2	2	6	2	2														
15-Phosphorus	2	2	6	2	3														
16-Sulfur	2	2	6	2	4														
17-Chlorine	2	2	6	2	5														
18-Argon	2	2	6	2	6														
19-Potassium	2	2	6	2	6		1												
20-Calcium	2	2	6	2	6		2												
21-Scandium	2	2	6	2	6	1	2												
22-Titanium	2	2	6	2	6	2	2												
23-Vanadium	2	2	6	2	6	3	2												
24-Chromium	2	2	6	2	6	5	1												
25-Manganese	2	2	6	2	6	5	2												
26-Iron	2	2	6	2	6	6	2												
27-Cobalt	2	2	6	2	6	7	2												
28-Nickle	2	2	6	2	6	8	2												
29-Copper	2	2	6	2	6	10	1												
30-Zinc	2	2	6	2	6	10	2												
31-Gallium	2	2	6	2	6	10	2	1											
32-Germanium	2	2	6	2	6	10	2	2											
33-Arsenic	2	2	6	2	6	10	2	3											
34-Selenium	2	2	6	2	6	10	2	4											
35-Bromium	2	2	6	2	6	10	2	5											
36-Krypton	2	2	6	2	6	10	2	6											
37-Rubidium	2	2	6	2	6	10	2	6			1								
38-Strontium	2	2	6	2	6	10	2	6			2								
39-Yttrium	2	2	6	2	6	10	2	6	1		2								
40-Zirconium	2	2	6	2	6	10	2	6	2		2								

	Element Eiectronic Configurations																		
Elements									sul	blev	els								
Elements	1s	2s	2p	3s	3р	3d	4s	4p	4d	4f	5s	5p	5d	5f	6s	6p	6d	7s	7p
41-Niobium	2	2	6	2	6	10	2	6	4		1								
42-Molybdenum	2	2	6	2	6	10	2	6	5		1								
43-Technetium	2	2	6	2	6	10	2	6	5		2								
44-Ruthenum	2	2	6	2	6	10	2	6	7		1								
45-Rhodium	2	2	6	2	6	10	2	6	8		1								
46-Palladium	2	2	6	2	6	10	2	6	10										
47-Silver	2	2	6	2	6	10	2	6	10		1								
48-Cadmium	2	2	6	2	6	10	2	6	10		2								
49-Indium	2	2	6	2	6	10	2	6	10		2	1							
50-Tin	2	2	6	2	6	10	2	6	10		2	2							
51-Antimony	2	2	6	2	6	10	2	6	10		2	3							
52-Tellurium	2	2	6	2	6	10	2	6	10		2	4							
53-Iodine	2	2	6	2	6	10	2	6	10		2	5							
54-Xenon	2	2	6	2	6	10	2	6	10		2	6							
55-Cesium	2	2	6	2	6	10	2	6	10		2	6			1				
56-Barium	2	2	6	2	6	10	2	6	10		2	6			2				
57-Lanthanum	2	2	6	2	6	10	2	6	10		2	6	1		2				
58-Cerium	2	2	6	2	6	10	2	6	10	1	2	6	1		2				
59-Prasedoymium	2	2	6	2	6	10	2	6	10	3	2	6			2				
60-Neodymium	2	2	6	2	6	10	2	6	10	4	2	6			2				
61-Promethium	2	2	6	2	6	10	2	6	10	5	2	6			2				
62-Samarium	2	2	6	2	6	10	2	6	10	6	2	6			2				
63-Europium	2	2	6	2	6	10	2	6	10	7	2	6			2				
64-Gadolinum	2	2	6	2	6	10	2	6	10	7	2	6	1		2				
65-Terbium	2	2	6	2	6	10	2	6	10	9	2	6			2				
66-Dysprosium	2	2	6	2	6	10	2	6	10	10	2	6			2				
67-Holmium	2	2	6	2	6	10	2	6	10	11	2	6			2				
68-Erbium	2	2	6	2	6	10	2	6	10	12	2	6			2				
69-Thulium	2	2	6	2	6	10	2	6	10	13	2	6			2				
70-Yetterbium	2	2	6	2	6	10	2	6	10	14	2	6			2				
71-Luteium	2	2	6	2	6	10	2	6	10	14	2	6	1		2				
72-Hafanium	2	2	6	2	6	10	2	6	10	14	2	6	2		2				
73-Tantalium	2	2	6	2	6	10	2	6	10	14	2	6	3		2				
74-Tungesten	2	2	6	2	6	10	2	6	10	14	2	6	4		2				
75-Rhenium	2	2	6	2	6	10	2	6	10	14	2	6	5		2				
76-Osmium	2	2	6	2	6	10	2	6	10	14	2	6	6		2				
77-Iridium	2	2	6	2	6	10	2	6	10	14	2	6	7		2				
78-Platinum	2	2	6	2	6	10	2	6	10	14	2	6	9		1				
79-Gold	2	2	6	2	6	10	2	6	10	14	2	6	10		1				
80-Mercury	2	2	6	2	6	10	2	6	10	14	2	6	10		2				