# THE p -BLOCK ELEMENTS

# **Objectives**

After studying this unit, you will be able to

- appreciate the general trends in the chemistry of p-block elements;
- describe the trends in physical and chemical properties of group 13 and 14 elements:
- explain anomalous behaviour of boron and carbon;
- describe allotropic forms of carbon;
- know the chemistry of some important compounds of boron, carbon and silicon;
- list the important uses of group 13 and 14 elements and their compounds.

The variation in properties of the p-block elements due to the influence of d and f electrons in the inner core of the heavier elements makes their chemistry interesting

In *p*-block elements the last electron enters the outermost p orbital. As we know that the number of p orbitals is three and, therefore, the maximum number of electrons that can be accommodated in a set of p orbitals is six. Consequently there are six groups of p-block elements in the periodic table numbering from 13 to 18. Boron, carbon, nitrogen, oxygen, fluorine and helium head the groups. Their valence shell **electronic configuration is**  $ns^2np^{1-6}$  (except for He). The **inner core** of the electronic configuration may, however, differ. The difference in inner core of elements greatly influences their physical properties (such as atomic and ionic radii, ionisation enthalpy, etc.) as well as chemical properties. Consequently, a lot of variation in properties of elements in a group of *p*-block is observed. The *maximum* oxidation state shown by a p-block element is equal to the total number of valence electrons (i.e., the sum of the sand *p*-electrons). Clearly, the number of possible oxidation states increases towards the right of the periodic table. In addition to this so called group oxidation state, p-block elements may show other oxidation states which normally, but not necessarily, differ from the total number of valence electrons by unit of two. The important oxidation states exhibited by p-block elements are shown in Table 11.1. In boron, carbon and nitrogen families the group oxidation state is the most stable state for the lighter elements in the group. However, the oxidation state two unit less than the group oxidation state becomes progressively more stable for the heavier elements in each group. The occurrences of oxidation states two unit less than the group oxidation states are sometime attributed to the 'inert pair effect'.

Group	13	14	15	16	17	18
General electronic configuration	ns²np¹	ns²np²	ns²np³	ns²np⁴	$ns^2np^5$	ns²np <sup>6</sup> (1 s² for He)
First member of the group	В	С	N	0	F	Не
Group oxidation state	+3	+4	+5	+6	+7	+8
Other oxidation states	+1	+2, -4	+3, - 3	+4, +2, -2	+5, + 3, +1, -1	+6, +4, +2

Table 11.1 General Electronic Configuration and Oxidation States of p-Block Elements

The relative stabilities of these two oxidation states – group oxidation state and two unit less than the group oxidation state – may vary from group to group and will be discussed at appropriate places.

It is interesting to note that the *non-metals* and *metalloids* exist only in the p-block of the periodic table. The non-metallic character of elements decreases down the group. In fact the heaviest element in each p-block group is the most metallic in nature. This change from non-metallic to metallic character brings diversity in the chemistry of these elements depending on the group to which they belong.

In general, non-metals have higher ionisation enthalpies and higher electronegativities than the metals. Hence, in contrast to metals which readily form cations, non-metals readily form anions. The compounds formed by highly reactive non-metals with highly reactive metals are generally ionic because of large differences in their electronegativities. On the other hand, compounds formed between non-metals themselves are largely covalent in character because of small differences in their electronegativities. The change of non-metallic to metallic character can be best illustrated by the nature of oxides they form. The non-metal oxides are acidic or neutral whereas metal oxides are basic in nature.

The first member of p-block differs from the remaining members of their corresponding group in two major respects. First is the size and all other properties which depend on size. Thus, the lightest *p*-block elements show the same kind of differences as the lightest s-block elements, lithium and beryllium. The second important difference, which applies only to the p-block elements, arises from the effect of dorbitals in the valence shell of heavier elements (starting from the third period onwards) and their lack in second period elements. The second period elements of p-groups starting from boron are restricted to a maximum covalence of four (using 2s and three 2p orbitals). In contrast, the third period elements of *p*-groups with the electronic configuration  $3s^23p^n$  have the vacant 3d orbitals lying between the 3p and the 4s levels of energy. Using these d-orbitals the third period elements can expand their covalence above four. For example, while boron forms only  $[BF_4]^-$ , aluminium gives  $[AlF_6]^{3-}$  ion. The presence of these d-orbitals influences the chemistry of the heavier elements in a number of other ways. The combined effect of size and availability of d orbitals considerably influences the ability of these elements to form  $\pi$  bonds. The first member of a group differs from the heavier members in its ability to form  $p\pi$  -  $p\pi$  multiple bonds to itself (e.g., C=C, C=C,

N≡N) and to other second row elements (e.g., C=O, C=N, C=N, N=O). This type of  $\pi$  - bonding is not particularly strong for the heavier p-block elements. The heavier elements do form  $\pi$  bonds but this involves d orbitals  $(d\pi - p\pi)$ or  $d\pi - d\pi$ ). As the d orbitals are of higher energy than the p orbitals, they contribute less to the overall stability of molecules than does  $p\pi$  -  $p\pi$  bonding of the second row elements. However, the coordination number in species of heavier elements may be higher than for the first element in the same oxidation state. For example, in +5 oxidation state both N and P form oxoanions :  $NO_3^-$  (three-coordination with  $\pi$  – bond involving one nitrogen p-orbital) and  $PO_4^{3-}$  (four-coordination involving s, p and d orbitals contributing to the  $\pi$  – bond). In this unit we will study the chemistry of group 13 and 14 elements of the periodic table.

# 11.1 GROUP 13 ELEMENTS: THE BORON FAMILY

This group elements show a wide variation in properties. Boron is a typical non-metal, aluminium is a metal but shows many chemical similarities to boron, and gallium, indium and thallium are almost exclusively metallic in character.

Boron is a fairly rare element, mainly occurs as orthoboric acid, (H3BO3), borax,  $Na_2B_4O_7 \cdot 10H_2O$ , and kernite,  $Na_2B_4O_7 \cdot 4H_2O$ . In India borax occurs in Puga Valley (Ladakh) and Sambhar Lake (Rajasthan). The abundance of boron in earth crust is less than 0.0001% by mass. There are two isotopic forms of boron  $^{10}{\rm B}$  (19%) and  $^{11}{\rm B}$  (81%). Aluminium is the most abundant metal and the third most abundant element in the earth's crust (8.3% by mass) after oxygen (45.5%) and Si (27.7%). Bauxite, Al<sub>2</sub>O<sub>3</sub>. 2H<sub>2</sub>O and cryolite, Na<sub>3</sub>AlF<sub>6</sub> are the important minerals of aluminium. In India it is found as mica in Madhya Pradesh, Karnataka, Orissa and Jammu. Gallium, indium and thallium are less abundant elements in nature.

The atomic, physical and chemical properties of these elements are discussed below.

### 11.1.1 Electronic Configuration

The outer electronic configuration of these elements is  $ns^2np^1$ . A close look at the electronic configuration suggests that while boron and aluminium have noble gas core, gallium and indium have noble gas plus 10~d-electrons, and thallium has noble gas plus 14~f- electrons plus 10~d-electrons cores. Thus, the electronic structures of these elements are more complex than for the first two groups of elements discussed in unit 10. This difference in electronic structures affects the other properties and consequently the chemistry of all the elements of this group.

### 11.1.2 Atomic Radii

On moving down the group, for each successive member one extra shell of electrons is added and, therefore, atomic radius is expected to increase. However, a deviation can be seen. Atomic radius of Ga is less than that of Al. This can be understood from the variation in the inner core of the electronic configuration. The presence of additional  $10 \, d$ -electrons offer only poor screening effect (Unit 2) for the outer electrons from the increased nuclear charge in gallium. Consequently, the atomic radius of gallium (135 pm) is less than that of aluminium (143 pm).

### 11.1.3 Ionization Enthalpy

The ionisation enthalpy values as expected from the general trends do not decrease smoothly down the group. The decrease from B to Al is associated with increase in size. The observed discontinuity in the ionisation enthalpy values between Al and Ga, and between In and Tl are due to inability of *d*- and *f*-electrons, which have low screening effect, to compensate the increase in nuclear charge.

The order of ionisation enthalpies, as expected, is  $\Delta_i H_1 < \Delta_i H_2 < \Delta_i H_3$ . The sum of the first three ionisation enthalpies for each of the elements is very high. Effect of this will be apparent when you study their chemical properties.

### 11.1.4 Electronegativity

Down the group, electronegativity first decreases from B to Al and then increases

marginally (Table 11.2). This is because of the discrepancies in atomic size of the elements.

### 11.1.5 Physical Properties

Boron is non-metallic in nature. It is extremely hard and black coloured solid. It exists in many allotropic forms. Due to very strong crystalline lattice, boron has unusually high melting point. Rest of the members are soft metals with low melting point and high electrical conductivity. It is worthwhile to note that gallium with unusually low melting point (303 K), could exist in liquid state during summer. Its high boiling point (2676 K) makes it a useful material for measuring high temperatures. Density of the elements increases down the group from boron to thallium.

### 11.1.6 Chemical Properties

# Oxidation state and trends in chemical reactivity

Due to small size of boron, the sum of its first three ionization enthalpies is very high. This prevents it to form +3 ions and forces it to form

only covalent compounds. But as we move from B to Al, the sum of the first three ionisation enthalpies of Al considerably decreases, and is therefore able to form Al<sup>3+</sup> ions. In fact, aluminium is a highly electropositive metal. However, down the group, due to poor shielding effect of intervening d and forbitals, the increased effective nuclear charge holds ns electrons tightly (responsible for inter pair effect) and thereby, restricting their participation in bonding. As a result of this, only p-orbital electron may be involved in bonding. In fact in Ga, In and Tl, both +1 and +3 oxidation states are observed. The relative stability of +1 oxidation state progressively increases for heavier elements: Al<Ga<In<Tl. In thallium +1 oxidation state is predominant whereas the +3 oxidation state is highly oxidising in character. The compounds in +1 oxidation state, as expected from energy considerations, are more ionic than those in +3 oxidation state.

In trivalent state, the number of electrons around the central atom in a molecule

Table 11.2 Atomic and Physical Properties of Group 13 Elements

Property		Element					
		Boron B	Aluminium Al	Gallium Ga	Indium In	Thallium Tl	
Atomic num	ıber	5	13	31	49	81	
Atomic mass(g mol <sup>-1</sup> )		10.81	26.98	69.72	114.82	204.38	
Electronic		[He] $2s^22p^1$	[Ne] $3s^23p^1$	$[Ar]3d^{10}4s^24p^1$	$[Kr]4d^{10}5s^25p^1$	$[Xe]4f^{14}5d^{10}6s^{2}6p^{1}$	
Atomic radi	us/pmª	(85)	143	135	167	170	
Ionic radius M³+/pmb		(27)	53.5	62.0	80.0	88.5	
Ionic radius M+/pm	3	-	-	120	140	150	
Ionization enthalpy (kJ mol <sup>-1</sup> )	$egin{array}{ll} \Delta_i H_1 \ \Delta_i H_2 \ \Delta_i H_3 \end{array}$	801 2427 3659	577 1816 2744	579 1979 2962	558 1820 2704	589 1971 2877	
Electronega	tivity <sup>c</sup>	2.0	1.5	1.6	1.7	1.8	
Density /g of at 298 K	cm <sup>-3</sup>	2.35	2.70	5.90	7.31	11.85	
Melting point / K		2453	933	303	430	576	
Boiling point / K		3923	2740	2676	2353	1730	
$E^{\ominus}$ / V for (M <sup>3+</sup> /M)		-	-1.66	-0.56	-0.34	+1.26	
E <sup>⊕</sup> / V for (N	/м'/M)	_	+0.55	-0.79(acid) -1.39(alkali)	-0.18	-0.34	

<sup>&</sup>lt;sup>a</sup>Metallic radius, <sup>b</sup> 6-coordination, <sup>c</sup> Pauling scale,

of the compounds of these elements (e.g., boron in  $BF_3$ ) will be only six. Such **electron deficient** molecules have tendency to accept a pair of electrons to achieve stable electronic configuration and thus, behave as Lewis acids. The tendency to behave as Lewis acid decreases with the increase in the size down the group.  $BCl_3$  easily accepts a lone pair of electrons from ammonia to form  $BCl_3 \cdot NH_3$ .

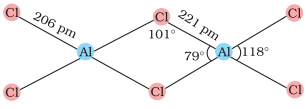
$$\begin{array}{c} Cl \\ B - Cl + NH_3 \end{array} \longrightarrow \begin{array}{c} NH_3 \\ Cl \end{array}$$

$$Cl Cl$$

Planar

Tetrahedral

AlCl<sub>3</sub> achieves stability by forming a dimer



**Tetrahedral** 

In trivalent state most of the compounds being covalent are hydrolysed in water. For example, the trichlorides on hyrolysis in water form tetrahedral  $[M(OH)_4]^-$  species; the hybridisation state of element M is  $sp^3$ . Aluminium chloride in acidified aqueous solution forms octahedral  $[Al(H_2O)_6]^{3^+}$  ion. In this complex ion, the 3d orbitals of Al are involved and the hybridisation state of Al is  $sp^3d^2$ .

#### Problem 11.1

Standard electrode potential values,  $E^{\odot}$  for  $Al^{3+}/Al$  is -1.66 V and that of  $Tl^{3+}/Tl$  is +1.26 V. Predict about the formation of  $M^{3+}$  ion in solution and compare the electropositive character of the two metals.

### **Solution**

Standard electrode potential values for two half cell reactions suggest that aluminium has high tendency to make Al<sup>3+</sup>(aq) ions, whereas Tl<sup>3+</sup> is not only unstable in

solution but is a powerful oxidising agent also. Thus Tl<sup>+</sup> is more stable in solution than Tl<sup>3+</sup>. Aluminium being able to form +3 ions easily, is more electropositive than thallium.

### (i) Reactivity towards air

Boron is unreactive in crystalline form. Aluminium forms a very thin oxide layer on the surface which protects the metal from further attack. Amorphous boron and aluminium metal on heating in air form  $B_2O_3$  and  $Al_2O_3$  respectively. With dinitrogen at high temperature they form nitrides.

$$2E(s) + 3O_2(g) \xrightarrow{\Delta} 2E_2O_3(s)$$
$$2E(s) + N_2(g) \xrightarrow{\Delta} 2EN(s)$$

(E = element)

The nature of these oxides varies down the group. Boron trioxide is acidic and reacts with basic (metallic) oxides forming metal borates. Aluminium and gallium oxides are amphoteric and those of indium and thallium are basic in their properties.

# (ii) Reactivity towards acids and alkalies

Boron does not react with acids and alkalies even at moderate temperature; but aluminium dissolves in mineral acids and aqueous alkalies and thus shows amphoteric character.

Aluminium dissolves in dilute HCl and liberates dihydrogen.

2Al(s) + 6HCl (aq) 
$$\rightarrow$$
 2Al<sup>3+</sup> (aq) + 6Cl<sup>-</sup> (aq) + 3H<sub>2</sub> (g)

However, concentrated nitric acid renders aluminium passive by forming a protective oxide layer on the surface.

Aluminium also reacts with aqueous alkali and liberates dihydrogen.

$$2 \text{Al (s)} + 2 \text{NaOH(aq)} + 6 \text{H}_2 \text{O(l)} \\ \downarrow \\ 2 \text{ Na}^{^{+}} [\text{Al(OH)}_4]^{^{-}} (\text{aq}) + 3 \text{H}_2 (\text{g}) \\ \text{Sodium}$$

tetrahydroxoaluminate(III)

# (iii) Reactivity towards halogens

These elements react with halogens to form trihalides (except  $TII_3$ ).

$$2E(s) + 3 X_2(g) \rightarrow 2EX_3(s)$$
 (X = F, Cl, Br, I)

#### Problem 11.2

White fumes appear around the bottle of anhydrous aluminium chloride. Give reason.

### Solution

Anhydrous aluminium chloride is partially hydrolysed with atmospheric moisture to liberate HCl gas. Moist HCl appears white in colour.

# 11.2 IMPORTANT TRENDS AND ANOMALOUS PROPERTIES OF BORON

Certain important trends can be observed in the chemical behaviour of group 13 elements. The tri-chlorides, bromides and iodides of all these elements being covalent in nature are hydrolysed in water. Species like tetrahedral  $[M(OH)_4]^-$  and octahedral  $[M(H_2O)_6]^{3+}$ , except in boron, exist in aqueous medium.

The monomeric trihalides, being electron deficient, are strong Lewis acids. Boron trifluoride easily reacts with Lewis bases such as  $\,\mathrm{NH}_3$  to complete octet around boron.

$$F_3B + :NH_3 \rightarrow F_3B \leftarrow NH_3$$

It is due to the absence of d orbitals that the maximum covalence of B is 4. Since the d orbitals are available with Al and other elements, the maximum covalence can be expected beyond 4. Most of the other metal halides (e.g., AlCl<sub>3</sub>) are dimerised through halogen bridging (e.g., Al<sub>2</sub>Cl<sub>6</sub>). The metal species completes its octet by accepting electrons from halogen in these halogen bridged molecules.

#### Problem 11.3

Boron is unable to form  $BF_6^{3-}$  ion. Explain.

### Solution

Due to non-availability of d orbitals, boron is unable to expand its octet. Therefore, the maximum covalence of boron cannot exceed 4.

# 11.3 SOME IMPORTANT COMPOUNDS OF BORON

Some useful compounds of boron are borax, orthoboric acid and diborane. We will briefly study their chemistry.

### 11.3.1 Borax

It is the most important compound of boron. It is a white crystalline solid of formula  $\mathrm{Na_2B_4O_7\cdot 10H_2O}$ . In fact it contains the tetranuclear units  $\left[\mathrm{B_4O_5}\left(\mathrm{OH}\right)_4\right]^{2^-}$  and correct formula; therefore, is  $\mathrm{Na_2[B_4O_5}\left(\mathrm{OH}\right)_4].8\mathrm{H_2O}$ . Borax dissolves in water to give an alkaline solution.

$$Na_2B_4O_7 + 7H_2O \rightarrow 2NaOH + 4H_3BO_3$$
  
Orthoboric acid

On heating, borax first loses water molecules and swells up. On further heating it turns into a transparent liquid, which solidifies into glass like material known as borax bead.

The metaborates of many transition metals have characteristic colours and, therefore, borax bead test can be used to identify them in the laboratory. For example, when borax is heated in a Bunsen burner flame with CoO on a loop of platinum wire, a blue coloured  $Co(BO_2)_2$  bead is formed.

### 11.3.2 Orthoboric acid

Orthoboric acid,  $H_3BO_3$  is a white crystalline solid, with soapy touch. It is sparingly soluble in water but highly soluble in hot water. It can be prepared by acidifying an aqueous solution of borax.

$$Na_2B_4O_7 + 2HCl + 5H_2O \rightarrow 2NaCl + 4B(OH)_3$$

It is also formed by the hydrolysis (reaction with water or dilute acid) of most boron compounds (halides, hydrides, etc.). It has a layer structure in which planar  $BO_3$  units are

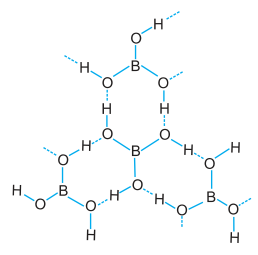
joined by hydrogen bonds as shown in Fig. 11.1.

Boric acid is a weak monobasic acid. It is not a protonic acid but acts as a Lewis acid by accepting electrons from a hydroxyl ion:

$$B(OH)_3 + 2HOH \rightarrow [B(OH)_4]^- + H_3O^+$$

On heating, orthoboric acid above 370K forms metaboric acid,  $HBO_2$  which on further heating yields boric oxide,  $B_2O_3$ .

$$H_3BO_3 \xrightarrow{\Delta} HBO_2 \xrightarrow{\Delta} B_2O_3$$



**Fig. 11. 1** Structure of boric acid; the dotted lines represent hydrogen bonds

### Problem 11.4

Why is boric acid considered as a weak acid?

#### Solution

Because it is not able to release  $H^+$  ions on its own. It receives  $OH^-$ ions from water molecule to complete its octet and in turn releases  $H^+$  ions.

# 11.3.3 Diborane, B<sub>2</sub>H<sub>6</sub>

The simplest boron hydride known, is diborane. It is prepared by treating boron trifluoride with LiAlH<sub>4</sub> in diethyl ether.

$$4\mathrm{BF}_3 + 3\ \mathrm{LiAlH}_4 \rightarrow 2\mathrm{B}_2\mathrm{H}_6 + 3\mathrm{LiF} + 3\mathrm{AlF}_3$$

A convenient laboratory method for the preparation of diborane involves the oxidation of sodium borohydride with iodine.

$$2NaBH_4 + I_2 \rightarrow B_2H_6 + 2NaI + H_2$$

Diborane is produced on an industrial scale by the reaction of  $BF_3$  with sodium hydride.

$$2BF_3 + 6NaH \xrightarrow{450K} B_2H_6 + 6NaF$$

Diborane is a colourless, highly toxic gas with a b.p. of 180 K. Diborane catches fire spontaneously upon exposure to air. It burns in oxygen releasing an enormous amount of energy.

$$B_2H_6 + 3O_2 \rightarrow B_2O_3 + 3H_2O;$$
  
 $\Delta_2H^{\odot} = -1976 \text{ kJ mol}^{-1}$ 

Most of the higher boranes are also spontaneously flammable in air. Boranes are readily hydrolysed by water to give boric acid.

$${\rm B_2H_6(g)} + 6{\rm H_2O(l)} \rightarrow 2{\rm B(OH)_3(aq)} + 6{\rm H_2(g)}$$

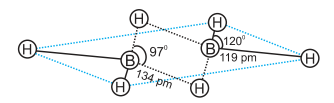
Diborane undergoes cleavage reactions with Lewis bases(L) to give borane adducts,  $\mathrm{BH_3}\text{-}\mathrm{L}$ 

$$B_2H_6 + 2 \text{ NMe}_3 \rightarrow 2BH_3 \cdot NMe_3$$
  
 $B_2H_6 + 2 \text{ CO} \rightarrow 2BH_3 \cdot CO$ 

Reaction of ammonia with diborane gives initially  $B_2H_6.2NH_3$  which is formulated as  $[BH_2(NH_3)_2]^+$   $[BH_4]^-$ ; further heating gives borazine,  $B_3N_3H_6$  known as "inorganic benzene" in view of its ring structure with alternate BH and NH groups.

$$3B_2H_6+6NH_3 \rightarrow 3[BH_2(NH_3)_2]^{\dagger}[BH_4]^{-}$$
  
 $-\frac{Heat}{2} \rightarrow 2B_3N_3H_6+12H_2$ 

The structure of diborane is shown in Fig.11.2(a). The four terminal hydrogen atoms and the two boron atoms lie in one plane. Above and below this plane, there are two bridging hydrogen atoms. The four terminal B-H bonds are regular two centre-two electron bonds while the two bridge (B-H-B) bonds are different and can be described in terms of three

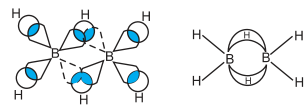


**Fig.11.2(a)** The structure of diborane,  $B_2H_6$ 

centre-two electron bonds shown in Fig.11.2 (b).

Boron also forms a series of hydridoborates; the most important one is the tetrahedral  $[BH_4]^{-}$  ion. Tetrahydridoborates of several metals are known. Lithium and sodium tetrahydridoborates, also known as **borohydrides**, are prepared by the reaction of metal hydrides with  $B_9H_6$  in diethyl ether.

 $2MH + B_2H_6 \rightarrow 2 M^+ [BH_4]^-$  (M = Li or Na)



**Fig.11.2(b)** Bonding in diborane. Each B atom uses sp³ hybrids for bonding. Out of the four sp³ hybrids on each B atom, one is without an electron shown in broken lines. The terminal B-H bonds are normal 2-centre-2-electron bonds but the two bridge bonds are 3-centre-2-electron bonds. The 3-centre-2-electron bridge bonds are also referred to as banana bonds.

Both LiBH<sub>4</sub> and NaBH<sub>4</sub> are used as reducing agents in organic synthesis. They are useful starting materials for preparing other metal borohydrides.

# 11.4 USES OF BORON AND ALUMINIUM AND THEIR COMPOUNDS

Boron being extremely hard refractory solid of high melting point, low density and very low electrical conductivity, finds many applications. Boron fibres are used in making bullet-proof vest and light composite material for aircraft. The boron-10 (<sup>10</sup>B) isotope has high ability to absorb neutrons and, therefore, metal borides are used in nuclear industry as protective shields and control rods. The main industrial application of borax and boric acid is in the manufacture of heat resistant glasses (e.g., Pyrex), glass-wool and fibreglass. Borax is also used as a flux for soldering metals, for heat, scratch and stain resistant glazed coating to earthenwares and as constituent of medicinal soaps. An aqueous solution of orthoboric acid is generally used as a mild antiseptic.

Aluminium is a bright silvery-white metal, with high tensile strength. It has a high electrical and thermal conductivity. On a weight-to-weight basis, the electrical conductivity of aluminium is twice that of copper. Aluminium is used extensively in industry and every day life. It forms alloys with Cu, Mn, Mg, Si and Zn. Aluminium and its alloys can be given shapes of pipe, tubes, rods, wires, plates or foils and, therefore, find uses in packing, utensil making, construction, aeroplane and transportation industry. The use of aluminium and its compounds for domestic purposes is now reduced considerably because of their toxic nature.

# 11.5 GROUP 14 ELEMENTS: THE CARBON FAMILY

Carbon (C), silicon (Si), germanium (Ge), tin (Sn) and lead (Pb) are the members of group 14. Carbon is the seventeenth most abundant element by mass in the earth's crust. It is widely distributed in nature in free as well as in the combined state. In elemental state it is available as coal, graphite and diamond; however, in combined state it is present as metal carbonates, hydrocarbons and carbon dioxide gas (0.03%) in air. One can emphatically say that carbon is the most versatile element in the world. Its combination with other elements such as dihydrogen, dioxygen, chlorine and sulphur provides an astonishing array of materials ranging from living tissues to drugs and plastics. Organic chemistry is devoted to carbon containing compounds. It is an essential constituent of all living organisms. Naturally occurring carbon contains two stable isotopes: 12 C and <sup>13</sup>C. In addition to these, third isotope, <sup>14</sup>C is also present. It is a radioactive isotope with halflife 5770 years and used for radiocarbon dating. Silicon is the second (27.7 % by mass) most abundant element on the earth's crust and is present in nature in the form of silica and silicates. Silicon is a very important component of ceramics, glass and cement.

Germanium exists only in traces. Tin occurs mainly as cassiterite,  ${\rm SnO_2}$  and lead as galena, PbS.

Ultrapure form of germanium and silicon are used to make transistors and semiconductor devices.

The important atomic and physical properties of the group 14 elements along with their electronic configuration are given in Table 11.2 Some of the atomic, physical and chemical properties are discussed below:

### 11.5.1 Electronic Configuration

The valence shell electronic configuration of these elements is  $ns^2np^2$ . The inner core of the electronic configuration of elements in this group also differs.

#### 11.5.2 Covalent Radius

There is a considerable increase in covalent radius from C to Si, thereafter from Si to Pb a small increase in radius is observed. This is due to the presence of completely filled d and f orbitals in heavier members.

### 11.5.3 Ionization Enthalpy

The first ionization enthalpy of group 14 members is higher than the corresponding members of group 13. The influence of inner core electrons is visible here also. In general the ionisation enthalpy decreases down the group. Small decrease in  $\Delta_l H$  from Si to Ge to Sn and slight increase in  $\Delta_l H$  from Sn to Pb is the consequence of poor shielding effect of intervening d and f orbitals and increase in size of the atom.

## 11.5.4 Electronegativity

Due to small size, the elements of this group are slightly more electronegative than group 13 elements. The electronegativity values for elements from Si to Pb are almost the same.

# 11.5.5 Physical Properties

All group 14 members are solids. Carbon and silicon are non-metals, germanium is a metalloid,

Table 11.3 Atomic and Physical Properties of Group 14 Elements

		Element					
Property		Carbon C	Silicon Si	Germanium Ge	Tin Sn	Lead Pb	
Atomic Number		6	14	32	50	82	
Atomic mass	s (g mol <sup>-1</sup> )	12.01	28.09	72.60	118.71	207.2	
Electronic configuration		$[\text{He}]2s^22p^2$	$[\text{Ne}]3s^23p^2$	$[Ar]3d^{10}4s^24p^2$	$[Kr]4d^{10}5s^25p^2$	$[Xe]4f^{14}5d6s^26p^2$	
Covalent radius/pm <sup>a</sup>		77	118	122	140	146	
Ionic radius M <sup>4+</sup> /pm <sup>b</sup>		_	40	53	69	78	
Ionic radius M <sup>2+</sup> /pm <sup>b</sup>		-	-	73	118	119	
Ionization	$\Delta_i H_1$	1086	786	761	708	715	
enthalpy/	$\Delta_{_{\mathbf{i}}}\!H_{_{2}}$	2352	1577	1537	1411	1450	
kJ mol <sup>-1</sup>	$\Delta_i H_3$	4620	3228	3300	2942	3081	
	$\Delta_i H_4$	6220	4354	4409	3929	4082	
Electronegativity <sup>c</sup>		2.5	1.8	1.8	1.8	1.9	
Density <sup>d</sup> /g cm <sup>-3</sup>		$3.51^{\rm e}$	2.34	5.32	$7.26^{\mathrm{f}}$	11.34	
Melting point/K		4373	1693	1218	505	600	
Boiling point/K		-	3550	3123	2896	2024	
Electrical resistivity/ ohm cm (293 K)		10 <sup>14</sup> -10 <sup>16</sup>	50	50	10 <sup>-5</sup>	$2 \times 10^{-5}$	

<sup>&</sup>lt;sup>a</sup> for M <sup>IV</sup> oxidation state; <sup>b</sup> 6–coordination; <sup>c</sup> Pauling scale; <sup>d</sup> 293 K; <sup>e</sup> for diamond; for graphite, density is 2.22;  $^f\beta$ -form (stable at room temperature)

whereas tin and lead are soft metals with low melting points. Melting points and boiling points of group 14 elements are much higher than those of corresponding elements of group 13.

# 11.5.6 Chemical Properties

# Oxidation states and trends in chemical reactivity

The group 14 elements have four electrons in outermost shell. The common oxidation states exhibited by these elements are +4 and +2. Carbon also exhibits negative oxidation states. Since the sum of the first four ionization enthalpies is very high, compounds in +4 oxidation state are generally covalent in nature. In heavier members the tendency to show +2oxidation state increases in the sequence Ge<Sn<Pb. It is due to the inability of  $ns^2$ electrons of valence shell to participate in bonding. The relative stabilities of these two oxidation states vary down the group. Carbon and silicon mostly show +4 oxidation state. Germanium forms stable compounds in +4 state and only few compounds in +2 state. Tin forms compounds in both oxidation states (Sn in +2 state is a reducing agent). Lead compounds in +2 state are stable and in +4 state are strong oxidising agents. In tetravalent state the number of electrons around the central atom in a molecule (e.g., carbon in CCl<sub>4</sub>) is eight. Being electron precise molecules, they are normally not expected to act as electron acceptor or electron donor species. Although carbon cannot exceed its covalence more than 4, other elements of the group can do so. It is because of the presence of d orbital in them. Due to this, their halides undergo hydrolysis and have tendency to form complexes by accepting electron pairs from donor species. For example, the species like,  $SiF_6^{2-}$ ,  $[GeCl_6]^{2-}$ ,  $[Sn(OH)_6]^{2-}$  exist where the hybridisation of the central atom is  $sp^3d^2$ .

### (i) Reactivity towards oxygen

All members when heated in oxygen form oxides. There are mainly two types of oxides, *i.e.*, monoxide and dioxide of formula MO and  $\mathrm{MO}_2$  respectively. SiO only exists at high temperature. Oxides in higher oxidation states of elements are generally more acidic than

those in lower oxidation states. The dioxides —  $\mathrm{CO}_2$ ,  $\mathrm{SiO}_2$  and  $\mathrm{GeO}_2$  are acidic, whereas  $\mathrm{SnO}_2$  and  $\mathrm{PbO}_2$  are amphoteric in nature. Among monoxides, CO is neutral, GeO is distinctly acidic whereas SnO and PbO are amphoteric.

### Problem 11.5

Select the member(s) of group 14 that (i) forms the most acidic dioxide, (ii) is commonly found in +2 oxidation state, (iii) used as semiconductor.

### Solution

(i) carbon (ii) lead

(iii) silicon and germanium

# (ii) Reactivity towards water

Carbon, silicon and germanium are not affected by water. Tin decomposes steam to form dioxide and dihydrogen gas.

$$Sn + 2H_2O \xrightarrow{\Delta} SnO_2 + 2H_2$$

Lead is unaffected by water, probably because of a protective oxide film formation.

## (iii) Reactivity towards halogen

These elements can form halides of formula  $MX_2$  and  $MX_4$  (where X = F, Cl, Br, I). Except carbon, all other members react directly with halogen under suitable condition to make halides. Most of the  $MX_4$  are covalent in nature. The central metal atom in these halides undergoes  $sp^3$  hybridisation and the molecule is tetrahedral in shape. Exceptions are SnF<sub>4</sub> and PbF<sub>4</sub>, which are ionic in nature. PbI<sub>4</sub> does not exist because Pb-I bond initially formed during the reaction does not release enough energy to unpair  $6s^2$  electrons and excite one of them to higher orbital to have four unpaired electrons around lead atom. Heavier members Ge to Pb are able to make halides of formula MX<sub>2</sub>. Stability of dihalides increases down the group. Considering the thermal and chemical stability,  $GeX_4$  is more stable than  $GeX_2$ , whereas  $PbX_2$  is more than  $PbX_4$ . Except  $CCl_4$ , other tetrachlorides are easily hydrolysed by water because the central atom can THE p-BLOCK ELEMENTS 317

accommodate the lone pair of electrons from oxygen atom of water molecule in d orbital.

Hydrolysis can be understood by taking the example of  $\mathrm{SiCl}_4$ . It undergoes hydrolysis by initially accepting lone pair of electrons from water molecule in d orbitals of Si, finally leading to the formation of  $\mathrm{Si}(\mathrm{OH})_4$  as shown below:

### Problem 11.6

 $[SiF_6]^{2-}$  is known whereas  $[SiCl_6]^{2-}$  not. Give possible reasons.

#### Solution

The main reasons are:

- (i) six large chloride ions cannot be accommodated around Si<sup>4+</sup> due to limitation of its size.
- (ii) interaction between lone pair of chloride ion and Si<sup>4+</sup> is not very strong.

# 11.6 IMPORTANT TRENDS AND ANOMALOUS BEHAVIOUR OF CARBON

Like first member of other groups, carbon also differs from rest of the members of its group. It is due to its smaller size, higher electronegativity, higher ionisation enthalpy and unavailability of d orbitals.

In carbon, only *s* and *p* orbitals are available for bonding and, therefore, it can accommodate only four pairs of electrons around it. This would limit the maximum covalence to four whereas other members can expand their covalence due to the presence of *d* orbitals.

Carbon also has unique ability to form  $p\pi-p\pi$  multiple bonds with itself and with other atoms of small size and high electronegativity. Few examples of multiple bonding are: C=C, C=C, C=O, C=S, and C=N. Heavier elements do not form  $p\pi-p\pi$  bonds because their atomic orbitals are too large and diffuse to have effective overlapping.

Carbon atoms have the tendency to link with one another through covalent bonds to form chains and rings. This property is called **catenation**. This is because C—C bonds are very strong. Down the group the size increases and electronegativity decreases, and, thereby, tendency to show catenation decreases. This can be clearly seen from bond enthalpies values. The order of catenation is  $C >> Si > Ge \approx Sn$ . Lead does not show catenation.

Bond	Bond enthalpy / kJ mol <sup>-1</sup>
С—С	348
Si —Si	297
Ge—Ge	260
Sn—Sn	240

Due to property of catenation and  $p\pi$ –  $p\pi$  bond formation, carbon is able to show allotropic forms.

### 11.7 ALLOTROPES OF CARBON

Carbon exhibits many allotropic forms; both crystalline as well as amorphous. Diamond and graphite are two well-known crystalline forms of carbon. In 1985, third form of carbon known as **fullerenes** was discovered by H.W.Kroto, E.Smalley and R.F.Curl. For this discovery they were awarded the Nobel Prize in 1996.

### 11.7.1 **Diamond**

It has a crystalline lattice. In diamond each carbon atom undergoes  $sp^3$  hybridisation and linked to four other carbon atoms by using hybridised orbitals in tetrahedral fashion. The C–C bond length is 154 pm. The structure extends in space and produces a rigid three-dimensional network of carbon atoms. In this

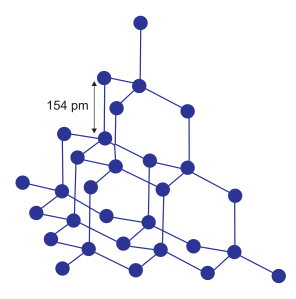


Fig. 11.3 The structure of diamond

structure (Fig. 11.3) directional covalent bonds are present throughout the lattice.

It is very difficult to break extended covalent bonding and, therefore, diamond is a hardest substance on the earth. It is used as an abrasive for sharpening hard tools, in making dies and in the manufacture of tungsten filaments for electric light bulbs.

### Problem 11.7

Diamond is covalent, yet it has high melting point. Why?

### Solution

Diamond has a three-dimensional network involving strong C—C bonds, which are very difficult to break and, in turn has high melting point.

### 11.7.2 Graphite

Graphite has layered structure (Fig.11.4). Layers are held by van der Waals forces and distance between two layers is 340 pm. Each layer is composed of planar hexagonal rings of carbon atoms. C—C bond length within the layer is 141.5 pm. Each carbon atom in hexagonal ring undergoes  $sp^2$  hybridisation and makes three sigma bonds with three neighbouring carbon atoms. Fourth electron forms a  $\pi$  bond. The electrons are delocalised over the whole sheet. Electrons are mobile and.

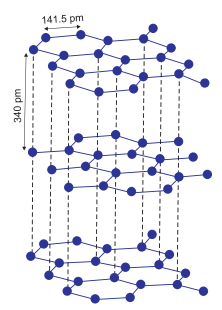


Fig 11.4 The structure of graphite

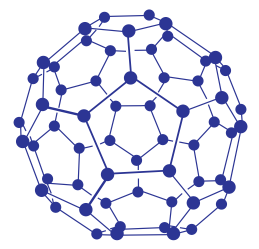
therefore, graphite conducts electricity along the sheet. Graphite cleaves easily between the layers and, therefore, it is very soft and slippery. For this reason graphite is used as a dry lubricant in machines running at high temperature, where oil cannot be used as a lubricant.

### 11.7.3 Fullerenes

Fullerenes are made by the heating of graphite in an electric arc in the presence of inert gases such as helium or argon. The sooty material formed by condensation of vapourised  $C^n$  small molecules consists of mainly  $C_{60}$  with smaller quantity of  $C_{70}$  and traces of fullerenes consisting of even number of carbon atoms up to 350 or above. Fullerenes are the only pure form of carbon because they have smooth structure without having 'dangling' bonds. Fullerenes are cage like molecules.  $C_{60}$  molecule has a shape like soccer ball and called **Buckminsterfullerene** (Fig. 11.5).

It contains twenty six- membered rings and twelve five membered rings. A six membered ring is fused with six or five membered rings but a five membered ring can only fuse with six membered rings. All the carbon atoms are equal and they undergo  $sp^2$  hybridisation. Each carbon atom forms three sigma bonds with other three carbon atoms. The remaining electron at each carbon is delocalised in

molecular orbitals, which in turn give aromatic character to molecule. This ball shaped molecule has 60 vertices and each one is occupied by one carbon atom and it also contains both single and double bonds with C–C distances of 143.5 pm and 138.3 pm respectively. Spherical fullerenes are also called bucky balls in short.



**Fig.11.5** The structure of  $C_{60}$ , Buckminster-fullerene: Note that molecule has the shape of a soccer ball (football).

It is very important to know that graphite is thermodynamically most stable allotrope of carbon and, therefore,  $\Delta_f H^{\circ}$  of graphite is taken as zero.  $\Delta_f H^{\circ}$  values of diamond and fullerene,  $C_{60}$  are 1.90 and 38.1 kJ mol<sup>-1</sup>, respectively.

Other forms of elemental carbon like carbon black, coke, and charcoal are all impure forms of graphite or fullerenes. Carbon black is obtained by burning hydrocarbons in a limited supply of air. Charcoal and coke are obtained by heating wood or coal respectively at high temperatures in the absence of air.

### 11.7.4 Uses of Carbon

Graphite fibres embedded in plastic material form high strength, lightweight composites. The composites are used in products such as tennis rackets, fishing rods, aircrafts and canoes. Being good conductor, graphite is used for electrodes in batteries and industrial electrolysis. Crucibles made from graphite are inert to dilute acids and alkalies. Being highly porous, activated charcoal is used in adsorbing poisonous gases; also used in water

filters to remove organic contaminators and in airconditioning system to control odour. Carbon black is used as black pigment in black ink and as filler in automobile tyres. Coke is used as a fuel and largely as a reducing agent in metallurgy. Diamond is a precious stone and used in jewellery. It is measured in carats (1 carat = 200 mg).

# 11.8 SOME IMPORTANT COMPOUNDS OF CARBON AND SILICON

### Oxides of Carbon

Two important oxides of carbon are carbon monoxide, CO and carbon dioxide,  $CO_2$ .

### 11.8.1 Carbon Monoxide

Direct oxidation of C in limited supply of oxygen or air yields carbon monoxide.

$$2C(s) + O_2(g) \xrightarrow{\Delta} 2CO(g)$$

On small scale pure CO is prepared by dehydration of formic acid with concentrated  ${\rm H_2SO_4}$  at 373 K

$$\text{HCOOH} \xrightarrow{373\text{K}} \text{H}_2\text{O} + \text{CO}$$

On commercial scale it is prepared by the passage of steam over hot coke. The mixture of CO and  $\rm H_2$  thus produced is known as **water** gas or synthesis gas.

$$C(s) + H_2O(g) \xrightarrow{473-1273K} CO(g) + H_2(g)$$
Water gas

When air is used instead of steam, a mixture of CO and  $\rm N_2$  is produced, which is called **producer gas**.

$$2C(s) + O_2(g) + 4N_2(g) \xrightarrow{1273 \text{ K}} 2CO(g) + 4N_2(g)$$

Producer gas

Water gas and producer gas are very important industrial fuels. Carbon monoxide in water gas or producer gas can undergo further combustion forming carbon dioxide with the liberation of heat.

Carbon monoxide is a colourless, odourless and almost water insoluble gas. It is a powerful reducing agent and reduces almost all metal oxides other than those of the alkali and alkaline earth metals, aluminium and a few transition metals. This property of

CO is used in the extraction of many metals from their oxides ores.

$$Fe_2O_3(s) + 3CO(g) \xrightarrow{\Delta} 2Fe(s) + 3CO_2(g)$$

$$ZnO(s) + CO(g) \xrightarrow{\Delta} Zn(s) + CO_2(g)$$

In CO molecule, there are one sigma and two  $\pi$  bonds between carbon and oxygen,  $:C \equiv O:$ . Because of the presence of a lone pair on carbon, CO molecule acts as a donor and reacts with certain metals when heated to form **metal carbonyls**. The highly poisonous nature of CO arises because of its ability to form a **complex with haemoglobin**, which is about 300 times more stable than the oxygen-haemoglobin complex. This prevents haemoglobin in the red blood corpuscles from carrying oxygen round the body and ultimately resulting in death.

#### 11.8.2 Carbon Dioxide

It is prepared by complete combustion of carbon and carbon containing fuels in excess of air.

$$C(s) + O_2(g) \xrightarrow{\Delta} CO_2(g)$$
  
 $CH_4(g) + 2O_2(g) \xrightarrow{\Delta} CO_2(g) + 2H_2O(g)$ 

In the laboratory it is conveniently prepared by the action of dilute HCl on calcium carbonate.

$$\label{eq:CaCO3} \text{CaCO}_3(\text{s}) + 2\text{HCl (aq)} \rightarrow \text{CaCl}_2 \text{ (aq)} + \text{CO}_2 \text{ (g)} + \\ \text{H}_2\text{O(l)}$$

On commercial scale it is obtained by heating limestone.

It is a colourless and odourless gas. Its low solubility in water makes it of immense biochemical and geo-chemical importance. With water, it forms carbonic acid,  $\rm H_2CO_3$  which is a weak dibasic acid and dissociates in two steps:

$$H_2CO_3(aq) + H_2O(l) \rightleftharpoons HCO_3^-(aq) + H_3O^+(aq)$$
  
 $HCO_3^-(aq) + H_2O(l) \rightleftharpoons CO_3^{2-}(aq) + H_3O^+(aq)$ 

 ${
m H_2CO_3/HCO_3^-}$  buffer system helps to maintain pH of blood between 7.26 to 7.42. Being acidic in nature, it combines with alkalies to form metal carbonates.

Carbon dioxide, which is normally present to the extent of  $\sim 0.03$  % by volume in the

atmosphere, is removed from it by the process known as **photosynthesis**. It is the process by which green plants convert atmospheric  $\mathrm{CO}_2$  into carbohydrates such as glucose. The overall chemical change can be expressed as:

$$6\text{CO}_2 + 12\text{H}_2\text{O} \xrightarrow{\text{Chlorphyll}} \text{C}_6\text{H}_{12}\text{O}_6 + 6\text{O}_2 + 6\text{H}_2\text{O}_6$$

By this process plants make food for themselves as well as for animals and human beings. Unlike CO, it is not poisonous. But the increase in combustion of fossil fuels and decomposition of limestone for cement manufacture in recent years seem to increase the  ${\rm CO}_2$  content of the atmosphere. This may lead to increase in **green house effect** and thus, raise the temperature of the atmosphere which might have serious consequences.

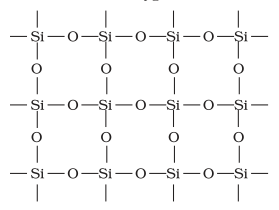
Carbon dioxide can be obtained as a solid in the form of **dry ice** by allowing the liquified  $CO_2$  to expand rapidly. Dry ice is used as a refrigerant for ice-cream and frozen food. Gaseous  $CO_2$  is extensively used to carbonate soft drinks. Being heavy and non-supporter of combustion it is used as fire extinguisher. A substantial amount of  $CO_2$  is used to manufacture urea.

In  $\mathrm{CO}_2$  molecule carbon atom undergoes sp hybridisation. Two sp hybridised orbitals of carbon atom overlap with two p orbitals of oxygen atoms to make two sigma bonds while other two electrons of carbon atom are involved in  $p\pi$ –  $p\pi$  bonding with oxygen atom. This results in its linear shape [with both C–O bonds of equal length (115 pm)] with no dipole moment. The resonance structures are shown below:

Resonance structures of carbon dioxide

# 11.8.3 Silicon Dioxide, SiO<sub>2</sub>

95% of the earth's crust is made up of silica and silicates. Silicon dioxide, commonly known as silica, occurs in several crystallographic forms. Quartz, cristobalite and tridymite are some of the crystalline forms of silica, and they are interconvertable at suitable temperature. Silicon dioxide is a covalent, three-dimensional network solid in which each silicon atom is covalently bonded in a tetrahedral manner to four oxygen atoms. Each oxygen atom in turn covalently bonded to another silicon atoms as shown in diagram (Fig 11.6). Each corner is shared with another tetrahedron. The entire crystal may be considered as giant molecule in which eight membered rings are formed with alternate silicon and oxygen atoms.



**Fig. 11.6** Three dimensional structure of  $SiO_2$ 

Silica in its normal form is almost non-reactive because of very high Si—O bond enthalpy. It resists the attack by halogens, dihydrogen and most of the acids and metals even at elevated temperatures. However, it is attacked by HF and NaOH.

$$SiO_2 + 2NaOH \rightarrow Na_2SiO_3 + H_2O$$
  
 $SiO_2 + 4HF \rightarrow SiF_4 + 2H_2O$ 

Quartz is extensively used as a piezoelectric material; it has made possible to develop extremely accurate clocks, modern radio and television broadcasting and mobile radio communications. Silica gel is used as a drying agent and as a support for chromatographic materials and catalysts. Kieselghur, an amorphous form of silica is used in filtration plants.

### 11.8.4 Silicones

They are a group of organosilicon polymers, which have  $\{R_2SiO\}$  as a repeating unit. The starting materials for the manufacture of silicones are alkyl or aryl substituted silicon chlorides,  $R_nSiCl_{(4-n)}$ , where R is alkyl or aryl group. When methyl chloride reacts with silicon in the presence of copper as a catalyst at a temperature 573K various types of methyl

substituted chlorosilane of formula MeSiCl $_3$ , Me $_2$ SiCl $_2$ , Me $_3$ SiCl with small amount of Me $_4$ Si are formed. Hydrolysis of dimethyldichlorosilane, (CH $_3$ ) $_2$ SiCl $_2$  followed by condensation polymerisation yields straight chain polymers.

The chain length of the polymer can be controlled by adding (CH<sub>3</sub>)<sub>3</sub>SiCl which blocks the ends as shown below:

Silicones being surrounded by non-polar alkyl groups are water repelling in nature. They have in general high thermal stability, high dielectric strength and resistance to oxidation and chemicals. They have wide applications. They are used as sealant, greases, electrical insulators and for water proofing of fabrics. Being biocompatible they are also used in surgical and cosmetic plants.

### Problem: 11.8

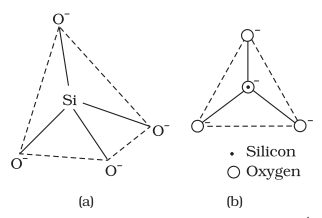
What are silicones?

### Solution

Simple silicones consist of  $\begin{pmatrix} 1 \\ Si - O \end{pmatrix}_n$  chains in which alkyl or phenyl groups occupy the remaining bonding positions on each silicon. They are hydrophobic (water repellant) in nature.

### 11.8.5 Silicates

A large number of silicates minerals exist in nature. Some of the examples are feldspar, zeolites, mica and asbestos. The basic structural unit of silicates is  $\mathrm{SiO_4}^{4^+}$  (Fig.11.7) in which silicon atom is bonded to four oxygen atoms in tetrahedron fashion. In silicates either the discrete unit is present or a number of such units are joined together via corners by sharing 1,2,3 or 4 oxygen atoms per silicate units. When silicate units are linked together, they form chain, ring, sheet or three-dimensional structures. Negative charge on silicate structure is



**Fig. 11.7** (a) Tetrahedral structure of  $SiO_4^{4-}$  anion; (b) Representation of  $SiO_4^{4-}$  unit

neutralized by positively charged metal ions. If all the four corners are shared with other tetrahedral units, three-dimensional network is formed.

Two important man-made silicates are glass and cement.

#### 11.8.6 Zeolites

If aluminium atoms replace few silicon atoms in three-dimensional network of silicon dioxide, overall structure known as aluminosilicate, acquires a negative charge. Cations such as Na<sup>+</sup>, K<sup>+</sup> or Ca<sup>2+</sup> balance the negative charge. Examples are feldspar and zeolites. Zeolites are widely used as a catalyst in petrochemical industries for cracking of hydrocarbons and isomerisation, e.g., ZSM-5 (A type of zeolite) used to convert alcohols directly into gasoline. Hydrated zeolites are used as ion exchangers in softening of "hard" water.

### SUMMARY

**p-Block** of the periodic table is unique in terms of having all types of elements – **metals**, **non-metals** and **metalloids**. There are six groups of p-block elements in the periodic table numbering from 13 to 18 Their valence shell electronic configuration is  $ns^2np^{1-6}$  (except for Ne). Differences in the **inner core** of their electronic configuration greatly influence their physical and chemical properties. As a consequence of this, a lot of variation in properties among these elements is observed. In addition to the *group exidation* state, these elements show other electrons by unit of two. While the group oxidation state is the most stable for the lighter elements of the group, lower oxidation states become progressively more stable for the heavier elements. The combined effect of size and availability of d orbitals considerably

influences the ability of these elements to form  $\pi$ -bonds. While the lighter elements form  $p\pi$ - $p\pi$  bonds, the heavier ones form  $d\pi$ - $p\pi$  or  $d\pi$ - $d\pi$  bonds. Absence of d orbital in second period elements limits their maximum covalence to 4 while heavier ones can exceed this limit.

Boron is a typical non-metal and the other members are metals. The availability of 3 valence electrons  $(2s^22p^1)$  for covalent bond formation using four orbitals  $(2s, 2p_x, 2p_y)$  and  $2p_z$  leads to the so called **electron deficiency** in boron compounds. This deficiency makes them good electron acceptor and thus boron compounds behave as **Lewis acids**. Boron forms covalent molecular compounds with dihydrogen as boranes, the simplest of which is **diborane**, B.H. Diborane contains two bridging hydrogen atoms between two boron atoms; these bridge bonds are considered to be **three-centre two-electron bonds**. The important compounds of boron with dioxygen are **boric acid** and **borax**. Boric acid, B(OH)3 is a weak monobasic acid; it acts as a Lewis acid by accepting electrons from hydroxyl ion. Borax is a white crystalline solid of formula  $Na_2[B_4O_5(OH)_2] \cdot 8H_2O$ . The **borax bead test** gives characteristic colours of transition metals.

Aluminium exhibits +3 oxidation state. With heavier elements +1 oxidation state gets progressively stabilised on going down the group. This is a consequence of the so called **inert pair effect**.

Carbon/is a typical non-metal forming covalent bonds employing all its four valence electrons  $(2/s^2)^2$ . It shows the property of **catenation**, the ability to form chains or rings, not only with C-C single bonds but also with multiple bonds (C $\neq$ C or C=C). The tendency/to catenation decreases as C>>Si>Ge  $\simeq$  Sn > Pb. Carbon provides one of the best examples of allotropy. Three important allotropes of carbon are diamond, graphite and **fullerenes**. The members of the carbon family mainly exhibit + A and +2 oxidation states / compouds in +4 oxidation states are generally opvalent in nature. The tendency to show +2 oxidation state increases among heavier elements. Lead in +2 state is stable whereas in +4 oxidation state it is a strong oxidising agent. Carbon also exhibits negative oxidation states. It forms two important oxides: CO and CO<sub>2</sub>. Carlyon monoxide is neutral whereas CO<sub>2</sub> is acidic in nature. Carbon monoxide having lone pair of electrons on C forms metal **carbonyls**. It is dealdly poisonous due to higher stability of its **haemoglobin ¢omplex** as compared to that of oxyhaemoglobin complex. Carbon dioxide as such is not toxic. However, increased content of CO, in atmosphere the to combustion of fossil fuels and decomposition of limestone is feared to cause increase in green house effect'. This, in turn, raises the temperalure of the atmosphere and causes serious complications. Silica, silicates and silicones are important class of compounds and find applications in industry and technology.

#### **EXERCISES**

- Discuss the pattern of variation in the oxidation states of (i) B to Tl and (ii) C to Pb.
- 11.2 How can you explain higher stability of BCl<sub>3</sub> as compared to TlCl<sub>3</sub>?
- 11.3 Why does boron triflouride behave as a Lewis acid?
- 11.4 Consider the compounds, BCl<sub>3</sub> and CCl<sub>4</sub>. How will they behave with water? Justify.
- 11.5 Is boric acid a protic acid? Explain.
- 11.6 Explain what happens when boric acid is heated.
- 11.7 Describe the shapes of  $\mathrm{BF_3}$  and  $\mathrm{BH_4}$ . Assign the hybridisation of boron in these species.
- 11.8 Write reactions to justify amphoteric nature of aluminium.

11.9 What are electron deficient compounds ? Are  $\mathrm{BCl_3}$  and  $\mathrm{SiCl_4}$  electron deficient species ? Explain.

- 11.10 Write the resonance structures of  $CO_3^2$  and  $HCO_3$ .
- 11.11 What is the state of hybridisation of carbon in (a)  $CO_3^{2-}$  (b) diamond (c) graphite?
- 11.12 Explain the difference in properties of diamond and graphite on the basis of their structures.
- 11.13 Rationalise the given statements and give chemical reactions:
  - Lead(II) chloride reacts with Cl<sub>2</sub> to give PbCl<sub>4</sub>.
  - Lead(IV) chloride is highly unstable towards heat.
  - Lead is known not to form an iodide, PbI<sub>4</sub>.
- 11.14 Suggest reasons why the B-F bond lengths in  $BF_3$  (130 pm) and  $BF_4^-$  (143 pm) differ.
- 11.15 If B–Cl bond has a dipole moment, explain why  $\mathrm{BCl_3}$  molecule has zero dipole moment.
- 11.16 Aluminium trifluoride is insoluble in anhydrous HF but dissolves on addition of NaF. Aluminium trifluoride precipitates out of the resulting solution when gaseous  $BF_3$  is bubbled through. Give reasons.
- 11.17 Suggest a reason as to why CO is poisonous.
- 11.18 How is excessive content of CO<sub>2</sub> responsible for global warming?
- 11.19 Explain structures of diborane and boric acid.
- 11.20 What happens when
  - (a) Borax is heated strongly,
  - (b) Boric acid is added to water,
  - (c) Aluminium is treated with dilute NaOH,
  - (d) BF<sub>3</sub> is reacted with ammonia?
- 11.21 Explain the following reactions
  - (a) Silicon is heated with methyl chloride at high temperature in the presence of copper;
  - (b) Silicon dioxide is treated with hydrogen fluoride;
  - (c) CO is heated with ZnO;
  - (d) Hydrated alumina is treated with aqueous NaOH solution.
- 11.22 Give reasons:
  - (i) Conc. HNO<sub>3</sub> can be transported in aluminium container.
  - (ii) A mixture of dilute NaOH and aluminium pieces is used to open drain.
  - (iii) Graphite is used as lubricant.
  - (iv) Diamond is used as an abrasive.
  - (v) Aluminium alloys are used to make aircraft body.
  - (vi) Aluminium utensils should not be kept in water overnight.
  - (vii) Aluminium wire is used to make transmission cables.
- 11.23 Explain why is there a phenomenal decrease in ionization enthalpy from carbon to silicon?
- 11.24 How would you explain the lower atomic radius of Ga as compared to Al?
- 11.25 What are allotropes? Sketch the structure of two allotropes of carbon namely diamond and graphite. What is the impact of structure on physical properties of two allotropes?

(a) Classify following oxides as neutral, acidic, basic or amphoteric: CO, B<sub>2</sub>O<sub>3</sub>, SiO<sub>2</sub>, CO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, PbO<sub>2</sub>, Tl<sub>2</sub>O<sub>3</sub> (b) Write suitable chemical equations to show their nature. 11.27 In some of the reactions thallium resembles aluminium, whereas in others it resembles with group I metals. Support this statement by giving some evidences. 11.28 When metal X is treated with sodium hydroxide, a white precipitate (A) is obtained, which is soluble in excess of NaOH to give soluble complex (B). Compound (A) is soluble in dilute HCl to form compound (C). The compound (A) when heated strongly gives (D), which is used to extract metal. Identify (X), (A), (B), (C) and (D). Write suitable equations to support their identities. 11.29 What do you understand by (a) inert pair effect (b) allotropy and (c) catenation? 11.30 A certain salt X, gives the following results. Its aqueous solution is alkaline to litmus. (ii) It swells up to a glassy material Y on strong heating. (iii) When conc. H<sub>o</sub>SO<sub>4</sub> is added to a hot solution of X, white crystal of an acid Z separates out. Write equations for all the above reactions and identify X, Y and Z. 11.31 Write balanced equations for: (i)  $BF_3 + LiH \rightarrow$ (ii)  $B_2H_6 + H_2O \rightarrow$ (iii) NaH +  $B_9H_6 \rightarrow$ (iv)  $H_aBO_a \xrightarrow{\Delta}$ (v) Al + NaOH  $\rightarrow$ (vi)  $B_2H_6 + NH_3 \rightarrow$ 11.32. Give one method for industrial preparation and one for laboratory preparation of CO and CO<sub>2</sub> each. 11.33 An aqueous solution of borax is (a) neutral (b) amphoteric (d) acidic (c) basic 11.34 Boric acid is polymeric due to (a) its acidic nature (b) the presence of hydrogen bonds (c) its monobasic nature (d) its geometry The type of hybridisation of boron in diborane is 11.35 (b)  $sp^{2}$ (a) sp (c)  $sp^3$ (d)  $dsp^2$ 11.36 Thermodynamically the most stable form of carbon is (a) diamond (b) graphite (c) fullerenes (d) coal 11.37 Elements of group 14 (a) exhibit oxidation state of +4 only (b) exhibit oxidation state of +2 and +4(c) form  $M^{2-}$  and  $M^{4+}$  ion

If the starting material for the manufacture of silicones is RSiCl<sub>3</sub>, write the

(d) form  $M^{2+}$  and  $M^{4+}$  ions

structure of the product formed.

11.38