

Unit

7

The *p*-Block Elements

Objectives

After studying this Unit, you will be able to

- appreciate general trends in the chemistry of elements of groups 15, 16, 17 and 18;
- learn the preparation, properties and uses of dinitrogen and phosphorus and some of their important compounds;
- describe the preparation, properties and uses of dioxygen and ozone and chemistry of some simple oxides;
- know allotropic forms of sulphur, chemistry of its important compounds and the structures of its oxoacids;
- describe the preparation, properties and uses of chlorine and hydrochloric acid;
- know the chemistry of interhalogens and structures of oxoacids of halogens;
- enumerate the uses of noble gases;
- appreciate the importance of these elements and their compounds in our day to day life.

*Diversity in chemistry is the hallmark of *p*-block elements manifested in their ability to react with the elements of *s*-, *d*- and *f*-blocks as well as with their own.*

In Class XI, you have learnt that the *p*-block elements are placed in groups 13 to 18 of the periodic table. Their valence shell electronic configuration is ns^2np^{1-6} (except He which has $1s^2$ configuration). The properties of *p*-block elements like that of others are greatly influenced by atomic sizes, ionisation enthalpy, electron gain enthalpy and electronegativity. The absence of *d*-orbitals in second period and presence of *d* or *d* and *f* orbitals in heavier elements (starting from third period onwards) have significant effects on the properties of elements. In addition, the presence of all the three types of elements; metals, metalloids and non-metals bring diversification in chemistry of these elements.

Having learnt the chemistry of elements of Groups 13 and 14 of the *p*-block of periodic table in Class XI, you will learn the chemistry of the elements of subsequent groups in this Unit.

71 Group 15 Elements

Group 15 includes nitrogen, phosphorus, arsenic, antimony and bismuth. As we go down the group, there is a shift from non-metallic to metallic through metalloidal character. Nitrogen and phosphorus are non-metals, arsenic and antimony metalloids and bismuth is a typical metal.

7.1.1 Occurrence

Molecular nitrogen comprises 78% by volume of the atmosphere. In the earth's crust, it occurs as sodium nitrate, NaNO_3 (called Chile saltpetre) and potassium nitrate (Indian saltpetre). It is found in the form of proteins in plants and animals. Phosphorus occurs in minerals

of the apatite family, $\text{Ca}_9(\text{PO}_4)_6 \cdot \text{CaX}_2$ ($\text{X} = \text{F}, \text{Cl}$ or OH) (e.g., fluorapatite $\text{Ca}_9(\text{PO}_4)_6 \cdot \text{CaF}_2$) which are the main components of phosphate rocks. Phosphorus is an essential constituent of animal and plant matter. It is present in bones as well as in living cells. Phosphoproteins are present in milk and eggs. Arsenic, antimony and bismuth are found mainly as sulphide minerals.

The important atomic and physical properties of this group elements along with their electronic configurations are given in Table 7.1.

Table 7.1: Atomic and Physical Properties of Group 15 Elements

Property	N	P	As	Sb	Bi
Atomic number	7	15	33	51	83
Atomic mass/g mol ⁻¹	14.01	30.97	74.92	121.75	208.98
Electronic configuration	[He]2s ² 2p ³	[Ne]3s ² 3p ³	[Ar]3d ¹⁰ 4s ² 4p ³	[Kr]4d ¹⁰ 5s ² 5p ³	[Xe]4f ¹⁴ 5d ¹⁰ 6s ² 6p ³
Ionisation enthalpy ($\Delta_i H/(\text{kJ mol}^{-1})$)	I	1402	1012	947	834
	II	2856	1903	1798	1595
	III	4577	2910	2736	2443
Electronegativity	3.0	2.1	2.0	1.9	1.9
Covalent radius/pm ^a	70	110	121	141	148
Ionic radius/pm	171 ^b	212 ^b	222 ^b	76 ^c	103 ^c
Melting point/K	63*	317 ^d	1089 ^e	904	544
Boiling point/K	77.2*	554 ^d	888 ^f	1860	1837
Density/[g cm ⁻³ (298 K)]	0.879 ^g	1.823	5.778 ^h	6.697	9.808

^a E^{III} single bond (E = element); ^b E^{3-} ; ^c E^{3+} ; ^d White phosphorus; ^e Grey α -form at 38.6 atm; ^f Sublimation temperature; ^g At 63 K; ^h Grey α -form; * Molecular N_2 .

Trends of some of the atomic, physical and chemical properties of the group are discussed below.

7.1.2 Electronic Configuration

The valence shell electronic configuration of these elements is ns^2np^3 . The s orbital in these elements is completely filled and p orbitals are half-filled, making their electronic configuration extra stable.

7.1.3 Atomic and Ionic Radii

Covalent and ionic (in a particular state) radii increase in size down the group. There is a considerable increase in covalent radius from N to P. However, from As to Bi only a small increase in covalent radius is observed. This is due to the presence of completely filled d and/or f orbitals in heavier members.

7.1.4 Ionisation Enthalpy

Ionisation enthalpy decreases down the group due to gradual increase in atomic size. Because of the extra stable half-filled p orbitals electronic configuration and smaller size, the ionisation enthalpy of the group 15 elements is much greater than that of group 14 elements in the corresponding periods. The order of successive ionisation enthalpies, as expected is $\Delta_i H_1 < \Delta_i H_2 < \Delta_i H_3$ (Table 7.1).

7.1.5 Electronegativity

The electronegativity value, in general, decreases down the group with increasing atomic size. However, amongst the heavier elements, the difference is not that much pronounced.

7.1.6 Physical Properties

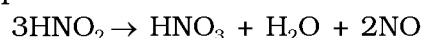
All the elements of this group are polyatomic. Dinitrogen is a diatomic gas while all others are solids. Metallic character increases down the group. Nitrogen and phosphorus are non-metals, arsenic and antimony metalloids and bismuth is a metal. This is due to decrease in ionisation enthalpy and increase in atomic size. The boiling points, in general, increase from top to bottom in the group but the melting point increases upto arsenic and then decreases upto bismuth. Except nitrogen, all the elements show allotropy.

7.1.7 Chemical Properties

Oxidation states and trends in chemical reactivity

The common oxidation states of these elements are -3, +3 and +5. The tendency to exhibit -3 oxidation state decreases down the group due to increase in size and metallic character. In fact last member of the group, bismuth hardly forms any compound in -3 oxidation state. The stability of +5 oxidation state decreases down the group. The only well characterised Bi (V) compound is BiF_5 . The stability of +5 oxidation state decreases and that of +3 state increases (due to inert pair effect) down the group. Nitrogen exhibits +1, +2, +4 oxidation states also when it reacts with oxygen. Phosphorus also shows +1 and +4 oxidation states in some oxoacids.

In the case of nitrogen, all oxidation states from +1 to +4 tend to disproportionate in acid solution. For example,



Similarly, in case of phosphorus nearly all intermediate oxidation states disproportionate into +5 and -3 both in alkali and acid. However +3 oxidation state in case of arsenic, antimony and bismuth becomes increasingly stable with respect to disproportionation.

Nitrogen is restricted to a maximum covalency of 4 since only four (one s and three p) orbitals are available for bonding. The heavier elements have vacant d orbitals in the outermost shell which can be used for bonding (covalency) and hence, expand their covalence as in PF_6^- .

Anomalous properties of nitrogen

Nitrogen differs from the rest of the members of this group due to its small size, high electronegativity, high ionisation enthalpy and non-availability of d orbitals. Nitrogen has unique ability to form **$p\pi-p\pi$ multiple** bonds with itself and with other elements having small size and high electronegativity (e.g., C, O). Heavier elements of this group do not form $p\pi-p\pi$ bonds as their atomic orbitals are so large and diffuse that they cannot have effective overlapping. Thus, nitrogen exists as a diatomic molecule with a triple bond (one s and two p) between the two atoms. Consequently, its bond enthalpy ($941.4 \text{ kJ mol}^{-1}$) is very high. On the contrary, phosphorus, arsenic and antimony form single bonds as P-P, As-As and Sb-Sb while bismuth forms metallic bonds in elemental state. However, the single N-N bond is weaker than the single P-P bond because of high interelectronic repulsion of the non-bonding electrons, owing to the small bond length. As a result the catenation tendency is weaker in

nitrogen. Another factor which affects the chemistry of nitrogen is the absence of *d* orbitals in its valence shell. Besides restricting its covalency to four, nitrogen cannot form ***d*π–*p*π bond** as the heavier elements can e.g., $R_3P = O$ or $R_3P = CH_2$ (*R* = alkyl group). Phosphorus and arsenic can form ***d*π–*d*π bond** also with transition metals when their compounds like $P(C_2H_5)_3$ and $As(C_6H_5)_3$ act as ligands.

- (i) *Reactivity towards hydrogen*: All the elements of Group 15 form hydrides of the type EH_3 where *E* = N, P, As, Sb or Bi. Some of the properties of these hydrides are shown in Table 7.2. The hydrides show regular gradation in their properties. The stability of hydrides decreases from NH_3 to BiH_3 which can be observed from their bond dissociation enthalpy. Consequently, the reducing character of the hydrides increases. Ammonia is only a mild reducing agent while BiH_3 is the strongest reducing agent amongst all the hydrides. Basicity also decreases in the order $NH_3 > PH_3 > AsH_3 > SbH_3 \geq BiH_3$.

Table 7.2: Properties of Hydrides of Group 15 Elements

Property	NH_3	PH_3	AsH_3	SbH_3	BiH_3
Melting point/K	195.2	139.5	156.7	185	–
Boiling point/K	238.5	185.5	210.6	254.6	290
(E–H) Distance/pm	101.7	141.9	151.9	170.7	–
HEH angle (°)	107.8	93.6	91.8	91.3	–
$\Delta_f H^\circ / kJ\ mol^{-1}$	–46.1	13.4	66.4	145.1	278
$\Delta_{diss} H^\circ (E-H) / kJ\ mol^{-1}$	389	322	297	255	–

- (ii) *Reactivity towards oxygen*: All these elements form two types of oxides: E_2O_3 and E_2O_5 . The oxide in the higher oxidation state of the element is more acidic than that of lower oxidation state. Their acidic character decreases down the group. The oxides of the type E_2O_3 of nitrogen and phosphorus are purely acidic, that of arsenic and antimony amphoteric and those of bismuth predominantly basic.
- (iii) *Reactivity towards halogens*: These elements react to form two series of halides: EX_3 and EX_5 . Nitrogen does not form pentahalide due to non-availability of the *d* orbitals in its valence shell. Pentahalides are more covalent than trihalides. All the trihalides of these elements except those of nitrogen are stable. In case of nitrogen, only NF_3 is known to be stable. Trihalides except BiF_3 are predominantly covalent in nature.
- (iv) *Reactivity towards metals*: All these elements react with metals to form their binary compounds exhibiting –3 oxidation state, such as, Ca_3N_2 (calcium nitride) Ca_3P_2 (calcium phosphide), Na_3As_2 (sodium arsenide), Zn_3Sb_2 (zinc antimonide) and Mg_3Bi_2 (magnesium bismuthide).

Though nitrogen exhibits +5 oxidation state, it does not form pentahalide. Give reason.

[Example 7.1](#)

Nitrogen with $n = 2$, has s and p orbitals only. It does not have d orbitals to expand its covalence beyond four. That is why it does not form pentahalide.

[Solution](#)

PH_3 has lower boiling point than NH_3 . Why?

[Example 7.](#)

Unlike NH_3 , PH_3 molecules are not associated through hydrogen bonding in liquid state. That is why the boiling point of PH_3 is lower than NH_3 .

[Solution](#)

Inte t uestions

7.1 Why are pentahalides more covalent than trihalides ?

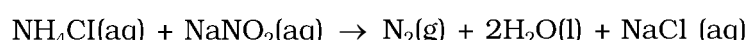
7.2 Why is BiH_3 the strongest reducing agent amongst all the hydrides of Group 15 elements ?

7. Nitrogen

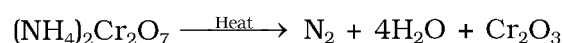
Preparation

Dinitrogen is produced commercially by the liquefaction and fractional distillation of air. Liquid dinitrogen (b.p. 77.2 K) distils out first leaving behind liquid oxygen (b.p. 90 K).

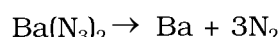
In the laboratory, dinitrogen is prepared by treating an aqueous solution of ammonium chloride with sodium nitrite.



Small amounts of NO and HNO_3 are also formed in this reaction; these impurities can be removed by passing the gas through aqueous sulphuric acid containing potassium dichromate. It can also be obtained by the thermal decomposition of ammonium dichromate.



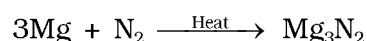
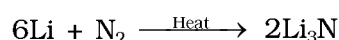
Very pure nitrogen can be obtained by the thermal decomposition of sodium or barium azide.



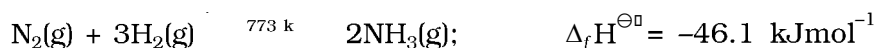
Properties

Dinitrogen is a colourless, odourless, tasteless and non-toxic gas. Nitrogen atom has two stable isotopes: ^{14}N and ^{15}N . It has a very low solubility in water (23.2 cm^3 per litre of water at 273 K and 1 bar pressure) and low freezing and boiling points (Table 7.1).

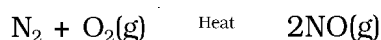
Dinitrogen is rather inert at room temperature because of the high bond enthalpy of $\text{N}\equiv\text{N}$ bond. Reactivity, however, increases rapidly with rise in temperature. At higher temperatures, it directly combines with some metals to form predominantly ionic nitrides and with non-metals, covalent nitrides. A few typical reactions are:



It combines with hydrogen at about 773 K in the presence of a catalyst (Haber's Process) to form ammonia:



Dinitrogen combines with dioxygen only at very high temperature (at about 2000 K) to form nitric oxide, NO.



Uses The main use of dinitrogen is in the manufacture of ammonia and other industrial chemicals containing nitrogen, (e.g., calcium cyanamide). It also finds use where an inert atmosphere is required (e.g., in iron and steel industry, inert diluent for reactive chemicals). Liquid dinitrogen is used as a refrigerant to preserve biological materials, food items and in cryosurgery.

Example 7. Write the reaction of thermal decomposition of sodium azide.

Solution Thermal decomposition of sodium azide gives dinitrogen gas.

$$2\text{NaN}_3 \rightarrow 2\text{Na} + 3\text{N}_2$$

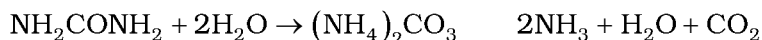
Inte t uestion

7.3 Why is N_2 less reactive at room temperature?

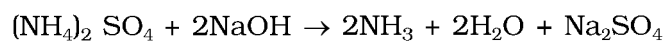
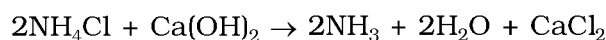
7. Ammonia

Preparation

Ammonia is present in small quantities in air and soil where it is formed by the decay of nitrogenous organic matter e.g., urea.



On a small scale ammonia is obtained from ammonium salts which decompose when treated with caustic soda or calcium hydroxide.



On a large scale, ammonia is manufactured by Haber's process.



In accordance with Le Chatelier's principle, high pressure would favour the formation of ammonia. The optimum conditions for the production of ammonia are a pressure of 200×10^5 Pa (about 200 atm), a temperature of ~ 700 K and the use of a catalyst such as iron oxide with small amounts of K_2O and Al_2O_3 to increase the rate of attainment of equilibrium. The flow chart for the production of ammonia is shown in Fig. 7.1. Earlier, iron was used as a catalyst with molybdenum as a promoter.

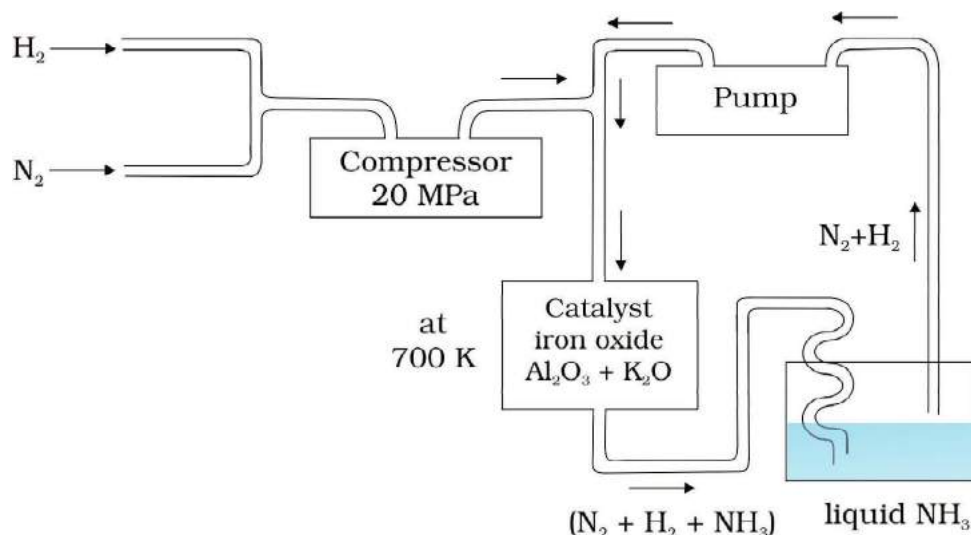
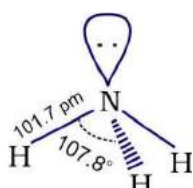


Fig. 7.1
Flow chart for the
manufacture of
ammonia



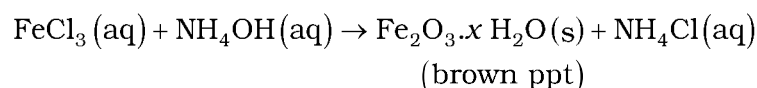
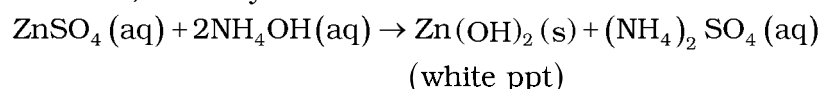
Properties

Ammonia is a colourless gas with a pungent odour. Its freezing and boiling points are 198.4 and 239.7 K respectively. In the solid and liquid states, it is associated through hydrogen bonds as in the case of water and that accounts for its higher melting and boiling points than expected on the basis of its molecular mass. The ammonia molecule is trigonal pyramidal with the nitrogen atom at the apex. It has three bond pairs and one lone pair of electrons as shown in the structure.

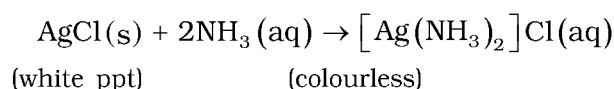
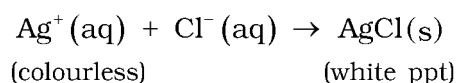
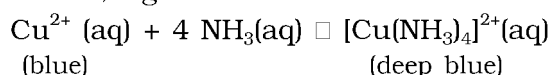
Ammonia gas is highly soluble in water. Its aqueous solution is weakly basic due to the formation of OH^- ions.



It forms ammonium salts with acids, e.g., NH_4Cl , $(\text{NH}_4)_2\text{SO}_4$, etc. As a weak base, it precipitates the hydroxides (hydrated oxides in case of some metals) of many metals from their salt solutions. For example,



The presence of a lone pair of electrons on the nitrogen atom of the ammonia molecule makes it a Lewis base. It donates the electron pair and forms linkage with metal ions and the formation of such complex compounds finds applications in detection of metal ions such as Cu^{2+} , Ag^+ :



Uses Ammonia is used to produce various nitrogenous fertilisers (ammonium nitrate, urea, ammonium phosphate and ammonium sulphate) and in the manufacture of some inorganic nitrogen compounds, the most important one being nitric acid. Liquid ammonia is also used as a refrigerant.

Example 7. Why does NH_3 act as a Lewis base ?
Solution Nitrogen atom in NH_3 has one lone pair of electrons which is available for donation. Therefore, it acts as a Lewis base.

Inte t uestions
7.4 Mention the conditions required to maximise the yield of ammonia.
7.5 How does ammonia react with a solution of Cu^{2+} ?

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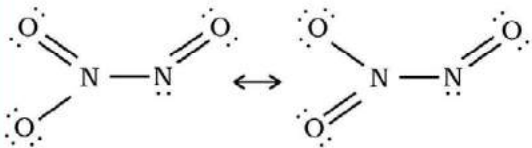
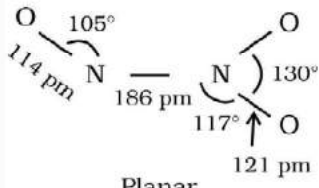
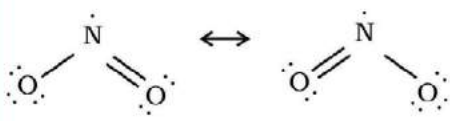
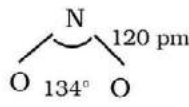
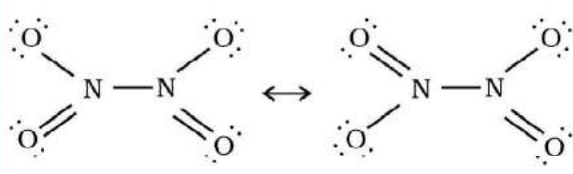
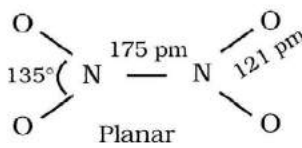
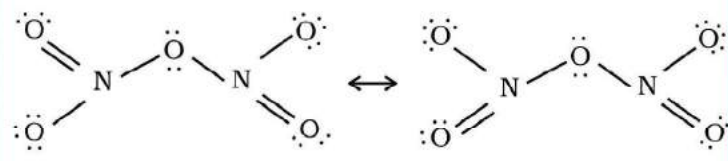
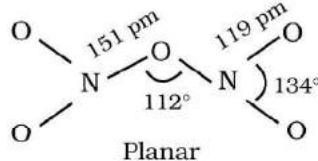
Nitrogen forms a number of oxides in different oxidation states. The names, formulas, preparation and physical appearance of these oxides are given in Table 7.3.

Table 7.3: Oxides of Nitrogen

Name	Formula	Oxidation state of nitrogen	Common methods of preparation	Physical appearance and chemical nature
Dinitrogen oxide [Nitrogen(I) oxide]	N_2O	+ 1	$\text{NH}_4\text{NO}_3 \xrightarrow{\text{Heat}} \text{N}_2\text{O} + 2\text{H}_2\text{O}$	colourless gas, neutral
Nitrogen monoxide [Nitrogen(II) oxide]	NO	+ 2	$2\text{NaNO}_2 + 2\text{FeSO}_4 + 3\text{H}_2\text{SO}_4 \rightarrow \text{Fe}_2(\text{SO}_4)_3 + 2\text{NaHSO}_4 + 2\text{H}_2\text{O} + 2\text{NO}$	colourless gas, neutral
Dinitrogen trioxide [Nitrogen(III) oxide]	N_2O_3	+ 3	$2\text{NO} + \text{N}_2\text{O}_4 \xrightarrow{250\text{K}} 2\text{N}_2\text{O}_3$	blue solid, acidic
Nitrogen dioxide [Nitrogen(IV) oxide]	NO_2	+ 4	$2\text{Pb}(\text{NO}_3)_2 \xrightarrow{673\text{K}} 4\text{NO}_2 + 2\text{PbO} + \text{O}_2$	brown gas, acidic
Dinitrogen tetroxide [Nitrogen(IV) oxide]	N_2O_4	+ 4	$2\text{NO}_2 \xrightleftharpoons[\text{Heat}]{\text{Cool}} \text{N}_2\text{O}_4$	colourless solid/liquid, acidic
Dinitrogen pentoxide [Nitrogen(V) oxide]	N_2O_5	+5	$4\text{HNO}_3 + \text{P}_4\text{O}_{10} \rightarrow 4\text{HPO}_3 + 2\text{N}_2\text{O}_5$	colourless solid, acidic

Lewis dot main resonance structures and bond parameters of oxides are given in Table 7.4.

Table 7.4: Structures of Oxides of Nitrogen

Formula	Resonance Structures	Bond Parameters
N ₂ O	$\ddot{\text{N}}=\ddot{\text{N}}=\ddot{\text{O}} \longleftrightarrow :\text{N}\equiv\text{N}-\ddot{\text{O}}:$	$\text{N}-\text{N}-\text{O}$ 113 pm 119 pm Linear
NO	$:\text{N}=\ddot{\text{O}}: \longleftrightarrow :\ddot{\text{N}}=\ddot{\text{O}}:$	$\text{N}-\text{O}$ 115 pm
N ₂ O ₃		 Planar
NO ₂		 Angular
N ₂ O ₄		 Planar
N ₂ O ₅		 Planar

Why does NO₂ dimerise ?

NO₂ contains odd number of valence electrons. It behaves as a typical odd molecule. On dimerisation, it is converted to stable N₂O₄ molecule with even number of electrons.

[Example 7.5](#)
[Solution](#)

Exercise 7.6

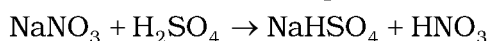
7.6 What is the covalence of nitrogen in N₂O₅ ?

7.5 Nitric acid

Nitrogen forms oxoacids such as H₂N₂O₂ (hyponitrous acid), HNO₂ (nitrous acid) and HNO₃ (nitric acid). Amongst them HNO₃ is the most important.

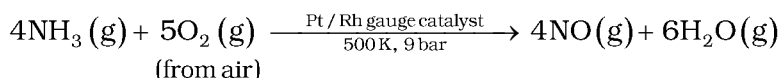
Preparation

In the laboratory, nitric acid is prepared by heating KNO_3 or NaNO_3 and concentrated H_2SO_4 in a glass retort.

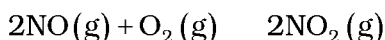


On a large scale it is prepared mainly by Ostwald's process.

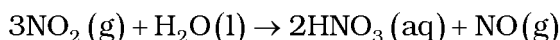
This method is based upon catalytic oxidation of NH_3 by atmospheric oxygen.



Nitric oxide thus formed combines with oxygen giving NO_2 .



Nitrogen dioxide so formed, dissolves in water to give HNO_3 .



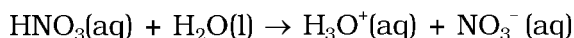
NO thus formed is recycled and the aqueous HNO_3 can be concentrated by distillation upto ~ 68% by mass. Further concentration to 98% can be achieved by dehydration with concentrated H_2SO_4 .

Properties

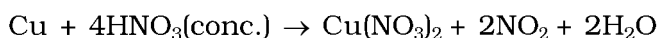
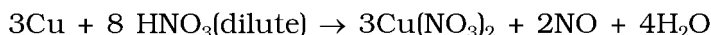
It is a colourless liquid (f.p. 231.4 K and b.p. 355.6 K). Laboratory grade nitric acid contains ~ 68% of the HNO_3 by mass and has a specific gravity of 1.504.

In the gaseous state, HNO_3 exists as a planar molecule with the structure as shown.

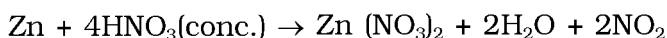
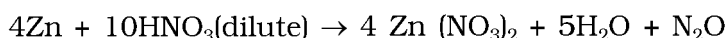
In aqueous solution, nitric acid behaves as a strong acid giving hydronium and nitrate ions.



Concentrated nitric acid is a strong oxidising agent and attacks most metals except noble metals such as gold and platinum. The products of oxidation depend upon the concentration of the acid, temperature and the nature of the material undergoing oxidation.

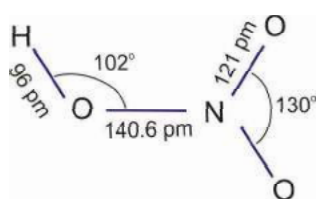


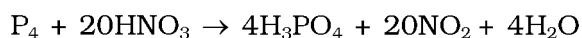
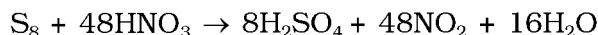
Zinc reacts with dilute nitric acid to give N_2O and with concentrated acid to give NO_2 .



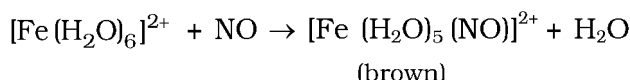
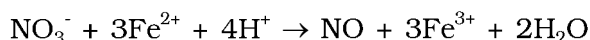
Some metals (e.g., Cr, Al) do not dissolve in concentrated nitric acid because of the formation of a passive film of oxide on the surface.

Concentrated nitric acid also oxidises non-metals and their compounds. Iodine is oxidised to iodic acid, carbon to carbon dioxide, sulphur to H_2SO_4 , and phosphorus to phosphoric acid.





Brown Ring Test: The familiar brown ring test for nitrates depends on the ability of Fe^{2+} to reduce nitrates to nitric oxide, which reacts with Fe^{2+} to form a brown coloured complex. The test is usually carried out by adding dilute ferrous sulphate solution to an aqueous solution containing nitrate ion, and then carefully adding concentrated sulphuric acid along the sides of the test tube. A brown ring at the interface between the solution and sulphuric acid layers indicates the presence of nitrate ion in solution.



Uses The major use of nitric acid is in the manufacture of ammonium nitrate for fertilisers and other nitrates for use in explosives and pyrotechnics. It is also used for the preparation of nitroglycerin, trinitrotoluene and other organic nitro compounds. Other major uses are in the *pickling of stainless steel*, etching of metals and as an oxidiser in rocket fuels.

7. Phosphorus Allotropic forms

Phosphorus is found in many allotropic forms, the important ones being white, red and black.

White phosphorus is a translucent white waxy solid. It is poisonous, insoluble in water but soluble in carbon disulphide and glows in dark (chemiluminescence). It dissolves in boiling NaOH solution in an inert atmosphere giving PH_3 .

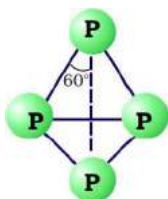
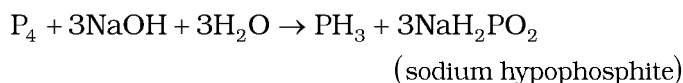
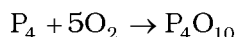


Fig. 7.2
White phosphorus

White phosphorus is less stable and therefore, more reactive than the other solid phases under normal conditions because of angular strain in the P_4 molecule where the angles are only 60° . It readily catches fire in air to give dense white fumes of P_4O_{10} .



It consists of discrete tetrahedral P_4 molecule as shown in Fig. 7.2.

Red phosphorus is obtained by heating white phosphorus at 573K in an inert atmosphere for several days. When red phosphorus is heated under high pressure, a series of phases of black phosphorus is formed. Red phosphorus possesses iron grey lustre. It is odourless, non-poisonous and insoluble in water as well as in carbon disulphide. Chemically, red phosphorus is much less reactive than white phosphorus. It does not glow in the dark.

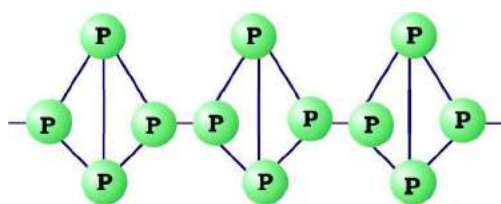


Fig.7.3: Red phosphorus

It is polymeric, consisting of chains of P₄ tetrahedra linked together in the manner as shown in Fig. 7.3.

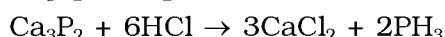
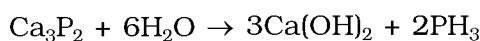
Black phosphorus has two forms α -black phosphorus and β -black phosphorus. α -Black phosphorus is formed when red phosphorus is heated in a sealed tube at 803K. It can be sublimed

in air and has opaque monoclinic or rhombohedral crystals. It does not oxidise in air. β -Black phosphorus is prepared by heating white phosphorus at 473 K under high pressure. It does not burn in air upto 673 K.

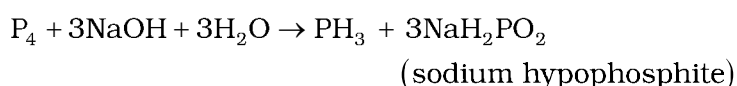
7.7 phosphine

Preparation

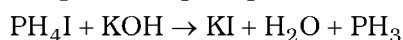
Phosphine is prepared by the reaction of calcium phosphide with water or dilute HCl.



In the laboratory, it is prepared by heating white phosphorus with concentrated NaOH solution in an inert atmosphere of CO₂.



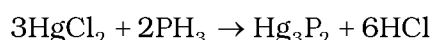
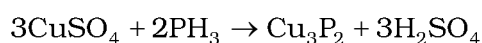
When pure, it is non inflammable but becomes inflammable owing to the presence of P₂H₄ or P₄ vapours. To purify it from the impurities, it is absorbed in HI to form phosphonium iodide (PH₄I) which on treating with KOH gives off phosphine.



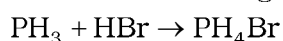
Properties

It is a colourless gas with rotten fish smell and is highly poisonous. It explodes in contact with traces of oxidising agents like HNO₃, Cl₂ and Br₂ vapours.

It is slightly soluble in water. The solution of PH₃ in water decomposes in presence of light giving red phosphorus and H₂. When absorbed in copper sulphate or mercuric chloride solution, the corresponding phosphides are obtained.



Phosphine is weakly basic and like ammonia, gives phosphonium compounds with acids e.g.,



Uses □ The spontaneous combustion of phosphine is technically used in *Holme's signals*. Containers containing calcium carbide and calcium phosphide are pierced and thrown in the sea when the gases evolved burn and serve as a signal. It is also used in *smoke screens*.

In what way can it be proved that PH_3 is basic in nature? [Example 7.](#)

PH_3 reacts with acids like HI to form PH_4I which shows that it is basic in nature. [Solution](#)



Due to lone pair on phosphorus atom, PH_3 is acting as a Lewis base in the above reaction.

Exercise Questions

7.7 Bond angle in PH_4^+ is higher than that in PH_3 . Why?

7.8 What happens when white phosphorus is heated with concentrated NaOH solution in an inert atmosphere of CO_2 ?

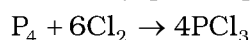
7. Phosphorus Halides

Phosphorus forms two types of halides, PX_3 ($\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$) and PX_5 ($\text{X} = \text{F}, \text{Cl}, \text{Br}$).

7.8.1 Phosphorus Trichloride

Preparation

It is obtained by passing dry chlorine over heated white phosphorus.

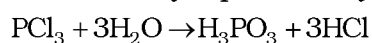


It is also obtained by the action of thionyl chloride with white phosphorus.

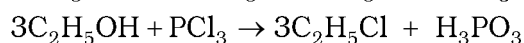
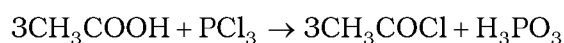


Properties

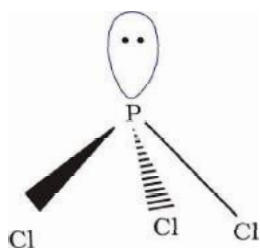
It is a colourless oily liquid and hydrolyses in the presence of moisture.



It reacts with organic compounds containing $-\text{OH}$ group such as CH_3COOH , $\text{C}_2\text{H}_5\text{OH}$.



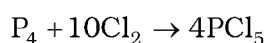
It has a pyramidal shape as shown, in which phosphorus is sp^3 hybridised.



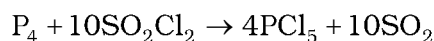
7.8.2 Phosphorus Pentachloride

Preparation

Phosphorus pentachloride is prepared by the reaction of white phosphorus with excess of dry chlorine.

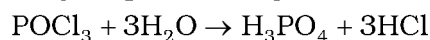
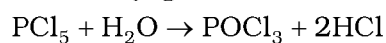


It can also be prepared by the action of SO_2Cl_2 on phosphorus.

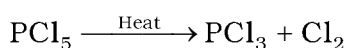


Properties

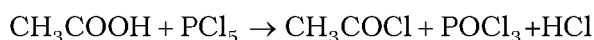
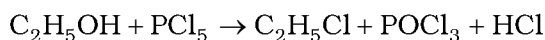
PCl_5 is a yellowish white powder and in moist air, it hydrolyses to POCl_3 and finally gets converted to phosphoric acid.



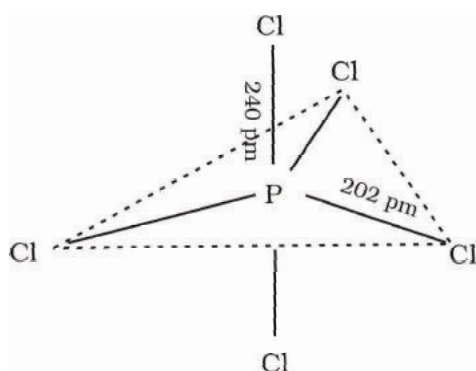
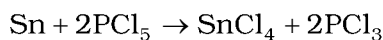
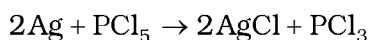
When heated, it sublimes but decomposes on stronger heating.



It reacts with organic compounds containing –OH group converting them to chloro derivatives.



Finely divided metals on heating with PCl_5 give corresponding chlorides.



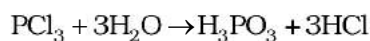
It is used in the synthesis of some organic compounds, e.g., $\text{C}_2\text{H}_5\text{Cl}$, CH_3COCl .

In gaseous and liquid phases, it has a trigonal bipyramidal structure as shown. The three equatorial P–Cl bonds are equivalent, while the two axial bonds are longer than equatorial bonds. This is due to the fact that the axial bond pairs suffer more repulsion as compared to equatorial bond pairs.

In the solid state it exists as an ionic solid, $[\text{PCl}_4]^+[\text{PCl}_6]^-$ in which the cation, $[\text{PCl}_4]^+$ is tetrahedral and the anion, $[\text{PCl}_6]^-$ octahedral.

Example 7.7 Why does PCl_3 fume in moisture ?

Solution PCl_3 hydrolyses in the presence of moisture giving fumes of HCl .



Example 7. Are all the five bonds in PCl_5 molecule equivalent? Justify your answer.

Solution PCl_5 has a trigonal bipyramidal structure and the three equatorial P–Cl bonds are equivalent, while the two axial bonds are different and longer than equatorial bonds.

Inte t uestions

7.9 What happens when PCl_5 is heated?

7.10 Write a balanced equation for the hydrolytic reaction of PCl_5 in heavy water.

7. Oxoacids of phosphorus

Phosphorus forms a number of oxoacids. The important oxoacids of phosphorus with their formulas, methods of preparation and the presence of some characteristic bonds in their structures are given in Table 7.5.

Table 7.5: Oxoacids of Phosphorus

Name	Formula	Oxidation state of phosphorus	Characteristic bonds and their number	Preparation
Hypophosphorous (Phosphinic)	H_3PO_2	+1	One P – OH Two P – H One P = O	white P_4 + alkali
Orthophosphorous (Phosphonic)	H_3PO_3	+3	Two P – OH One P – H One P = O	$\text{P}_2\text{O}_3 + \text{H}_2\text{O}$
Pyrophosphorous	$\text{H}_4\text{P}_2\text{O}_5$	+3	Two P – OH Two P – H Two P = O	$\text{PCl}_3 + \text{H}_3\text{PO}_3$
Hypophosphoric	$\text{H}_4\text{P}_2\text{O}_6$	+4	Four P – OH Two P = O One P – P	red P_4 + alkali
Orthophosphoric	H_3PO_4	+5	Three P – OH One P = O	$\text{P}_4\text{O}_{10} + \text{H}_2\text{O}$
Pyrophosphoric	$\text{H}_4\text{P}_2\text{O}_7$	+5	Four P – OH Two P = O One P – O – P	heat phosphoric acid
Metaphosphoric*	$(\text{HPO}_3)_n$	+5	Three P – OH Three P = O Three P – O – P	phosphorus acid + Br_2 , heat in a sealed tube

* Exists in polymeric forms only. Characteristic bonds of $(\text{HPO}_3)_3$ have been given in the Table.

The compositions of the oxoacids are interrelated in terms of loss or gain of H_2O molecule or O-atom.

The structures of some important oxoacids are given below:

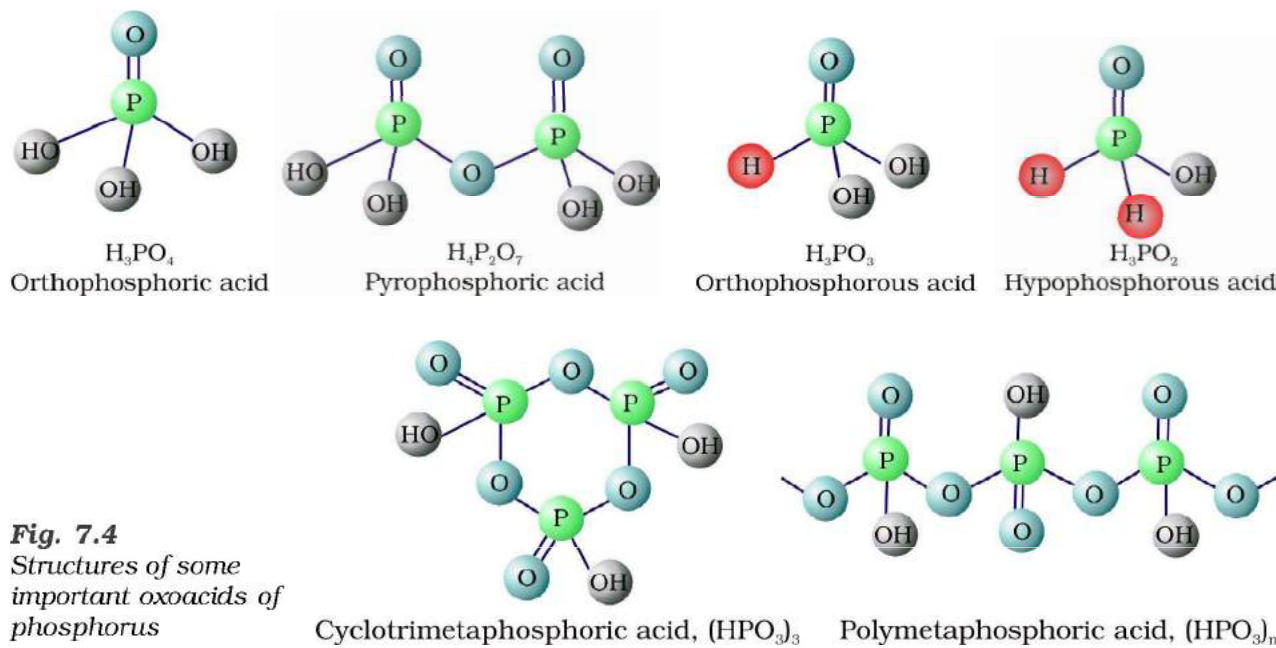
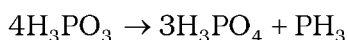
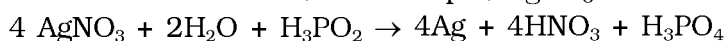


Fig. 7.4
Structures of some important oxoacids of phosphorus

In oxoacids phosphorus is tetrahedrally surrounded by other atoms. All these acids contain at least one P=O bond and one P–OH bond. The oxoacids in which phosphorus has lower oxidation state (less than +5) contain, in addition to P=O and P–OH bonds, either P–P (e.g., in $\text{H}_4\text{P}_2\text{O}_6$) or P–H (e.g., in H_3PO_2) bonds but not both. These acids in +3 oxidation state of phosphorus tend to disproportionate to higher and lower oxidation states. For example, orthophosphorous acid (or phosphorous acid) on heating disproportionates to give orthophosphoric acid (or phosphoric acid) and phosphine.



The acids which contain P–H bond have strong reducing properties. Thus, hypophosphorous acid is a good reducing agent as it contains two P–H bonds and reduces, for example, AgNO_3 to metallic silver.



These P–H bonds are not ionisable to give H^+ and do not play any role in basicity. Only those H atoms which are attached with oxygen in P–OH form are ionisable and cause the basicity. Thus, H_3PO_3 and H_3PO_4 are dibasic and tribasic, respectively as the structure of H_3PO_3 has two P–OH bonds and H_3PO_4 three.

Example 7. How do you account for the reducing behaviour of H_3PO_2 on the basis of its structure?

Solution In H_3PO_2 , two H atoms are bonded directly to P atom which imparts reducing character to the acid.

Inte t uestions

7.11 What is the basicity of H_3PO_4 ?

7.12 What happens when H_3PO_3 is heated?

7.1 Group 16 Elements

Oxygen, sulphur, selenium, tellurium and polonium constitute Group 16 of the periodic table. This is sometimes known as group of *chalcogens*. The name is derived from the Greek word for brass and points to the association of sulphur and its congeners with copper. Most copper minerals contain either oxygen or sulphur and frequently the other members of the group.

7.10.1 Occurrence

Oxygen is the most abundant of all the elements on earth. Oxygen forms about 46.6% by mass of earth's crust. Dry air contains 20.946% oxygen by volume.

However, the abundance of sulphur in the earth's crust is only 0.03-0.1%. Combined sulphur exists primarily as sulphates such as *gypsum* $\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$, *epsom salt* $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, *baryte* BaSO_4 and sulphides such as *galena* PbS , *zinc blende* ZnS , *copper pyrites* CuFeS_2 . Traces of sulphur occur as hydrogen sulphide in volcanoes. Organic materials such as eggs, proteins, garlic, onion, mustard, hair and wool contain sulphur.

Selenium and tellurium are also found as metal selenides and tellurides in sulphide ores. Polonium occurs in nature as a decay product of thorium and uranium minerals.

The important atomic and physical properties of Group 16 along with electronic configuration are given in Table 7.6. Some of the atomic, physical and chemical properties and their trends are discussed below.

7.10.2 Electronic Configuration	The elements of Group 16 have six electrons in the outermost shell and have ns^2np^4 general electronic configuration.
7.10.3 Atomic and Ionic Radii	Due to increase in the number of shells, atomic and ionic radii increase from top to bottom in the group. The size of oxygen atom is, however, exceptionally small.
7.10.4 Ionisation Enthalpy	Ionisation enthalpy decreases down the group. It is due to increase in size. However, the elements of this group have lower ionisation enthalpy values compared to those of Group 15 in the corresponding periods. This is due to the fact that Group 15 elements have extra stable half-filled p orbitals electronic configurations.
7.10.5 Electron Gain Enthalpy	Because of the compact nature of oxygen atom, it has less negative electron gain enthalpy than sulphur. However, from sulphur onwards the value again becomes less negative upto polonium.
7.10.6 Electronegativity	Next to fluorine, oxygen has the highest electronegativity value amongst the elements. Within the group, electronegativity decreases with an increase in atomic number. This implies that the metallic character increases from oxygen to polonium.

Elements of Group 16 generally show lower value of first ionisation enthalpy compared to the corresponding periods of group 15. Why?

Example 7.1

Due to extra stable half-filled p orbitals electronic configurations of Group 15 elements, larger amount of energy is required to remove electrons compared to Group 16 elements.

Solution

7.10.7 Physical Properties	Some of the physical properties of Group 16 elements are given in Table 7.6. Oxygen and sulphur are non-metals, selenium and tellurium metalloids, whereas polonium is a metal. Polonium is radioactive and is short lived (Half-life 13.8 days). All these elements exhibit allotropy. The melting and boiling points increase with an increase in atomic number down the group. The large difference between the melting and boiling points of oxygen and sulphur may be explained on the basis of their atomicity; oxygen exists as diatomic molecule (O_2) whereas sulphur exists as polyatomic molecule (S_8).
7.10.8 Chemical Properties	<i>Oxidation states and trends in chemical reactivity</i> The elements of Group 16 exhibit a number of oxidation states (Table 7.6). The stability of -2 oxidation state decreases down the group. Polonium hardly shows -2 oxidation state. Since electronegativity of oxygen is very high, it shows only negative oxidation state as -2 except

Table 7.6: Some Physical Properties of Group 16 Elements

Property	O	S	Se	Te	Po
Atomic number	8	16	34	52	84
Atomic mass/g mol ⁻¹	16.00	32.06	78.96	127.60	210.00
Electronic configuration	[He]2s ² 2p ⁴	[Ne]3s ² 3p ⁴	[Ar]3d ¹⁰ 4s ² 4p ⁴	[Kr]4d ¹⁰ 5s ² 5p ⁴	[Xe]4f ¹⁴ 5d ¹⁰ 6s ² 6p ⁴
Covalent radius/(pm) ^a	66	104	117	137	146
Ionic radius, E ²⁻ /pm	140	184	198	221	230 ^b
Electron gain enthalpy, /Δ _{eg} H kJ mol ⁻¹	-141	-200	-195	-190	-174
Ionisation enthalpy (Δ _i H ₁) /kJ mol ⁻¹	1314	1000	941	869	813
Electronegativity	3.50	2.58	2.55	2.01	1.76
Density /g cm ⁻³ (298 K)	1.32 ^c	2.06 ^d	4.19 ^e	6.25	–
Melting point/K	55	393 ^f	490	725	520
Boiling point/K	90	718	958	1260	1235
Oxidation states*	-2,-1,1,2	-2,2,4,6	-2,2,4,6	-2,2,4,6	2,4

^aSingle bond; ^bApproximate value; ^cAt the melting point; ^d Rhombic sulphur; ^eHexagonal grey; ^fMonoclinic form, 673 K.
* Oxygen shows oxidation states of +2 and +1 in oxygen fluorides OF₂ and O₂F₂ respectively.

in the case of OF₂ where its oxidation state is + 2. Other elements of the group exhibit + 2, + 4, + 6 oxidation states but + 4 and + 6 are more common. Sulphur, selenium and tellurium usually show + 4 oxidation state in their compounds with oxygen and + 6 with fluorine. The stability of + 6 oxidation state decreases down the group and stability of + 4 oxidation state increases (inert pair effect). Bonding in +4 and +6 oxidation states is primarily covalent.

Anomalous behaviour of oxygen

The anomalous behaviour of oxygen, like other members of *p*-block present in second period is due to its small size and high electronegativity. One typical example of effects of small size and high electronegativity is the presence of strong hydrogen bonding in H₂O which is not found in H₂S.

The absence of *d* orbitals in oxygen limits its covalency to four and in practice, rarely exceeds two. On the other hand, in case of other elements of the group, the valence shells can be expanded and covalence exceeds four.

- (i) *Reactivity with hydrogen:* All the elements of Group 16 form hydrides of the type H₂E (E = O, S, Se, Te, Po). Some properties of hydrides are given in Table 7.7. Their acidic character increases from H₂O to H₂Te. The increase in acidic character can be explained in terms of decrease in bond enthalpy for the dissociation of H–E bond down the group. Owing to the decrease in enthalpy for the dissociation of H–E bond down the group, the thermal stability of hydrides also decreases from H₂O to H₂Po. All the hydrides except water possess reducing property and this character increases from H₂S to H₂Te.

Table 7.7: Properties of Hydrides of Group 16 Elements

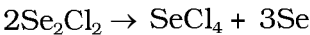
Property	H ₂ O	H ₂ S	H ₂ Se	H ₂ Te
m.p/K	273	188	208	222
b.p/K	373	213	232	269
H-E distance/pm	96	134	146	169
HEH angle (°)	104	92	91	90
$\Delta_f H/\text{kJ mol}^{-1}$	-286	-20	73	100
$\Delta_{\text{diss}} H (H-E)/\text{kJ mol}^{-1}$	463	347	276	238
Dissociation constant ^a	1.8×10^{-16}	1.3×10^{-7}	1.3×10^{-4}	2.3×10^{-3}

^a Aqueous solution, 298 K

- (ii) *Reactivity with oxygen*: All these elements form oxides of the EO₂ and EO₃ types where E = S, Se, Te or Po. Ozone (O₃) and sulphur dioxide (SO₂) are gases while selenium dioxide (SeO₂) is solid. Reducing property of dioxide decreases from SO₂ to TeO₂; SO₂ is reducing while TeO₂ is an oxidising agent. Besides EO₂ type, sulphur, selenium and tellurium also form EO₃ type oxides (SO₃, SeO₃, TeO₃). Both types of oxides are acidic in nature.
- (iii) *Reactivity towards the halogens*: Elements of Group 16 form a large number of halides of the type, EX₆, EX₄ and EX₂ where E is an element of the group and X is a halogen. The stability of the halides decreases in the order F⁻ > Cl⁻ > Br⁻ > I⁻. Amongst hexahalides, hexafluorides are the only stable halides. All hexafluorides are gaseous in nature. They have octahedral structure. Sulphur hexafluoride, SF₆ is exceptionally stable for steric reasons.

Amongst tetrafluorides, SF₄ is a gas, SeF₄ a liquid and TeF₄ a solid. These fluorides have sp³d hybridisation and thus, have trigonal bipyramidal structures in which one of the equatorial positions is occupied by a lone pair of electrons. This geometry is also regarded as *see-saw* geometry.

All elements except oxygen form dichlorides and dibromides. These dihalides are formed by sp³ hybridisation and thus, have tetrahedral structure. The well known monohalides are dimeric in nature. Examples are S₂F₂, S₂Cl₂, S₂Br₂, Se₂Cl₂ and Se₂Br₂. These dimeric halides undergo disproportionation as given below:



H₂S is less acidic than H₂Te. Why?

[Example 7.11](#)

Due to the decrease in bond (E-H) dissociation enthalpy down the group, acidic character increases.

[Solution](#)

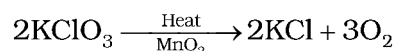
Practice Questions

- 7.13** List the important sources of sulphur.
- 7.14** Write the order of thermal stability of the hydrides of Group 16 elements.
- 7.15** Why is H₂O a liquid and H₂S a gas ?

Preparation

Dioxygen can be obtained in the laboratory by the following ways:

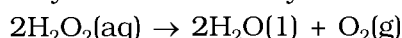
- (i) By heating oxygen containing salts such as chlorates, nitrates and permanganates.



- (ii) By the thermal decomposition of the oxides of metals low in the electrochemical series and higher oxides of some metals.



- (iii) Hydrogen peroxide is readily decomposed into water and dioxygen by catalysts such as finely divided metals and manganese dioxide.



On large scale it can be prepared from water or air. Electrolysis of water leads to the release of hydrogen at the cathode and oxygen at the anode.

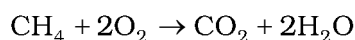
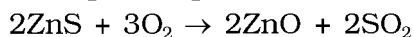
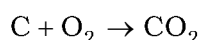
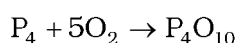
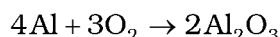
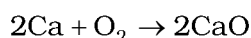
Industrially, dioxygen is obtained from air by first removing carbon dioxide and water vapour and then, the remaining gases are liquefied and fractionally distilled to give dinitrogen and dioxygen.

Properties

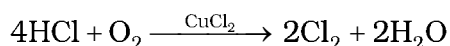
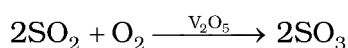
Dioxygen is a colourless and odourless gas. Its solubility in water is to the extent of 3.08 cm^3 in 100 cm^3 water at 293 K which is just sufficient for the vital support of marine and aquatic life. It liquefies at 90 K and freezes at 55 K . Oxygen atom has three stable isotopes: ^{16}O , ^{17}O and ^{18}O . Molecular oxygen, O_2 is unique in being paramagnetic inspite of having even number of electrons (see Class XI Chemistry Book, Unit 4).

Dioxygen directly reacts with nearly all metals and non-metals except some metals (e.g., Au, Pt) and some noble gases. Its combination with other elements is often strongly exothermic which helps in sustaining the reaction. However, to initiate the reaction, some external heating is required as bond dissociation enthalpy of oxygen-oxygen double bond is high ($493.4 \text{ kJ mol}^{-1}$).

Some of the reactions of dioxygen with metals, non-metals and other compounds are given below:



Some compounds are catalytically oxidised. For example,



Uses In addition to its importance in normal respiration and combustion processes, oxygen is used in oxyacetylene welding, in the manufacture of many metals, particularly steel. Oxygen cylinders are widely used in hospitals, high altitude flying and in mountaineering. The combustion of fuels, e.g., hydrazines in liquid oxygen, provides tremendous thrust in rockets.

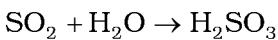
Exercise Questions

- 7.16** Which of the following does not react with oxygen directly?
Zn, Ti, Pt, Fe
- 7.17** Complete the following reactions:
(i) $\text{C}_2\text{H}_4 + \text{O}_2 \rightarrow$
(ii) $4\text{Al} + 3 \text{O}_2 \rightarrow$

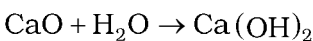
7.1 Simple oxides

A binary compound of oxygen with another element is called oxide. As already stated, oxygen reacts with most of the elements of the periodic table to form oxides. In many cases one element forms two or more oxides. The oxides vary widely in their nature and properties.

Oxides can be classified as acidic (e.g., SO_2 , Cl_2O_7 , CO_2 , N_2O_5) or mixed (Pb_3O_4 , Fe_3O_4). Simple oxides can be classified on the basis of their acidic, basic or amphoteric character. An oxide that combines with water to give an acid is termed acidic oxide (e.g., SO_2 , Cl_2O_7 , CO_2 , N_2O_5). For example, SO_2 combines with water to give H_2SO_3 , an acid.

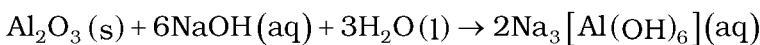
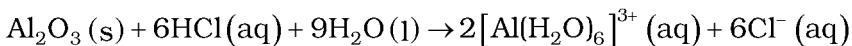


As a general rule, only non-metal oxides are acidic but oxides of some metals in high oxidation state also have acidic character (e.g., Mn_2O_7 , CrO_3 , V_2O_5). The oxides which give a base with water are known as basic oxides (e.g., Na_2O , CaO , BaO). For example, CaO combines with water to give Ca(OH)_2 , a base.



In general, metallic oxides are basic.

Some metallic oxides exhibit a dual behaviour. They show characteristics of both acidic as well as basic oxides. Such oxides are known as amphoteric oxides. They react with acids as well as alkalies. For example, Al_2O_3 reacts with acids as well as alkalies.



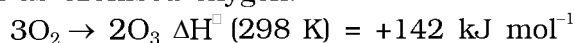
There are some oxides which are neither acidic nor basic. Such oxides are known as neutral oxides. Examples of neutral oxides are CO , NO and N_2O .

Ozone is an allotropic form of oxygen. It is too reactive to remain for long in the atmosphere at sea level. At a height of about 20 kilometres, it is formed from atmospheric oxygen in the presence of sunlight. This ozone layer protects the earth's surface from an excessive concentration of ultraviolet (UV) radiations.

7.1 One

Preparation

When a slow dry stream of oxygen is passed through a silent electrical discharge, conversion of oxygen to ozone (10%) occurs. The product is known as ozonised oxygen.



Since the formation of ozone from oxygen is an endothermic process, it is necessary to use a silent electrical discharge in its preparation to prevent its decomposition.

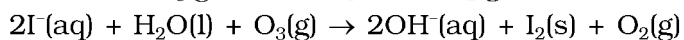
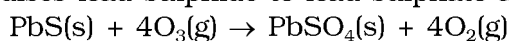
If concentrations of ozone greater than 10 per cent are required, a battery of ozonisers can be used, and pure ozone (b.p. 101.1K) can be condensed in a vessel surrounded by liquid oxygen.

Properties

Pure ozone is a pale blue gas, dark blue liquid and violet-black solid. Ozone has a characteristic smell and in small concentrations it is harmless. However, if the concentration rises above about 100 parts per million, breathing becomes uncomfortable resulting in headache and nausea.

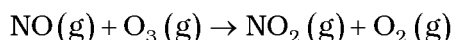
Ozone is thermodynamically unstable with respect to oxygen since its decomposition into oxygen results in the liberation of heat (ΔH is negative) and an increase in entropy (ΔS is positive). These two effects reinforce each other, resulting in large negative Gibbs energy change (ΔG) for its conversion into oxygen. It is not really surprising, therefore, high concentrations of ozone can be dangerously explosive.

Due to the ease with which it liberates atoms of nascent oxygen ($\text{O}_3 \rightarrow \text{O}_2 + \text{O}$), it acts as a powerful oxidising agent. For example, it oxidises lead sulphide to lead sulphate and iodide ions to iodine.

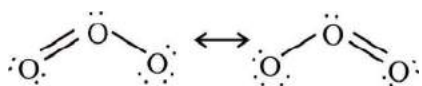


When ozone reacts with an excess of potassium iodide solution buffered with a borate buffer (pH 9.2), iodine is liberated which can be titrated against a standard solution of sodium thiosulphate. This is a quantitative method for estimating O_3 gas.

Experiments have shown that nitrogen oxides (particularly nitric oxide) combine very rapidly with ozone and there is, thus, the possibility that nitrogen oxides emitted from the exhaust systems of supersonic jet aeroplanes might be slowly depleting the concentration of the ozone layer in the upper atmosphere.



Another threat to this ozone layer is probably posed by the use of freons which are used in aerosol sprays and as refrigerants.



The two oxygen-oxygen bond lengths in the ozone molecule are identical (128 pm) and the molecule is angular as expected with a bond angle of about 117° . It is a resonance hybrid of two main forms:

Uses It is used as a germicide, disinfectant and for sterilising water. It is also used for bleaching oils, ivory, flour, starch, etc. It acts as an oxidising agent in the manufacture of potassium permanganate.

7.18 Why does O_3 act as a powerful oxidising agent?

7.19 How is O_3 estimated quantitatively?

7.1 Sulphur Allotropic Forms

Sulphur forms numerous allotropes of which the **yellow rhombic** (α -sulphur) and **monoclinic** (β -sulphur) forms are the most important. The stable form at room temperature is rhombic sulphur, which transforms to monoclinic sulphur when heated above 369 K.

Rhombic sulphur (α -sulphur)

This allotrope is yellow in colour, m.p. 385.8 K and specific gravity 2.06. Rhombic sulphur crystals are formed on evaporating the solution of roll sulphur in CS_2 . It is insoluble in water but dissolves to some extent in benzene, alcohol and ether. It is readily soluble in CS_2 .

Monoclinic sulphur (β -sulphur)

Its m.p. is 393 K and specific gravity 1.98. It is soluble in CS_2 . This form of sulphur is prepared by melting rhombic sulphur in a dish and cooling, till crust is formed. Two holes are made in the crust and the remaining liquid poured out. On removing the crust, colourless needle shaped crystals of β -sulphur are formed. It is stable above 369 K and transforms into α -sulphur below it. Conversely, α -sulphur is stable below 369 K and transforms into β -sulphur above this. At 369 K both the forms are stable. This temperature is called transition temperature.

Both rhombic and monoclinic sulphur have S_8 molecules. These S_8 molecules are packed to give different crystal structures. The S_8 ring in both the forms is puckered and has a crown shape. The molecular dimensions are given in Fig. 7.5(a).

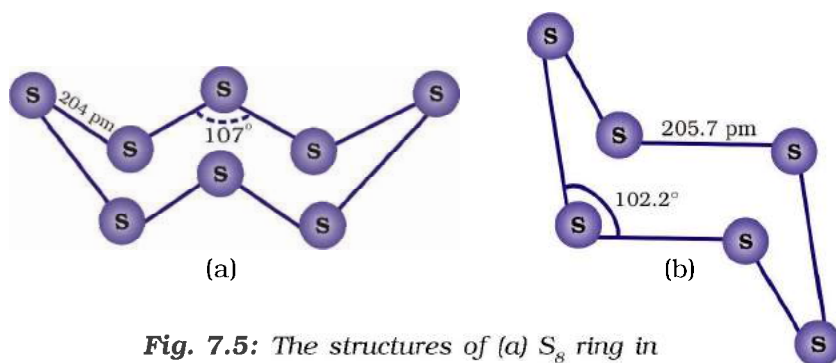


Fig. 7.5: The structures of (a) S_8 ring in rhombic sulphur and (b) S_6 form

Several other modifications of sulphur containing 6-20 sulphur atoms per ring have been synthesised in the last two decades. In cyclo- S_6 , the ring adopts the chair form and the molecular dimensions are as shown in Fig. 7.5 (b). At elevated temperatures (~ 1000 K), S_2 is the dominant species and is paramagnetic like O_2 .

Which form of sulphur shows paramagnetic behaviour ?

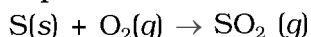
In vapour state sulphur partly exists as S_2 molecule which has two unpaired electrons in the antibonding π^* orbitals like O_2 and, hence, exhibits paramagnetism.

Example 7.1

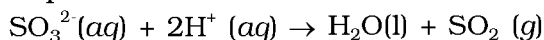
Solution

Preparation

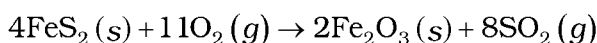
Sulphur dioxide is formed together with a little (6-8%) sulphur trioxide when sulphur is burnt in air or oxygen:



In the laboratory it is readily generated by treating a sulphite with dilute sulphuric acid.



Industrially, it is produced as a by-product of the roasting of sulphide ores.

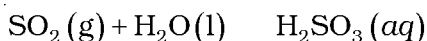


The gas after drying is liquefied under pressure and stored in steel cylinders.

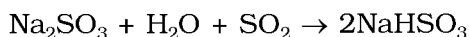
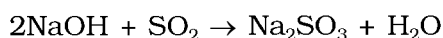
Properties

Sulphur dioxide is a colourless gas with pungent smell and is highly soluble in water. It liquefies at room temperature under a pressure of two atmospheres and boils at 263 K.

Sulphur dioxide, when passed through water, forms a solution of sulphurous acid.

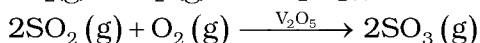
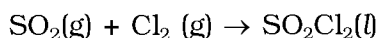


It reacts readily with sodium hydroxide solution, forming sodium sulphite, which then reacts with more sulphur dioxide to form sodium hydrogen sulphite.

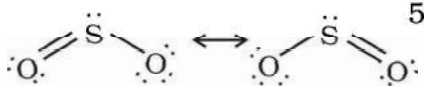
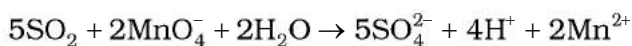
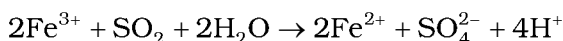


In its reaction with water and alkalis, the behaviour of sulphur dioxide is very similar to that of carbon dioxide.

Sulphur dioxide reacts with chlorine in the presence of charcoal (which acts as a catalyst) to give sulphuryl chloride, SO_2Cl_2 . It is oxidised to sulphur trioxide by oxygen in the presence of vanadium(V) oxide catalyst.



When moist, sulphur dioxide behaves as a reducing agent. For example, it converts iron(III) ions to iron(II) ions and decolourises acidified potassium permanganate(VII) solution; the latter reaction is a convenient test for the gas.



The molecule of SO_2 is angular. It is a resonance hybrid of the two canonical forms:

Uses Sulphur dioxide is used (i) in refining petroleum and sugar (ii) in bleaching wool and silk and (iii) as an anti-chlor, disinfectant and preservative. Sulphuric acid, sodium hydrogen sulphite and calcium hydrogen sulphite (industrial chemicals) are manufactured from sulphur dioxide. Liquid SO_2 is used as a solvent to dissolve a number of organic and inorganic chemicals.

- 7.20** What happens when sulphur dioxide is passed through an aqueous solution of Fe(III) salt?
- 7.21** Comment on the nature of two S–O bonds formed in SO₂ molecule. Are the two S–O bonds in this molecule equal ?
- 7.22** How is the presence of SO₂ detected ?

7.1 Oxoacids of Sulphur

Sulphur forms a number of oxoacids such as H₂SO₃, H₂S₂O₃, H₂S₂O₄, H₂S₂O₅, H₂S_xO₆ (x = 2 to 5), H₂SO₄, H₂S₂O₇, H₂SO₅, H₂S₂O₈. Some of these acids are unstable and cannot be isolated. They are known in aqueous solution or in the form of their salts. Structures of some important oxoacids are shown in Fig. 7.6.

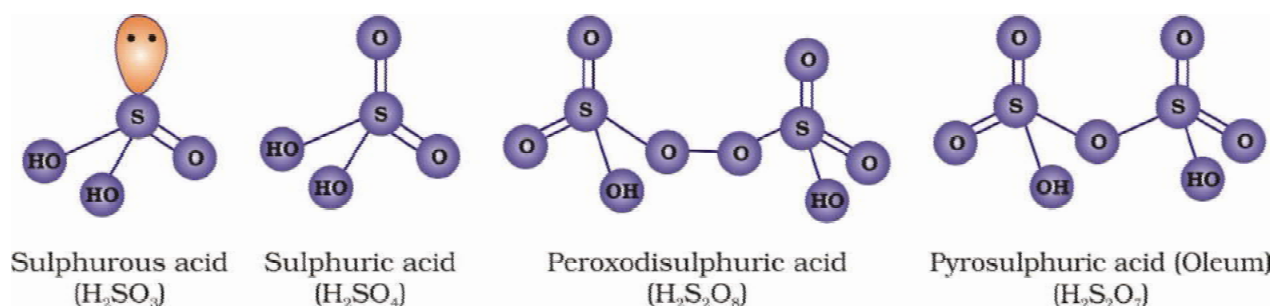


Fig. 7.6: Structures of some important oxoacids of sulphur

7.17 Sulphuric Acid

Manufacture

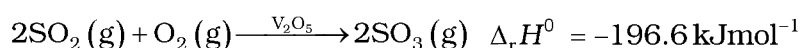
Sulphuric acid is one of the most important industrial chemicals worldwide.

Sulphuric acid is manufactured by the **Contact Process** which involves three steps:

- burning of sulphur or sulphide ores in air to generate SO₂.
- conversion of SO₂ to SO₃ by the reaction with oxygen in the presence of a catalyst (V₂O₅), and
- absorption of SO₃ in H₂SO₄ to give *Oleum* (H₂S₂O₇).

A flow diagram for the manufacture of sulphuric acid is shown in (Fig. 7.7). The SO₂ produced is purified by removing dust and other impurities such as arsenic compounds.

The key step in the manufacture of H₂SO₄ is the catalytic oxidation of SO₂ with O₂ to give SO₃ in the presence of V₂O₅ (catalyst).



The reaction is exothermic, reversible and the forward reaction leads to a decrease in volume. Therefore, low temperature and high pressure are the favourable conditions for maximum yield. But the temperature should not be very low otherwise rate of reaction will become slow.

In practice, the plant is operated at a pressure of 2 bar and a temperature of 720 K. The SO₃ gas from the catalytic converter is

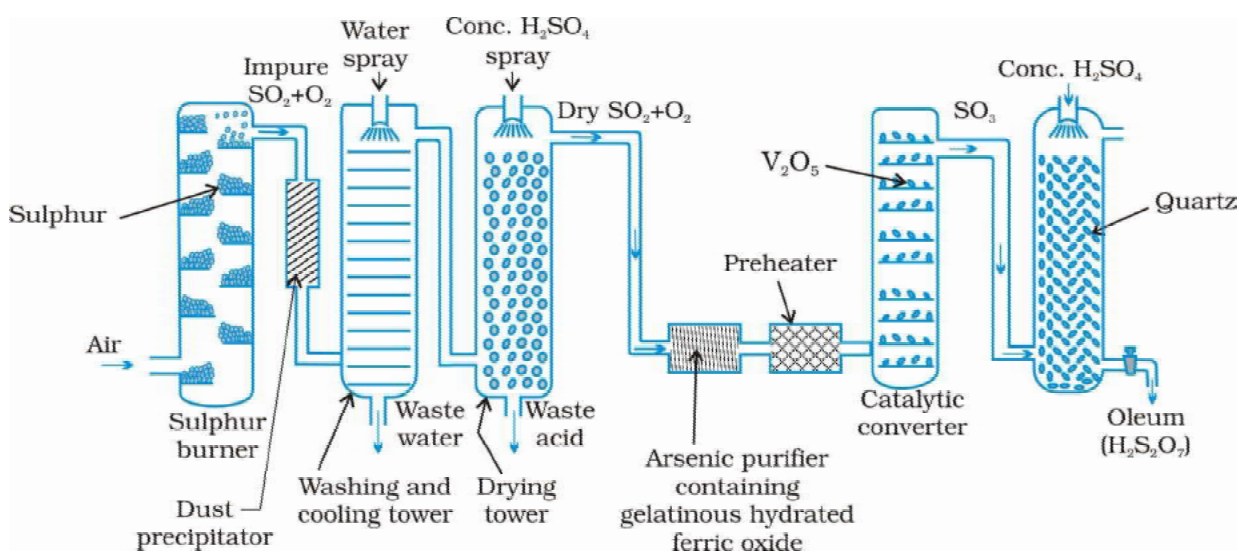
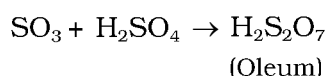


Fig. 7.7: Flow diagram for the manufacture of sulphuric acid

absorbed in concentrated H_2SO_4 to produce *oleum*. Dilution of oleum with water gives H_2SO_4 of the desired concentration. In the industry two steps are carried out simultaneously to make the process a continuous one and also to reduce the cost.

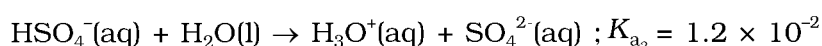


The sulphuric acid obtained by Contact process is 96-98% pure.

Properties

Sulphuric acid is a colourless, dense, oily liquid with a specific gravity of 1.84 at 298 K. The acid freezes at 283 K and boils at 611 K. It dissolves in water with the evolution of a large quantity of heat. Hence, care must be taken while preparing sulphuric acid solution from concentrated sulphuric acid. The concentrated acid must be added slowly into water with constant stirring.

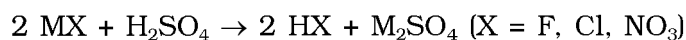
The chemical reactions of sulphuric acid are as a result of the following characteristics: (a) low volatility (b) strong acidic character (c) strong affinity for water and (d) ability to act as an oxidising agent. In aqueous solution, sulphuric acid ionises in two steps.



The larger value of K_{a_1} ($K_{a_1} > 10$) means that H_2SO_4 is largely dissociated into H^+ and HSO_4^- . Greater the value of dissociation constant (K_a), the stronger is the acid.

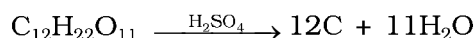
The acid forms two series of salts: normal sulphates (such as sodium sulphate and copper sulphate) and acid sulphates (e.g., sodium hydrogen sulphate).

Sulphuric acid, because of its low volatility can be used to manufacture more volatile acids from their corresponding salts.

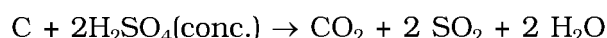
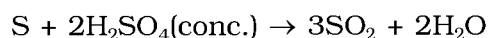
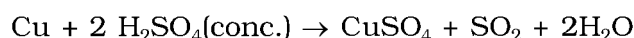


(M = Metal)

Concentrated sulphuric acid is a strong dehydrating agent. Many wet gases can be dried by passing them through sulphuric acid, provided the gases do not react with the acid. Sulphuric acid removes water from organic compounds; it is evident by its charring action on carbohydrates.



Hot concentrated sulphuric acid is a moderately strong oxidising agent. In this respect, it is intermediate between phosphoric and nitric acids. Both metals and non-metals are oxidised by concentrated sulphuric acid, which is reduced to SO_2 .

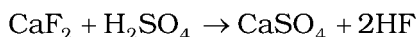


Uses □ Sulphuric acid is a very important industrial chemical. A nation's industrial strength can be judged by the quantity of sulphuric acid it produces and consumes. It is needed for the manufacture of hundreds of other compounds and also in many industrial processes. The bulk of sulphuric acid produced is used in the manufacture of fertilisers (e.g., ammonium sulphate, superphosphate). Other uses are in: (a) petroleum refining (b) manufacture of pigments, paints and dyestuff intermediates (c) detergent industry (d) metallurgical applications (e.g., cleansing metals before enameling, electroplating and galvanising (e) storage batteries (f) in the manufacture of nitrocellulose products and (g) as a laboratory reagent.

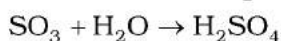
What happens when

- (i) Concentrated H_2SO_4 is added to calcium fluoride
- (ii) SO_3 is passed through water?

- (i) It forms hydrogen fluoride



- (ii) It dissolves SO_3 to give H_2SO_4 .



Example 7.13

Solution

Exercise 7.23

7.23 Mention three areas in which H_2SO_4 plays an important role.

7.24 Write the conditions to maximise the yield of H_2SO_4 by Contact process.

7.25 Why is $K_{a2} \ll K_{a1}$ for H_2SO_4 in water ?

7.1 Group 17
Elements

Fluorine, chlorine, bromine, iodine and astatine are members of Group 17. These are collectively known as the **halogens** (Greek *halo* means salt and *genes* means born i.e., salt producers). The halogens are highly reactive non-metallic elements. Like Groups 1 and 2, the elements of Group 17 show great similarity amongst themselves. That much similarity is not found in the elements of other groups of the periodic table. Also, there is a regular gradation in their physical and chemical properties. Astatine is a radioactive element.

7.18.1 Occurrence Fluorine and chlorine are fairly abundant while bromine and iodine less so. Fluorine is present mainly as insoluble fluorides (fluorspar CaF_2 , cryolite Na_3AlF_6 and fluoroapatite $3\text{Ca}_3(\text{PO}_4)_2 \cdot \text{CaF}_2$) and small quantities are present in soil, river water plants and bones and teeth of animals. Sea water contains chlorides, bromides and iodides of sodium, potassium, magnesium and calcium, but is mainly sodium chloride solution (2.5% by mass). The deposits of dried up seas contain these compounds, e.g., sodium chloride and carnallite, $\text{KCl} \cdot \text{MgCl}_2 \cdot 6\text{H}_2\text{O}$. Certain forms of marine life contain iodine in their systems; various seaweeds, for example, contain upto 0.5% of iodine and Chile saltpetre contains upto 0.2% of sodium iodate.

The important atomic and physical properties of Group 17 elements along with their electronic configurations are given in Table 7.8.

Table 7.8: Atomic and Physical Properties of Halogens

Property	F	Cl	Br	I	At ^a
Atomic number	9	17	35	53	85
Atomic mass/g mol ⁻¹	19.00	35.45	79.90	126.90	210
Electronic configuration	[He]2s ² 2p ⁵	[Ne]3s ² 3p ⁵	[Ar]3d ¹⁰ 4s ² 4p ⁵	[Kr]4d ¹⁰ 5s ² 5p ⁵	[Xe]4f ¹⁴ 5d ¹⁰ 6s ² 6p ⁵
Covalent radius/pm	64	99	114	133	–
Ionic radius X ⁻ /pm	133	184	196	220	–
Ionisation enthalpy/kJ mol ⁻¹	1680	1256	1142	1008	–
Electron gain enthalpy/kJ mol ⁻¹	–333	–349	–325	–296	–
Electronegativity ^b	4	3.2	3.0	2.7	2.2
Δ _{hyd} H(X ⁻)/kJ mol ⁻¹	515	381	347	305	–
	F ₂	Cl ₂	Br ₂	I ₂	–
Melting point/K	54.4	172.0	265.8	386.6	–
Boiling point/K	84.9	239.0	332.5	458.2	–
Density/g cm ⁻³	1.5 (85) ^c	1.66 (203) ^c	3.19(273) ^c	4.94(293) ^d	–
Distance X – X/pm	143	199	228	266	–
Bond dissociation enthalpy /(kJ mol ⁻¹)	158.8	242.6	192.8	151.1	–
E ^o /V ^e	2.87	1.36	1.09	0.54	–

^a Radioactive; ^b Pauling scale; ^c For the liquid at temperatures (K) given in the parentheses; ^d solid; ^e The half-cell reaction is $\text{X}_2(\text{g}) + 2\text{e}^- \rightarrow 2\text{X}(\text{aq})$.

The trends of some of the atomic, physical and chemical properties are discussed below.

- 7.18.2 Electronic Configuration** All these elements have seven electrons in their outermost shell ($ns^2 np^5$) which is one electron short of the next noble gas.
- 7.18.3 Atomic and Ionic Radii** The halogens have the smallest atomic radii in their respective periods due to maximum effective nuclear charge. The atomic radius of fluorine like the other elements of second period is extremely small. Atomic and ionic radii increase from fluorine to iodine due to increasing number of quantum shells.
- 7.18.4 Ionisation Enthalpy** They have little tendency to lose electron. Thus they have very high ionisation enthalpy. Due to increase in atomic size, ionisation enthalpy decreases down the group.
- 7.18.5 Electron Gain Enthalpy** Halogens have maximum negative electron gain enthalpy in the corresponding periods. This is due to the fact that the atoms of these elements have only one electron less than stable noble gas configurations. Electron gain enthalpy of the elements of the group becomes less negative down the group. However, the negative electron gain enthalpy of fluorine is less than that of chlorine. It is due to small size of fluorine atom. As a result, there are strong interelectronic repulsions in the relatively small $2p$ orbitals of fluorine and thus, the incoming electron does not experience much attraction.
- 7.18.6 Electronegativity** They have very high electronegativity. The electronegativity decreases down the group. Fluorine is the most electronegative element in the periodic table.

Halogens have maximum negative electron gain enthalpy in the [Example 7.1](#) respective periods of the periodic table. Why?

Halogens have the smallest size in their respective periods and therefore [solution](#) high effective nuclear charge. As a consequence, they readily accept one electron to acquire noble gas electronic configuration.

- 7.18.7 Physical Properties** Halogens display smooth variations in their physical properties. Fluorine and chlorine are gases, bromine is a liquid and iodine is a solid. Their melting and boiling points steadily increase with atomic number. All halogens are coloured. This is due to absorption of radiations in visible region which results in the excitation of outer electrons to higher energy level. By absorbing different quanta of radiation, they display different colours. For example, F_2 , has yellow, Cl_2 , greenish yellow, Br_2 , red and I_2 , violet colour. Fluorine and chlorine react with water. Bromine and iodine are only sparingly soluble in water but are soluble in various organic solvents such as chloroform, carbon tetrachloride, carbon disulphide and hydrocarbons to give coloured solutions.
- One curious anomaly we notice from Table 7.8 is the smaller enthalpy of dissociation of F_2 compared to that of Cl_2 whereas X-X bond dissociation enthalpies from chlorine onwards show the expected

trend: $\text{Cl} - \text{Cl} > \text{Br} - \text{Br} > \text{I} - \text{I}$. A reason for this anomaly is the relatively large electron-electron repulsion among the lone pairs in F_2 molecule where they are much closer to each other than in case of Cl_2 .

Example 7.15

Solution

Although electron gain enthalpy of fluorine is less negative as compared to chlorine, fluorine is a stronger oxidising agent than chlorine. Why?

It is due to

- (i) low enthalpy of dissociation of F-F bond (Table 7.8).
- (ii) high hydration enthalpy of F^- (Table 7.8).

7.18.8 Chemical Properties

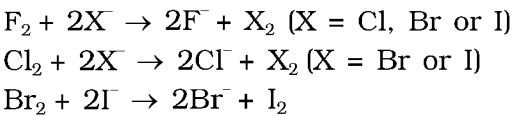
Oxidation states and trends in chemical reactivity
 All the halogens exhibit -1 oxidation state. However, chlorine, bromine and iodine exhibit $+1$, $+3$, $+5$ and $+7$ oxidation states also as explained below:

Halogen atom in ground state (other than fluorine)	ns	np	nd	
	$\uparrow\downarrow$	$\uparrow\downarrow\uparrow\downarrow\uparrow$		1 unpaired electron accounts for -1 or $+1$ oxidation states
First excited state	$\uparrow\downarrow$	$\uparrow\downarrow\uparrow\uparrow$	$\uparrow$	3 unpaired electrons account for $+3$ oxidation states
Second excited state	$\uparrow\downarrow$	$\uparrow\uparrow\uparrow$	$\uparrow\uparrow$	5 unpaired electrons account for $+5$ oxidation state
Third excited state	$\uparrow$	$\uparrow\uparrow\uparrow$	$\uparrow\uparrow\uparrow$	7 unpaired electrons account for $+7$ oxidation state

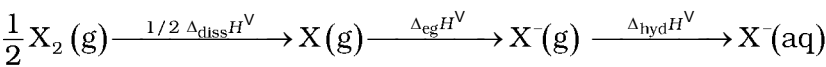
The higher oxidation states of chlorine, bromine and iodine are realised mainly when the halogens are in combination with the small and highly electronegative fluorine and oxygen atoms. e.g., in interhalogens, oxides and oxoacids. The oxidation states of $+4$ and $+6$ occur in the oxides and oxoacids of chlorine and bromine. The fluorine atom has no d orbitals in its valence shell and therefore cannot expand its octet. Being the most electronegative, it exhibits only -1 oxidation state.

All the halogens are highly reactive. They react with metals and non-metals to form halides. The reactivity of the halogens decreases down the group.

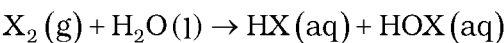
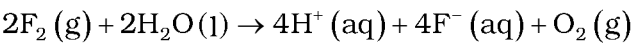
The ready acceptance of an electron is the reason for the strong oxidising nature of halogens. F_2 is the strongest oxidising halogen and it oxidises other halide ions in solution or even in the solid phase. In general, a halogen oxidises halide ions of higher atomic number.



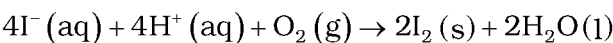
The decreasing oxidising ability of the halogens in aqueous solution down the group is evident from their standard electrode potentials (Table 7.8) which are dependent on the parameters indicated below:



The relative oxidising power of halogens can further be illustrated by their reactions with water. Fluorine oxidises water to oxygen whereas chlorine and bromine react with water to form corresponding hydrohalic and hypohalous acids. The reaction of iodine with water is non-spontaneous. In fact, I⁻ can be oxidised by oxygen in acidic medium; just the reverse of the reaction observed with fluorine.



(where X = Cl or Br)



Anomalous behaviour of fluorine

Like other elements of *p*-block present in second period of the periodic table, fluorine is anomalous in many properties. For example, ionisation enthalpy, electronegativity, and electrode potentials are all higher for fluorine than expected from the trends set by other halogens. Also, ionic and covalent radii, m.p. and b.p., enthalpy of bond dissociation and electron gain enthalpy are quite lower than expected. The anomalous behaviour of fluorine is due to its small size, highest electronegativity, low F-F bond dissociation enthalpy, and non availability of *d* orbitals in valence shell.

Most of the reactions of fluorine are exothermic (due to the small and strong bond formed by it with other elements). It forms only one oxoacid while other halogens form a number of oxoacids. Hydrogen fluoride is a liquid (b.p. 293 K) due to strong hydrogen bonding. Other hydrogen halides are gases.

- (i) *Reactivity towards hydrogen*: They all react with hydrogen to give hydrogen halides but affinity for hydrogen decreases from fluorine to iodine. Hydrogen halides dissolve in water to form hydrohalic acids. Some of the properties of hydrogen halides are given in Table 7.9. The acidic strength of these acids varies in the order: HF < HCl < HBr < HI. The stability of these halides decreases down the group due to decrease in bond (H-X) dissociation enthalpy in the order: H-F > H-Cl > H-Br > H-I.

Table 7.9: Properties of Hydrogen Halides

Property	HF	HCl	HBr	HI
Melting point/K	190	159	185	222
Boiling point/K	293	189	206	238
Bond length (H – X)/pm	91.7	127.4	141.4	160.9
$\Delta_{\text{diss}} H^{\circ} / \text{kJ mol}^{-1}$	574	432	363	295
pK_{a}	3.2	–7.0	–9.5	–10.0

- (ii) *Reactivity towards oxygen*: Halogens form many oxides with oxygen but most of them are unstable. Fluorine forms two oxides OF₂ and O₂F₂. However, only OF₂ is thermally stable at 298 K. These oxides

are essentially oxygen fluorides because of the higher electronegativity of fluorine than oxygen. Both are strong fluorinating agents. O_2F_2 oxidises plutonium to PuF_6 and the reaction is used in removing plutonium as PuF_6 from spent nuclear fuel.

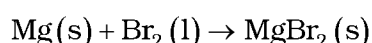
Chlorine, bromine and iodine form oxides in which the oxidation states of these halogens range from +1 to +7. A combination of kinetic and thermodynamic factors lead to the generally decreasing order of stability of oxides formed by halogens, $I > Cl > Br$. The higher oxides of halogens tend to be more stable than the lower ones.

Chlorine oxides, Cl_2O , ClO_2 , Cl_2O_6 and Cl_2O_7 are highly reactive oxidising agents and tend to explode. ClO_2 is used as a bleaching agent for paper pulp and textiles and in water treatment.

The bromine oxides, Br_2O , BrO_2 , BrO_3 are the least stable halogen oxides (middle row anomaly) and exist only at low temperatures. They are very powerful oxidising agents.

The iodine oxides, I_2O_4 , I_2O_5 , I_2O_7 are insoluble solids and decompose on heating. I_2O_5 is a very good oxidising agent and is used in the estimation of carbon monoxide.

- (iii) *Reactivity towards metals*: Halogens react with metals to form metal halides. For example, bromine reacts with magnesium to give magnesium bromide.



The ionic character of the halides decreases in the order $MF > MCl > MBr > MI$ where M is a monovalent metal. If a metal exhibits more than one oxidation state, the halides in higher oxidation state will be more covalent than the one in lower oxidation state. For example, $SnCl_4$, $PbCl_4$, $SbCl_5$ and UF_6 are more covalent than $SnCl_2$, $PbCl_2$, $SbCl_3$ and UF_4 respectively.

- (iv) *Reactivity of halogens towards other halogens*: Halogens combine amongst themselves to form a number of compounds known as interhalogens of the types XX' , XX_3 , XX_5 and XX_7 where X is a larger size halogen and X' is smaller size halogen.

Example 7.1

Fluorine exhibits only -1 oxidation state whereas other halogens exhibit +1, +3, +5 and +7 oxidation states also. Explain.

Solution

Fluorine is the most electronegative element and cannot exhibit any positive oxidation state. Other halogens have *d* orbitals and therefore, can expand their octets and show +1, +3, +5 and +7 oxidation states also.

Text Questions

- 7.26** Considering the parameters such as bond dissociation enthalpy, electron gain enthalpy and hydration enthalpy, compare the oxidising power of F_2 and Cl_2 .
- 7.27** Give two examples to show the anomalous behaviour of fluorine.
- 7.28** Sea is the greatest source of some halogens. Comment.

Chlorine was discovered in 1774 by Scheele by the action of HCl on MnO_2 . In 1810 Davy established its elementary nature and suggested the name chlorine on account of its colour (Greek, *chloros* = greenish yellow).

Preparation

It can be prepared by any one of the following methods:

- (i) By heating manganese dioxide with concentrated hydrochloric acid.



However, a mixture of common salt and concentrated H_2SO_4 is used in place of HCl.

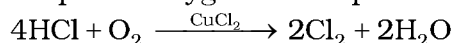


- (ii) By the action of HCl on potassium permanganate.



Manufacture of chlorine

- (i) *Deacon's process*: By oxidation of hydrogen chloride gas by atmospheric oxygen in the presence of CuCl_2 (catalyst) at 723 K.



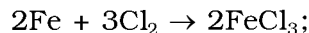
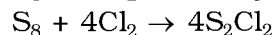
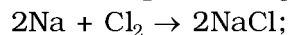
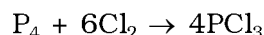
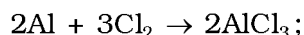
- (ii) *Electrolytic process*: Chlorine is obtained by the electrolysis of brine (concentrated NaCl solution). Chlorine is liberated at anode.

It is also obtained as a by-product in many chemical industries.

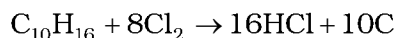
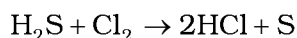
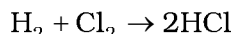
Properties

It is a greenish yellow gas with pungent and suffocating odour. It is about 2-5 times heavier than air. It can be liquefied easily into greenish yellow liquid which boils at 239 K. It is soluble in water.

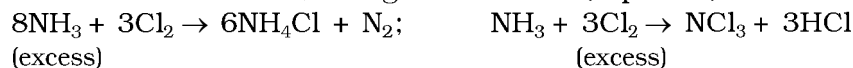
Chlorine reacts with a number of metals and non-metals to form chlorides.



It has great affinity for hydrogen. It reacts with compounds containing hydrogen to form HCl.



With excess ammonia, chlorine gives nitrogen and ammonium chloride whereas with excess chlorine, nitrogen trichloride (explosive) is formed.



With cold and dilute alkalies chlorine produces a mixture of chloride and hypochlorite but with hot and concentrated alkalies it gives chloride and chlorate.

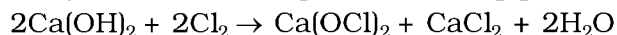


(cold and dilute)



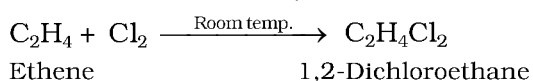
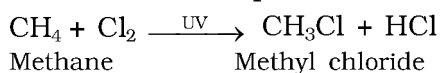
(hot and conc.)

With dry slaked lime it gives bleaching powder.



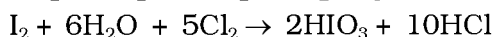
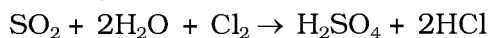
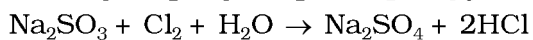
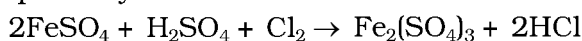
The composition of bleaching powder is $\text{Ca}(\text{OCl})_2 \cdot \text{CaCl}_2 \cdot \text{Ca}(\text{OH})_2 \cdot 2\text{H}_2\text{O}$.

Chlorine reacts with hydrocarbons and gives substitution products with saturated hydrocarbons and addition products with unsaturated hydrocarbons. For example,

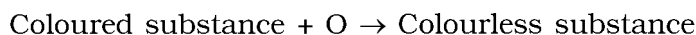
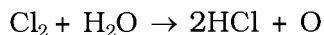


Chlorine water on standing loses its yellow colour due to the formation of HCl and HOCl. Hypochlorous acid (HOCl) so formed, gives nascent oxygen which is responsible for oxidising and bleaching properties of chlorine.

- (i) It oxidises ferrous to ferric and sulphite to sulphate. Chlorine oxidises sulphur dioxide to sulphur trioxide and iodine to iodate. In the presence of water they form sulphuric acid and iodic acid respectively.



- (ii) It is a powerful bleaching agent; bleaching action is due to oxidation.



Uses It is used (i) for bleaching woodpulp (required for the manufacture of paper and rayon), bleaching cotton and textiles, (ii) in the extraction of gold and platinum (iii) in the manufacture of dyes, drugs and organic compounds such as CCl_4 , CHCl_3 , DDT, refrigerants, etc. (iv) in sterilising drinking water and (v) preparation of poisonous gases such as phosgene (COCl_2), tear gas (CCl_3NO_2), mustard gas ($\text{ClCH}_2\text{CH}_2\text{SCH}_2\text{CH}_2\text{Cl}$).

Example 7.17 Write the balanced chemical equation for the reaction of Cl_2 with hot and concentrated NaOH. Is this reaction a disproportionation reaction? Justify.

Solution $3\text{Cl}_2 + 6\text{NaOH} \rightarrow 5\text{NaCl} + \text{NaClO}_3 + 3\text{H}_2\text{O}$
Yes, chlorine from zero oxidation state is changed to -1 and +5 oxidation states.

Inte t uestions

7.29 Give the reason for bleaching action of Cl_2 .

7.30 Name two poisonous gases which can be prepared from chlorine gas.

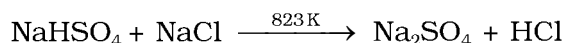
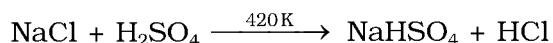
7. Chlorine Chloride

It bleaches vegetable or organic matter in the presence of moisture. Bleaching effect of chlorine is permanent.

Glauber prepared this acid in 1648 by heating common salt with concentrated sulphuric acid. Davy in 1810 showed that it is a compound of hydrogen and chlorine.

Preparation

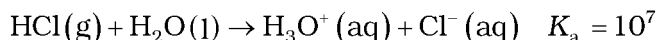
In laboratory, it is prepared by heating sodium chloride with concentrated sulphuric acid.



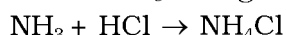
HCl gas can be dried by passing through concentrated sulphuric acid.

Properties

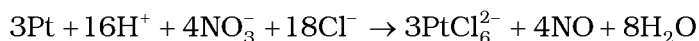
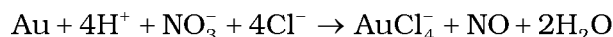
It is a colourless and pungent smelling gas. It is easily liquefied to a colourless liquid (b.p. 189 K) and freezes to a white crystalline solid (f.p. 159 K). It is extremely soluble in water and ionises as follows:



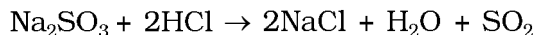
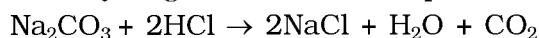
Its aqueous solution is called hydrochloric acid. High value of dissociation constant (K_a) indicates that it is a strong acid in water. It reacts with NH_3 and gives white fumes of NH_4Cl .



When three parts of concentrated HCl and one part of concentrated HNO_3 are mixed, aqua regia is formed which is used for dissolving noble metals, e.g., gold, platinum.



Hydrochloric acid decomposes salts of weaker acids, e.g., carbonates, hydrogencarbonates, sulphites, etc.



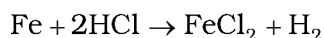
Uses It is used (i) in the manufacture of chlorine, NH_4Cl and glucose (from corn starch), (ii) for extracting glue from bones and purifying bone black, (iii) in medicine and as a laboratory reagent.

When HCl reacts with finely powdered iron, it forms ferrous chloride and not ferric chloride. Why?

[Example 7.1](#)

Its reaction with iron produces H_2 .

[Solution](#)



Liberation of hydrogen prevents the formation of ferric chloride.

7.1 Oxoacids of halogens

Due to high electronegativity and small size, fluorine forms only one oxoacid, HOF known as fluoric (I) acid or hypofluorous acid. The other halogens form several oxoacids. Most of them cannot be isolated in pure state. They are stable only in aqueous solutions or in the form of their salts. The oxoacids of halogens are given in Table 7.10 and their structures are given in Fig. 7.8.

Table 7.10: Oxoacids of Halogens

Halic (I) acid (Hypohalous acid)	HOX (Hypofluorous acid)	HOX (Hypochlorous acid)	HOX (Hypobromous acid)	HOX (Hypoiodous acid)
Halic (III) acid (Halous acid)	– –	HOX (chlorous acid)	– –	– –
Halic (V) acid (Halic acid)	– –	HOX ₂ (chloric acid)	HOX ₂ (bromic acid)	HOX ₂ (iodic acid)
Halic (VII) acid (Perhalic acid)	– –	HOX ₃ (perchloric acid)	HOX ₃ (perbromic acid)	HOX ₃ (periodic acid)

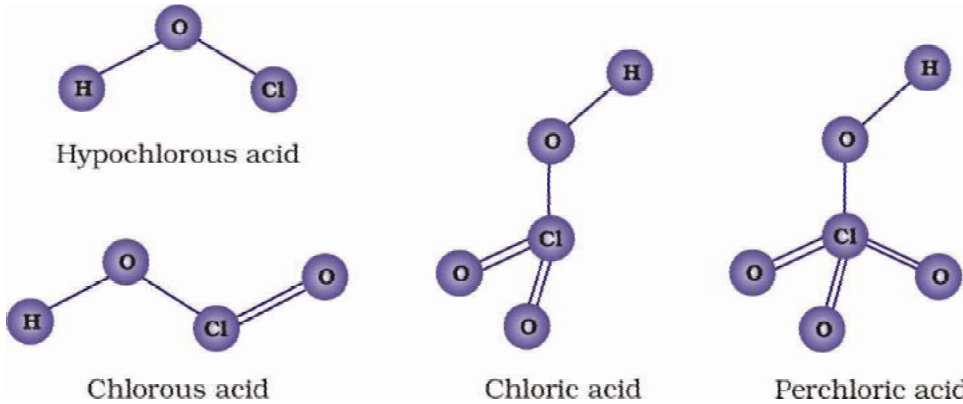


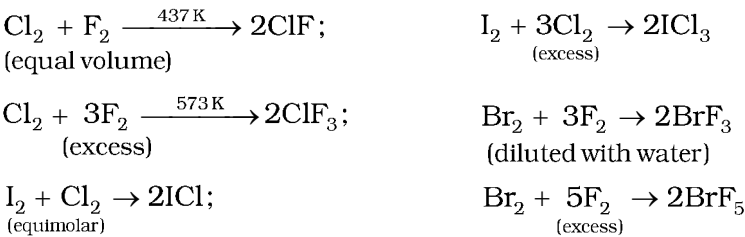
Fig. 7.8
The structures of
oxoacids of chlorine

7. Interhalogen compounds

When two different halogens react with each other, interhalogen compounds are formed. They can be assigned general compositions as XX' , XX'_3 , XX'_5 and XX'_7 where X is halogen of larger size and X' of smaller size and X is more electropositive than X'. As the ratio between radii of X and X' increases, the number of atoms per molecule also increases. Thus, iodine (VII) fluoride should have maximum number of atoms as the ratio of radii between I and F should be maximum. That is why its formula is IF_7 (having maximum number of atoms).

Preparation

The interhalogen compounds can be prepared by the direct combination or by the action of halogen on lower interhalogen compounds. The product formed depends upon some specific conditions, For e.g.,



Properties

Some properties of interhalogen compounds are given in Table 7.11.

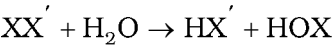
Table 7.11: Some Properties of Interhalogen Compounds

Type	Formula	Physical state and colour	Structure
XX' ₁	ClF	colourless gas	–
	BrF	pale brown gas	–
	IF ^a	detected spectroscopically	–
	BrCl ^b	gas	–
	ICl	ruby red solid (α-form)	–
		brown red solid (β-form)	–
	IBr	black solid	–
XX' ₃	ClF ₃	colourless gas	Bent T-shaped
	BrF ₃	yellow green liquid	Bent T-shaped
	IF ₃	yellow powder	Bent T-shaped (?)
	ICl ₃ ^c	orange solid	Bent T-shaped (?)
XX' ₅	IF ₅	colourless gas but solid below 77 K	Square pyramidal
	BrF ₅	colourless liquid	Square pyramidal
	ClF ₅	colourless liquid	Square pyramidal
XX' ₇	IF ₇	colourless gas	Pentagonal bipyramidal

^aVery unstable; ^bThe pure solid is known at room temperature; ^cDimerises as Cl-bridged dimer (I₂Cl₆)

These are all covalent molecules and are diamagnetic in nature. They are volatile solids or liquids at 298 K except ClF which is a gas. Their physical properties are intermediate between those of constituent halogens except that their m.p. and b.p. are a little higher than expected.

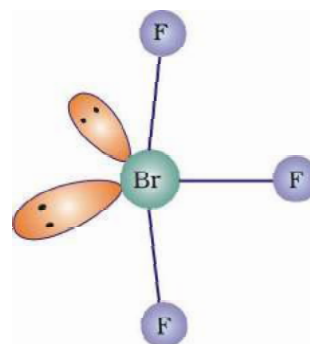
Their chemical reactions can be compared with the individual halogens. In general, interhalogen compounds are more reactive than halogens (except fluorine). This is because X-X' bond in interhalogens is weaker than X-X bond in halogens except F-F bond. All these undergo hydrolysis giving halide ion derived from the smaller halogen and a hypohalite (when XX'), halite (when XX'₃), halate (when XX'₅) and perhalate (when XX'₇) anion derived from the larger halogen.



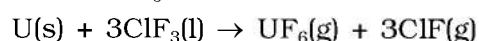
Their molecular structures are very interesting which can be explained on the basis of VSEPR theory (Example 7.19). The XX₃ compounds have the bent ‘T’ shape, XX₅ compounds square pyramidal and IF₇ has pentagonal bipyramidal structures (Table 7.11).

Example 7.1 Discuss the molecular shape of BrF_3 on the basis of VSEPR theory.

Solution The central atom Br has seven electrons in the valence shell. Three of these will form electron-pair bonds with three fluorine atoms leaving behind four electrons. Thus, there are three bond pairs and two lone pairs. According to VSEPR theory, these will occupy the corners of a trigonal bipyramid. The two lone pairs will occupy the equatorial positions to minimise lone pair-lone pair and the bond pair-lone pair repulsions which are greater than the bond pair-bond pair repulsions. In addition, the axial fluorine atoms will be bent towards the equatorial fluorine in order to minimise the lone-pair-lone pair repulsions. The shape would be that of a slightly bent 'T'.



Uses These compounds can be used as non aqueous solvents. Interhalogen compounds are very useful fluorinating agents. ClF_3 and BrF_3 are used for the production of UF_6 in the enrichment of ^{235}U .



Inte t uestion

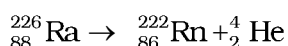
7.31 Why is ICl more reactive than I_2 ?

7. Group 18 Elements

Group 18 consists of six elements: helium, neon, argon, krypton, xenon and radon. All these are gases and chemically unreactive. They form very few compounds. Because of this they are termed noble gases.

7.23.1 Occurrence

All the noble gases except radon occur in the atmosphere. Their atmospheric abundance in dry air is ~ 1% by volume of which argon is the major constituent. Helium and sometimes neon are found in minerals of radioactive origin e.g., pitchblende, monazite, cleveite. The main commercial source of helium is natural gas. Xenon and radon are the rarest elements of the group. Radon is obtained as a decay product of ^{226}Ra .



Example 7. Why are the elements of Group 18 known as noble gases ?

Solution The elements present in Group 18 have their valence shell orbitals completely filled and, therefore, react with a few elements only under certain conditions. Therefore, they are now known as noble gases.

The important atomic and physical properties of the Group 18 elements along with their electronic configurations are given in Table 7.12. The trends in some of the atomic, physical and chemical properties of the group are discussed here.

Table 7.12: Atomic and Physical Properties of Group 18 Elements

Property	He	Ne	Ar	Kr	Xe	Rn*
Atomic number	2	10	18	36	54	86
Atomic mass/ g mol ⁻¹	4.00	20.18	39.95	83.80	131.30	222.00
Electronic configuration	1s ²	[He]2s ² 2p ⁶	[Ne] 3s ² 3p ⁶	[Ar]3d ¹⁰ 4s ² 4p ⁶	[Kr]4d ¹⁰ 5s ² 5p ⁶	[Xe]4f ¹⁴ 5d ¹⁰ 6s ² 6p ⁶
Atomic radius/pm	120	160	190	200	220	–
Ionisation enthalpy /kJmol ⁻¹	2372	2080	1520	1351	1170	1037
Electron gain enthalpy /kJmol ⁻¹	48	116	96	96	77	68
Density (at STP)/gcm ⁻³	1.8×10 ⁻⁴	9.0×10 ⁻⁴	1.8×10 ⁻³	3.7×10 ⁻³	5.9×10 ⁻³	9.7×10 ⁻³
Melting point/K	–	24.6	83.8	115.9	161.3	202
Boiling point/K	4.2	27.1	87.2	119.7	165.0	211
Atmospheric content (% by volume)	5.24×10 ⁻⁴	–	1.82×10 ⁻³	0.934	1.14×10 ⁻⁴	8.7×10 ⁻⁶

* radioactive

- 7.23.2 Electronic Configuration

All noble gases have general electronic configuration ns^2np^6 except helium which has $1s^2$ (Table 7.12). Many of the properties of noble gases including their inactive nature are ascribed to their closed shell structures.
- 7.23.3 Ionisation Enthalpy

Due to stable electronic configuration these gases exhibit very high ionisation enthalpy. However, it decreases down the group with increase in atomic size.
- 7.23.4 Atomic Radii

Atomic radii increase down the group with increase in atomic number.
- 7.23.5 Electron Gain Enthalpy

Since noble gases have stable electronic configurations, they have no tendency to accept the electron and therefore, have large positive values of electron gain enthalpy.

Physical Properties

All the noble gases are monoatomic. They are colourless, odourless and tasteless. They are sparingly soluble in water. They have very low melting and boiling points because the only type of interatomic interaction in these elements is weak dispersion forces. Helium has the lowest boiling point (4.2 K) of any known substance. It has an unusual property of diffusing through most commonly used laboratory materials such as rubber, glass or plastics.

Noble gases have very low boiling points. Why?

Noble gases being monoatomic have no interatomic forces except weak dispersion forces and therefore, they are liquefied at very low temperatures. Hence, they have low boiling points.

Example 7. 1
Solution

Chemical Properties

In general, noble gases are least reactive. Their inertness to chemical reactivity is attributed to the following reasons:

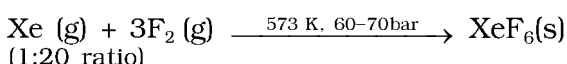
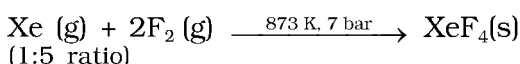
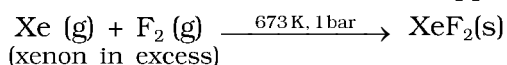
- The noble gases except helium ($1s^2$) have completely filled ns^2np^6 electronic configuration in their valence shell.
- They have high ionisation enthalpy and more positive electron gain enthalpy.

The reactivity of noble gases has been investigated occasionally, ever since their discovery, but all attempts to force them to react to form the compounds, were unsuccessful for quite a few years. In March 1962, Neil Bartlett, then at the University of British Columbia, observed the reaction of a noble gas. First, he prepared a red compound which is formulated as $O_2^+PtF_6^-$. He, then realised that the first ionisation enthalpy of molecular oxygen (1175 kJ mol^{-1}) was almost identical with that of xenon (1170 kJ mol^{-1}). He made efforts to prepare same type of compound with Xe and was successful in preparing another red colour compound $Xe^+PtF_6^-$ by mixing PtF_6 and xenon. After this discovery, a number of xenon compounds mainly with most electronegative elements like fluorine and oxygen, have been synthesised.

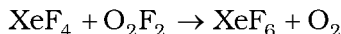
The compounds of krypton are fewer. Only the difluoride (KrF_2) has been studied in detail. Compounds of radon have not been isolated but only identified (e.g., RnF_2) by radiotracer technique. No true compounds of Ar, Ne or He are yet known.

(a) Xenon-fluorine compounds

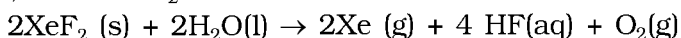
Xenon forms three binary fluorides, XeF_2 , XeF_4 and XeF_6 by the direct reaction of elements under appropriate experimental conditions.



XeF_6 can also be prepared by the interaction of XeF_4 and O_2F_2 at 143K.

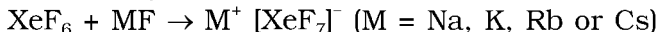
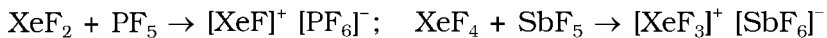


XeF_2 , XeF_4 and XeF_6 are colourless crystalline solids and sublime readily at 298 K. They are powerful fluorinating agents. They are readily hydrolysed even by traces of water. For example, XeF_2 is hydrolysed to give Xe, HF and O_2 .



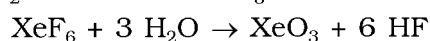
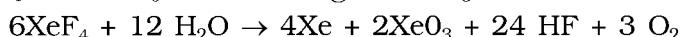
The structures of the three xenon fluorides can be deduced from VSEPR and these are shown in Fig. 7.9. XeF_2 and XeF_4 have linear and square planar structures respectively. XeF_6 has seven electron pairs (6 bonding pairs and one lone pair) and would, thus, have a distorted octahedral structure as found experimentally in the gas phase.

Xenon fluorides react with fluoride ion acceptors to form cationic species and fluoride ion donors to form fluoroanions.

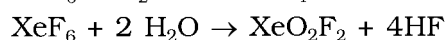
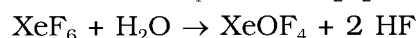


(b) Xenon-oxygen compounds

Hydrolysis of XeF_4 and XeF_6 with water gives XeO_3 .



Partial hydrolysis of XeF_6 gives oxyfluorides, XeOF_4 and XeO_2F_2 .



XeO_3 is a colourless explosive solid and has a pyramidal molecular structure (Fig. 7.9). XeOF_4 is a colourless volatile liquid and has a square pyramidal molecular structure (Fig. 7.9).

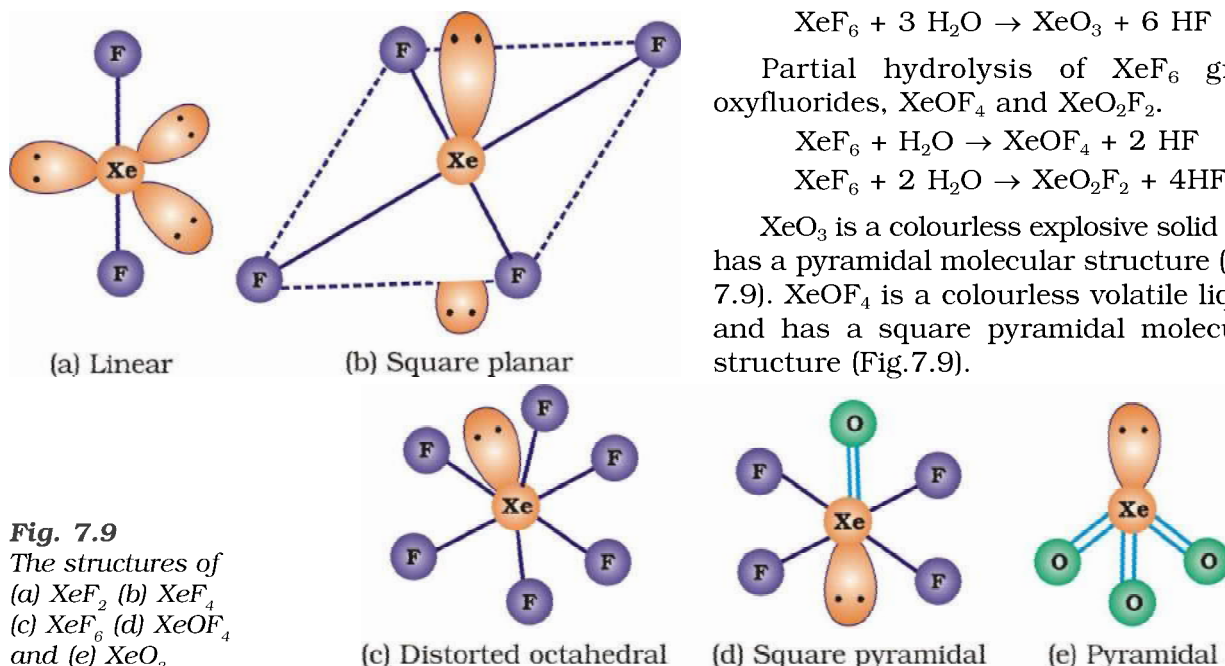


Fig. 7.9
The structures of
(a) XeF_2 (b) XeF_4
(c) XeF_6 (d) XeOF_4
and (e) XeO_3

Does the hydrolysis of XeF_6 lead to a redox reaction?

No, the products of hydrolysis are XeOF_4 and XeO_2F_2 where the oxidation states of all the elements remain the same as it was in the reacting state.

Example 7.

Solution

Uses Helium is a non-inflammable and light gas. Hence, it is used in filling balloons for meteorological observations. It is also used in gas-cooled nuclear reactors. Liquid helium (b.p. 4.2 K) finds use as cryogenic agent for carrying out various experiments at low temperatures. It is used to produce and sustain powerful superconducting magnets which form an essential part of modern NMR spectrometers and Magnetic Resonance Imaging (MRI) systems for clinical diagnosis. It is used as a diluent for oxygen in modern diving apparatus because of its very low solubility in blood.

Neon is used in discharge tubes and fluorescent bulbs for advertisement display purposes. Neon bulbs are used in botanical gardens and in green houses.

Argon is used mainly to provide an inert atmosphere in high temperature metallurgical processes (arc welding of metals or alloys) and for filling electric bulbs. It is also used in the laboratory for handling substances that are air-sensitive. There are no significant uses of Xenon and Krypton. They are used in light bulbs designed for special purposes.

Exercises

7.32 Why is helium used in diving apparatus?

7.33 Balance the following equation: $\text{XeF}_6 + \text{H}_2\text{O} \rightarrow \text{XeO}_2\text{F}_2 + \text{HF}$

7.34 Why has it been difficult to study the chemistry of radon?

Groups 13 to 18 of the periodic table consist of **p-block elements** with their valence shell electronic configuration ns^2np^{1-6} . Groups 13 and 14 were dealt with in Class XI. In this Unit remaining groups of the p-block have been discussed.

Group 15 consists of five elements namely, N, P, As, Sb and Bi which have general electronic configuration ns^2np^3 . Nitrogen differs from other elements of this group due to small size, formation of **$p\pi-p\pi$ multiple bonds** with itself and with highly electronegative atom like O or C and **non-availability of d orbitals** to expand its valence shell. Elements of group 15 show gradation in properties. They react with oxygen, hydrogen and halogens. They exhibit two important oxidation states, +3 and +5 but +3 oxidation is favoured by heavier elements due to 'inert pair effect'.

Dinitrogen can be prepared in laboratory as well as on industrial scale. It forms oxides in various oxidation states as N_2O , NO, N_2O_3 , NO_2 , N_2O_4 and N_2O_5 . These oxides have **resonating structures** and have multiple bonds. Ammonia can be prepared on large scale by **Haber's process**. HNO_3 is an important industrial chemical. It is a strong monobasic acid and is a powerful oxidising agent. Metals and non-metals react with HNO_3 under different conditions to give NO or NO_2 .

Phosphorus exists as P_4 in elemental form. It exists in several **allotropic forms**. It forms hydride, PH_3 which is a highly poisonous gas. It forms two types of halides as PX_3 and PX_5 . PCl_3 is prepared by the reaction of white phosphorus with dry chlorine while PCl_5 is prepared by the reaction of phosphorus with SO_2Cl_2 . Phosphorus forms a number of oxoacids. Depending upon the number of P-OH groups, their basicity varies. The oxoacids which have P-H bonds are good reducing agents.

The Group 16 elements have general electronic configuration ns^2np^4 . They show maximum oxidation state, +6. Gradation in physical and chemical properties is observed in the group 16 elements. In laboratory, dioxygen is prepared by heating $KClO_3$ in presence of MnO_2 . It forms a number of oxides with metals. Allotropic form of oxygen is O_3 which is a highly oxidising agent. Sulphur forms a number of allotropes. Of these, α - and β - forms of sulphur are the most important. Sulphur combines with oxygen to give oxides such as SO_2 and SO_3 . SO_2 is prepared by the direct union of sulphur with oxygen. SO_2 is used in the manufacture of H_2SO_4 . Sulphur forms a number of oxoacids. Amongst them, the most important is H_2SO_4 . It is prepared by **contact process**. It is a dehydrating and oxidising agent. It is used in the manufacture of several compounds.

Group 17 of the periodic table consists of the following elements F, Cl, Br, I and At. These elements are extremely reactive and as such they are found in the combined state only. The common oxidation state of these elements is -1. However, highest oxidation state can be +7. They show regular gradation in physical and chemical properties. They form oxides, hydrogen halides, interhalogen compounds and oxoacids. Chlorine is conveniently obtained by the reaction of HCl with $KMnO_4$. HCl is prepared by heating NaCl with concentrated H_2SO_4 . Halogens combine with one another to form **interhalogen compounds** of the type XX'_n ($n = 1, 3, 5, 7$) where X' is lighter than X. A number of oxoacids of halogens are known. In the structures of these oxoacids, halogen is the central atom which is bonded in each case with one OH bond as X-OH. In some cases X = O bonds are also found.

Group 18 of the periodic table consists of **noble gases**. They have ns^2np^6 valence shell electronic configuration except He which has $1s^2$. All the gases except Rn occur in atmosphere. Rn is obtained as the decay product of ^{226}Ra .

Due to complete octet of outermost shell, they have less tendency to form compounds. The best characterised compounds are those of xenon with fluorine and oxygen only under certain conditions. These gases have several uses. Argon is used to provide inert atmosphere, helium is used in filling balloons for meteorological observations, neon is used in discharge tubes and fluorescent bulbs.

- 7.1 Discuss the general characteristics of Group 15 elements with reference to their electronic configuration, oxidation state, atomic size, ionisation enthalpy and electronegativity.
- 7.2 Why does the reactivity of nitrogen differ from phosphorus?
- 7.3 Discuss the trends in chemical reactivity of group 15 elements.
- 7.4 Why does NH_3 form hydrogen bond but PH_3 does not?
- 7.5 How is nitrogen prepared in the laboratory? Write the chemical equations of the reactions involved.
- 7.6 How is ammonia manufactured industrially?
- 7.7 Illustrate how copper metal can give different products on reaction with HNO_3 .
- 7.8 Give the resonating structures of NO_2 and N_2O_5 .
- 7.9 The HNH angle value is higher than HPH, HAsH and HSbH angles. Why? [Hint: Can be explained on the basis of sp^3 hybridisation in NH_3 and only s-p bonding between hydrogen and other elements of the group].
- 7.10 Why does $\text{R}_3\text{P} = \text{O}$ exist but $\text{R}_3\text{N} = \text{O}$ does not (R = alkyl group)?
- 7.11 Explain why NH_3 is basic while BiH_3 is only feebly basic.
- 7.12 Nitrogen exists as diatomic molecule and phosphorus as P_4 . Why?
- 7.13 Write main differences between the properties of white phosphorus and red phosphorus.
- 7.14 Why does nitrogen show catenation properties less than phosphorus?
- 7.15 Give the disproportionation reaction of H_3PO_3 .
- 7.16 Can PCl_5 act as an oxidising as well as a reducing agent? Justify.
- 7.17 Justify the placement of O, S, Se, Te and Po in the same group of the periodic table in terms of electronic configuration, oxidation state and hydride formation.
- 7.18 Why is dioxygen a gas but sulphur a solid?
- 7.19 Knowing the electron gain enthalpy values for $\text{O} \rightarrow \text{O}^-$ and $\text{O} \rightarrow \text{O}^{2-}$ as -141 and 702 kJ mol^{-1} respectively, how can you account for the formation of a large number of oxides having O^{2-} species and not O^- ? [Hint: Consider lattice energy factor in the formation of compounds].
- 7.20 Which aerosols deplete ozone?
- 7.21 Describe the manufacture of H_2SO_4 by contact process?
- 7.22 How is SO_2 an air pollutant?
- 7.23 Why are halogens strong oxidising agents?
- 7.24 Explain why fluorine forms only one oxoacid, HOF.
- 7.25 Explain why inspite of nearly the same electronegativity, nitrogen forms hydrogen bonding while chlorine does not.
- 7.26 Write two uses of ClO_2 .
- 7.27 Why are halogens coloured?
- 7.28 Write the reactions of F_2 and Cl_2 with water.
- 7.29 How can you prepare Cl_2 from HCl and HCl from Cl_2 ? Write reactions only.
- 7.30 What inspired N. Bartlett for carrying out reaction between Xe and PtF_6 ?
- 7.31 What are the oxidation states of phosphorus in the following:
 (i) H_3PO_3 (ii) PCl_3 (iii) Ca_3P_2 (iv) Na_3PO_4 (v) POF_3 ?

- 7.32** Write balanced equations for the following:
 (i) NaCl is heated with sulphuric acid in the presence of MnO_2 .
 (ii) Chlorine gas is passed into a solution of NaI in water.
- 7.33** How are xenon fluorides XeF_2 , XeF_4 and XeF_6 obtained?
- 7.34** With what neutral molecule is ClO^- isoelectronic? Is that molecule a Lewis base?
- 7.35** How are XeO_3 and XeOF_4 prepared?
- 7.36** Arrange the following in the order of property indicated for each set:
 (i) F_2 , Cl_2 , Br_2 , I_2 - increasing bond dissociation enthalpy.
 (ii) HF , HCl , HBr , HI - increasing acid strength.
 (iii) NH_3 , PH_3 , AsH_3 , SbH_3 , BiH_3 - increasing base strength.
- 7.37** Which one of the following does not exist?
 (i) XeOF_4 (ii) NeF_2 (iii) XeF_2 (iv) XeF_6
- 7.38** Give the formula and describe the structure of a noble gas species which is isostructural with:
 (i) ICl_4^- (ii) IBr_2^- (iii) BrO_3^-
- 7.39** Why do noble gases have comparatively large atomic sizes?
- 7.40** List the uses of neon and argon gases.

Answers to Some Intext Questions

- 7.1** Higher the positive oxidation state of central atom, more will be its polarising power which, in turn, increases the covalent character of bond formed between the central atom and the other atom.
- 7.2** Because BiH_3 is the least stable among the hydrides of Group 15.
- 7.3** Because of strong $p\pi$ - $p\pi$ overlap resulting into the triple bond, $\text{N}\equiv\text{N}$.
- 7.6** From the structure of N_2O_5 it is evident that covalence of nitrogen is four.
- 7.7** Both are sp^3 hybridised. In PH_4^+ all the four orbitals are bonded whereas in PH_3 there is a lone pair of electrons on P, which is responsible for lone pair-bond pair repulsion in PH_3 reducing the bond angle to less than $109^\circ 28'$.
- 7.10** $\text{PCl}_5 + \text{D}_2\text{O} \rightarrow \text{POCl}_3 + 2\text{DCl}$
- 7.11** Three P-OH groups are present in the molecule of H_3PO_4 . Therefore, its basicity is three.
- 7.15** Because of small size and high electronegativity of oxygen, molecules of water are highly associated through hydrogen bonding resulting in its liquid state.
- 7.21** Both the S-O bonds are covalent and have equal strength due to resonating structures.
- 7.25** H_2SO_4 is a very strong acid in water largely because of its first ionisation to H_3O^+ and HSO_4^- . The ionisation of HSO_4^- to H_3O^+ and SO_4^{2-} is very very small. That is why $K_{a_2} \ll K_{a_1}$.
- 7.31** In general, interhalogen compounds are more reactive than halogens due to weaker X-X^1 bonding than X-X bond. Thus, ICl is more reactive than I_2 .
- 7.34** Radon is radioactive with very short half-life which makes the study of chemistry of radon difficult.

