HYDROGEN AND ITS COMPOUNDS •

INTRODUCTION

Hydrogen is the lightest element and also the lightest gas in the periodic table.

It is the lightest non-metal. It is the simplest element in periodic table having only 1e⁻, 1p & no neutron.

Electronic configuration of H is 1s1

Number of $e^- = 1$

Number of orbital = 1

Number of shells = 1Number of subshell =1

It is discovered by Henry Cavendish & it was called inflammable element.

The name hydrogen was given by Lavoisier

HYDROGEN

Hydro (Water)

Gene (Producer) or generator

means according to Lavoisier the gas which produce water on burning with oxygen.

$$H_2 + \frac{1}{2}O_2 \rightarrow H_2O$$

Order of Abundance of H

Hydrogen is the most abundant element in the universe (70% of the total mass of the universe.)

Order of abundance of H

Universe > Sun atm > Earth

The planet Jupiter & Saturn consist mainly of H₂. Similarly about half the mass of the sun & some other stars is made up of hydrogen.

In Sun's atmosphere & in universe, It is found in atomic form. While in earth it is generally found in molecular form. At Sun, the stratosphere is made up of H (atomic hydrogen) & they undergoes fusion & converted into He nuclei & this reaction is exothermic so lot of amount of energy is liberated.

It is the ninth element on earth in order of abundance.

Earth does not posses enough gravitational force to retain live hydrogen molecule i.e. why it is not found in earth atmosphere in atomic form.

Hydrogen is the most reactive elements in atomic form but it is less reactive in molecular form because of very high bond dissociation energy due to 1s -1s overlapping.

POSITION OF H, IN PERIODIC TABLE

Hydrogen is the first element of Periodic table but still it could not be assigned a proper position either in Mendeleef periodic table or in Modern periodic table because of following reasons.

It may kept in 1st/IA or 17th/VIIA group due to following reason.

Resembles with 1st / IA Alkali metals.

ELECTRONIC CONFIGURATION : Like alkali metals hydrogen also has only one electron in outer most shell. **(i)**

 ns^1

 \Rightarrow \rightarrow

 \rightarrow

$H = 1 s^1$	$K = 4s^1$	▲
$Li = 2s^1$	$Rb = 5s^1$	
$Na = 3s^1$		₊



(ii) ELECTROPOSITIVE CHARACTERS: Like alkali metals hydrogen also have the tendency to loose one electron to form cation.

(iii) **OXIDATION NUMBERS**: Like alkali metals hydrogen can also exhibit the oxidation number of +1 in most of its compound.

Ex. Na⁺Cl⁻ K⁺Cl H⁺ Cl⁻ =+1

Ex.

(iv) **REACTION WITH ELECTRONEGATIVE ELEMENTS (NON-METALS)**: Like alkali metals H also reacts with Oxygen, Sulphur, Halogens to form oxides, sulphides and halides respectively.

 $O \xrightarrow{gives} H_2O$, like Na₂O, K₂O

S
$$\xrightarrow{gives}$$
 H₂S, like Na₂S, K₂S

with halogen \xrightarrow{gives} HX, like NaX, KX

(v) LIBERATIONAT CATHODE

2Na ⁺ Cl	electrolysis >	2Na	+	Cl_2
		(liberated at cath	node)(liberate	d at anode)

$2H^+$	CF	electrolysis	(H	+	(1)

like alkali metals H also get liberated at cathode on electrolysis.

(vi) **REDUCING NATURE :** Like alkali metals H, also have reducing nature.

 $CuO + H_2 \longrightarrow Cu + H_2O$

Due to resembling of these properties with alkali metals H can be placed in IA/group/alkali metals group. **Resembles with halogen/** VIIA/17th

(i) ELECTRONIC CONFIGURATION

Electronic configuration of halogen $H \longrightarrow 1s^1$

 \longrightarrow ns²np⁵

both these requires one electron to attain the stable configuration of their nearest inert gas.

$$H \longrightarrow 1s^{1} \xrightarrow{1e^{-} \text{ less than}} \text{He}(1s^{2})$$

 $F \longrightarrow 2p^5 \xrightarrow{1e^- \text{ less than}} \text{Ne}(2p^6)$

 $Cl \longrightarrow 3p^5 \xrightarrow{1e^{-} less than} Ar(3p^6)$

(ii) **ELECTRONEGATIVE CHARACTER**: Both halogen & Hydrogen has the tendency to gain one e⁻.

$$F + e^- \longrightarrow F^-$$

 $H + e^{-} \longrightarrow H^{-}$

i.e. why both will act as electronegative species.

- (iii) IONIZATION POTENTIAL : Ionization potential of hydrogen is almost similar to halogen.
- (iv) OXIDATION NUMBER : Both halogen & hydrogen can exhibit -1 oxidation number.

Hydrogen in metal hydride shows -1 oxidation state.

 $NaH \longrightarrow Na^+ + H^ NaCl \longrightarrow Na^+ + Cl^-$



- (v) DIATOMIC MOLECULE : Both halogen & hydrogen have the tendency to exist as diatomic molecule.
 Ex. F₂, Cl₂ Br₂, I₂ & H₂.
- (vi) LIBERATION AT ANODE : Whenever Alkali metals halides & hydrides undergoes electrolysis, both halogen and hydrogen will liberate at anode.

 $2Na^+Cl^- \longrightarrow 2Na+Cl_2$ (anode)

 $2Na^+H^- \longrightarrow 2Na + H_2$ (anode)

(vii) **REACTION WITH HIGHLY ELECTROPOSITIVE METALS**: Hydrogen reacts with highly electropositive element (i.e. s-block elements) & forms their hydrides like their halides.

 $2Na + H_2 \longrightarrow 2NaH$ $2Na + Cl_2 \longrightarrow 2NaCl$

This reaction shows oxidising character of hydrogen.

(viii) FORMATION OF COVALENT COMPOUND : Both Hydrogen & halogen on reaction with non metals to form covalent compounds like.

 CH_4 , NH_3 , H_2O CCl_4 , NCl_2 , Cl_2O

Conclusion : The position of hydrogen is still in controversy & no proper position is assigned to H in periodic table i.e. why it is also called notorious or rogue elements.

ISOTOPES OF HYDROGEN

There are 3 isotopes of H,

	Protium or ordinary hydrogen	Deuterium/ Heavy hydrogen	Tritium
Symbol	,H ¹ H	$_{1}\mathrm{H}^{2}/_{1}\mathrm{D}^{2}$	1H ³ /1T ³

Property	Protium or ordinary hydrogen	Deuterium or heavy hydrogen	Tritium
Atomic number	1	1	1
Mass number	1	2	3
Exact atomic mass	1.008123	2.0142	3.0170
Symbol	$^{1}_{1}H$	2_1 H or 2_1 D	${}^{3}_{1}$ H or ${}^{3}_{1}$ T
Molecular formula	H ₂	D_2	Τ ₂
No. of protons in		-	-
the nucleus	1	1	1
No. of neutrons in			
the nucelus	Nil	1	2
No. of electron	1	1	1
Electronic configu.	$1s^1$	$1 s^{1}$	$1s^1$
Relative abundance	99.984%	0.016%	10 ⁻¹⁵ %
Stability	Stable	Stable	Unstable(Radioactive)



HYDROGEN AND ITS COMPOUNDS

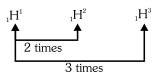
Properties	${ m H_2}$	D ₂	T ₂
M.P.	-259°C	-254.3°C	-252.4ºC
B.P.	$-252.6^{\circ}C$	-249.3°C	-248.0°C
Bond length (H–H)	74 pm	74 pm	74 pm
Bond energy (H—H)	436.0 KJ mol ⁻¹	443.3 KJ mol ⁻¹	446.9 KJ mol ⁻¹
Heat of fusion & vaporisation	Minimum	H <d<t< td=""><td>Maximum</td></d<t<>	Maximum

Isotopic effect

The effect which can change the physical & chemical properties of isotopes is called isotopic effect. It is because of difference in mass.

In isotopic effect maximum changes occurs in physical properties like melting point, boiling point, bond energy, while minimum changes occurs in chemical properties like state of chemical reaction etc.

Imp. Isotopic effect is found only in hydrogen isotopes. Because there is large difference in mass.



While in other isotopes like ${}_{7}N^{14}$ ${}_{7}N^{15}$ only a fractional mass is increased. ${}_{6}C^{14}$ ${}_{6}C^{13}$ ${}_{6}C^{12}$ \longrightarrow Here also only a fraction is increased

But in case of hydrogen mass increased to 2 to 3 times in their isotopes.

Ex. Isotopic effect is found in : (1) H (2) N

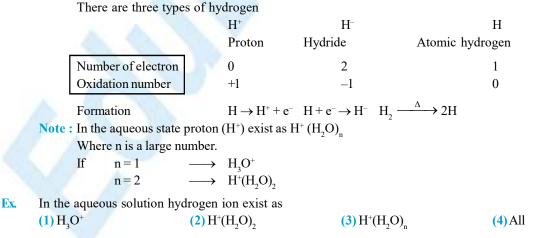
(3) C

(4) All

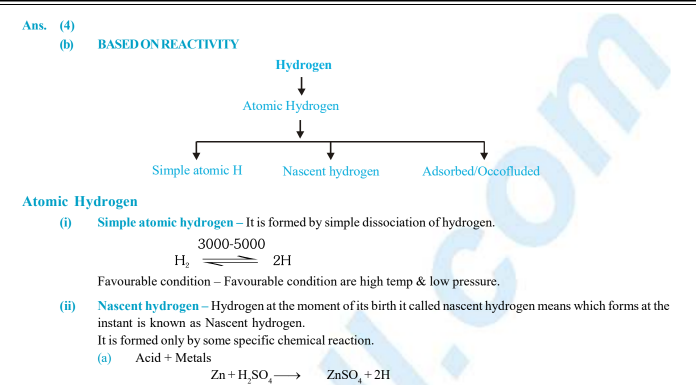
- Ans. (1)
- Ex. Which of the following reaction is fast & why? (i) $CH_4 + Cl_2 \longrightarrow CH_3Cl + HCl$ (ii) $CD_4 + Cl_2 \longrightarrow CD_3Cl + DCl$
- Ans. (i) because C–H bond energy is less in comparison to C–D bond energy.

DIFFERENT FORMS OF HYDROGEN

(a) BASED ON OXIDATION NUMBER





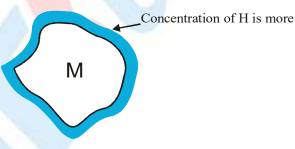


(b) Base + element
$$2NaOH + Be \longrightarrow Na_2BeO_2 + 2H$$

(c)
$$C_2H_5OH + Alkali metal$$

 $C_2H_5OH + Na \longrightarrow C_2H_5ONa + H$

(iii) Adsorbed/Occluded hydrogens

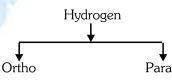


Adsorbed H is hydrogen present at the outer surface of metal. Occlusion – The property of metal to adsorb any gas is called occlusion.

Reactivity order

Atomic hydrogen > Nascent hydrogen > Molecular hydrogen

(c) BASED ON NUCLEAR SPIN (NUCLEAR ISOMERS)



(i) Ortho hydrogen – The molecular form of hydrogen having same spin of proton is called ortho hydrogen.



(ii) Para hydrogen – The molecular form of hydrogen having opposite spin of proton is called para hydrgen.

In ortho hydrogen spin of proton is same, so they will repel each other & because of this repulsion, internal energy of ortho hydrogen increases. So ortho hydrogen has more internal energy.

Stability of ortho & para hydrogen

Stability of ortho & para hydrogen depends upon temperature condition.

At low temp : para hydrogen is more stable than ortho hydrogen while at high temp ortho hydrogen is more stable than para hydrogen.

	$P_{H_2} \xrightarrow{T\uparrow} O_{H_2}$	
	Ortho	Para
At 25°C	75%	25%
At-253°C/20K	0	100%

EDUBULL KEY POINTS

- (i) We can obtain 100% pure para hydrogen at low temp but can't ortho because at high temp parahydrogen will dissociate into atomic hydrogen.
- (ii) Ortho & Para hydrogen differs only in physical properties but have same chemical properites.

METHOD OF PREPARATION

(i) **FROM ACIDS**: The metal which are placed about H₂ in electrochemical series react with dil acids they liberate H₂.

Ex.
$$Fe + H_2SO_4 \rightarrow FeSO_4 + H_2$$

(dil)
 $Cu + H_2SO_4 \rightarrow \times (No reaction)$

Lab preparation : When impure Zn reacts with dil H,SO₄ it forms H,

Zn	+	$H_2SO_4 \rightarrow ZnSO_4 + H_2$
(impure)		(dil)

IN OLI N 7 O LUT

- **Ex.** Why we use impure Zn.
- **Ans.** Because the rate of reaction with pure Zn is very slow.

(2) **BYALKALIES**: Only (Be, Zn, Al, Sn, Pb, Si) (Amphoteric metal) react with boiling NaOH or KOH they evolve H₂.

$$2h + 2NaOH \rightarrow Na_{2}ZhO_{2} + H_{2} + (sodium zincate)$$

$$2Al + 2NaOH + 2H_{2}O \longrightarrow NaAlO_{2} + 3H_{2}^{\uparrow} (sodium meta aluminat e)$$

$$Sn + 2NaOH + H_{2}O \longrightarrow Na_{2}SnO_{3} + 2H_{2}^{\uparrow} (sodium stannate)$$

$$Pb + 2NaOH + H_{2}O \longrightarrow Na_{2}PbO_{3} + 2H_{2}^{\uparrow} (sodium plumbate)$$

$$Si + 2NaOH + H_{2}O \longrightarrow Na_{2}SiO_{3} + 2H_{2}^{\uparrow} (sodium silicate)$$

$$Be + 2NaOH \longrightarrow Na_{2}BeO_{2} + H_{2} (sodium beryliate)$$



(3) FROM WATER

All the metals which are placed above than H, when react with water the evolve H₂.

 $Zn + H_2O \longrightarrow ZnO + H_2\uparrow$

Three type of water is used

(i) Cold water : The temperature of cold water is 7 to 25°C this water is used for highly reactive metals.

Such as Li, K, Ba, Sr, Ca, Na, means alkali metals of alkaline earth metals.

The reaction with alkali metals are vigorous to minimum the rate of reaction these metals are used in the form of amalgam.

 $2Na(Hg) + H_2O \longrightarrow 2NaOH + H_2\uparrow$

- (ii) Hot water : The temperature of hot water is 25°C to 90°C. This water is used for reactive metals, such as Mg, Al, Mn, Zn, Cr.
- (iii) Steam : The temperature of steam is more than 100°C. This form of water is used for very less reactive metals like Fe, Cd, Co, Ni, Sn, Pb.

Condition for best yield of H,

- (i) Cold water \longrightarrow With highly reactive metals.
- (ii) Hot water \longrightarrow With reactive metals
- (iii) Steam \longrightarrow With less reactive metals.

(4) **ON IONIC HYDRIDE**

Whenever ionic hydride reacts with water then form H₂.

 $CaH_2 + 2H_2O \longrightarrow Ca(OH)_2 + H_2$

(Hydrolith)

$$NaBH_4 + H_2O \longrightarrow NaBO_2 + 4H_2$$

METHOD TO PREPARE PURE HYDROGEN

1. ELECTROLYSIS OF WATER : To prepare pure hydrogen we use impure water (i.e. having 15-20% solution of alkali

or acid)

 $4H_2O \downarrow 4H^+ + 4OH^ 4H^+ + 4e^- \longrightarrow 2H_2$

at cathode

at anode

 $4OH^- \longrightarrow 2H_2O + O_2 + 4e^-$

The SO_4^{-2} or K⁺ ion present in acid or alkali does not move towards anode or cathode as their discharge potential is higher than of OH⁻ ions or H⁺ ions respectively.

2. BY REACTION OF MAGNESIUM WITH DIL. H₂SO₄

 $Mg + H_2SO_4 \longrightarrow MgSO_4 + H_2$

3. BREACTION OF NAH WITH WATER

 $NaH + H_2O \longrightarrow NaOH + H_2$

4. UYENO METHOD : This method is used for military purpose

 $2A1 + 2KOH + 2H_2O \longrightarrow 2KAIO_2 + 3H_2\uparrow$

(Potassium meta aluminate)

Because by this reaction we can prepare H₂ in a rapid manner.



COMMERCIAL OR INDUSTRIAL METHOD TO PREPARE H,

(i) **LANE PROCESS :** Steam is passes over hot iron it converts into $Fe_3O_4 \& H_2$.

$$3\text{Fe} + 4\text{H}_2\text{O} \longrightarrow \text{Fe}_3\text{O}_4 + 4\text{H}_2\uparrow$$

Iron is regenerated by reducing Fe_3O_4 into Fe by water gas $(CO + H_2)$

$$2Fe_{3}O_{4} + 4CO + 4H_{2} \longrightarrow 6Fe + 4CO_{2} + 4H_{2}O$$

 $\operatorname{Fe_3O_4} + 2\operatorname{CO} + 2\operatorname{H_2} \longrightarrow 3\operatorname{Fe} + 2\operatorname{CO_2} + 2\operatorname{H_2O}$

This process is a continuous process.

(ii) **BOSCH PROCESS :** In this process initially steam is passed over red hot coke then water gas is formed. $C+H_2O\longrightarrow CO+H_2$

Then water gas is mixed with more stem in presence of Fe_2O_3/Cr_2O_3 then CO will convert into O_2 & we can obtain more H_2 .

 $CO + H_2 + H_2O \longrightarrow CO_2 + 2H_2$

(iii) FROM NATURALGAS

$$C_nH_{2n+2} + nH_2O \longrightarrow nCO + (2n+1)H_2$$

PHYSICAL PROPERTIES OF HYDROGEN

- (i) Hydrogen is a lightest, colourless, odourless and tasteless gas. It is sparingly soluble in water. It is inflammable and less reactive gas.
- (ii) Its f.p. (-259.2°C) and b.p. (-252°C) are very low indicating less intermolecular attraction. Due to low f.p. liquid hydrogen is used as a cryogenic fluid (to produce low temperature).
- (iii) H—H bond energy [$104 \text{ Kcal mol}^{-1}$] and 436 KJ/m
- (iv) H H bond length [74 pm] so H_2 is less reactive and require high temp for reaction.

CHEMICAL PROPERTIES OF H,

- (i) H_2 is neutral in nature i.e. why it does not react with acids & bases.
- (ii) LESS REACTIVE : Hydrogen is very less reactive in nature because of very high bond dissociation energy.
- (iii) **COMBUSTIBLE NATURE :** H₂ is highly combustible in nature & it burns with oxygen or air with pale blue flame to give water.

$$H_2 + \frac{1}{2}O_2 \longrightarrow H_2O$$

(iv) REACTION WITH HIGHLY ELECTROPOSITIVE METALS

IA & IIA group elements are called highly electropositive metal. Whenever H reacts with these metals they form ionic hydrides.

$$2Na+H_2 \longrightarrow 2NaH$$
$$CO+H_2 \longrightarrow CaH_2$$

(vi) REACTION WITH NON-METAL

(a) Reaction with halogen

$$H_2 + X_2 \longrightarrow 2HX$$

[F₂, Cl₂, Br₂, I₂]

order of reactivity of halogen with hydrogen

$$F_2 > Cl_2 > Br_2 > I$$

(b) Reaction with sulphur

 $H_2 + S \longrightarrow H_2S$

(c) With nitrogen – (Haber process)

$$N_2 + 3H_2 \xrightarrow{683 \text{ K}} 2\text{NH}_3$$



(vi) **REDUCINGNATURE**

 $CuO + H_2 \longrightarrow Cu + H_2O$

(vii) REACTION WITH CARBON MONO OXIDE

$$2H_2 + CO \xrightarrow{ZnO / Cr_2O_3} CH_3OH (Methanol)$$

(viii) HYDROGENATION OF UNSATURATED HYDROCARBON

$$C_{2}H_{2} + 2H_{2} \longrightarrow C_{2}H_{6}$$

$$C_{2}H_{4} + H_{2} \xrightarrow{Ni/Pb/Pt}{473K} \rightarrow$$

When unsaturated hydrocarbon having C = C or C = C reacts with hydrogens in the presence of Ni/Pd/Pt forms saturated hydrocarbons.

EDUBULL KEY POINTS

C,H

Hydrogenation of vegetable oil.

Vegetable oils are also called polyunsaturated oils because they contain many C = C bond. When these oil are exposed to air for a long time then double bond will get oxidized and the oil becomes. Rancid (having fowl. smell or unpleasant test) in nature. So to avoid this vegetable oil are converted into edible fats (Vanaspati Ghee)

Vegetable oils +
$$H_2 \xrightarrow{Ni}$$
 edible fats solid
(liquid)

(liquid)

This whole process is known as hydrogenation or hardening of oil.

USES OF HYDROGEN

- 1. Hydrogenation of vegetable oil to form solid fats i.e. vanaspati ghee.
- 2. In liquid form as a rocket fuel. (Liquid $H_2 + Liquid O_2$)
- 3. In a air ship of balloons as a mixture of Hydrogen & Helium $[15\% H_2 + 85\% He]$
- 4. Formation of different compounds.

Like \rightarrow NH₃, (Haber process) alkane, alcohol and other hydrocarbon

- **Ex.** Write the names of isotopes of hydrogen. What is the mass ratio of these isotopes ?
- Sol. The various, isotopes of hydrogen are :

¹₁H (Protium); ²₁H or D(Deuterium); ³₁H or T(Tritium)

The mass ratio of ${}_{1}^{1}H : {}_{1}^{2}H : {}_{1}^{3}H$ is 1 : 2 : 3

- Ex. Why does hydrogen occur in a diatomic form rather than in a monoatomic form under normal conditions?
- **Sol.** Hydrogen atom has only one electron and thus, to achieve stable inert gas configuration of helium, it shares its single electron with electron of other hydrogen atom to form a stable diatomic molecule. The stability of H_2 is further confirmed by the fact, that formation of one mole of gaseous H_2 molecules results in the release of 435.8 kJ of energy

 $H(g) + H(g) \longrightarrow H_2(g); \Delta H = -435.8 \text{ kJ mol}^{-1}$

- **Ex.** Which of the following can adsorb largest volume of hydrogen gas?
 - (1) Finely divided platinum (2) Finely divided nickel
 - (3) Colloidal palladium

(4) Colloidal platinum

Ans. (3)

Sol. Order of adsorption of $H_2($ occlusion) is :

Colloidal Palladium > Palladium > Platinum > Gold > Nickel.



- Ex. Describe the bulk preparation of hydrogen by electrolytic method. What is the role of an electrolyte in this process
- **Sol.:** The electrolyte (15–20% NaOH solution) increases conductivity of water.

Cathode (iron) : Reduction of water occurs. $2H_2O + 2e^- \longrightarrow 2H_2 + 2OH^-$

Anode (nickel coated iron) : Oxidation of OH⁻ occurs.

 $2OH^{-} \longrightarrow H_2O + 1/2O_2 + 2e^{-}$

Ex. Complete the following reactions :

(i)
$$H_2(g) + M_m O_n(s) \xrightarrow{\Delta}$$

(ii) $CO(g) + H_2(g) \xrightarrow{\Delta}$
(iii) $C_3H_8(g) + 3H_2O(g) \xrightarrow{\Delta}$
(iv) $Zn(s) + NaOH(aq)$
(iv) $Zn(s) + NaOH(aq)$

Ans. (i)
$$nH_2(g) + M_mO_n(s) \xrightarrow{\Delta} mM(s) + nH_2O(\bullet)$$

(ii) $CO(g) + 2H_2(g) \xrightarrow{\Delta} CH_3OH(\bullet)$ (methanol) (iii) $CU(g) + 2H_2(g) \xrightarrow{N_1} CH_3OH(\bullet)$ (methanol)

(iii) $C_3H_8(g) + 3H_2O(g) \xrightarrow{Ni} 3CO(g) + 7H_2(g)$

(iv) $Zn(s) + 2NaOH(aq) \xrightarrow{heat} Na_2ZnO_2(aq) + H_2(g)$

- **Ex.** Can conc. H_2SO_4 be used for drying H_2 gas ? Justify.
- Sol. Conc. H_2SO_4 cannot be used for drying H_2 gas because H_2SO_4 does absorb moisture from moist H_2 , but the process is highly exothermic. The heat so produced causes hydrogen to catch fire because of its inflammable nature.
- Ex. Can dihydrogen act as oxidising agent ? If so give chemical reactions to support the statement.
- Sol. Dihydrogen can act as oxidising agent when it forms metal hydrides.

 $2Li + H_2 \longrightarrow 2LiH$

HYDRIDES

The compounds of hydrogen with different elements are called hydrides.

These are of three types :

(1) IONIC/SALT LIKE/SALINE HYDRIDES

Compounds of hydrogen with s-block elements except berylium & magnesium are called ionic hydrides.

LiH, NaH, KH, RbH, CsH, CaH₂, SrH₂, BaH₂

 BeH_2 , MgH_2 are covalent polymeric hydride.

- Structure of these hydrides are similar to rock salt, so they are also called salt like/saline hydrides.
- Down the group size \uparrow Lattice energy \downarrow stability \downarrow Melting point \downarrow Boiling point \downarrow
- On electrolysis of these hydrides, hydrogen is liberated at anode.
- On reaction with water these hydrides will form hydrogen

 $NaH + H_2O \longrightarrow NaOH + H_2$

These hydrides forms complex hydrides which are very good reducing agents.

 $4LiH + AlCl_{3} \longrightarrow LiAlH_{4} + 3LiCl$

 $NaBH_4 \longrightarrow Sodium borohydride$

 $LiAlH_4 \longrightarrow Lithium aluminium hydride.$



(2) METTALIC/INTERSTITIAL HYDRIDES

They are the compounds of d & f-block elements.

In these hydrogen occupies interstitial sites present in metallic lattice, so they are called interstitial hydrides.

- Properties of these hydrides are similar to parent metals, so they are also known as metallic hydrides.
 - These hydrides are non. stoichiometric in nature (i.e. having variable composition)

ZrHx (x = 1.3 - 1.75)

TiHx (x = 1.8 - 2)

• Metals of group 7,8,9 donot form any hydrides so this particular part of periodic table is known as hydride gap.

(3) COVALENT/MOLECULES HYDRIDES

- They are the compounds of hydrogen with p-block elements CH₄, NH₃, H₂O, HF, etc.
- These hydrides exist as molecules, so they are also known as molecular hydrides. There hydrides are nonconductor of electricity.

Nomenclature - element + Suffix (ane)

 $PH_3 \longrightarrow Phosphane$

 $NH_3 \longrightarrow Azane$

 $H_2O \longrightarrow Oxidane$

These hydrides are again divides into 3 categories.

- (a) Electron deficient hydrides :
- They are the hydrides of group 13 elements. BH₃, AlH₃, GaH₃ – In these hydrides central element does not have complete octet. i.e. why they are called electron deficient compounds.
- (b) Electron precise hydrides They are the hydrides of group 14 element. Ex. CH_4 , SiH_4 , GeH_4

In these type of hydrides central elements has 8e⁻ in their outer most shell.

(c) Electron rich hydrides : These are the hydrides of group 15, 16, 17

Ex. NH₃, H₂O, HF: etc.

In these hydrides lone pair are present on central dement which can be given to others. So they are called electron rich hydrides.

Ex. What characteristics do you expect from electron deficient hydrides with respect to their structure and chemical reactivity?

Sol. Electron deficient hydrides do not have sufficient number of electrons to form normal covalent bonds. They generally exist in polymeric forms such as B_2H_6 , B_4H_{10} , $(AIH_3)_p$, etc.

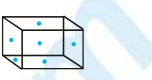
Due to deficiency of electrons, these hydrides act as Lewis acids and thus, form complex entities with Lewis bases such as : NH_{2} , H^{-} ions, etc.

 $B_{2}H_{6} + 2NH_{3} \longrightarrow [BH_{2}(NH_{3})_{2}]^{+}[BH_{4}]^{-}$ $B_{2}H_{6} + 2NaH \longrightarrow 2Na^{+}[BH_{4}]^{-} (sodium borohydride)$

Ex. Do you expect the carbon hydride of the type (C_nH_{2n+2}) to act as Lewis acid or base ? Justify your answer.

Sol. Carbon hydride of the type (C_nH_{2n+2}) are electron precise hydrides. In other words, they have exact numbers of electrons required to form covalent bonds. Therefore, they do not have tendency to either gain or lose electrons and hence, they do not act as Lewis acids or Lewis bases.





Ex. Arrange the following :

- (i) CaH₂, BeH₂ and TiH₂ in order of increasing electrical conductance.
- (ii) LiH, NaH and CsH in order of increasing ionic character
- (iii) H–H, D–D and F–F in order of increasing bond dissociation enthalpy.
- (iv) NaH, MgH₂ and H₂O in order of increasing reducing property.
- Sol. (i) BeH_2 is a covalent hydride, therefore, it does not conduct electricity at all. CaH_2 conducts electricity in the fused state while TiH₂ conducts electricity at room temperature. Thus, the order of increasing electrical conductance is : $BeH_2 < CaH_2 < TiH_2$.
 - (ii) Electronegativity decreases down the group from Li to Cs, therefore, the ionic character of their hydrides also increases in the same order, i.e., LiH, < NaOH < CsH.
 - (iii) F F bond dissociation enthalpy is the minimum. This is due to high concentration of electron density around each F atom in the form of 3 lone pairs which cause significant repulsive interactions. The bond enthalpy of D-D bond is slightly higher than that of H-H bond. It is due to marginally smaller size of D. Thus, the bond dissociation enthalpy increases in the order : F-F < H-H < D-D.
 - (iv) Ionic hydrides are powerful reducing agents. Both MgH₂ and H₂O are covalent hydrides but the bond dissociation of O–H bonds in H₂O is much higher than that of Mg–H bond in MgH₂. Therefore, the reducing character increases in the order : $H_2O < MgH_2 < NaH$.
- Ex. How do you expect the metallic hydrides to be useful hydrogen storage ? Explain.
- Sol. In some of the transition metal hydrides, hydrogen is absorbed as H atoms. Due to the inclusion of H-atoms, the metal lattice expands and thus becomes less stable. Therefore, when such metallic hydride is heated, it decomposes to release hydrogen gas and very finely divided metal. The hydrogen evolved in this manner can be used as a fuel. Thus, transition metals or their alloys can act as sponge and can be used to store and transport hydrogen to be used as a fuel.
- **Ex.** Explain the meaning of term hydride gap.
- Ans. Elements of group 7, 8, 9 of d-block do not form hydrides at all. This inability of metals of group 7, 8, 9 of periodic table to form hydrides is referred to as hydride gap of d-block.
- **Ex.** Explain the following :
 - (a) Water is excellent solvent for ionic compounds.
 - (b) Lakes freeze from top to bottom.
- Sol. (a) Water has a high dielectric constant (78.39) due to the polar character of its molecule. Water is an excellent solvent for many ionic as well as covalent compounds. Dissolution of ionic compounds takes place because of ion-dipole interactions. Dissolution of molecular compounds such as alcohols, amides, urea, sugar, glucose, honey, etc., in water takes place because of the tendency of these substances to form hydrogen bonds with water molecules.
 - (b) This is due to the fact that the frozen water does not sink to the bottom but keeps floating at the surface due to its lesser density. This provides thermal insulation to the water below it. The lesser density of ice can be attributed to open cage-like structure on account of hydrogen bonding.

WATER (H_2O)

PROPERTIES OF PURE WATER

(a) **Physical properties**

(i) Pure water is colourless, tasteless and odourless. It gives bluish tinge in thick layers. (ii) It freezes at 0°C and boils at 100°C. (iii) Its maximum density is 1.00 at 4°C. (iv) It is a polar molecule and has V-shaped structure. The bond angle is 104.5°. (v) It has a high dielectric constant. The polar character of water makes it an excellent solvent for polar and ionic substances. (vi) It is a poor conductor of electricity. (vii) It has the tendency to associate. It exists in the liquid state not as a single H₂O molecule but as associated molecules through hydrogen bonding. The existence of hydrogen bonding is responsible for high values of specific heat, the latent heat of fusion and latent heat of vaporisation.



(b) Chemical properties

(i) Water is neutral in nature. pH of the pure water is 7. It is a weak electrolyte and feebly ionises into H^+ and OH^- ions.

 $H_{2}O \Longrightarrow H^{+} + OH^{-}$

In pure water $[H^+] = [OH^-] = 10^{-7}$ at 25°C.

(ii) WITH METALS : it reacts with active metals and evolves hydrogen. The reaction is exothermic in the case of alkali and alkaline earth metals.

 $2Na+2H_2O \longrightarrow 2NaOH + H_2$

 $Ca + 2H_2O \longrightarrow Ca(OH)_2 + H_2$

(iii) REACTION WITH NON METALS

Chlorine decomposes cold water forming HCl and HClO.

$$Cl_2 + H_2O \longrightarrow HCl + HClO (Hypochlorous acid)$$

When steam is passed over red hot coke (1000°C), water gas is formed.

$$C + H_2O \rightarrow CO_2 + H_{3^2}$$

_{Water gas}

(iv) ACTION ON NONMETALLIC OXIDES : Acidic oxides combine with water to form acids.

$CO_2 + H_2O f H_2CO_3$	(Carbonic acid)
$SO_2 + H_2O \longrightarrow H_2SO_3$	(Sulphurous acid)
$SO_3 + H_2O \longrightarrow H_2SO_4$	(Sulphuric acid)
$P_2O_5 + 3H_2O \longrightarrow 2H_3PO_4$	(Orthophosphoric acid)
$N_2O_5 + H_2O \longrightarrow 2HNO_3$	(Nitric acid)
$Cl_2O_7 + H_2O \longrightarrow 2HClO_4$	(Perchloric acid)

(v) ACTION ON METALLIC OXIDES : basic oxides combine with water to form alkalies.

 $Na_2O + H_2O \longrightarrow 2NaOH$

 $CaO + H_2O \longrightarrow Ca(OH)_2$

(vi) ACTION ON HYDRIDES, CARBIDES, NITRIDES, PHOSPHIDES: Water decomposes these compounds with liberation of hydrogen, acetylene (or methane), ammonia, phosphine respectively.

$$CaH_{2} + 2H_{2}O \longrightarrow Ca(OH)_{2} + 2H_{2}$$

$$CaC_{2} + 2H_{2}O \longrightarrow Ca(OH)_{2} + C_{2}H_{2}$$

$$Al_{4}C_{3} + 12H_{2}O \longrightarrow 4Al(OH)_{3} + 3CH_{4}$$

$$Mg_{3}N_{2} + 6H_{2}O \longrightarrow 3Mg(OH)_{2} + 2NH_{3}$$

$$Ca_{2}P_{3} + 6H_{2}O \longrightarrow 3Ca(OH)_{3} + 2PH_{3}$$

HYDROLYSIS: Many salts specially the salts of strong bases with weak acids, weak bases with strong acids and weak bases with weak acids undergo hydrolysis with water.

 $CH_3COONa + H_2O \rightarrow CH_3COOH + NaOH$

$$CH_3COONH_4 + H_2O \rightarrow CH_3COOH + NH_4OH$$

Halides of nonmetals are decomposed by water.

 $PCl_{5} + 4H_{2}O \longrightarrow H_{3}PO_{4} + 5HCl$ $PCl_{3} + 3H_{2}O \longrightarrow H_{3}PO_{3} + 3HCl$



(vii)

HARD AND SOFT WATER

A water is said to be a soft water if it produces sufficient lather with the soap and water is described as being hard if it forms an insoluble scum before it forms a lather with soap. The hardness of natural water is generally caused by presence of bicarbonates, chlorides and sulphates of calcium and magnesium but infact soluble salts that form a scum with soap cause hardness.

$$Ca^{2+}(aq) + 2C_{17}H_{35}COO^{-}(aq) \longrightarrow (C_{17}H_{35}COO)_{2}Ca$$

$$Mg^{2+}(aq) + 2C_{17}H_{35}COO^{-}(aq) \longrightarrow (C_{17}H_{35}COO)_{2}Mg$$
Anion of soap (Lag) Inso lub le precipitates

Soap will not produce lather with water until all the calcium and magnesium ions have been precipitated. Hard water thus wastes soap. Hardness of water is of two types :

- (a) Temporary hardness
- (b) Permanent hardness
- (a) **Temporary hardness :** This is due to the presence of bicarbonates of calcium and magnesium.

Temporary hardness in water is easily removed by boiling, as the bicarbonates decompose readily and the insoluble carbonates are precipitated.

$$Ca(HCO_3)_2 \xrightarrow{Boil} CaCO_3 + H_2O + CO_2$$

(Inso lub le)

$$Mg(HCO_3)_2 \xrightarrow{\text{Boil}} MgCO_3 + H_2O + CO_2$$
(Inso lub k)

Temporary hardness can also be removed by Clark's process which involves the addition of slaked lime [Ca(OH),].

$$Ca(HCO_3)_2 + Ca(OH)_2 \longrightarrow 2CaCO_{(Inso huble)} + 2H_2O$$

It is essential to add only the calculated amount of Ca(OH), because excess will cause artificial hardness.

(b) Permanent hardness : Permanent hardness is introduced when water passes over rocks containing the sulphates or chlorides of both of calcium and magnesium. This type of hardness cannot be removed by boiling or by the addition of slaked lime.

The various water softeners are :

(i) Washing soda : It removes both the temporary and permanent hardness by converting soluble calcium and magnesium compounds into insoluble carbonates.

$$CaCl_{2} + NaCO_{3} = CaCO_{3} + 2NaCl$$
$$CaSO_{4} + Na_{2}CO_{3} = CaCO_{3} + Na_{2}SO_{4}$$
$$Ca(HCO_{3})_{2} + Na_{2}CO_{3} = CaCO_{3} + 2NaHCO_{3}$$
(So lab le)

In place of sodium carbonate, caustic soda or sodium phosphate can also be used.

$$MgCl_2 + 2NaOH \longrightarrow Mg(OH)_2 + 2NaCl_{(Inso lub le)}$$

$$3MgSO_4 + 2Na_3PO_4 \longrightarrow Mg_3(PO_4)_2 + 3Na_2SO_4$$



(ii) Calgon : The complex salt of metaphosphoric acid, sodium hexametaphosphate $(NaPO_3)_6$, is known as calgon. It is represented as $Na_2[Na_4(PO_3)_6]$. Calcium and magnesium salts present in hard water react with calgon to give complex salts.

$$2CaSO_4 + Na_2[Na_4(PO_3)_6] \longrightarrow Na_2[Ca_2(PO_3)_6] + 2Na_2SO_4$$
$$2MgSO_4 + Na_2[Na_4(PO_3)_6] \longrightarrow Na_2[Mg_2(PO_3)_6] + 2Na_2SO_4$$

(iii) Permutit process : Permutit is hydrated sodium alumino silicate

[Na₂Al₂Si₂O₈. xH₂O] [Na₂O.Al₂O₃.2SiO₂.xH₂O]

Permutit also known as sodium zeolite (Na_2Z) . means Zeolite is $Al_2Si_2O_8.xH_2O$.

In this process when hard water is poured into chamber, it may contain organic impurities like plant. This impurities can be removed by gravel. On moving upwards hard water will react with Na_2Z during this reaction Na^+ ions of Na_2Z will replace Mg^{+2} & Ca^{+2} of impurities.

$$Na_2Z + CaCl_2 \longrightarrow 2NaCl + CaZ$$

NaCl is dissolved in water & water becomes soft.

This soft water is fit for washing purpose but not fit for drinking purpose.

After some time when Na_2Z is completely converted into CaZ. Process is stopped and for regeneration of Na_2Z . We use NaCl solution.

 $2NaCl + CaZ \longrightarrow CaCl_2 + Na_2Z$

(iv) Ion exchange resin : By this process we can remove both cation & anion of hardness.

This process contain two chamber.

- (a) Cation exchange resin : This resin contains granular insoluble organic acid having giant molecules with –COOH group.
- (b) Anion exchange resin : This resin contain giant organic molecules with basic groups derived from amines.

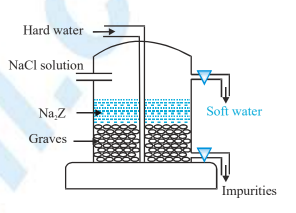
R-NH3+OH-

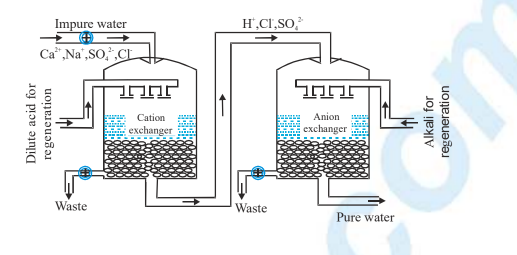
Process : When hard water is poured into first chamber the cation of hardness (Mg⁺². Ca⁺²) removed by H⁺ ions of organic acid.

$$2R-COO^{-}H^{+}+CaCl_{2} \longrightarrow (R-COO)_{2}Ca+2H^{+}+2Cl^{-}$$

$$+ MgSO_4 \longrightarrow (R-COO)_2Mg + 2H^+ + SO_4^{-2}$$

3





This water becomes soften but not used for drinking purpose because this water contain the impurity of acid. To remove anion of hardness this acidic water then passed through another bed containing anion exchanger. This exchanger removes anion like Cl^- , $SO_4^{-2} \& HCO_3^{-2}$.

 $R-NH_{3}^{+}OH^{-}+H^{+}+CI^{-}\longrightarrow R-NH_{3}^{+}CI^{-}+H^{+}OH$

This water is free from impurities & can be used for drinking purpose.

After some times when both resin gets exhausted process is stopped.

Regeneration of resin

(i) Cation exchange resin : We use dil acid.

 $2H^+Cl^- + (R-COO)_2Ca \longrightarrow 2R^-COO^-H^+ + CaCl_2$

(ii) Anion exchange resin : We use dil NaOH solution

 $R-NH_3+Cl^-+Na^+OH^-\longrightarrow Na^+Cl^-+R-NH_3+OH^-$

HEAVY WATER (D_2O)

METHOD OF PREPARATION

Repeated electrolysis of H_2O: On electrolysis of water (impure) H_2O dissociate into H^+ & OH⁻ while a fractional part of D_2O will dissociate into D^+ & OD⁻

 $H_2O f H^+ + OH^-$

 $D_2O f D^+ + OD^-$

 D^+ & OD^- due to more mass have less mobility i.e. why H^+ & OH^- will move towards cathode & anode respectively while D^+ & OD^- will be in solution.

This process will repeated for six time.

PROPERTIES OF HEAVY WATER

Physical properties : (a) Heavy water is a colourless, odourless and tasteless mobile liquid, (b) Nearly all the physical constants are higher than the corresponding values of ordinary water.

Chemical properties : Heavy water is chemically similar to ordinary water. However, D_2O reacts more slowly than H_2O in chemical reactions.



(a) ACTION OF METALS : D_2O reacts with alkali and alkaline earth metals liberates heavy hydrogen.

$$2D_2O + 2Na \longrightarrow 2NaOD_{\text{Sodium deuteroxide}} + D_2$$

$$2D_2O + Ca \longrightarrow Ca(OD)_2 + D_2$$

(b) ACTION WITH METALLIC OXIDES : D₂O reacts slowly with basic oxides to form heavy alkalies.

 $Na_2O + D_2O \longrightarrow 2NaOD$

 $CaO + D_2O \longrightarrow Ca(OD)_2$

(c) ACTION WITH NONMETALLIC OXIDES : D₂O reacts slowly with acidic oxides to form deutero acids.

$$N_2O_5 + D_2O \longrightarrow 2DNO_3$$

Deutero nitric acid

$$P_2O_5 + 3D_2O \longrightarrow 2D_3PO_4$$

Deutero phosphoric aci

$$SO_3 + D_2O \longrightarrow D_2SO_4$$

Deutero sulphuric ac

(d) **ACTION WITH METALLIC CARBIDES, PHOSPHIDES, NITRIDES, ARSENIDES, ETC.**: Like H₂O heavy water reacts with carbides, phosphide nitrides, arsenides, etc. to form corresponding deutero compounds.

$$CaC_2 + 2D_2O \longrightarrow Ca(OD)_2 + C_2D_2$$

Deutero acetylene

$$Al_4C_3 + 12D_2O \longrightarrow 4Al(OD)_3 + 3CD_4$$

Deutero methan

$$Mg_{3}N_{2} + 6D_{2}O \longrightarrow 3Mg(OD)_{2} + 2ND_{3}$$

Deutero ammoni

$$AIN + 3D_2O \longrightarrow AI(OD)_3 + ND_3$$

Deutero ammo

$$Ca_3P_2 + 6D_2O \longrightarrow 3Ca(OD)_2 + 2PD_3$$

Deutero phosphir

$$Na_3As + 3D_2O \longrightarrow 3NaOD + AsD_3$$

Deutero ar sin o

(e) **ELECTROLYSIS**: A solution of heavy water containing Na_2CO_3 when electrolysed evolve heavy hydrogen at cathode.

$$2D_2O \xrightarrow{\text{Electrolysis}} 2D_2 + O_2$$

(Cathode) (Anode)

USES : As a neutron moderator : Fission in uranium-235 is brought by slow speed neutrons. The substances which are used for slowing down the speed of neutrons are called moderators. Heavy water is used for this purpose in nuclear reactors.



H₂O₂ (Hydrogen Peroxide)

LABORATORY METHOD

In laboratory, H_2O_2 is prepared by adding calculated amounts of sodium peroxide to ice cod dilute (20%) solution of H_2SO_4 .

$$Na_2O_2 + H_2SO_4 \longrightarrow Na_2SO_4 + H_2O_2$$

By the action of sulphuric acid or phosphoric acid on hydrated barium peroxide BaO₂.8H₂O (Merck process)

(a) $BaO_2.8H_2O + H_2SO_4 \longrightarrow BaSO_4 \downarrow + H_2O_2 + 8H_2O_3$

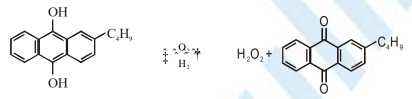
Anhydrous barium peroxide does not react readily with sulphuric acid (because a coating of insoluble barium sulphate is formed on its surface which stops further action of the acid). Therefore, hydrated barium peroxide, BaO₂.8H₂O must be used.

(b) $3BaO_2 + 2H_3PO_4 \longrightarrow Ba_3(PO_4)_2 + 3H_2O$ $Ba_3(PO_4)_2 + 3H_2SO_4 \longrightarrow 3BaSO_4 + 2H_3PO_4$

Phosphoric acid is preferred over H_2SO_4 because soluble impurities like barium persulphate (BaO₂.8H₂O -H₂SO₄) tends to decompose H_2O_2 while H_3PO_4 acts as preservative (negative catalyst) for H_2O_2 .

INDUSTRIAL METHOD

(i) Auto oxidation of 2 butyl anthraquinol (cyclic process)



(ii) Oxidation of isopropyl alcohol

(iii) Electrolytic Process : Used 50% H₂SO₄ in electrolytic cell using Pt as anode graphite as cathode.

$$2H_2SO_4 \longrightarrow 2H^+ + 2HSO_4^{\Theta}$$
At Cathode $2H^+ + 2e^- \longrightarrow H_2^{\uparrow}$
At Anode $2HSO_4^- \longrightarrow H_2S_2O_8 + 2e^-$
Peroxo disulphhuric acid
$$H_2S_2O_8 + H_2O \longrightarrow H_2SO_4 + H_2SO_5$$

$$H_2SO_5 + H_2O \longrightarrow H_2SO_4 + H_2O_2$$

Physical property

- (i) Pure H₂O₂ is colour less, odourless liquid and impure with bluish layer.
- (ii) It has more Hydrogen bonding then H_2O , So order of boiling point. $H_2O_2 > D_2O > H_2O$
- (iii) It is soluble in H₂O, alcohol and ether
- (iv) It has bitter test and harmful for skin
- (v) It is a dibasic weak acid

- (vi) It has a oxidising as well as reducing property
- (vii) H₂O₂ easily decompose in presence of light and temperature, So H₂O₂ always kept in dark bottles and kept at cool places.
- (viii) 30% solution of H_2O_2 is called Perhydrol.

Chemical property

1. **STABILITY**: It is unstable in nature decompose on standing and heating. It is an example of auto oxidation-reduction Reaction; $H_2O_2 \rightarrow H_2O + O$

NOTE : Nascent oxygen working as colourless agent

Decomposition of H_2O_2 ; $2H_2O_2$; $2H_2O_2$; $2H_2O + O_2$

Note : This is retarded by R–OH, acetanilide, glycerol

2. **OXIDIZING NATURE :** H_2O_2 is a powerful oxidant in acidic as well as in alkaline medium.

(In acid)
$$H_2O_2 + 2H^+ + 2e \xrightarrow{Fast} 2H_2O = E_{RP}^0 = 1.77V$$

(In alkali)
$$H_2O + HO_2^- + 2e \xrightarrow{\text{Slow}} 3HO^- E_{RP}^0 = 0.87 \text{ V}$$

Thus H_2O_2 is more powerful oxidant in acidic medium. The simple interpretation of H_2O_2 as oxidant can be shown by the equation.

 $H_2O_2 \longrightarrow H_2O + O$

Following are some important examples of oxidant action of H₂O₂:

(a) H_2O_2 oxidises black lead sulphide (PbS) to white lead sulphate (PbSO₄). This reaction is used in restoring the white colour of old paintings which have blackened due to the formation of lead sulphide by the action of H_2S present in air.

$$4[H_2O_2 \longrightarrow H_2O + O]$$

PbS+4O
$$\longrightarrow PbSO_4$$

 $PbS + 4H_2O_2 \longrightarrow PbSO_4 + 4H_2O$

(b) H_2O_2 oxidises sulphites into sulphates.

$$H_2O_2 \longrightarrow H_2O + O$$
$$Na_2SO_3 + O \longrightarrow Na_2SO_4$$

$$Na_2SO_3 + H_2O_2 \longrightarrow Na_2SO_4 + H_2O$$

(c) H_2O_2 oxidises nitrites to nitrates.

$$H_2O_2 \longrightarrow H_2O+O$$
$$Na_2NO_2 + O \longrightarrow NaNO_3$$

$$Na_2NO_2 + H_2O_2 \longrightarrow Na_2NO_3 + H_2O$$

(d) H_2O_2 oxidises H_2S into sulphur.



3. Reducing nature : It can also act as a reducing agent towards powerful oxidising agents.

$$H_2O_2 \longrightarrow 2H^+ + O_2 + 2e^-$$

(a) It reduces Ag_2O to silver. $Ag_2O + H_2O_2 \longrightarrow$

$$Ag_2O + H_2O_2 \longrightarrow 2Ag + H_2O + O_2$$

(b) It reduces ozone to oxygen.

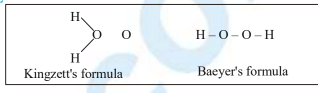
$$H_2O_2 + O_3 \longrightarrow H_2O + 2O_2$$

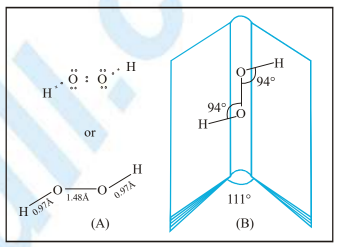
STRUCTURE OF HYDROGEN PEROXIDE

The vapour density as determined by Victor Meyer method at 90°C is 17. Hence, the molecular mass of H_2O_2 is 34. Two formulae have been suggested for hydrogen peroxide.

The calculated value of the single bond O–O

distance is 1.48Å and X-ray measurements shows what in hydrogen peroxide, O–O bond distance is 1.46 ± 0.03 Å. The value of dipole moment of H₂O₂ is 2.1 D. This suggests that all the four atoms do not lie in the same plane. The molecule can be pictured as lying on the spine of a book open to an angle of 111°. The hydrogen atoms are present one on each cover and H–O bonds making angles of 94° with the O–O bond as shown in fig. The bond distance between O–H is 0.97Å.





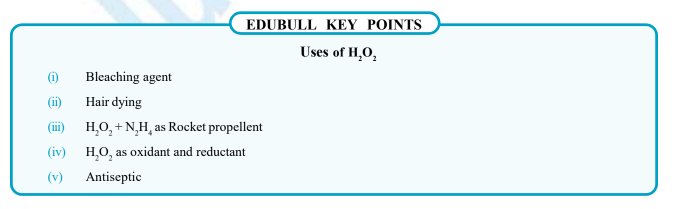
Precautions

(i) H_2O_2 can not be stored in simple glass bottles since rough surface of glass [alkali oxides present in it] excited by light and decomposed H_2O_2 .

So the H₂O₂ usually stored in coloured, paraffin wax coated, plastic bottle.

(ii) Always with H_2O_2 add small quantity of inhibitor or negative catalyst to stay decomposition of H_2O_2 .

like H_3PO_4 , acetanilides etc.





- **Ex.** What causes the temporary and permanent hardness of water ?
- Sol. Temporary hardness is caused by presence of soluble bicarbonates of calcium and magnesium, i.e. $Ca(HCO_3)_2$ and $Mg(HCO_3)_2$ in water whereas permanent hardness is caused by presence of soluble chlorides and sulphates of calcium and magnesium, i.e. $CaCl_2$, $CaSO_4$, $MgCl_2$ and $MgSO_4$ in water.
- **Ex.** What is 'demineralised water' and how can it be obtained ?
- **Sol.** Water which is free from all cations and anions is called demineralised water. It is obtained by passing hard water first through cation exchange resin and then through anion exchange resin.
- **Ex.** Explain the following :
 - (i) Soft water lathers with soap but not hard water.
 - (ii) Temporary hard water becomes soft on boiling.
 - (iii) Water can extinguish most fires but not petrol fire.
 - (iv) Hard water is softened before use in boilers.
- Sol. (i) Hard water contains calcium and magnesium salts. These react with soap to form insoluble calcium and magnesium salts of fatty acids, i.e., form scum and not lather.

 $2RCOONa (soap) + Ca^{2+} \longrightarrow (RCOO), Ca (insoluble) + 2Na^{+}$

 $2RCOONa (soap) + Mg^{2+} \longrightarrow (RCOO), Mg (insoluble) + 2Na^{+}$

(ii) Temporary hardness of water is due to the presence of soluble bicarbonates of calcium and magnesium. On boiling, the bicarbonates are converted into insoluble carbonates which can be removed by filtration.

 $Ca(HCO_3)_2(soluble) \longrightarrow CaCO_3 \downarrow + H_2O + CO_2$

 $Mg(HCO_3)_2$ (soluble) $\longrightarrow MgCO_3 \downarrow + H_2O + CO_2$

- (iii) When water poured over petrol fire, petrol being lighter than water floats over water and thus fire spreads instead being extinguished.
- (iv) Hard water consists bicarbonates and other soluble salts of calcium and magnesium. When hard water is evaporated in boilers, scales of precipitated carbonates of calcium and magnesium along with other salts such as sulphates, chlorides, etc., are formed. In order to avoid the formation of scales hard water is first softened before use in boilers.
- **Ex.** What happens when ?
 - (i) Heavy water reacts with magnesium nitride.
 - (ii) Heavy water reacts with sodium.

Ans. (i) $Mg_3N_2 + 6D_2O \longrightarrow Mg(OD)_2 + 2ND_3$ (Deutrammonia)

(ii) Na + 2D₂O \longrightarrow 2NaOD + D₂

Ex. Knowing the properties of H₂O and D₂O, do you think that D₂O can be used for drinking purposes?

Sol. Heavy water is injurious to human beings, plants and animals since it slows down the rates of reactions occuring in them. Thus, heavy water does not support life. In fact it retards certain cellular process, such as mitosis, cell division, etc, Thus, prolonged use of D₂O leads to degeneration of tissues.



Ex. Assertion : Anhydrous BaO_2 is not used for preparing H_2O_2 .

Reason : H_2O_2 is prepared on large scale by air oxidation of 2-Ethyl anthraquinol.

(1) If both Assertion and Reason are true and Reason is a correct explanation of Assertion.

(2) If both Assertion and Reason are true and Reason is not a correct explanation of Assertion.

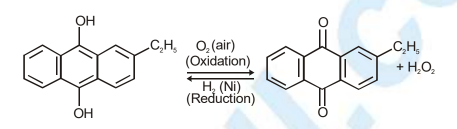
(3) If Assertion is true but Reason is false.

(4) If Assertion is false but Reason is true.

Sol. Assertion : $BaO_2 + H_2SO_4 \longrightarrow BaSO_4 \downarrow$ (white) $+ H_2O_2$

Insoluble $BaSO_4$ forms a thin layer around BaO_2 and therefore reaction occurs slowly and finally stops. So anhydrous BaO_2 , is not used.

Reason : Industrial method (Auto oxidation)



2- Ethyl anthraquinol

2-Ehtylanthraquinone

It is cheaper method as only O_2 from air and H_2 are consumed.

Ans. (2)

- Ex. Write chemical reactions to justify that hydrogen peroxide can function as an oxidising as well as a reducing agent.
- Sol. H_2O_2 can act as an oxidising as well as a reducing agent both in acidic and basic media as illustrated below. (i) Oxidising agent in acidic medium

$$2Fe^{2+}(aq) + 2H^{+}(aq) + H_{2}O_{2}(aq) \longrightarrow 2Fe^{3+}(aq) + 2H_{2}O(\bullet)$$

(ii) Oxidising agent in basic medium

 $Mn^{2+}(aq) + H_2O_2(aq) + 2HO^{-}(aq) \longrightarrow MnO_2(s) + 2H_2O(\bullet)$ manganese dioxide

(iii) Reducing agent in acidic medium

$$2MnO_4(aq) + 6H^+(aq) + 5H_2O_2(aq) \longrightarrow 2Mn^{2+}(aq) + 8H_2O(\bullet) + 5O_2(g)$$

(iv) Reducing agent in basic medium

Ex. Explain the following :

(i) Why hydrated barium peroxide is used in the preparation of hydrogen peroxide instead of the anhydrous variety ?

- (ii) Phosphoric acid is preferred to sulphuric acid in the preparation of H₂O₂ from barium peroxide.
- (iii) Statues coated with white lead on long exposure to atmosphere turns black and the original colour can be restored on treatment with H_2O_2 .
- Sol.

(i) If anhydrous barium peroxide is used in the preparation, the barium sulphate, thus formed, forms an insoluble protective coating on the surface of solid barium peroxide hydrated.

This prevents the further reaction of the acid, i.e., causing the reaction to stop. If, however, hydrated barium peroxide (in the form of the paste) is used, the water causes to dislodge the insoluble $BaSO_4$ from the surface of BaO_2 . Hence $BaSO_4$ settles at the bottom of the reaction vessel and the reaction continues without any difficulty.



- (ii) When phosphoric acid is used in the preparation of H_2O_2 from BaO_2 , it plays the dual role. It liberates H_2O_2 and also acts as a preservator by retarding its decomposition.
- (iii) White lead is used as a pigment. The statues coated with white lead get blackened due to the action of H_2S present in atmosphere in traces.

$$Pb(OH)_2$$
 (white lead).2PbCO₃ + 3H₂S \longrightarrow 3PbS + 2CO₂ + 4H₂O

DIHYDROGEN AS A FUEL

Dihydrogen releases large quantities of heat on combustion. The data on energy released by combustion of fuels like dihydrogen, methane, LPG etc. are compared in terms of the same amounts in mole, mass and volume, are shown in Table

THE ENERGY RELEASED BY COMBUSTION OF VARIOUS FUELS IN MOLES, MASS AND VOLUME

Energy released on combustion in kJ state)	Dihydrogen (in gaseous state)	Dihydrogen (in liquid)	LPG	CH ₄ gas	Octane (in liquid state)
per mole	286	285	2220	880	5511
per gram	143	142	50	53	47
per litre	12	9968	25590	35	34005

From this table it is clear that on a mass for mass basis dihydrogen can release more energy than petrol (about three times). Moreover, pollutants in combustion of dihydrogen will be less than petrol. The only pollutants will be the oxides of dinitrogen (due to the presence of dinitrogen as impurity with dihydrogen). This, of course, can be minimised by injecting a small amount of water into the cylinder to lower the temperature so that the reaction between dinitrogen and dioxygen may not take place. However, the mass of the containers in which dihydrogen will be kept must be taken into consideration. A cylinder of compressed dihydrogen weighs about 30 times as much as a tank of petrol containing the same amount of energy. Also, dihydrogen gas is converted into liquid state by cooling to 20K. This would require expensive insulated tanks. Tanks of metal alloy like NaNi₅, Ti–TiH₂, Mg–MgH₂ etc. are in use for storage of dihydrogen in small quantities. These limitations have prompted researchers to search for alternative techniques to use dihydrogen in an efficient way.

In this view Hydrogen Economy is an alternative. The basic principle of hydrogen economy is the transportation and storage of energy in the form of liquid or gaseous dihydrogen. Advantage of hydrogen economy is that energy is transmitted in the form of dihydrogen and not as electric power. It is for the first time in the history of India that a pilot project using dihydrogen as fuel was launched in October 2005 for running automobiles. Initially 5% dihydrogen has been mixed in CNG for use in four-wheeler vehicles. The percentage of dihydrogen would be gradually increased to reach the optimum level.

Nowadays, it is also used in fuel cells for generation of electric power. It is expected that economically viable and safe sources of dihydrogen will be identified in the years to come, for its usage as a common source of energy.



HYDROGEN AND ITS COMPOUNDS

COORCELE, THIS & FORMELIAS

Preparationof hydrogen

(f) Passing steam over hot iron (Lane process)

 $3Fe = 4H_sO \rightarrow Fe_3P_4 = H_s \uparrow$

(II) By the action of water on hydrolith.

$$C_{0}H_{2} + H_{2}O \longrightarrow Cu(OH)_{2} + 3H_{2}$$

(III) By the electrolysis of water

(IV) Bosch process

$$C(s_1 - H_sO(g) \longrightarrow CO - H_g$$

$$-$$

$$\Pi_sO(g) \xrightarrow{(o_1, o_2)} CO, \quad \Pi,$$

$$\downarrow$$

$$25 a.m.$$

$$CO, absurbed$$

Properties of molecular hydrogen

Metuls like Pd. Pt. Ni, Collett, adsorb large quantities of the gas due to vacancies between these atoms. This is known as "occlusion".

Reaction with vegetable oils

Vegetable oil – $H_2 \xrightarrow{127.6}{13} \Rightarrow$ Vegetable gheet (Slid Iat)

This process is known as "hydrogenation." or "hardening of eils"

Isotopes of hydrogen

(a) Hydrogen (Protium)	(b) Deuterium	(c) Tritium1
1 H' or H	H [^] er D	H ³ or T
(P 1,* 1.u 0)	(P 1.e 1.u 1	(P 1.e 1.n 2)

Different forms of hydrogen

Atomic hydrogen

H ______ 2H

(Molecular hydrogen) (Atomic hydrogen)

Nuscent hydrogen

Zo_dil.11.80. > Zn80, 1211

Ortho and Para hydrogen

If two nuclei have same spin then it is called "Ortho H_3 ". If two nuclei have different spin then it is called "Para H_2 ". HYDRIDES



- Ionic or salt like hydrides : s block
- Li H, NaH, KH, RbH, SrH₂, BaH₂ etc.

Be & Mg hydrides are covalent in nature

Molecular or covalent hydrides : p block

NH₃, PH₃, H₂O, CH₄ etc.

Metallic or Interstitial hydrides : trasition elements

In these hydrides, hydrogen atoms occupies the interstitial position of metallic latice.

 ${\rm La\,H}_{2.87}, {\rm Ti\,H}_{1.8}, {\rm Zr\,H}_{1.9}$

Hydride gap = 7, 8, 9

WATER

- (I) The existence of hydrogen bonding is responsible for high values o specific heat, the latent of fusion and latent heat of vapourisation and high boiling point.
- (II) Some of the covalent compounds such as alcohols, carboxylic acids and carbohydrates are soluble in water due to formation of hydrogen bonding.

Hardness of Water

Temporary hardness of water is due to bicarbonates of Ca and Mg. It can be removed by -

- (a) Boiling
- (b) Clark method using CaO

Permanent hardness of water is due to chloride/sulphate of Ca and Mg. It can be removed by -

- (a) Adding washing soda, Na_2CO_3
- (b) Calgon $[Na6(PO_3)_6]$
- (c) Permutit, $Na_2 Al_2 Si_2 O_8 xH_2 O$
- (d) Artificial resins, $RSO_3H \& RNH_2OH$

Heavy water or Deuterium oxide (D,O)

- (a) It is produced by repeated electrolysis of ordinary water containing alkali.
- (b) Most important use of heavy water is in nuclear reactors for slow down the speed of neutron (i.e. as a moderator)

HYDROGEN PEROXIDE

Preparation

- (I) BaO_2 . $\operatorname{H}_2O + \operatorname{H}_2SO_4 \longrightarrow \operatorname{BaSO}_4 + \operatorname{H}_2O_2 + 8\operatorname{H}_2O$
- (II) $\operatorname{Na_2O_2} + 2 \operatorname{H_2SO_4} \longrightarrow 2 \operatorname{Na_2SO_4} + \operatorname{H_2O_2}$
- (III) $BaO_2 + H_2O + CO_2 \longrightarrow BaCO_3 + H_2O_2$
- (IV) Oxidation of 2 Alkyl anthraquinol
- (V) Electrolysis of 50% H₂SO₄

Chemical properties

(a) **Decomposition**

$$H_2O_2 \longrightarrow H_2O + O_2$$

 $\downarrow \qquad reduction \qquad \uparrow$

- (b) H_2O_2 can accept as well can donate electrons and thus it can acts as an oxidising and reducing agtent in acidic as well as alkaline medium
- (c) **Oxidising properties** : It is a strong oxidising agent under acidic and alkaline conditions. Oxidation in



acidic medium is generally slow while rapid in alkaline solution.

$$H_2O_2 + 2H^+ + 2e \longrightarrow 2H_2O$$

In alkaline solution

 $H_2O_2 + 2OH^- + 2e \longrightarrow 2H_2O + 2O^{2-}$

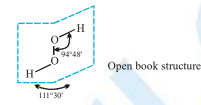
(d) **Reducing properties**

It acts as a reducing agent towards strong oxidising agents in acidic as well as alkaline medium. Reducing action in acidic solution is slower than in alkaline solution.

(e) **Bleaching action**

$$H_2O_2 \longrightarrow H_2O + [O]$$

Structure



Structure of H_2O_2 in the gas phase.

Tests of H₂O₂

- (a) It liberates iodien from KI solutin in the presence of ferrous sulphate.
- (b) It gives orange red colour with acidified titanium oxide due to formation of pertitianic acid.

 $\mathrm{Ti}(\mathrm{SO}_4)_2 + \mathrm{H}_2\mathrm{O}_2 + 2\mathrm{H}_2\mathrm{O} \longrightarrow \mathrm{H}_2\mathrm{TiO}_4 + 2\mathrm{H}_2\mathrm{SO}_4$

(c) It gives blue colour with acidic field $K_2Cr_2O_7$ and ether. The blue colour of chromium peroxide is stabilized by ether.

 $\operatorname{CrO}_4^{2-} + 2\operatorname{H} + 2\operatorname{H}_2\operatorname{O}_2 \longrightarrow \operatorname{CrO}_5 + 3\operatorname{H}_2\operatorname{O}_5$

