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## **Inorganic Chemistry**



Haydrogen O Hydrogen choices similar characteristics to both alleali <sup>∞</sup> Graup  $\rightarrow$  1 (Alkali metals) ©Period  $\rightarrow$  1 Non metal metals (Can.1) and halogens (Can. 17) Dihydrogen (Hz): Host abundant molecule in the universe (70%) Isotopes: profium (:H), deuterium (?H), tràtium (3H,) Only nachicachive 0.01567. in terrestrial Hydroger, emits & particle most abundant mostly in form of HD ty = 12.33 years. • Granulated In and dilute HCI: Zn + 2HCI -> ZnCl2 + H2 -> [Lab prep] [Mina Ashido] • In and aquous alkali -> In + 2NaOH -> Na, In O2 + H2 • Electro-lysis of acidified water:  $2H_2O(1) \xrightarrow{7} 2H_2(9) + O_2(9)$  [Undercoater Kaminari] · High purity (>99.95%) H2 is obtained by electrolysing coarm aquious barium hydroxide solution w/ nickel electrodes. • Electrolysis of brine solution; 201 2H2O(1) + 20 - Mgol H2(3) + 20H (aq) [As a by product] • Steam on hydrocarbon/coke at high T & catalyst: Cn Hzn+2 + nH20 -1270K > nCO + (2n+1) Hz  $C H_4 (9) + H_{10}(9) \xrightarrow{1270K} CO(8) + 3B H 2(9)$ > water gas / synthesis gas/ & Prep of Syngas from Coal (Coal Garification): syngors Knep of Hydrogen  $C(s) + H_2O(g) \xrightarrow{1270k} CO(g) + H_2(g)$ tlina coalked in the lab heith a piece of Zinc Colourless, Odourless, tasteless to make H2 and found Kaminara sacimming → comburtible, insoluble in H2O in a cidified water. After drying him for - (Phynical - ) Lighter than air, PROPERTIES Chemical -> (H-H bond dissociation enthalpy is the highert for single high purity bakugo and tetsu tetsu were bond betaseen two atoms of any a element.) called. they took him to Doctor Brown. Can Lose 1 c to form Ht, gain to form H- or There was steam coming out of hisroom, in such hotness he was drinking coke form covalent bond. → 2HX(8) Reaction with Halogens: H2(9) + X2(9) with his cat in lap. MP Reaction with f occurs even in dark . Io dine needs catalyst.  $\rightarrow 2H_{2O}(l) + \Delta H$ Reaction with O2: 2H2(4+ O2 (3) - cat/A Reaction with N2: 3H2 + N2 673K, 2000th 2NH3 + All [Habers Process] Reaction with metal ion and metal oxids: H2(9) + Pd 2+(aq) → Pd(s) + 2H+ (ar) +yH20(1)  $y_{H_2}(g) + M_{x}O_{y}(s) \longrightarrow x M(s)$ @ <u>Reaction with Organic compounds</u> Hydrogenation of negetable oils using nickel as catalyst gives edikle fats. · Hydroformylation of olefins yields aldebydes celvich further undergo reduction to give alcohols.  $\rightarrow$  RCH<sub>2</sub>CH<sub>2</sub>-CHO ; H<sub>2</sub>+RCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CHO  $\rightarrow$  RCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH > Ionic or saline or sattlike by duides  $H_2 + CO + RCH = CH_2 -$ Thes of H Amelia takes a feit organo-metallic > Covalent or molecular hydrides @ Uses of Hydrogen>> pipe to weld the rocket full · synthesis of ammonia tank lifted by hydrolic, but > Metallic or non-stoichiometric hydride · manufacture of vanaspatifat the electricity ready out · manufacture of organic compounds @Hydrides (EHx → E\*+ +xH-) particularly methan of -> Ionic or Soline Hydrides» . stoichiometric compounds with highly electropositive elen · manufacture of metal hydrides, • Greverally ionic but covalent character found in light metal hydrides · preparation of HCI · Ionic ones are crystalline, non-volatile, non · Reducing heavy metal oxides to · BeH2, Mg H2 are polymeric · But melts of ionic ones a conduct electricity and likerate Hz solid state. metal · Saline hydrides react violently with H, O producing H, gas. · Atomic II and ony-hydrogen torches used for cutting and welding. NaH(st H2O(99)  $\longrightarrow$  NaOH (09) + H2(9) [Atomic It are alloced to recombines · Lit is nather unreactive at moderate temperatures with 0, or C/2. on the surface generating 4000k] It is therefore, used in the synthesis of other useful hydrides. · Rocket fuel. . . Used in fuel cells for generating 8Liti + Al, Cl6  $\longrightarrow$  2LiAlHg + 6LiCI electrical energy  $2LiH + B_2H_6 \longrightarrow 2LiBH_4$ 🚹 EXAMSROADOFFICIAL < EXAMSROAD 🔟 EXAMSROAD 💟 EXAMSROAD

@ Covalent/Holecular Hydries . covalent molecules formed with the p block elements. [CH4, NH3, H, O, HF etc] · Being covalent, they are volatile Form Hydrogen bond → electron-deficient: Incomplete octate, electrion que acceptors, acts as lessis acid, group (13) > electron-precise: complete octate, All group 14 hydridus -> electron-ruick: More than 8 e-s in last shell, 10 graup (15-17), electron donor, Lewis base Metallic/Non-stoichiometric Hydrides . d compounds with d block and f block [7,8,9th Crowp elements don't] • From 6th group only Chromium forms Crtt · Good conductors of heat and electricity. Pd Ho, 6-0.2 •Almost always non-stoichiometric e.g. La H2.87, Yb H2.55, Ti H1.5-1.8, Zr H 1.3-1.75, VH 0.50, Ni H0.6-0.7 · Some metals absorbs He into their lattice, they are called interstitial hydrides. This property is used is acidely used in catalytic reduction / hydrogenation. · Pd, Pt can accommodate a very large volume of Hydrogen tarteless, colourless liquid I high mp freezing point, b.p., heat of vap , heat of funion → high thermal conductivity, sweface terrion, high specific heat capacity @WATER . Human body has about 65%. PROPS dipole moment and dielectric corut. 00 95.7pm-Structure - Anamelous behaviour 104.5° · highly polar molecule · sp3 hybridised Structure of Ice Each O atom is sorrounded tetrahedrally by form O atoms. . In liquid face, there exist intermolecular H-bond. · Dencity of ice is less than Water. · Redox involving H2O H2O is easily reduced to I H2 by Chemical · Amphoteric Nature: Reacts with as an acid with highly electropositive metals. NH3 and as a base with H2S.  $2H_20 + 2N_a \longrightarrow 2N_a0H$ + H2 H20 + NH3 == 0H- + NH4 Oxidised to Oz during photo synthesis and with F2 too. H20+H2S = H30+ +HS- $\rightarrow 4H^+ + 4F^- + O_2$ 2f2 +21+20 -· Auto-protolysis(self-ionization) · Hy drates formation From aqueous solution many salts  $H_{20}(l) + H_{20}(l) \rightleftharpoons H_{30}^{+} + 0H^{-}$ can be crystallined as hydrated salts, callification · Hydrolyiis Reaction Due to high dielectric constant, -> coordinated coates [Cr (H20)6] 3+ 3CT it has a very strong hydrating tendency. → interstitial water Bac1, .2 H20  $P_4 D_{10} + 6H_2 O \longrightarrow 4H_3 PO_4$ -> Hydrogen-bonded water [Gu(H20)4]2+SO42-, CuSO4.5H20  $S_iCl_4 + 2H_{20} \longrightarrow S_iO_2 + 4HCl$ @ Soft Water : Barically pure from minerals and readily  $N^{3-} + 3H_2O \longrightarrow NH_3 + 3OH^$ forms lather with soap. @ Hard Water Contains carbonate, bicarbonates, and chlarides and sulphates. Basen't readily form > @ Temporary Hardness [Due to Mg/Ca bicarbonate] can be nemoved by lather. (forms scum/precipitate with soap) So Boiling By boiling, soluble Mg (HCO)2 and Ca (HCO2)2 are [soap containing sodium stearate (C17 H35 (OO Na) reacts converted into insoluble Mg (OH), and Ca(HCOJ) Ca (03 with hard water to precipite out Ca/mg stearate] respectively, which forms precipitate. " Clark's Method Calculated amount of added lime precipitates 2C17 H35 COO Na (aa) + M 2+ (aq) out en Cacos and Mg (OH) (C1, H35 COO)2 M & + 2 Na+ (aq) [M → ca/Mg]  $Ca(HCO_3)_1 + Ca(OH)_2 \longrightarrow 2CaCO_3\downarrow + 2H_2O$ Classification - Temporary Hardness  $Mg(H(0_3)_2 + 2G(0H)_2 \longrightarrow 2Ga(0_3)_+ Mg(0H)_1 + 2H_20$ @ Permanent Hardness @ Permanent Hardness [ Due to Hg/Ca chlorides/sulphates] > Ocalgon's method Sodium hexameta phosphate (Na6 P6 O18) commercially can be removed by. called ' calgon'. The following reactions occurs. So Treating w/ washing sola (Na, CO3) Converts the solube salts into insoluble carbonates. Nac Pe O13 - 2Na + Na4 Pe O13 - $MCl_2 + Na_2CO_3 \longrightarrow MCO_3 \downarrow + 2NaCl_3$  $M^{2+} + Na_4 P_6 O_{13}^{2-} \longrightarrow \left[ Na_2 M P_6 O_{13} \right]^{2-} + 2 Na_{10}^{+} H$  $MSO_4 + Na_2CO_3 \longrightarrow MCO_3 \downarrow + Na_2SO_4$ This complex anion keeps the Mg 8Lit + AbCls -> 2Litilly + 6Lici and Ca2+ ions in solution offit + Bits - slink

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	S-Block
· Gro	ups 1 & 2 Group 1: Li, Na, K, Rb, Cs, Fr [ns] @ Group 2: Be, Mg, Ca, Sr, Ba, Ra [ns <sup>2</sup> ] <u>Alkali metals</u> <u>Alkali metals</u>
M From	it radioactive, 223 Fr has a ch = 21 minutes of bolger and configuration of the first of the fir
OLi sk	news similarities to Mg and Be to AI [Diagonal relationship] -> Use to similar size and I'm ratio,
	oup 1: Alkali metals @ Higly electropositive metals and lose 1 electron to form M+ ions.
II TRE	NDSI wic/Ioiric Radii: Increase down the group [Li < Na < K< Rb < Cs < Fr]. Ions are smaller than parent atom. Withium has mor hydration enthalpy, thus liselth are morely hydrated [LiCI-2H20]
> Hur	ration Enthology: Decrease down the group LLiman Marine -
<b>IIPHYS</b>	ICAL PROPERTIES DU
@ All o	ne silvery white, soft and light metallic bonding.
OH.P O O & Impl	und B.P. s are low due to weak metallic bonding. and B.P. s are low due to weak metallic bonding. art characteric flame in oxidising flame due to shifting of excited electron and coming back.
Metal	
Li Na	Crimson Red 670.8 Yellow 589.2
K	Violet 766.5
Rb	Red ruidet 780.0
Cs	Blue 455.5 Blue 455.5 Blue 455.5 I TO - is stable w/ large
Real Real	clion with Air: • They ternish in dry air the borns nonoxide, bitness form superoxides (K, Rb, cs) -
	Blue 455.5 YICAL PROPERTIES II ction <u>cuith</u> Air: • They ternish in dry air <del>coh</del> i due to formation of oxides, which in turn reacts with moisture to form hydroxide. • Na forms peroxide, Li forms monoxide, bthens form superoxides [0] is stable w/lange to form hydroxide. • Na forms peroxide, Li forms monoxide, bthens form superoxides [M = K, Rb, Cs] +02
411	+02 2L120 UNION / ZITU OF II Hu with No of air to form nitride [L131]
04 4	10 ces exception w between our these metals are kept tub merged in Neroscie (2M+2H20 -> 2M++20H++H2)
. Deed	in with histor @ Alkali metais to
€Li→	most -ve c the tradaction energy, on etc.
This Also	is due to small size and very high Hydrotion energy, oner terminal alloynes. react with proton donors like alcohol, gaseous ammonia and terminal alloynes. (400°C) (200°C) (20
-> Read	is due to small size and very high right amonia and terminal allynes. react with proton donors like alcohol, gaseous ammonia and terminal allynes. (400°C) (800°C) (800°C) (100°C) (10
2M+	ion with H2: They react with H2 at about on the form control of the form ionic holides. H2 $\longrightarrow 2M^+H^-$ cn with Halogens; React vigorously wto form ionic holides. (LiI is most covalent due to large anion) (LiI is most covalent due to large anion)
-> Reacti	on with Halogens; @ React vigorously wto form ionic houses, e thep
nucle.	Shang adving agents, Li stronger, the comment internet in the stronger the devices the devices the devices in t
-> Solutio	ing Nature: © Strong reducing agents, Li strongert, Na ceeclest. $M(g) \rightarrow H^+(g) + e^-$ Liousanton on enthalpy] ing Nature: © Strong reducing agents, Li strongert, Na ceeclest. $M(g) \rightarrow H^+(g) + e^-$ Liousanton on enthalpy] ons in liquid Ammonia: © Dissolve in liq NHs giving deep blue solution $H^+(g) + H_2O \longrightarrow M^+(aq)$ [Hydrotion enthalpy] which are conducting in nature. which are conducting in nature. (shich are conducting in nature. (shich are conducting in nature. (shich are conducting in nature. (shich are conducting in formation of the solution © Solution is paramagnetic. (H(NH_3) x]^+ + [e(NH_3)y]^- thus impaires blue colour to the solution © Solution is paramagnetic. (H) (NH_3) + [e(NH_3)y]^- thus impaires blue colour to the solution © Solution is paramagnetic. (shich are colour dranges to bronze colour and becomes dramagnetic. SII SII
M+(x	++)NH2→ [M(NH2)2]+ + (NH3)2] - The blue colour is due to the solution @ Solution is paramagnetic ++)NH2→ [M(NH2)2]+ + ((NH3)2) - thus imparts blue colour to the solution @ Solution is paramagnetic
o On s	tanding slowly, liberates hydrogen resulting in formation of amide. [M <sup>+</sup> cam) + e <sup>-</sup> + NH3(1) - + NH12(am) + 21/2 -
	$C_{nn} = \frac{1}{2} 1$
OLi With	SID 1 Pb makes 'celuite metal' ball bearings for motor engine. @ PLi ceith AI is used to make air craft parts. 1 Pb makes 'celuite metal' ball bearings for motor engine. @ PLi ceith AI is used to make air craft parts. 1 Pb makes 'celuite metal' ball bearings for motor engine. @ PLi ceith AI is used to make air craft parts.
🖉 Li avit	h Mg alloy is used to make armous process of the and Phile. When to be used as anti-concurry and
1 tiouid	No is used as apple this fast breader nuclear reactors. OK has with role for biological systems. OKCI > fertilized
KOH is	s used in the manufacture of soft soap, Also used as an excellent absorbent of CO2
Cs is	used in devising photoelectric cells.
	The start balls not up of the of open to have considered a starting of the start of

VI COMPOUNDS OF ALKALI METALOD	and the second
-> Oxides & Hychroxides @ Li form oxide (Lizo) and penovide (Lizo), Na Jorms peroplat ( 1)	レート
Increasing stability of perovide and hydroxide, as the size of the metal ion increases, is due to the stability of perovide and hydroxide, as the size of the metal ion increases, is due to the stability formation of hydroxide by hydroxide.	2 84
$M_2O + H_2O \rightarrow 2H^+ + 2OH^- / H_2O_2 + 2H_2O \rightarrow 2H^+ + 2OH^- + H_2O_2 / 2H_2O =$	
$M_2O + H_2O \rightarrow 2M^+ + 2OH^- / H_2O_2 + 2H_3O \rightarrow 2H^+ + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2H_2O - 2H^- + 2OH^- + H_2O_3 / 2HO_2 + 2H_2O - 2H^- + 2$	*
@ Alkalimetal hydroxides are strongest of all bases and dissolve freely in H,O, releases heat on incluse regarderies	5
-> <u>Halides</u> @ Gystalline colourless solids with high melting point.	- 1 · ·
<ul> <li>C All these holides have high regative enthalpies of formation: Δ<sub>c</sub>H<sup>O</sup> values for fluorides become less negative as the go do can the group but that for chlorides, Bromides and Iodicles are neverse.</li> <li>O for a given metal Δ<sub>f</sub>H<sup>O</sup> always becomes less negative from flow ride to iodicle</li> <li>M.p. 8 b.p → Fluoride &gt; Chloride &gt; Bromide &gt; Iodicle → All are soluble in water.</li> </ul>	ローショー
@EXC Low solubity of Lif in water is due to its high lattice enthalpy cohereas low solubility of CSI is due	1
to smaller Hydration enthalize of its top ions	「日日」
<ul> <li>Other halides of Li are soluble in ethanol, actione and ethylacetate; Lici is soluble in pyreidine also.</li> <li>→ Salts of Oxo-Acids @ Acidic proton and oxo group both are attached to the same hydroxyl group. H+203 → occol</li> <li>@ Alkali metals form salts with all the oxo acids (Generally soluble).</li> </ul>	シュリ
-> Salts of Oxo-Acids @ Acidic proton and oxo group both are attached to the same hydroxy group. [""	1 13
42 C3 is not statue, earling small control in pointer C3 for watering is	010
© Litto; doern't exist as a solid.	AN
Anomalous Properties of Li	M
<ul> <li>→ Cause: @ 2xceptionally small size. @ High polarising power (\$a) (2/n ratio).</li> <li>→ Cause: @ 2xceptionally small size. @ High polarising power (\$a) (2/n ratio).</li> <li>→ Points: @ Increased covalent character which nexults in solubility in organic conpounds. @ Diagonal relationship with Mg.</li> <li>@ hander and higher m.p., b.p than other alkali metals @ Li is least neactive @ strongert neducing agent among Gap:</li> <li>@ Forms nitride on compusition like no other metal. alkali metal.</li> <li>@ Forms nitride on compusition like no other metal. alkali metal.</li> <li>@ Lici is deliquescent and crystallises as a hydrate (Lici.2420), othe alkali vnetal khlorides dort</li> <li>@ Lici is deliquescent and crystallises as a hydrate (Lici.2420), othe alkali vnetal khlorides dort</li> <li>@ Li khlor ides dont form achill all other elements form 'solid hydriogen carbonate.</li> <li>@ Li htco; is not obtained in the solid form achill all other elements form 'solid hydriogen carbonate.</li> <li>@ Li while other alkali metals dont form ethynide on reaction ceith ethyne.</li> <li>@ Li No; gives Li,0 on heating other doenpose [4LiNo; → 2Li,0+4No; +0, /2NaNo; → 2NaNO; +02]</li> <li>@ LiNo; gives Li,0 on heating other due to covalency.</li> <li>@ Li f and Li;0 are niuch less soluble. due to covalency.</li> <li>@ Li f and Li;0 are niuch less soluble. due to covalency.</li> </ul>	1
O horder and higher m.p., b.p. than other alkali metals @ Li is least reactive ostrongest reality of	1
© Forms nitride on compusition like no other mean hydrate (Lic1.24,0); othe alkali metal khlorides don't @ Lici is it deliguescent and crystallises as a hydrate (Lic1.24,0); othe alkali metal khlorides don't	
@ LiHCO3 is not obtained in the solid form achill all other elanews form studies of reaction ceith allyne.	,Ó
@ Li unlike other clical metals the formal acompose [4LiNo3 -> 2Li20 + 4NO2 + 02 / 2NaNO3 -> 2NaNO2 + 02]	5
· Lif and lizo are nuch less soluble. due to covalency.	· ···
@ Both Li and My are harder, and in a start of her	1º
Both neact slowly with water . Orides and theme hudganister and much adulate the trite	0
an heating.	14
The oxides as Lizo and 20 as not compared by 21 and propaging the second the noit	113
Both react stocky, with water . Orides and theme hydroxides are, much soluble. hydroxides die decompose an heading. The oxides as Lizo and M20 do not combine with excess of as Coth Lizo are soluble in ethanol as Both Lizo are soluble in ethanol a Both Lizo and MgCl3 are delequeost and , crystalline.	T
10 COMPUNDS OF Nall Of Generally prepared by soviey process.	10
10 COMPUNDS OF Na 11 -> Warling Soda (Na CO3. 10 H, of @ Generally prepared by soviey process. -> Warling Soda (Na CO3. 10 H, of @ Generally prepared by soviey process. -> Warling Soda (Na CO3. 10 H, of @ Generally prepared by soviey process. -> Warling Soda (Na CO3. 10 H, of @ Generally prepared by soviey process.	U.
· Advantage of Low something of the parting CO2 to a conc solution of NaCl stasaturated with NH3	)
$2NH_3 + H_2O + CO_2 \longrightarrow (NH_4)_2 CO_3 / (NH_4)_2 CO_3 + H_2O + CO_2 \longrightarrow 2NH_4 HCO_3 / NH_4 HCO_3 + Nact \longrightarrow NH_4 CO_4 + H_2O_7$	
(NH4)HCO3. (NH4) HCO3 IS OCCURRENT OF O 2NH3 + H2O + CO2 → (NH4)2 CO3 / (NH4)2 CO3 + H2O + CO2 → 2NH4 HCO3 / NH4 HCO3 + NaCI → NH4CI + Na HCO3 @ NaHCO3 crystal separates. These are heated to give @ Na2CO3 [2Na HCO3 → Na, CO3 + CO2 + H2O] @ NaHCO3 crystal separates. These are heated to give @ Na2CO3 [2Na HCO3 → Na, CO3 + CO2 + H2O] [2 Na HCO3 crystal separates. These are heated to give @ Na2CO3 [2Na HCO3 → Na, CO3 + CO2 + H2O] [2 Na HCO3 crystal separates. These are heated to give @ Na2CO3 [2Na HCO3 → Na, CO3 + CO2 + H2O] [2 Na HCO3 crystal separates. These are heated to give @ Na2CO3 [2Na HCO3 → Na, CO3 + CO2 + H2O] [2 Na HCO3 crystal separates. These are heated to give @ Na2CO3 [2Na HCO3 → Na, CO3 + CO2 + H2O]	5
10 In this process NH3 is recovered when the other of 0	1
@ Solvay process can't be extended to the manufacture of K2CO3 bez KHCO3 is too soluble to be	,
precipitated by the addition of NHaHCO, to a satured solution of KCI.	1.7
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-> Sadium Chloride (Nacl) • Most abundant source is soa water containing 2.7 to 2.9% by mars of Nacl.
• In tropical counties, it is prepared by sclar evapowration [50 latch tons/year in India] as impurities,
• On Crude satium chloride, generally obtained from crystallisation of brine solution, contains Marsol, (asol, (asol, (ac)), 19-2
<ul> <li>In tropical counties, it is prepared by sclar evapouration [50 latch tors/year in India] as impurities.</li> <li>Or Crucle satium chloride, generally obtained from crystallisation of brine solution, contains Na, 504, Cas04, Cac12, MgC12<sup>*</sup></li> <li>Cac12, MgC12 are impurities because they are deliquesed (absorbs moisture)</li> <li>To obtain pure Nac1, caude salt is dissolved in minimum amount of coaler and fillered to remove insoluble impurities. The solution is then saturated with HCI gas. Cystals of pure sodium chloride separate out. Calcium and mg MgC12.</li> </ul>
• To obtain pure Naci, caude salt is dissolved in minimum amount of exceed and fillened to remove insource and mg Mg Cl2.
being more solution, normans in solution. Sodium chlorido mells al 1081 K, solubility = 36g, solubility doesn't increase approciably with increase in tamp.
$ \underbrace{\text{Caustic Soda (NaOH)}}_{\text{Cathodo}: Na^{+} + e^{-\frac{119}{2}} \text{Na-amalgam / Anodo: Cl^{-} \rightarrow \frac{1}{2}Cl_2 + e^{-} $
-> Caustic Soda (NaOH) @ propared commercially by electrolysis of NaCl in Caster-Kellner cell, at controls or on Cl2 is avolved at anode
Cathodo: Na <sup>+</sup> + $e^{-\frac{1}{2}}$ Na-amalgam / Anodo: $c_{1}^{-} \rightarrow \frac{1}{2}c_{12}^{-} + e^{-\frac{1}{2}}$
© Treated with water, 2 Na-amolgam + 2H2O -> 2 NaOH + 2Hg + H2 © NaOH is white translucent solid, m.p = 591 K, readily soluble in water to give strong alkaline solution. © Orystals of NaOII are deliquescent, NaOII at the swepace reacts with CO2 in atmosphere to form NaCO3.
oughter of have the here a high when the here a fining a purification of pauxite
@ In textile industries for mercerising conten juenais @ for preparation of public, or as a final of a
-> Baking soda (Na HCO2) @ Decomposes on heating to redeare CO2 making cakes or partnies fluffy.
Made by saturating a solution of Na2CO3 with CO2, while crystalline Na HCO3 power, being less soluble, seperates out,
$Na_2CO_3 + H_2O + CO_2 \longrightarrow 2 NaHCO_3 @ Hild antiseptic and used in fire extinguishers.$
@ Biological Importance of Na and K
O Biological Importance of the and the outside of cells, being located in blood plasma and in the interestitial fluid ashich sorrounds the cells. These ions participate in the transmission of nerve signals, in regulating the Acro of coater across sorrounds the cells. These ions participate in the transmission of nerve signals, in regulating the Acro of coater across
cell membranes and in the transport of sugars and ammino acids into cells
cell membranes and in the transport of sugars and ammino acias into certain the transmissions of nerve signals. K <sup>+</sup> ions are the most abundant ations withing cell fluids, where they activate en zymes, participate in the oxidation of glucase to produce ATP and with Nat, are responsible for the transmissions of nerve signals. The ionic and light across variants blood cells demonstrate that a discriminatory mechanism called the sodium-
oxidation of glucase to produce ATP and with Nat, are responsible for the transmissions of view -
• The ionic gradiant across various blood cells demonstrate that a discriminatory menument of the ATP used
• The ionic gradiant across various blood cells demonstrate that a discriminatory mechanism called the sodium- -potassium pump, operates across the cell membrianes achich consumes more than one-thind of the ATP used by resting animal and about 15 kg por 24 h in human.
by resting animal and about 15 kg. per 24 h in human.
bigen de la la service de la
@ Electronic conf: ns <sup>2</sup> @ Elements: Be, Mg, Ca, Sr, Ba, Ra     [II] TREDIDSED → Atomic Radii/Ionic Radii: Be (Mg (Ca (Sr CBa (Ra CAll are smaller than corresponding Alkali metalistics)     [Ra > Ba]     [II] TREDIDSED → Atomic Radii/Ionic Radii: Be (Mg (Ca (Sr CBa (Ra CAll are smaller than corresponding Alkali metalistics)     [Ra > Ba]     [II] TREDIDSED → Atomic Radii/Ionic Radii: Be (Mg (Ca (Sr CBa (Ra CAll are smaller than corresponding Alkali metalistics)     [Ra > Ba]     [II] TREDIDSED → Atomic Radii/Ionic Radii: Be (Mg (Ca (Sr CBa (Ra CAll are smaller than corresponding Alkali metalistics)     [II] TREDIDSED → Atomic Radii/Ionic Radii: Be (Mg (Ca (Sr CBa (Ra CAll are smaller than corresponding Alkali metalistics)     [II] TREDIDSED → Atomic Radii/Ionic Radii: Be (Mg (Ca (Sr CBa (Ra CAll are smaller than corresponding Alkali metalistics)     [II]
DUTREDIDSD -> Atomic Radii/Ionic Radii: Be < My < Ca < Sr < Ba < Ra LAII are smaller man control of the smaller man control of th
→ IE: Inonation Ionisation Enthology decreases down the group. [Ra> Ba]
Hydration Entructures. Be Mg > Ca > Sr > Ba L [More than alkali ructure] hydrated linu ve unit of
ID PHYSICAL PROPERTIESDO: @ Gunerally are silvery cohite, lustrous and relatively soft but harder than alkalimetals.
@ Be, Mg are greyich @ M.Ps and B.Ps are higher than corresponding
· Electropositive charachter increases down the group from Be to Ba
© Ca, Sr, Ba import characteristic brick red, orimson and apple green colours respectively to the frame, the es in Be and My age too trapitly boy held to be excited and then release energy, hence no colour.
(tigh conductivity and typical metal properties.
M CHEMICAL PROPERTIES MILONS reacting than alkali mercus /
- Ren (of Air and H.O: @ Ro the see lingtically inert to air and water due to formation of a la film on their storgale.
But powdered Bo burns brillionty on ignation in air to give dec and ises 12
Mg is more electropositive and burns with dazzling brilliance in air to give MgO and Mg3N2.
Ca, sr, is a are readily attacked ky air to form on the and intride. React also with H2 0 with increasing vigour even
in cold to form by drawides.
And BO & Racin (H + X2 - MX2] O But Thermal decomposition of (NH2), Befa is the best route for the preparation of Befa,
and Bezel Beclz is conviniently made from the oxide. [BeO + C+C12 BeClz + CO]
-> Ran Ly H2: @ All encept Be combine to form by dride HH2. @ BeH2 can be prepared by rxn of BeCl2 with LiAlH2
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→ <u>Reducing Nature</u> : © Weaker neducing agent than corresponding alkali metals. © Doci has rather less negative reduction potential than other alkaline earth metals but its reducing nature is due to large hydration energy associated with small size of Be <sup>2+</sup> ion and relatively large value of the atomization enthalpy of the metal.
@ Bop has after the all a strated than other alkaline earth metals but its reducing normal is alle
to loss here the some to be adomition political price of Bo2+ ion and relatively large value of the adomitication
entholog - C U II
in the line forming amounted ions.
→ Soln in liquid NH3 @ dissolues in liq NH3 to give deep blue black solution forming ammoniated ions. [M+(x+y)NH3 → [M(NH3)x] <sup>2+</sup> + 2[e(NH3)y] [monthese soln, the ammoniates [M(NH3)6] <sup>2+</sup> can be nemoved.
(M+(x+y)NH) -> [H(NH2), ]2+ + 2[o(NH1),]- @ From these sol", the ammoniates [H(NH3)6] car
TITLICEC DD Q Q
III USES DO @ Be is used in manufacture of alloys. @ Copper-beryllium allogs are used in the prep of high strangth springs. @ Hatellia Qa is used in manufacture of alloys. @ May forms alloys with AI, Zn, Mn and Sn
enclaute be is used for making windows hing it is a the Co. to Co. to Co. the
<ul> <li>Hg-Al alloys being light are used in cir-oraft construction. Mg (roisday) reason used as Antacid.</li> <li>bubs, incendiary bombs and single signals. Surpention of Mg (014 in H20 (Hilk of Magnovia) used as Antacid.</li> <li>Mg (03 is used in toothpaste. Of a is used in extraction of metals from oxides ashich are difficult to reduce with C.</li> <li>Mg (03 is used in toothpaste. Of a is used in extraction of metals from oxides ashich are difficult to reduce of Canar)</li> <li>Of a and Ba metal used to remove oir from vaccum tubes. Of a salts are used in radiotherapy (Treatment of Canar)</li> </ul>
bulbs, incendiary bombs and single rignals. @ Surpention of My (04) in H20 (Hilk of magnitude to reduce with C.
My CO3 is used in toothparte. @ Ca is used in extraction of metals from oxides which we applied in the cancer]
@ Ca and Ba metal used to remove air from vaccum tukes. @ Ra salts are used in radiotherated
Specially covalent in nature
<ul> <li><sup>©</sup> Mg CO<sub>3</sub> is used in toothparte. <sup>©</sup> Ca is used in extraction of metals from oxides ceruch the topper (Treatment of Cancer)</li> <li><sup>©</sup> Ca and Ba metal used to remove air from vaccum tubes. <sup>©</sup> Ra salts are used in radiotherapy (Treatment of Cancer)</li> <li><sup>©</sup> Ca and Ba metal used to remove air from vaccum tubes. <sup>©</sup> Ra salts are used in radiotherapy (Treatment of Cancer)</li> <li><sup>©</sup> Ca and Ba metal used to remove air from vaccum tubes. <sup>©</sup> Ra salts are used in radiotherapy (Treatment of Cancer)</li> <li><sup>©</sup> Specially covalent in nature</li> <li><sup>©</sup> Specially covalent in nature</li> <li><sup>©</sup> Oxides &amp; Hudgenides: <sup>©</sup> Russ in O</li> <li><sup>©</sup> Except BeO, they have rock salt structure.</li> </ul>
UD COMPOUNDS OF ALKALINE EARTH METALUU → Oxides 8 Hydroxides: @ Runn in 02 to form MO (monoxide). @ Except BeO, they have rock salt structure. UD COMPOUNDS OF ALKALINE EARTH METALUU
Someruty, thermal stability and basic character increase with increase in the no film of the region of the
less basic and less stable than Corp. 1 @ Be(OH), being amphoteric [Be(OH), + 20H <sup>-</sup> → [Be(OH),] <sup>2</sup> → (Bery late ion)]     [Bery late ion)]
$Be(0H)_{L} + 2HCI + 2H_{20} \longrightarrow [Be(0H)_{4}]^{Cl_{2}}$
Heliday a Crust D X is when I are a set in the and the solvents.
-> Halides: Except Bex, all holides of corp. 2 are ionic in nature. @ Bex, is covalent and soluble in organic solvents.
BeCl2 has a chain structure in solid state [>Be Cl > Be Cl >
Control dissosicites at order of 1200K. Tendency toform halide hydrates gradually decreases down [MyCl: 8420, Calls: 6420, Shell 64200, Sh
"Instead of an dehydration hydrated halldes of Be and Mg on heating suffer hydrolysis;
and relatively less soluble than chlorides owning high lattice energies.
Carbonates @ Insoluble in water, can be precipitated by addition of a social of an animitation decompose on heating to give CQ and origine The solubility of carbonates decreases with increase in at no. • All the carbonates decompose on heating to give CQ and origine The solubility of carbonates decreases with increase in at no. • All the carbonates decompose on heating to give CQ and origine The solubility of carbonates decreases with increase in at no. • All the carbonates decompose on heating to give CQ and origine to give contain the increase of
Be Co; is unstable and kept in CO, atmosphere. Thermal stability increases with increasing cotionic size. Thus couble
Sulphoto: @ All while colids and while to have a Bas SO: and Ma SOs are readily soluble in water.
<ul> <li>Be Co; is unstable and kept in CO, atmosphere. Thermal Stability increases with increasing cationic size.</li> <li>Sulphate: All cohite solids and stable to heat. Be SO4, and Mg SO4 are radily soluble in water.</li> <li>Sulphate: All cohite solids and stable to heat. Be SO4, and Mg SO4 are radily soluble in water.</li> <li>Sulphate: All cohite solids and stable to heat. Be SO4, and Mg SO4 are radily soluble in water.</li> </ul>
-> <u>Nitrates</u> : @ Made by dissolution of the of the carbonates in dil nitric acids. @ hg (No3), crystallises with 6 H, @ molecules, Ba(No)
orgstallises as adhydrous : salt. @ All decompose on heating [24(NO3) -> 2NO. + 4NO, + 02]
1 L. marking
@ Encostionally small atom and ionic size, Rue to high ionisation entrangly and small size it forms largery control of
Due to choose of clochital does not exibit coordination no more than town. other members can have coordination no of 0.
All like there manhed and hydroxide of be are amphoteric.
<ul> <li>Diagonal Relationship blu Bo and A!: Be<sup>2+</sup> and Al<sup>3+</sup> have similar charge / radius vatio</li> <li>All and Be are not readily attacked by acids because of the presence of an oxide film on the surface of the metal</li> <li>All and Be are not readily attacked by acids because of the presence of an oxide film on the surface of the metal</li> </ul>
@ All and Be are not readily attacked by acids because of the presence of an oxide film on the surgice of the metal
( 130 (OH), dissolved in excess of alkali to aive Gergiane ion Lise (014)
Chlorides of Be and Al have CI bridged chloride structure in vapour state. Both chloridus we non polar, strong cears
base liked as Friedel Brott cotolists
OF COMPOUNDS OF Calcium 10 CaO + CO.]
(II COMPOUNDS OF Calcium II) $\rightarrow \underline{\text{Suick lime}(Ca0)}: @ Proped by heating lime stone (Ca(03) in a notary kiln at 1070 - 1270 K [Ca(03 - Ca0 + C02]) \rightarrow \underline{\text{Suick lime}(Ca0)}: @ Proped by heating lime stone (Ca(03) in a notary kiln at 1070 - 1270 K [Ca(03 - Ca0 + C02])) \rightarrow \underline{\text{Suick lime}(Ca0)}: @ Proped by heating lime stone (Ca(03) in a notary kiln at 1070 - 1270 K [Ca(03 - Ca0 + C02]))) \rightarrow \underline{\text{Suick lime}(Ca0)}: @ Proped by heating lime stone (Ca(03) in a notary kiln at 1070 - 1270 K [Ca(03 - Ca0 + C02]))))))))))))))))))))))))))))))))))))$
-> Quick line (Cao): @ Proped by heating time stone (Caus) in a carbon disxide. @ Cao is a cshite amorphous solid. mp = 2870K. On exposure to air, it absorbs moisture and carbon disxide. @ Cao is a cshite amorphous solid. mp = 2870K. On exposure to air, it absorbs moisture and carbon disxide.
$[f_{0} \circ f_{0}] = [f_{0} \circ f$
Suick lime slaked with sodo is known as sodalime @ Being basic oxide, it combines with acidic oxides at high Temp.
$\left[C_{a0} + Sio_{2} \longrightarrow C_{a}Sio_{3} / 6C_{a0} + P_{4}O_{10} \longrightarrow 2C_{a3}(Po_{4})_{2}\right]$
-> Uses: @ For manufacturing coment and is the cheapent form alkali @ Hanufacturing of sodium car bonate (Nagoz) and (Na 04),
@ In purification of sugar and in manufacture of dye stuffs
- Sloked lime Co (OH) . on Proparied by adding Ca0 to H20 withit amagehour analy
→ Slaked lime, Ca (OH); @ Preparied by adding Ca0 to H2O, ishite amorphous powder, sparingly soluble in water.
The aqueous solution is known as lime asater and a suspension of Ca(OH)2 in H2 O is known as milk of lime.
<ul> <li>On passing CO2, it turns milley due to formation of CaCO3 [CaCOH)2 in H2 O is known as mille of line.</li> <li>On passing excess CO2, the ppt dissolues to form Ca(HO3), [CaCOH)2 + CO2 -&gt; CaCO3 + H2 O]</li> <li>Bleaching Power.</li> </ul>
- 11222 a preparation of mentor at ill and if control of bleaching powder Balot, + 201, -> Call, + Calocu, +240
-> Uses: @ preparation of mortar, a building material. @ used in white ask due to its disinfectant nature. @ Used in glass making, in tanning industry, prep of bleaching poorder and in its disinfectant nature.
O Used in glass making, in tanning industry, prop of bleaching pocoder and purification of sugar
Let + WE + ME + ME + ME + ME
a second s

-> Calcium Carbonate. (CacO3): @ CaCO3 occurs in nature in several forms like lime stone, chalk, markle etc. @ Can be prepared by passing CO2 through Ca(OH)2 or by addition of Na2CO3 to CaCI2

 $\left[Ca(OH), + CO_2 \longrightarrow Ca(O_3 + H_2O / CaCl_2 + Na_2CO_3 \longrightarrow CaCO_3 + 2NaCl\right]$ 

@ CaCO3 is a cohite fluffy powder, almost insoluble in water. When heated at 1200 k, it decomposes. [COO3 -1700K Ca0 +CO2] @ Reacts with dilute acid to liberate CO2 [CaCO3 + 2HCI -- > CaCI1 + H20 + CO2 / CaCO3 + H2SO4 + H20 + CO2 -> Uses: @ Used as building material inform of markle and manufacture of quick lime.

@ Cacoz and Mg coz is used as a flux in the extraction of metals such as iron

@ specially ppt. CaO3 is used in making high quality paper.

@ used as antacid, mild abrasive in tooth paste, a constituent of cheasing yum and filler in cosmetics.

→ Calcium Sulphate ( Plaster of Paris) (CasO4·2H2O): @ Hemi-hydrate of EasO4, obtained by heating gyprum, CaSO4·2H2O at 393K  $\left[2(CaSO_4 \cdot 2H_20) \longrightarrow 2(CaSO_4) \cdot H_20 + 3H_20\right]$ 

- @ Above 393K, no water of erystallisation is left and anhydrous calcium sulphate, CasD4 is formed (dead burnt plaster) @ Has remarkable property of settling with water. On, mixing with adequate quantity of water it forms plastic mass that gets into a hard solid in 5 to 15 minutes.
- -> Uses: " In building industry as plaster. " Used to immobilise the affected part of organ where there is a bonde fracture of sprain, also employed in dentistry, in ornamental works and making casts of statues and burts
  - @ Cement: Comment is a product obtained by combining a meterial rich in lime, cao with other materials such as day which contains silica, Sioz along with the oxides of aluminium. The raw materials for the manufacture o are limestone and day. which are mixed together to form
    - "cement clinker" which is mixed with 2-3% by weight of gypsum (Cas04.24,20) to form cement.
    - Ingredients of Portland cement ( dical cium silicate (CarsiO4) 26% ( trical cium silicate (CarsiO5) 51%. and Tricalcium aluminate (Caz Alz 06) 11%
    - @ Settling of Cement: When mixed with water, it settles to a hard mars. This is due to the hydraction of the molecules of the constituents and rearrangements.
    - The purpose of adding gypsum is only to slow down the process of settling of the comment so that it gets sufficient hardening.

#### Biological Importance of Mg and Ca:

@ An adult body contains 25g of Mg and 1200g of Ca. @ All enzymes that utilise ATP in phosphate transfer require Mg as a cofactor. @ Main pigment for the absorption of light in plants is chlorophyll which contains Mg. @ About 99% of body Ca is present in bones and teeth, it also plays important roles in neuromascular function, interneuronal transmission, 'cell membrane integrity and blood coagulation. @ Ca concentration in blood plasma is regulated at about 100 mgL-', It is maintained by by two hormones: calcitonin and parathyroid.

Group 15/ (N, P, As, Sb, Gi) [NS <sup>2</sup> np <sup>3</sup> ]
Group 15 (N, P, As, Sb, Bi) [ns <sup>2</sup> np <sup>3</sup> ] elements: → down the group, there is a shift from non-metallic to metallic through metaloidic character. N, P are non metals, As, Sb is metalloids and Bi is a typical metal.
Occurance: → N <sub>2</sub> is 78% of the atmosphere by volume. it occurs in earth's crust as NaNO3, KNO3. It is also found in proteins. → P appears in members of the apatite family [Cag(PO4)6·CaX2(X=F,ClorOH)] it is an essential constituent of animal and plant matter (bones, cells). → As, Sb and Bi are found mainly as sulphide minerals.
$\begin{array}{llllllllllllllllllllllllllllllllllll$
ionisation: $\rightarrow$ decrases down the group. due to half filled p orbital, IE is enthalpy: $\rightarrow$ considerably high than group 14. $(\Delta H_1 < \Delta H_2 < \Delta H_3)$
electronegativity: -> decreases down the group, not profound in heavier element
$\begin{array}{rcl} physical & : & \rightarrow N_2 \text{ is solid gas, others are diatomic solids. metallic character increases} \\ properties & down the group. & \rightarrow b,p increases down the group. \\ & \rightarrow mp increases up to As then decreases up to Bi. \\ & \rightarrow all other than N show allotropy. \end{array}$
CHEMICAL
properties ox states and -> common oxidation states are -3, +3, +5. de tendency to exibit
trends in chemical reactivity of +5 ox state decreases down the group. Bi almost doesn't. A stability of +5 ox state decreases down the group only Bitucompound is BiFs, stability of +3 ox state increases down the group (due to inert pair effect)
$ \rightarrow N \text{ also exhibits } +1, +2, +4 \text{ ox. states reacting with 'O_2} $ $ \rightarrow P \text{ shows } +1 \text{ and } +4 \text{ in some } 0 \times 0 \text{ acids} $ $ \rightarrow \text{ all ox states of } N \text{ from } +1 \text{ to } +4 \text{ disproportionate in acid solution.} $
3HNO2 → HNO3 + H2O + 2NO → similarly in P, nearly all ox states disproportionate in alkali and acid. → N is stricted to a maximum covalency of 4 since only 4(1s + 3P) orbitals are available, others with d orbital can expand their octate.
$\begin{array}{llllllllllllllllllllllllllllllllllll$
→ P-P, As-As and Sb-Sb form single bond, Bi forms metallic bond, Single N-N is weaker than P-P because of high inter electronic repulsion of the non-bonding electrons. As a result, catenation tendency is weaker in N.
→ due to absense of d orbital in valence shell, it cannot form dπ-pπ por bond as the others. P and As can form dπ-dπ bond also with transition metals when their compounds like P(C2H5)3 and As(C6H5)3 act as ligands.
necessi and a fundas.

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reactivity $\rightarrow$ all form hydride of form $EH_3$ , stability of the hydrides decreases from
towards H2 NH3 to BiH3. the reducing nature of the hydrides increases down the group,
$\rightarrow$ basicity order; NH <sub>3</sub> > PH <sub>3</sub> > As H <sub>3</sub> > SbH <sub>3</sub> ≥ BIH <sub>3</sub>
reactivity. $\rightarrow$ all form two types of onides : $E_2O_3$ and $E_2O_5$ . Oxide with higher towards $O_2$ ox. state is more acidic. acidic character decreases down the group. $\rightarrow E_2O_3$ of N.P are purely acidic, As, 56 are amphoteric, Bi is basic.
reactivity $\rightarrow$ these form EX <sub>3</sub> and EX <sub>5</sub> . N doesn't form NX <sub>5</sub> due to absence of d.orb. Howards $\Rightarrow$ EX <sub>5</sub> are $\Xi \times_3$ more covalent than $\Xi \times_3$ . except NX <sub>3</sub> all tribalides halogens are stable (NF <sub>3</sub> is stable though )( $\Xi \times_3$ except BiF <sub>3</sub> are covalent).
reactivity → all form binary compounds exhibiting -3 ox state & e.g. Ca <sub>3</sub> N <sub>2</sub> , Ca <sub>3</sub> P <sub>2</sub> towards: metals
metals <b>DINITROGEN</b>
→ produced commercially by liquification and fractional distillation of air preparation: → treating an aqueous sol <sup>n</sup> of NH4CI with sodium nitrile (NaNO2)
$NH_4Cl(\alpha\alpha) + N\alpha NO_2(\alpha\alpha) \longrightarrow N_2(g) + 2H_2O(l) + N\alpha Cl(\alpha q)$
$\rightarrow$ by thermal decomposition of $(NH_4)_2 Cr_2 O_7$ [ammonium dichromate]
$(NH_4)_2 Cr_2 O_7 \longrightarrow N_2 + 4 H_2 O + Cr_2 O_3$ $\rightarrow$ very pure N_2 by thermal decomposition of sodium or barium azide.
$Ba(N_3)_2 \longrightarrow Ba + 3N_2$
[properties: -> has two stakle isotopes: "N and "N. very less soluble in water.
low freezing and boiling points. rather inert at room temp.
→ reactivity increases rapidly with temperature, at high Temp, it directly combines with some metals to form predominantly
ionic nitrides and covalent ones aeith non-metals.
$6Li + N_2 \longrightarrow 2Li_3N / 3Mg + N_2 \longrightarrow Mg_3N_2$
$\rightarrow$ it combines with H <sub>2</sub> at 500 °C in presence of catalyst (Haber's process) $N_2(g) + 3H_2(g) = \frac{500°C}{cat} 2NH_3(g)$
$\rightarrow N_2$ combines with $O_2$ at about 2000 K to form NO.
$N_2$ + $O_2(g) \stackrel{A}{\longrightarrow} 2NOCR$ dents
Annonia.
preparation: -> in nature, due to decay of nitrogenous compounds e.g. wrea.
preparation: $NH_2 CO NH_2 + 2H_2 O \longrightarrow (NH_4)_2 CO_3 \rightleftharpoons 2NH_3 + H_2 O + CO_2$
→ in small scale, ammonium salts decompose when treated w/ NaOH, Ca(OH),
$\sqrt{2NH_4Cl + Ca(OH)_2} \xrightarrow{2NH_3} + 2H_2O + CaCl_2$
$(NH_4)_2SO_4 + 2NAOH \longrightarrow 2NH_3 + 2H_2O + No. SO.$
H 106.8° = H -> On large scale, it is manufactured by Habers Process
$\frac{1}{N_2(g) + 3H_2(g)} \xrightarrow{200 atm}_{400^{\circ}c} 2NH_3(g)  [catalyst \rightarrow K_20, Al_2O_3]$
properties! → colourless, pungent odour. possesses hydrogen bonds causing high b.p. → highly soluble in water, #in ag sol <sup>h</sup> is weakly basic due to form OH -
$NH_3 \oplus H_2 O(l) \longrightarrow NH_4^+ (\alpha \alpha) + OH^- (\alpha \alpha)$
$\rightarrow$ It precipitates the hydroxides of many metals from their salt. e.g.
$\frac{Zn SO_4(aq) + 2NH_4OH(aq) \longrightarrow Zn(OH)_2(s) + (NH_4)_2SO_4(aq)}{\text{white ppt}}$
Fecis (ma) + NH4 OH (mar)> FezO3 · X H2O (SH NH4 CI (mag)
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 $\rightarrow$  Zn reacts with dilute nitric acid to give NO2 and with conc acid to give NO2  $4Z_n + 10HNO_3$  (dil)  $\longrightarrow 4Z_n (NO_3)_2 + 5H_2O + N_2O$  $Z_n + 4 HNO_3 (conc) \longrightarrow Z_n (NO_3)_2 + 2 H_2 O_2 + 2 NO_2$ -> some metals (e.g. Cr, Al) do not dissolve in conc nitric acid because of the formation of f a passive film of oxide on the surface. → conc HNO3 also oxidises non-metals and other compounds.  $I_2 + 10 H NO_3 \longrightarrow 2HIO_3 + 10 NO_2 + 4H_2O_3$  $C + 4 HNO_3 \longrightarrow CO_2 + 2H_{20} + 4 NO_2$  $S_8 + 48 HNO_3 \longrightarrow 8 H_2 SO_4 + 48 NO_2 + 16 H_2 O_4$  $P_4 + 20 H NO_3 \longrightarrow 4 H_3 PO_4 + 20 NO_2 + 4 H_2 O_4$  $\rightarrow$  the familiar Brown Ring Test for nitrates depends on the ability of Fe<sup>2+</sup> to reduce nitrates to nitric oxide, which reacts with Fe<sup>2+</sup> 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. NO3 + 3Fe2+ + 4H+ ---- NO + 3Fe3+ + 2H20  $\left[Fe(H_2O)_6\right]^{2+} + NO \longrightarrow \left[Fe(H_2O)_5(NO)\right]^{2+} + H_2O$ brown · Phosphorus Allotropic forms : white phosphorus: -> translucent white, waxy, poisonous, insoluble in water but soluble in CS2 and glows in dark. -> It dissolves in boiling NaOH solution in an inert atmosphere giving PH3.  $P_4 + 3N_{a}OH + 3H_{2}O \longrightarrow PH_3 + 3N_{a}H_2PO_2$ -> it is less stable and therefore, more reactive than others because of angular strain in the Py molecule (angles 60°). It readily chatches fire in air to give dense white fumes of PaDio  $P_4 + 50$ ,  $\longrightarrow P_6 O_{10}$ red phosphorus: → obtained by heating white phosphorus at 573K in inert atmosphere for several days, when it is headed under high pressure; a veries of phases of black P is formed. -> Red P posseses iron grey lusture. Odowrless, non-poisonous and insoluble in water as well as in carbon CS2. Red Pis less reactive than white P. doesn't glow in the dark.  $- P \left[ p - P \right] - P \left[ p - P$  $\rightarrow$  It is polymaru'c.

black phosphorus:  $\rightarrow$  it has two forms e.g.  $\alpha$ -black and  $\beta$ -black.  $\alpha$ -black phosphorus is formed when red php is heated in a sealed tube at 803k. it can be sublimed in air and has opaque mono clinic or rhombothedral crystals. does not oxidise in air,  $\rightarrow \beta$ -black P is prepared by heating white P at 473k under high P. It does not burn in air upto 673K. Phosphine preparation:  $\rightarrow$  by the reaction of Calcium phosphile (Ca<sub>3</sub>P<sub>2</sub>) with H<sub>2</sub>O or dilute HCI  $C_{a_3}P_3 + 6H_2O \longrightarrow 3C_a(OH)_2 + 2PH_3$  $Ca_3P_2 + GHCI \longrightarrow 3Ca Cl_2 + 2PH_3$ → lab prep: by heating white p with conc NaOH in inertatmosphere of CO2.  $P_4 + 3N_aOH + 3H_2O \longrightarrow PH_3 + 3N_aH_2PO_2$ - when pure, it is non inflammable but becomes inflammable owing to the presence of P2H4 of P4 vapours, to purify it from impurities, it is absorbed in HI to form phosphonium iodide (PH4I) which on treating with KOH. gives PH3.  $PH_{4}I + KOH \longrightarrow KI + H_{2}O + PH_{3}$ properties: -> colourless gas, rotten fish smell and is highly pisonous, it explodes in contact with traces of oxidising agent like HNO3, CI2 and Brz vapours.  $\rightarrow$  slightly solutile in water. solution of  $PH_3$  in water decomposes in presence of light giving red P and Hz.  $\rightarrow$  when absorbed in CuSO4 or HgCl<sub>2</sub> solution, the corresponding phosphides formed  $3 \operatorname{Cu} \operatorname{SO}_4 + 2 \operatorname{PH}_3 \longrightarrow \operatorname{Cu}_3 \operatorname{P}_2 + 3 \operatorname{H}_2 \operatorname{SO}_4$  $3Hg_{3}Cl_{2} + 2PH_{3} \longrightarrow Hg_{3}P_{2} + 6HCl$ -> phosphine is weakly basic and like ammonia, gives phosphonium compounds

Phosphorus Halides

auth acids e.g. "Brin PH3 + HBr ----> PH4Br ents"

 $P_4 + 8SOCI_2 \longrightarrow 4PCI_3 + 4SO_2 + 2S_2CI_2$ 

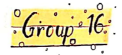
Properties:  $\rightarrow$  colourless oily liquid and hydrolyses in the pnesence of moisture.  $PCI_3 + 3H_2O \longrightarrow H_3PO_3 + 3HCI$ 

-> reacts with organic compounds containing -OH group.

 $3CH_3(00H + PCl_3 \longrightarrow 3CH_3COCl + H_3PO_3$ 

 $3C_2H_5OH + PCI_3 \longrightarrow 3C_2H_5CI + H_3PO_3$ 

properties: 
$$\rightarrow$$
 yellowith white powder, in moith air, it by dongues to POCI3 and Analy gets  
convected to phashoric acid.  
PCB + H\_DO  $\rightarrow$  POCI3 + 2HCI / POCI3 + 3H20  $\rightarrow$  H3, POA + 3HCI  
 $\rightarrow$  when heated, it sublimes but decomposes on stronger heating. PCI3  $\rightarrow$  PCI3 + CCC  
 $\rightarrow$  reacts with organic compounds containing.  $-OH$  group convecting to CI-denies.  
Chapter + PCI3  $\rightarrow$  CLUCCI + PCCI3 + HCI  
 $\rightarrow$  finally divided medals on hawding with PCIG glue corresponding delorides.  
2H39 + PCI3  $\rightarrow$  2H201 + PCI3  $/$  Sin Clus + 2PCI3  
 $\rightarrow$  it has trigonal biggranidal shauther and in solid state, it acids as an ionic  
solid [PCI3] + PCI6] in white the cotion, [PCI3] + bothedend, [PCI3] + octa.  
 $PCI3 + PCI3 = 2H201 + PCI3 / Sin + 2PCI3 = Sin Clus + 2PCI3
 $\rightarrow$  it has trigonal biggranidal shauther and in solid state, it acids as an ionic  
solid [PCI3] + PCC6] in white the cotion, [PCI3] + bothedend, [PCI3] + octa.  
 $PCI3 + PCI3 = 2H201 + PCI3 / Sin + 2PCI3 = H10 + H100 +$$ 



electronic configuration: ns2np4 elements: O, S, Se, Te, Po atomic and ionic radii: increases down the group. Oxygen is exceptionally small. Ionisation enthalpy; increases down the group. IE of grp 16 is lower compared to grp 15. electron gain enthalpy: because of compact nature, O atom has less negative electron gain enthalpy than S. from S onwards the value again becomes less negative up to Po. electronegativity: next to F, O has the highest EN value amongst the elements. EN decreases down the group. metallic character increases from 0 to Po. physical properties: -> O,S are non metals, Se, Te are metalloids, Po is a metal. → Po is radioactive and short lived. → mp and bp increase with atomic number. large difference b/w the mp and bp of → all these elements exhibits allotnopy O and S may be explained on the basis of atomicity [02 diatomic, S& polyctomic\_ CHEMICAL oridation state d: -> stability of -2 ox. state decreases down the group. Po doewit. O only shows -2. PROPERTIES exception : OF2 (+2) chemical reactivity → others show +2, +4, +6 ox states. +4 and +6 are more common. S, Se, Te generally show +4 with oxygen and +6 with F. stability of +6 oxidation state decreases down the group and that of +4 increases (inert pair effect). anomalous -> due to small size, high electronegativity. due to this H-bond exists in H2O but not H2S. behaviour: - absence of dorbital limits its covalency to 4 and in practice, it rarely exceeds 2. of oxygen reactivity of -> all form hydrides of the type H2E (E = O, S, Se, Te, Po), acidic character increases downer hydrogen: -> the increase in acidic character down the group can be explained in terms of decrease in bond enthalpy for dissosiation of H-E bond down the group. -> the thermal stability of hydries also decreases from H2O to H2Po due to this. -> all hydrides except water possesses reducing property and this character increases from H2S to H2Te. reactivity all -> all form oxides of EO2 and EO3 type. O3 and SO2 are gases while SeO2 is solid. -> reducing property of dioxide decreases from SO2 to TeO2, SO2 is reducing while B TeO2 is an oridising agent. > besides EOz, S, Se and Te form EO3. both are acidic in nature, reactivity wit > they form a large number of halides of type EX6, EX4 and EX2. halogens:  $\rightarrow$  stability of halides decreases in the order  $F^- > CI^- > B_r^- > I^-$ . among  $E X_6$ , ECI6 are only stable halides, all EF6 are gaseous in nature. SF6 is exceptionally stable for steric reasons. → among EX4, SF4 is a gas, SeF4 a liquid, TeF4 a solid.  $\rightarrow$  all except 0 forms dichlorides and dibromides. → the well known monohalides are dimeric in nature eg. 52F2, 52G2, 52Br2, Se2C12, these undergo disproportionation as given below. 2 Se2 Cl2 ---- Se Cl4 + 3 Se

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<pre>Classical Dioxygen DD</pre>
preparation: $\rightarrow$ lab props: by heating 0-containing salts such as chlorates, nitrates and permanganates. $\frac{2\kappa c_{10_3}}{M_{N_0}} \rightarrow 2\kappa c_{1} + 30_2$
→ lab prep: by thermal decomposition of the oxides of metals low in the electrochemical series and higher oxides of some metals.
$2Ag_{2O} \longrightarrow 4Ag + O_2$ $2Pb_3O_4 \longrightarrow 6PbO + O_2$
$2 \text{ Hg} \cup \longrightarrow 2 \text{Hg} + 0$ , $2 \text{ Pb} 0$ , $2 \text{ Pb} 0 + 0$
$   ab pnep: H_2O_2 \text{ is readily decomposed into } H_2O \text{ and } O_2 \text{ with finely diwided metal } \\ and MnO_2 \text{ as catalyst, } 2H_2O_2  2H_2O + O_2 } $
→ on large scale it can prepared from water or air via electrolysis.
-> industrially, Oz is obtained by tractional distilation of liquified air after
remowing CO2 and water vapour.
properties: $\rightarrow 2Ca + O_2 \rightarrow 2CaO$ ; $4AI + 3O_2 \rightarrow 2AI_2O_3$ ; $P_4 + 5O_2 \rightarrow P_4O_{10}$ $C + O_2 \rightarrow CO_1$ ; $O_2 = O_2O_3$ ; $P_4 + 5O_2 \rightarrow P_4O_{10}$
$2 \neq 1$ $2 \neq $
$CH_{4} + 20_{2} \longrightarrow CO_{2} + 2H_{2}O ; 2SO_{2} + O_{2} \xrightarrow{V_{2}O_{5}} 2SO_{3}$ $4HC_{1} + O_{2} \xrightarrow{CuCl_{2}} 2Cl_{2} + 2H_{2}O$
<u>Simple Oxides</u> : -> generally, non-metal oxides are acidic but oxides of some metals in high oxidation state have some acidic character (e.g. Mn207, Cr03, V205)
→ generally, metal oxides are basic. → some show dual behaviour known as amphoteric oxides. e.g.
$A_{1_2O_3} + 6HCI + 9H_2O \longrightarrow 2[AI(H_2O)_6]^{3+} + 6CI^{-1}$
$Alo \rightarrow CN GU + 2U = 2NG + A(OH)/Cl$
→ some oxides neither acidic nor basic known as neutral oxides. eg. CO, NO, N20.
°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°
-> too reactive to remain in atmosphere at sea level. at about 20 km, 0 zone layer is formed.
Toroparation: _ notaing day on through a vilent electrical discharge, conversion of D, to
$\begin{array}{cccccccccccccccccccccccccccccccccccc$
$30_2 \rightarrow 20_3$
→ as the conversion in endothermic, it is manning to use a main and
. O. Shan have a latter of ozon's care can be used and pure of the can be
(0)(d)(h)(h)(h)(h)(h)(h)(h)(h)(h)(h)(h)(h)(h)
properties: -> pale blue gas, dark blue liquid and violet-gues on the source however above characteristic smell and in small concentration is harmless. however above
characteristic smell and in small concentration is normanices. Now and a nausea, 100 ppm, breathing becomes uncomfortable resulting in headache and nausea, 100 ppm, breathing becomes uncomfortable cort On since in decomposition $\Delta H$ is -ve and
-> 020ne is thermody namically unitable who at on of O2 is explosive.
As is positive. $DG < O$ thus high content oxygen $(O_3 \rightarrow O_2 + O)$ , it acts as a $\rightarrow due^{\frac{1}{2}}$ tendency to liberate atoms of nascent oxygen $(O_3 \rightarrow O_2 + O)$ , it acts as a
$ \rightarrow due^{to} tendency to liberate atoms of nascent oxygen (03 - 02 + 0) + 02  powerful oxidising agent. e.g. Pbs + 10 403 - pbs04 + 402  21 - + 120 + 03 - 201 - + 12 + 02  21 - + 120 + 03 - 201 - + 12 + 02  21 - + 120 + 03 - 201 - + 12 + 02  21 - + 120 + 03 201 - + 12 + 02$
21 + 420 + 03 - 21 + 420 + 03 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 - 21 + 10 + 10 + 10 + 10 + 10 + 10 + 10 +
$\rightarrow$ nitrogen 0×ides (particularly & nitric acid) combine rapidly with O3. NO + O3 $\longrightarrow NO_2 + O_2$

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$\rightarrow$ freens used in aerosol sprays and as refrigerants damages ozone layer.
· Sulphin?
$\rightarrow$ forms numerous all othopes of which yellow rhombic (x-sulphur) and monodinic ( $\beta$ -sulphur) forms are most important. rhombic sulphur is stable at room temperature,
Thombic subplust: $\rightarrow$ yellow in colour, formed on evaporation of solution of roll subplus in $CS_2$ , $(\alpha - subplus) \rightarrow in soluble in water but dissolves to some extend in (\Omega, R-0) + and R-0-R'.(30) = (S_3) \rightarrow readily soluble in CS_2.$
monoclinic sulphus $\rightarrow$ soluble in CS <sub>2</sub> , prepared by melting rhombic sulphus in dish and cooling, ( $\beta$ -sulpher) till crust is formed. two holes are made in the crust and the remaining liquid is powed out. On removing the crust colougless needle co-changed
(S <sub>6</sub> ) → at 369K, both forms are stable, it is known as transition temperature. → both forms have S <sub>8</sub> molecules with different structure (puckered and crown)
→ in synthenised «cyclo-S6, it becomes chair form. above 100DK, S2 is dominant.
Sulphur Dioride
preparation: $\rightarrow SO_2$ is formed with a little $(6-8\%)$ SO3 when S is burnt in air or oxygen. S + $O_2 \longrightarrow SO_2$
$\rightarrow$ lab prep: treating a sulphite with dilute sulphuric acid $SO_3^{2^-} + 2H^+ \longrightarrow H_2O + SO_2$
$\rightarrow$ industrially, produced as a p by product of roasting of sulphide ores. $4 \operatorname{FeS}_2 + 110_2 \longrightarrow 2\operatorname{Fe}_20_3 + 8 \operatorname{SO}_2$
properties: $\rightarrow$ colourless, pungent smelling, soluble in water. forms $H_2SO_3$ with water.
$SO_2 + H_2O \rightleftharpoons H_2SO_3$ $SO_2 + H_2O \rightleftharpoons H_2SO_3$ $SO_2 + H_2O \rightleftharpoons H_2SO_3$ $SO_2$ reads neadily anith NaOH solution forming sodium sulphite which reads with more $SO_2$ forming NaHSO_3. $2NaOH + SO_2 \longrightarrow Na_2SO_3 + H_2O$
$Na_2SO_3 + H_2O + SO_2 \longrightarrow Na_2SO_3 + H_2O$ $Na_2SO_3 + H_2O + SO_2 \longrightarrow 2NaHSO_3.$
: → reacts ceith Cl <sub>2</sub> in presence of charcoal (catalyst) to give sulphuryl chloride(so <sub>2</sub> cl <sub>2</sub> ). it is oxidiaest to so3 by O <sub>2</sub> , SO <sub>2</sub> + Cl <sub>2</sub> charcoal > sO <sub>2</sub> Cl <sub>2</sub>
→ moist SO2 behaves as a reducing agent. it converts fe(II) ions to fe(II) and decolourises acidified KMnO4 solution.
$2fe^{3+} + SO_2 + 2H_2O \longrightarrow 2fe^{2+} + SO_4^{2-} + 4H^+$
$5SO_2 + 2MnO_4^- + 2H_2O \longrightarrow 5SO_4^{2^-} + 4H^+ + 2Mn^{2^+}$ Oxoacids of Sulpher,
140/
HO JS OF STORY CACH. OF PER OXO disulphunic acid Pyrosulphunic acid (Oleum)
sulphurous acid sulphuric acid, of per 0×0 disulphuric acid (0 leum) (H2SO3) (H2SO4) (H2S208) (H2S207)

Sulphuric Acid
manufacture: → by the Contact Process which involues 3 steps: i) burning of sulphur or sulphide ores in air to generate SO <sub>2</sub> . i) burning of sulphur or sulphide ores in air to generate SO <sub>2</sub> . ii) conversion of SO <sub>2</sub> to SO <sub>3</sub> by the reaction with O <sub>2</sub> with V <sub>2</sub> O <sub>5</sub> as catalyst.
ii) absorption of so3 in H2SO4 to give Oleum (H2S2O7) iii) absorption of so3 in H2SO4 to give Oleum (H2S2O7)
is exothermic, renormize and forward (generally 2 bar and 720K is who therefore, low T and high P are favourable. (generally 2 bar and 720K is who therefore, low T and high P are favourable. (generally 2 bar and 720K is who therefore, low T and high P are favourable. (generally 2 bar and 720K is who therefore, low T and high P are favourable. (generally 2 bar and 720K is who dilution of aleum gives H2SO4 of the desired concentration (96-98% pure).
properties: -> colourless, dissolves in water with large evolution of authority (ii) strong affinity -> chemical reactions are due to (i) low volatility (ii) strong acid (iii) strong offinity
$F = \{x \in \mathcal{L}, N \in \mathcal{L}\}$
→ it is strong deby drating agent (wet gases can be dried up pnowided they don't read) → it is strong deby drating agent (wet gases can be dried up pnowided they don't read) it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H2SO4 > 12 C + 11 H20 it removes water from organic compounds. C12 H22 O11 - H22 O11 - H22 O11 -
intermidiate between phosphoric and nitric acid). Hz SOn is reduced to SO2.
$(L_1 + 2H_2SO_4 \xrightarrow{\longrightarrow} L_4SO_4 + SO_2 + 2H_2O_4)$
C + 2H2SO4 (conc) → CO2 + 2SO2 + 2H2O C + 2H2SO4 (conc) → CO2 + 2SO2 + 2H2O C + 2H2SO4 (conc) → CO2 + 2SO2 + 2H2O in making fertilizers, petroleum refining, manufacture of pigments, paints, detergent in dustry
Group 17 Elements
occurance: → F, CI are fairly abundant. At is radioactive. F is normally present as insoluble fluorides (CaFz, NazArF6 etc) and small quantity is present in soil, river water, bones and teeth of animals.
river water, bones and teetre of unimum. → sea water contains chlorides, bromides and iodicles of Na, K, Mg and Ca (mainly Na → sea water contains chlorides, bromides and iodicles of Na, K, Mg and Ca (mainly Na
electronic config $\rightarrow$ ns <sup>2</sup> np <sup>5</sup> atomic and ionic radii': $\rightarrow$ smallest atomic radii' in respective group. F is extremely small. $\rightarrow$ atomic and ionic radii' increases down the grown.
→ atomic and ionic radii increases down the grown.
ionisation on the py: -> very high ionisation enhancing.
electron gain: -> maximum electron gain enthalpy in respective group periods because they need
electronegativity: -> highlyly electronegative, decreases down the group. F is most electronegative.
physical: $\rightarrow$ all halogens are coloured. (F <sub>2</sub> yellow, Cl <sub>2</sub> greenish yellow, B <sub>5</sub> red, L <sub>2</sub> violet). properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly soluble in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly soluble in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly soluble in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly soluble in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly collabele in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly collabele in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly collabele in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly collabele in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly collabele in H <sub>2</sub> O. but so luble in properties $\rightarrow$ F and Cl react with H <sub>2</sub> O, Br and I are spaningly collabele in H <sub>2</sub> O. but so luble in H <sub>2</sub> O. but so luble in H <sub>2</sub> O. Br and H <sub>2</sub> O. but so luble in H <sub>2</sub> O. but so luble in H <sub>2</sub> O. but so luble in H <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and H <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble is here span and h <sub>2</sub> O. but so luble span and h <sub>2</sub> O. but so lub
→ enthalpy of dissosiation of P2 13 with real of old control of old control of the control of t
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### CHEMICAL PROPERTIES

ox. state and : - > all hologens exhibit -1 ox state. however CI, Br and I exhibit +1, +3, +5 and +7 also. higher oxidation state of CI, Br, I are realised mainly when thend in chemically reactivity the halogens combine with small and highly electronegative For O atoms. - all halogens are highly reactive. reactivity decreases down the group. -> due to high acceptance of one electron, are strong oxidising nature

F is the strongest oxidising halogen, oxidises other halide ions in solution or even is solid state. a halogen oxidises halide ions of higher ating  $F_2 + 2X^- \longrightarrow 2F^- + X_2 \quad (X = CI, Br, I)$ 

 $\frac{Cl_2 + 2X}{2Cl} + \frac{X}{2} \left(X = Br, 1\right)$  $Br_2 + 21^- \longrightarrow 2Br^- + I_2$ 

-> relative oxidising power of halogens can further be illustrated by their reaction with water. F oxidises water to Oz whereas CI, Br form hydrohalic and hydrohalous acids. I doesn't react sponteneously, I2 can be oxidized by O2

$2F_2 + 1$	120	→ 4H+	+ 4F +	• O2
×2 + 11	1 <sub>2</sub> 0	→ H× ·	+ HOX ( *	= (1, Br)
41 +	4H <sup>+</sup> + 02	>	212 + 2H	20

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anomalous: -> due to small size, high electronegativity, low F-F bond dissociation enthalpy and non availability of d'orbital. -> IE, EN and electrode potential are highly-er than expected.

- -> most reactions of F are exothermic (due to small and strong bond formed with other elements).
- -> It forms only one execut while others from a number of them.
- → IF is a liquid (due to H bonding), while other by H× are gases.

reactivity:  $\rightarrow$  all react to give It × but affinity for Hz decreases from F to I. H× dissolute to wards Hz in H2O to form hydnohalic acid.

- rightingens -> acidic strength: HF<HCI<HBr<HI. stability decreases down the group.
- only OF, is there mally stable, both are strong fluorinating agents, O2F2 owands 02
  - oxidises Pu to Pufe (used in removing Pu from nuclear fuel)
  - -> CI, Br, I form oxides of +1 to +7. decrease in stability in order I>CI>Br.
  - $\rightarrow$  higher oxides tend to be more stable.
  - -> CI oxides, CI20, CIO2, CI206 and CI207 are highly reactive ox. agent and tend to explode. CI20 is used as bleaching agent for paper pulps, textiles, water treatment
  - $\rightarrow$  Br oxides, Br<sub>2</sub>O, BrO<sub>2</sub>, BrO<sub>3</sub> are least stable stable halogen oxides.
  - $\rightarrow$  1 oxides,  $I_2O_4$ ,  $I_2O_5$ ,  $I_2O_7$  are insoluble solids and decompose on heating.  $I_2O_5$ is a very good on agent and is a used in estimation of CO.

reactivity: -> forms metal halidesey. Mg + Brz -> Mg Brz

-> ionic character MF>HCI>MBr>MI. It me metals with higher or state is more towards metals stable and more covalent.

reactivity ---- they combine to form compounds of types XX', XX', XX's and XX', where X towards is a larger size and X' is smaller size halogens. hatogens



preparation: --> heating MnO2 with concentrated HCI. MnO2 + 4HCI ---> MnC12 + C12 + 2H2O however Nacl and conc H2SOg is used in steal of HCl. 4Nacl + MnO2 + 4H2 SO4 ----- MnCl2 + 4NaHSO4 + 2H2O + Cl2 → by the action of HCI on KMn04. 2KMnOg +16HC1 ---- 2KC1 + 2MnCl2 + 8H20 + 5C12 -> Electrolytic process: Cl2 is obtained by electrolysis of brine (conc. Nacl solution), Cl2 is liberated at anode. properties: -> greenish yellow gas with pungent and suffocating adour. It is about 2-5 times heavier than air. It is 2-5 times heavier than air, can be easily liquified. -> reacts with metals and non-metals to form chlori des.  $2AI + 3CI_2 \longrightarrow 2AICI_3$  $P_4 + 6Cl_2 \longrightarrow 4PCl_3$  $2N_{q} + CI_{2} \longrightarrow 2N_{q}CI \qquad S_{8} + 4CI_{2} \longrightarrow 4S_{2}CI_{2}$  $2Fe + 3Cl_2 \longrightarrow 2FeCl_3$ → It has great affinity for hydrogen, reacts awith H- compounds forming HCI  $H_2 + CI_2 \longrightarrow 2HCI \qquad H_2S + CI_2 \longrightarrow 2HCI + S \qquad C_{10} + BCI_2 \longrightarrow 16HCI + 10C$ -> with excess ammonia, C/2 gives N2 and NH4CI but with excess C/2, NC/3 (explosive) is formed.  $8NH_3 + 3CI_2 \longrightarrow 6NH_4CI + N_2$   $NH_3 + 3CI_2 \longrightarrow NCI_3 + 3HC/$ (excess) → with cold and dilutes, Cl2 produces a mixture of chloride and hypochlorite but not with hot and concentrated alkalies it gives choride + chlorate. 2NaOH + Cl2 ---- NaCI + NaOCI + H2O  $\frac{(\operatorname{cold} + \operatorname{dil})}{6\operatorname{NaOH}} + 3\operatorname{Cl}_2 \longrightarrow 5\operatorname{NaCl} + \operatorname{NaClO}_3 + 3\operatorname{H}_20$ (cold + dil) (hot + conc) -> with dry slaked lime it gives bleaching powder [Ca(04)2. Ca(12. (a(0H)2.2H22]  $2C_{\alpha}(OH), + 2CI_{2} \longrightarrow C_{\alpha}(OCI)_{2} + C_{\alpha}CI_{2} + 2H_{2}O$ -> C12 reacts with hydrocarbons and gives substitution products with saturated hydrocarbons and addition products with unsaturated hydrocarbons. CH4 + C12 - W CH3 CI + HCI C2H4 + C12 room temp C2H4C12 -> Chlorine water on standing loses its yellow colour due to the formation of HCI and HOCI. Hypochlorous acid (HOCI) so formed, gives na seent oxygen which is responsible for oxidising and bleaching properties of C/2, it oxidises ferrous to ferric and subphite to subphate, subphur to so3, todine to iodate. In presence of H2O, they form H2SO, and HIO3.  $2Fe SO_{4} + H_{2}SO_{4} + Cl_{2} \longrightarrow Fe_{2}(SO_{4})_{3} + 2HCl$   $Na_{2}SO_{3} + Cl_{2} + H_{2}O \longrightarrow Na_{2}SO_{4} + 2HCl$   $SO_{2} + 2H_{2}O + Cl_{2} \longrightarrow H_{2}SO_{4} + 2HCl$   $I_{2} + 6H_{2}O + 5Cl_{2} \longrightarrow 2H1O_{3} + 10Hcl$ -> It is a poweerful bleaching agent; bleaching action is due to oridation. C12 + 1420 ----> 2HCI + 0 Coloured substance + 0 --- colourless substance.

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·Hydrog		CLL.	100
o Hydrog	en.	Cruo	nae.
.0.0	0	. 0	°0

o Hydrogen. Crybride.	NoHSO + HCL
preapation: lab: by heating Nacl with concentrated sulphuric a	aid. $\frac{N_{aCl} + H_2 SO_4}{N_{all} SO_4 + H_Cl} \rightarrow N_{all} SO_4 + H_Cl}{N_{all} SO_4 + H_Cl}$
properties: -> bleaches veg and organic matter in presence of m	
permanent. → colourless ang pungert smelling gas, easily liquified.c	and extremely soluble in water and ionises.
$HCI(9) + H_2O(1) - H_3O(10) + C$	
→ when 3 parts of conc HCI and one part of concentration formed which is used for dissolving noble metals e.g.	gold, platinum
	$\rightarrow 3PtCl_6^{2-} + 4NO + 8H_2O$
we will be sails of weater acid e.g. carbonates, hydri	ogencarbonates, sulphites etc.
$Na_2(O_3 + 2HCI \longrightarrow 2NaCI + H_2O + CO_2$	
$N_{a}HCO_{3} + HCI \longrightarrow N_{a}CI + H_{2}O + CO_{2}$	
$Na_2 SO_3 + 2HCI \longrightarrow 2NaCI + H_2O + SO_2$	
• Oxoacids of Halogens	
-> due to high EN and small size, F forms only one oxoacid. HOF	known as fluoric (1) acid or
hypofluorous acid. Others form several oxoacids which cann.	of he isolated in pureform.
they one stable is their salts and in aqueous solution.	
Halic (I) acid (hypohalous) HOF HOCI HOBr (hypoflownows) (hypochlorows) (hypobrome	(hypoiodous)
Halic (II) acid (Halous) HOCIO (chlorous)	-
Halic (v) a cid (Halic) Ho C(O2 (chioroid) (chioroid) (bnomic)	(iodic)
Halic (VI) acid (Perhalic) (perchloric) (perchloric) (perchloric) (perchloric)	O HOIOZ (periodic)
HOCI HOCEOExcellero	$\frac{1}{2} \frac{1}{2} \frac{1}$
INTERHALOGEN COMPONNDS	
preparation: -> by direct combination or by the action of halogen	on lower interhalogen compounds.
$\begin{array}{c} Cl_2 + f_2 \xrightarrow{4374} 2CIF \\ aqual volume \\ \end{array} \qquad \qquad$	$\begin{array}{c} \Box_{3} & \hline{\Box_{2} + 3F_{2}} & \underline{573K} \\ \hline{\Box_{2} + 3F_{2}} & \underline{575K} \\ \hline{\Box_{2} + 3F_{2}} & \underline$
$B_{12} + 3F_2 \longrightarrow 2B_1F_3 \qquad I_2 + C_{12} \longrightarrow 2.$	$1CI  Br_2 + 5F_2 \longrightarrow 2BrFs$
(dil) (equimoion)	rally more a struct 1 1 1 ( ( ( ))
[properties: → covalent mole cules and dia magnetic in nature.gene cos r x - x' bond is weaker than X-× (ce exception)	F-F).
→ all undergo hydrolysis giving halide ion derived ; hypohalite (when ××'), halite (××'3), halate (w	from the smaller halogen and a
hypohalite (when xx'), halite (xx'3), halate (w	hen $XX_5$ ) and perhalate (when $XX_7$ )
$X X' + H_2 O \longrightarrow H X' + H O X$	

# GROUP 18 ELEMENTS

occurance: $\rightarrow$ all except kn occur in the atmosphere. He and sometimes Ne are found in minerals
of radioactive origin e.g. pitchblende, monazite, deveite.
-> commercial source of helium is natural gas. Ra is obtained as a decay product of 226 Ra
$ \begin{array}{c} 226\\ 83\\ 83\\ 83\\ 86\\ \end{array} \xrightarrow{222}_{86} R_{\rm H} + \frac{4}{2} {\rm He} \end{array} $
electron: $\rightarrow n5^2 np^6$ except helium which is $15^2$ .
atomic radii: $\rightarrow$ increases down the group ionisation enthalpy: $\rightarrow$ exhibit very high ionisation enthalpy. decreases down the group.
electron gain $\rightarrow$ no tendency to accept electron therefore, have large positive values of EGA.
eruna quy:
physical: → all monoatomic, colourless, o dourless and tasteless. Its sparingly soluble in H2O, properties; very low m.p and b.p. He has the lowest b.p. (4.2K) of any known substance. → it has unusual property of diffuring through most commonly used lab materials such as rubber or plastics.
chemical there are several Xe compounds. compounds of Kr are fearer. only Krfz has
properties: been studied in detail. —> compounds of Rn has Bnot keen is plated just identified.(e.g. RnFs.).
for X 0 X 0 X 6 H wells converted
Xenon-B $\longrightarrow$ forms XeF <sub>2</sub> , XeF <sub>4</sub> and XeF <sub>6</sub> but combination under appropriate experimental compounds: conditions. Xe(g) + F <sub>2</sub> (g) <u>-673K</u> , 1000, XeF <sub>2</sub> (S)
$\frac{(e \times en)}{\times e(\theta) + 2F_2(\theta)} \xrightarrow{873 \times 7bar} \times eF_4(s)$
(A. C. artic)
(1: 5 valle) $X_{e}(9) + 3F_{2}(9) - 573,60-70ker XeF_{6}(5)$ (1: 20 ratio)
-> Xefo can also be prepared by interaction of Xefo and O2F2 at 143K
$x_e F_4 + O_2 F_2 \longrightarrow x_e F_6 + O_2$
→ Xe Fz, Xe Fq, and Xe F6 are colourless crystalline solids and sublime readily at 25°C they are powerful fluor inating agents. readily hycholysed even by traces of H2O.
2×efy (s) + 2H2O (l)> 2×e(l) + 4HF (aq) + 02 (g)
Xe fluorides react with P ion acceptors to form cationic species and flucoide ion donors to form fluoranions,
$X'eF_2 + PF_g \longrightarrow [XeF]^+ [PF_6]^-$
$\times eF_4 + SbF_5 \longrightarrow [XeF_3]^+ [SbF_6]^-$
Xefo+MF M+ [Xefer] (M = No, K, Rbor Cs)
Xenon-07 -> Hydrolysis of XeF4 and XeF6 with H2O gives XeO3.
compounds: $6 \times ef_4 + 12H_20 \longrightarrow 4 \times e + 2 \times eO_3 + 24H_f + 3O_2$
$XeFe + 3H_2O \longrightarrow XeO_3 + 6HF$
$\rightarrow$ partial hydrolysis of XeFe gives oxyfluorides, XeOF4 and XeO <sub>2</sub> F <sub>2</sub>
-> XeOs is a colourless explosive solid and has a pyramidal moleculer. structure.
→ Xe OF4 is a colourless volatile liquid and has a square pyramidal molecular structure.