S Block Elements

S-Block Elements: compartive study 04

IA 1 H IA Li Be Na Mg K Ca Rb SΥ CS Bα Ra Fr

IA -> Alkali Metal (They react with 150 to form Hydroxides which are strong alkalies)

IT A → Alkaline earth Metal (Because their oxides are alkaline in Nature and existed in the earty)

Electronic Configuration

and Ionic Radii Atomic

II A Elements gre smaller in size than Elements.

But their ionic Radii are smaller than atomic Radii B/UZ After Removal & Zeft? significantly.

Ionisation potential:

in Periodic table
hence used in Photocell size 1

strong Electropositive ... Co have Lowest IP

(ii) IIA Elements have nigher value of 1. P b/(02 of 100 stable E.C.

: A -> strong Electropositive Nature: - due to Low Value of IP, they loose easily.

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Na -> 152 252 20 35 Na LMg Mg - 152 252 2P6 352 Bud Ind I.b Na> Mg

Alkali Metals are extremly reactive and strong Reducing agents give Reason.

ANS: Alkali metals have very low value of I.F. they have tendency to loose & readily, that's why they are extremly Reactive and Kept in Kerosenoil. They have high value of SOP (standard axidation? potential) so strong Reducing Na 2.71 2.92 2.99 3.02 SOP 3.04

Density, melting point and Boiling point: ->

(V)

size of alkalimetals are highest in their period so they doesn't have close packing in crystal. all alkeli metals the have Bcc str. so they have (CN 8, PF 68-1)

Low of Density. which I on Moving Down the Alkali Metals have weak force of attaraction So they are soft and have LOW M.P & B.P.

While alkaline earth metal are heavy and hard, high value of M.P & B.P. due to small size and high charge density

Metallic property: -

M. C. 8 L M. C. 1

Flame Test :>

All the alkalimetals give characteristic colour in bunsen flame. The reasonis that when an alkalimetal is heated in a bunsen flame, toway the electrons get excited to higher energy level. when these e's returns to their ground level, energy is realesed in form of radiation visible region of spectrum.

golden violet reddish yellow (liliae) violet Li Na Comson Red

Among II group Members, Be and Mg do not impact any colour to the flame b/coz their valence e's are so tightly held by their Nuclei that they require much greater amount of energy for their excitation and hence energy released during return Journey lies beyond visible regian (U.V. Regian)

> Mg Ca SY Ba Be comisar Apple crimson Colourless Colourless Brick Red green Red

chemical proposeties: ->
The alkali metals exhibit high

chemical reactivity which is due to their low I.E.

and low heat of Atomisation. Reactivity 1 on going

down two gp.

LIL NALK CROLCS

(i) Action of Air (Reaction with Atmospheric oxygen)

All the alkali metals tarnish rapidly on exposure
to air due to formation of exides at the surface.

The exides may further react with moisture and
atmospheric Co2 to form a layer of Cambonates
at the surface. Therefore they are always
kerosene or faraftins oil to protect them

From the Action of Air.

form mixt of oxide and superaride, rest form super oxide.

4 Li + 02 -> 2 Li20

4 & BNR+02 \longrightarrow 2 N920 \nearrow 2 NA + 02 \longrightarrow 2 N9202 Peroxide

M + 02 \longrightarrow M D2 Superoxide

(M = K, Rb, US)

Reason: The formation and stability of these oxides Can be explained on the basis of their lattice energy Lit im being a small in has a strong field around it and can stabilise only small anion 0⁻² where Nation being a large Cation stabilise a large anion.

2 Mg + 02 -> 2 Mg0

Reaction with water: >

due to low value of I'E

Reactivity 1 2Na+ 2H20 -> 2Na0H+ H11

2Na+ GH50H -> GH50Na+ H11

Na+ HC=CH-> HC=CNa+ H11

2 Mg + 2 H20 Boil SMg(0H)2 + HL

Ca + 2420 Cold (OH)2 + HL

Solubility in liquid NH3

All the alkali metals

dissolve in liquid NH3 giving deep Blue solution.

The colour deepens with 1 in conc. of the sol.

The concentrated solutions are Bronze in colour and Posses metallic lusture. The solutions are good conductor of electricity and are paramagnetic conductor of electricity and are paramagnetic in Nature. The paramagnetic character 1 with 1 in concentration. The dissolution of two metal is

occompanied by imisation. M = M+ E

whe alkalimetals are dissolved in liq. NHz, the NHz solvated the Cation as well as electron.

M + NHz

(in excess)

The e released by metals caused polarisation of the charges of twe technis of the

it is the ammoniated \(\tilde{e}\) which is responsible for the Blue colour of solution. The electrical conductivity is due to ammoniated \(\tilde{e}\) as well as ammoniated cation. The dilute dilute solution are paramagnetic in Nature The I in Paramagnetic Character with \(\tilde{f}\) in Conc.

Suggests that in Concentrated sol. ammoniated \(\tilde{e}\) associated to form \(\tilde{e}\) pairs.

The value of X and y depends upon extent of solvation

The alkali metal solution in lig. NHz are stable and decompose only Slowly liberating Hydrogen the presence of toansition Metal (Fe, Pt, Zn)

M+liq. NH3 MNH2 + 1 H2 1

M(NH) --- M(NH) 2 + 4NH + 12 1

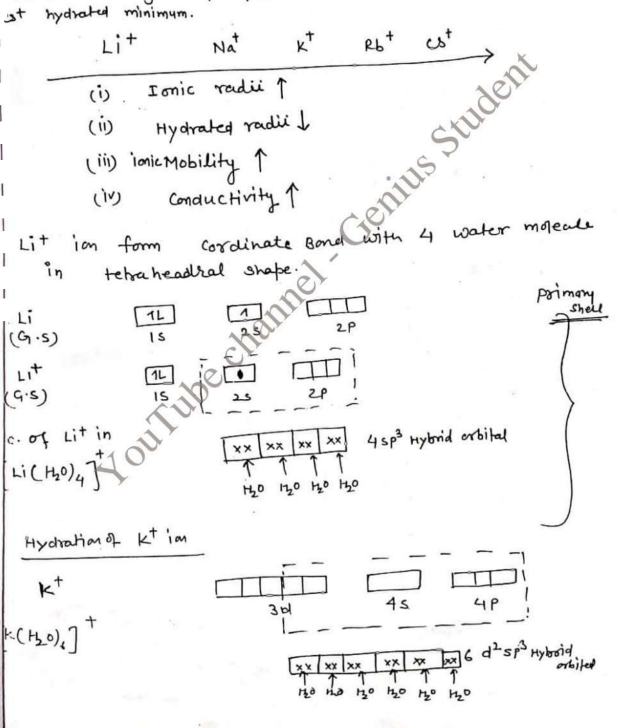
Due to presence of ammoniated & there solution behave as strong reducing Agent.

K2 [Ni(CN)4] + 2K = (NH3)4 K4[Ni(CN)4]

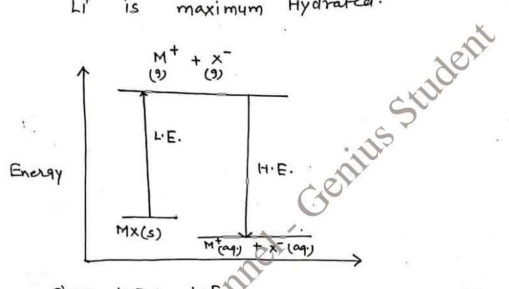
Fe ((0)s + 2Na = (Ny) Naz [Fe((0)4]

SOLVATION TENDENCY

simple salts of alkalimetals are solvable in water and give ions. The alkalimetals ions are extensively Hydrated. The smaller the size of the ion, the grater is the degree of hydration. Lit is hydrated max. while it hydrated minimum.



After this primary layer, a secondary layer of water morecule forms in which Morecule doesn't form Bond But they are Bonded with weak ion - dipose force of Altraction. and this Altraction force is inversely proportional to radius. I.e. smaller ion have strong force. that's why Lit is maximum Hydrated.



since H.E > L.E

Energy $\begin{array}{c}
M^{\dagger} + \chi^{\bullet} \\
(9) \\
(1)
\end{array}$ $\begin{array}{c}
M^{\dagger} + \chi^{\bullet} \\
(9) \\
(9)
\end{array}$ $\begin{array}{c}
M^{\dagger} + \chi^{\bullet} \\
(9) \\
(9)
\end{array}$ $\begin{array}{c}
M^{\dagger} + \chi^{\bullet} \\
(9) \\
(9)
\end{array}$

since L.E > H.E.

Salt is insolvable in water.

COMPLEXATION - TENDENCY

S-Block Elements have very less tendency to form complexes with ligands. Bloz-

(i) they do not have vacant orbitals.

(ii) large size

(111) small charge.

he Common Unidentate ligand with which the alkali metal ions form some what stable complexes are Hop, NHz,

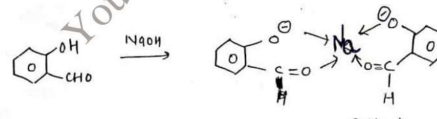
Py, Phy PO; Triphenyl phospine oxide

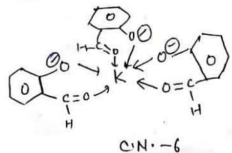
[M(H20)4] M = Lit, Nat, 100 [M'(H20)6] M' = Kt, Pbt, Cot

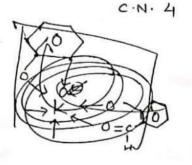
Some Examples of lithium Metal

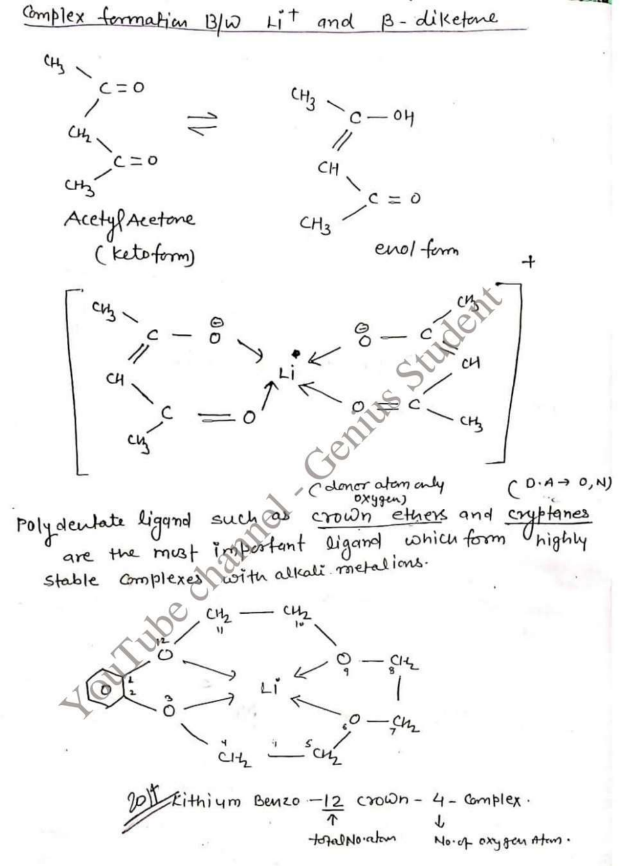
[Li(NH3)4] [Lic(150)(PY)2]

However alkali metals form reasonably reasonably stable complexes with some of the bidentate ligand and boly dentate ligand such as salicylablehyde, B-diketones, nitrophenols









hese ligands are also highly selective in complex formation. The cause of selectivity of lies in the size of the hole available in a Polydentate ligand. only the alkali metal ion which exactly fits into the holes form stable complexes. For example the hole in is Just of the right size for the (cryptate - 22,2) kt ion to fit in whearas Lit and Nat are too small while Pst and cot are too big. thus kt complex of cryptate -222 is Most stable. also don' doesn't have Alkaline earth Metals gre grater tendency to form complex But due to small size of Be it Can form complexes (Tetiahedral) Example Basic Berelium Acetate Be40 (CH3(W)6 Berilium oxalate [Be((204))] [Ca(EDTA)] Important complex of Ca is culorophyll Imperfant complex of Mg is

Functions of s-Block Elements in 1310 systems:

1. Transport of ions across the cell membrane

For the fuctioning of nerve and myscle cell, an electrical potential across the cell membrane is essential. This mainted by the different ratio of Nat and to kt inside and outside the cell. Nat are actively expelled from a cells, whereas kt are not. This ian transport is somtimes called a sodium pump, and it involves both the Active expulsion of Nat and the active take-up of kt. thus the Mechanism of ian transport involves polyether Complex formation.

- 2. Energy transfer: > Intu living beings energy is stored in the form of ATP which form Complex with Mg+2 in

chlorophyll is the pigment responsible for the green colour of the plants which is complex of Mg+2 ion.

4. Roll of Cath cath ion is present in bones as apatite (as Cas (PO4), OH, fluoro apalite in teeth enamales [3 Cas (PO4)2. Caf2].

It also important for Blood clotting and ambaction of Muscles.