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(5) Wavelength (
$$\lambda$$
) - linear distance blow two consecutive create or blow two  
consequive trough. It is generally expressed in terms of  
nu Angitrom( $\lambda$ ). [ $1A = 10^{10}$  m]  
(3) Frequency ( $y$ ) - No. of waves passing through a point in one second  
SI unit is hertz ( $H_2$ ).  
(3) Velocity (c) - linear distance travelled by more per second.  
(4) When the electromagnetic radiations are arranged in norder of their increating  
mavelengths or decreasing travents are arranged. In order of their increating  
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Electromagnetic spectrum.  
(S) Ric conflerer in units in site (207)  
wave number ( $\overline{y}$ ) =  $\int_{1}^{2} = \int_{1}^{200} + 2 \times 10^{4}$  m<sup>4</sup> with that st sheeted  
frequency ( $\overline{y}$ ) =  $\int_{2}^{2} = 3 \times 10^{2}$  m<sup>4</sup> for  $5 \times 10^{-1}$  or  $6 \times 10^{10}$  Hz  
(2) Particle Nature of Element's or  $6 \times 10^{10}$  Hz  
(2) Particle Nature of Element's constant and photoelectric effect.  
(2) Particle Nature of Element's or eating the constant action inclust but  
discontinuously but  
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discontinuously but  
 $f_{1-e}$  is a called or eating and ere entited from body not continuously but  
discontinuously but  
 $f_{1-e}$  is a planet's constant = 6.62 \times 10^{-3} Js  
at we know,  $y = \sum_{x = 1}^{2}$  hz planet's constant = 6.62 \times 10^{-3} Js  
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of sodium in terms of e.v.
sol:
                 Given, 2= 310 n-m , KE= 15ev ; Wo= ??
         We know, E= Wo + KE
                      hV = W_0 + kE \Rightarrow W_0 = hV - kE
                                          N_0 = h\left(\frac{c}{\lambda}\right) - k\epsilon
                                              = 12400 evA - 1.5 ev ⇒ 4-1.5ev
3100Å ⇒ 2.5ev
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(3) Dual Nature of EMR: To explain reflection, refraction, diffraction etc., light has been considered as a wave whereas to explain the photoelectric effect, Einstein considered it to be made up of tiny particles called photons. In other words, light is a kind of radiation exhibits dual behaviour i.e. waves as nell as particle behaviour. Such a wave like as well as particle like nature of radiation

is known as dual nature of radiation.

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(K3B)

(1) The splitting of light into series of colour bands is known as <u>dispersion</u> and the series of colour bands is called a <u>spectrum</u>. Two types of spectrum:
i) <u>Emission spectra</u>: when the radiations emitted from some source eg: from the sun

a: when the radiations emitted from some source eg: from the sun or by passing electric discharge through a gas at low pressure or by heating some substance to high temperature is passed directly through the prism and then recieved on the photographic plate, the spectrum obtained is called emission spectrum. Depending upon source of radiation, the emission spectra are of two types: Continuous and Line spectra.

(ii) <u>Absorption spectrum</u>. It is like the photographic negative of an emission spectrum. A continuum of radiation is passed through a sample which absorbs racliation of certain wavelength. The missing wavelength which corresponds to the radiation absorbed by the matter, leave dark spaces in the bright continuous spectrum.





tor eq. (i) lyman series. Let line 
$$\rightarrow n=2$$
 ton=1  
ind line  $\rightarrow n=3$  to  $n=1$   
ind line  $\rightarrow n=\infty$  to  $n=1$   
ind line  $\rightarrow n=\infty$  to  $n=2$   
ind line  $\rightarrow n=4$  to  $n=2$   
ind line  $\rightarrow n=4$  to  $n=2$   
ind line  $\rightarrow n=4$  to  $n=2$   
ind line  $\rightarrow n=0$  to  $n=2$   
last line  $\rightarrow n=\infty$  to  $n=2$   
ind line  $\rightarrow n=\infty$  to  $n=2$   
ind line  $\rightarrow n=\infty$  to  $n=2$   
ind line  $\rightarrow n=\infty$  to  $n=2$ 

10° The diff. Hw the V of 1st line of Balmer series and the last line of Parchen series for Litz Pan is how much?

soln Balmer: 1st line (n=3ton=2)  $\overline{\mathcal{V}} = \frac{1}{\lambda} = \mathcal{R}(3)^2 \left( \frac{1}{(2)^2} - \frac{1}{(3)^2} \right) \implies \mathcal{R}(9) \left( \frac{9-4}{36} \right) \implies \mathcal{R}(\frac{5}{4}) - (1)$ Paschen: last  $(n=\infty \text{ fo } n=3)$  (0) $\overline{\gamma} = \frac{1}{\lambda} = R(3)^2 (\frac{1}{\beta^2} - (1)) = R - (1)$ difference = 1 - 1 इस - R = R अब finally टम बात मरने जा रहे है Bohr पापा के Model के बारे में। BOHR Model (Applicable for single electronic species eg: H, Het, Litz, Betzete) Postulates: is Electron revolve around the nucleus in a fixed circular path of definite energy called stationary orbits. (i) Electron revolve only in those circular paths for which the value of angular momentum is equal to integral multiple of <u>h</u> i.e. 211 radius of <del>C</del>. mars of <u>mevo = n h</u> eog <u>velocity of eo</u> porbit no velocity of eo porbit no in Energy exchange takes place only when electron jumps blw the orbits.
 in frequency of radiation absorbed or emitted when transition occurs between two stationary states that differ in energy by AF is given by
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 in the state that the electron is not possible as long as the electron remains in one of its energy levels and hence. remains in one of its energy levels and hence , there is no cause of instability in n's model. # Calculation of Bohr Radius? According to Bohr's Model, radius of nth orbit is given by:  $T_n = \frac{0.0529 n^2}{z^2} nm$ Bohr radius: for hydrogen(z=1), the radius of first stationary state is called as Bohr Radius. fe. v=0.0529n-m is value of Bohr radius. # Energy of an electron:  $E = -13.6\left(\frac{z^2}{y_2}\right) = -V = 0^{3}$  $-2.18 \times 10^{-18} \left(\frac{z^2}{n^2}\right) J$ 

(B) Why energy of the electron in a hydrogen atom has negative sign for all

possible orbits? ans: This is because the energy of electron in the atom is lower than the energy of free electron at rest. A free electron at rest is an electron that is infinite for away from the nucleus i.e. free election at rest is an electron that is infinite far away from the nucleus i.e. free from influence of nucleus and is assigned the energy value of zero. Mathematically,  $n = \infty$  and thus  $E_{\infty} = 0$ . In this state hydrogen atomis called ionised H-atom. As the electron gets closer to the nucleus (as n decreases), En becomes larger in absolute value and more and more negative. The most negative energy value is given by n=1 which corresponds to the most stable orbit. We call this the ground state.

 +
 Calculation of frequency for a transition:

 S
 The frequency (v) associated with the absorption and emission of the photon can be evaluated by using equation:

 be evaluated by using equation:
 (v) = 2.20 × 10<sup>15</sup>

 $\sqrt{\gamma} = 3.29 \times 10^{15} \left( \frac{1}{\eta_1^{2}} - \frac{1}{\eta_4^{2}} \right) \text{Hz}$ 

and  $\overline{\nu} = \frac{1}{L} = Rz^2 \left( \frac{1}{n_1^2} - \frac{1}{n_2^2} \right)$  (calculated above)

(K<sup>3B)</sup>•In general, the number of emission lines when an electron jumps from n2 level to n1 level are given by the expression:-(n2-n1)(n2-n1+1) 2

(I·f·) Ionisation Energy: - Energy required to semore the e completely from the atom so as to convert it to a positive ion. Simply, The energy absorbed by electron in ground state (n=1) so as to jump to infinity. (n=a).  $f_{0} r eq, for H : I f := E_0 - E_1$   $= \left[ -\frac{2 \cdot 18 \times 10}{(0)^2} - [-2 \cdot 18 \times 10^{-18} (1)^2] - \left[ -2 \cdot 18 \times 10^{-18} (1)^2 \right] J$   $J_{TTE} = \left[ -\frac{2 \cdot 18 \times 10^{-18}}{(0)^2} - [-2 \cdot 18 \times 10^{-18} (1)^2] \right] J$ ईमी तरह की किसी की भी ⇒ 2.18×10-18J I.E. तिमाल समते हैं।

· Velocity of an electron in any Orbit:

$$V = \frac{2\pi K z e^2}{nh} m/sec = 2.18 \times 10^6 \left(\frac{z}{n}\right) m/sec$$

Frequency of revolutions/Number of revolutions =  $\frac{V}{2\pi r}$ substituting value of  $x_{3} = 2\pi m v Ze^{2}$ 

LP: Radius of two different orbits in H like sample is 4R f16R respectively. Find the ratio of frequency of revolution of electron in their 20rbits.

$$\underbrace{ \left\{ \underbrace{0} \right\}^{n}}_{1} : \underbrace{\frac{1}{1_{2}}}_{1} = \underbrace{\frac{V_{12}}{V_{2}} x_{1_{2}}^{n}}_{1} - \underbrace{2}_{1}^{n} \\ \xrightarrow{V_{2}} x_{1_{2}}^{n} + \underbrace{2}_{1}^{n} \\ \xrightarrow{V_{2}} x_{1_{2}}^{n} + \underbrace{2}_{1}^{n} \\ \xrightarrow{V_{2}} x_{1_{2}}^{n} + \underbrace{2}_{1}^{n} \\ \xrightarrow{V_{1}} x_{1_{2}}^{n} = \underbrace{\frac{V_{12}}{V_{2}} x_{1_{2}}^{n}}_{n} \\ \xrightarrow{N_{1}} = \frac{1}{2} \\ \xrightarrow{N_{1}} = \underbrace{N_{1}}_{n} \\ \xrightarrow{N_{1}}_{n} \\ \xrightarrow{N_{1}} = \underbrace{N_{1}}_{n} \\ \xrightarrow{N_{1}}_{n} \\ \xrightarrow{N_{1}} = \underbrace{N_{1}}_{n} \\ \xrightarrow{N_{1}}_{n} \\ \xrightarrow{N_$$

o could not explain the ability of atoms to form molecules by chemical bonds.

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Ĥψ=EΨ Ĥ is called Hamiltonian operator, Eard Y are obtained from solution of schrodinger wave equation. (K<sup>3</sup>B) ⇒ When Schrodinger equation is solved for hydrogen atom, the solution gives the possible energy levels the electron can occupy and the corresponding wave functions of V of electron associated with each energy level. These quantized energy states and corresponding wavefunctions which are characterized by a set of three quantum numbers - principal quantum number n, azimuthal quantum number l and magnetic quantum number m.

Atomic Orbitals: Ware functions of hydrogen or hydrogen like species with one electron. (Wave function is a mathematical function whose value depends upon the coordinates of the electron in the atom and cloes not carry any physical meaning.

An orbital is a region in space around the nucleus where the prob. of finding the electron is maximum.

ORBIT	ORBITAL
• It is well defined circular path around the	· It shows the 3 dimensional space around
nucleus in which electron revolve.	the nucleus within which the probability
• Max <sup>m</sup> no. of e <sup>®</sup> that an orbit can have is	of finding an et is maximum.
given by 2n² where n is the no. of orbit.	· Max <sup>m</sup> no. of electrons that can be occupied by
0 1	an orbital is always two.

Quantum Numbers

 Set of 4 numbers which is used to define an electron completely -n, l, m, s n→principal quantum number L→ Azimuthal quantum number. mL→ Magnetic quantum number. s→spin quantum number.
 (K<sup>3</sup>B) → Orbit Tri sigt subshell, The stip sigt orbital.
 (reular path around nucleus in which e<sup>Q</sup> revolve.
 (max<sup>m</sup> no. of e<sup>Q</sup> in any orbit is 2n<sup>2</sup>) in which e<sup>Q</sup> revolve.
 (orbital can have max<sup>m</sup> of 2e<sup>Q</sup>) where probability of finding an e<sup>Q</sup> is maximum.

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lobe → e<sup>0</sup> density





In notes ko padhkar is

• Spin angular momentum =  $\frac{h}{2\pi} \int s(s+1)$ 

# Energy of orbitals:	
Monoelectronic species	Multi electronic species
• There is only nuclear attractive force	• It is nuclear attractive force as well as interelectronic forces.
<ul> <li>Energy of all orbitals in same shell is same</li> </ul>	o Energy of different orbitals is different in same shell due to interelectronic
<ul> <li>Energy only depends on principal</li> <li>Quantum number.</li> </ul>	• Energy depends upon 'n' as well as
15 < 2s < = 2p < 3s = 3p = 3d	1s < 2s < 2p < 3s < 3p < 4s < 3d
Zeff (Effective Nuclear Charge):- Due to shielding of outer shell e <sup>⊕</sup> from nucleus by inner shell e <sup>⊕</sup> , net ⊕ve charge experienced by outer shell e <sup>⊕</sup> from nucleus.	
Zefft lf Shielding Power s>p7d>f	
for same subshell Energy + as atomic no. (Zefff)	
eg: $E_{2s}(H) > E_{2s}(L_1^\circ) > E_{2s}(Na) > E_{2s}(k)$	
means ding $\psi$ here $n \rightarrow const$ $F \propto -z^2$ $z = F + z^2$	
Autbau Principal	
· Acc to Authan principal, e are filled in orbitals in increasing	
<ul> <li>∑st e are filled in lower energy orbitals and then e are filled in higher energy orbitals.</li> </ul>	
Energy: 1s<2s<2p<3s<3p<4s<3d<4p<5s<4d<5p<6s<4f<5d <6p<7s	