

Maa Omwati Degree Collage

Hassanpur

NOTES

Subject:- Atomic Molecular And laser Physics

B.SC Sem:-VI

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A & M Physics

Quantum Analysis is the best study of A & M Physics

- NO e^- occurs in isolation. We need to study via a sample.

~~Similar atoms & molecules in sample~~

- ✓ Best way to reveal internal structure of nucleus, atom & molecules is spectrum.

Spectral line can be characterized by ν or ω or λ .

Every field of science has its own notations!!
 $\frac{\nu}{c} = \bar{\omega} = \frac{1}{\lambda_{vacuum}}$ * Called wave number, $\bar{\omega}$
 $\bar{\omega}$ is a replacement of frequency which is usually a large number. $\bar{\omega}$ = no. of waves per metre or centimeter
It remains const. irrespective of medium.

- ⊗ As long as particle (e^- or atom or molecule or nucleus)

~~is in fixed state, its energy is constant~~ Bohr's

When it goes to higher energy and it comes down and releases energy

$$\Delta E = h\nu = hc\bar{\omega} = \left(\frac{hc}{\lambda}\right)$$

- Same specimen can give spectra of e^- , atoms, molecules & nucleus.

- ~~Excitation~~ Excitation Mechanism : that causes particle to go to higher energy.

It has to be chosen in such a way that it is able to change:
Molecular Energy levels
& Atomic Energy levels
& Nucleus Energy levels

It could be suitable radiation or current or atomic collisions.

Typical Spectrometer



① If white light source, I require modulator to choose wavelength of our choice (filter)

Maximum ΔE : atoms several eV eg. $(-13.6 \text{ eV} - (-\frac{13.6}{4} \text{ eV}))$

Then ΔE : molecular

Then ΔE_{min} : nuclear $\approx 10^{-8} \text{ eV}$



② Note that there are two types of nuclear energy levels. 1 due to excitation of shell model states... they correspond to γ ray. While these states are excited due to external magnetic fields are Radio Frequency Regions that of radio frequency rays.

$$\lambda = \frac{hc}{\Delta E}$$

Hence, Radio Frequency Spectra of nuclear energy levels
Therefore suitable source is Radio Frequency Oscillator.

Example NMR.

$$\Delta E_{molecular} \approx 10^{-3} \text{ eV}$$

3 types of energy: ① Rotational $\sim$ Microwave SPECTRA
eg. Rhyphon Oscillator: Microwave source

② Vibration 10^{-4} eV $\sim$ Infrared Region

③ Electronic Energy Levels $\sim$ Several eV
ie. Visible or UV

Source

$\Delta E_{atomic} \approx$ many eV Balmer $\leftarrow$ Visible and Infrared Spectrum $\leftarrow$ Rosen, Balmer, Lyman, Paschen, etc.

eg. $(13.6 - \frac{13.6}{4}) \text{ eV}$ in H atom

If $T = 300 \text{ K} = 27^\circ \text{C}$

$$\text{Equivalent} = kT = 0.026 \text{ eV}$$

It is sufficient to cause rotation of molecules but not vibration.



Atomic Models.

Thomson Model (Plum Pudding Model)

Rutherford Model (Based on scattering experiments)

Bohr's Model (1913) (Angular Momentum Quantization Model)

$$\alpha = 0.53 \text{ \AA}$$

$$E_n = -\left(\frac{13.6}{n^2}\right) \text{ eV}$$

Schrodinger's Model



Heisenberg's Model

or

Vector Model of Atom

Boris Model

Bohr removed discrepancies in Rutherford Model by introducing quantization of Angular Momentum, thereby quantizing energy and saying that Energy will remain constant and energy will not be continuously emitted thereby e- will not collapse in nucleus.

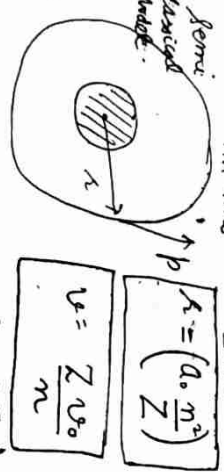
Bohr Quantized Angular Momentum $\vec{L} = |\vec{r} \times \vec{p}| = n\hbar$

$$E_n = -\left(\frac{13.6}{n^2}\right)$$

$$\Delta E = \left(\frac{hc}{\lambda}\right) \Rightarrow E_f - E_i = \frac{hc}{\lambda}$$

$$= 13.6 \left[\frac{1}{n_1^2} - \frac{1}{n_2^2} \right]$$

$\frac{mv^2}{r} = \frac{Zq^2}{4\pi\epsilon_0 r^2}$ $m\hbar v = \frac{Zq^2}{4\pi\epsilon_0}$ $v = \frac{Zq^2}{m\hbar 4\pi\epsilon_0}$



$$E = \frac{p^2}{2m} + V_{el}$$

$$= \frac{p^2}{2m} - \frac{Ze^2}{4\pi\epsilon_0 r}$$

$$= \frac{1}{2} m v^2 - \frac{Ze^2}{4\pi\epsilon_0 r}$$

$$= \frac{1}{2} m e \frac{Z^2 e^4}{n^2 \hbar^2 (4\pi\epsilon_0)^2} - \frac{e^4 Z^2 m e}{(4\pi\epsilon_0)^2 n^2 \hbar^2} = -\frac{1}{2} \frac{Z^2 m e^4}{n^2 \hbar^2 (4\pi\epsilon_0)^2}$$

$$\alpha = \frac{e^2}{4\pi\epsilon_0 \hbar c} \approx \frac{1}{137}$$

dimensionless number : FINE STRUCTURE CONSTANT.

$$E_n = -\frac{m e^4}{2 \hbar^2} \left(\frac{Z^2}{n^2} \right) = -R h c \left(\frac{Z^2}{n^2} \right) \quad R = \text{Rydberg Const.}$$

$$R = 1.09 \times 10^7 \text{ m}^{-1}$$

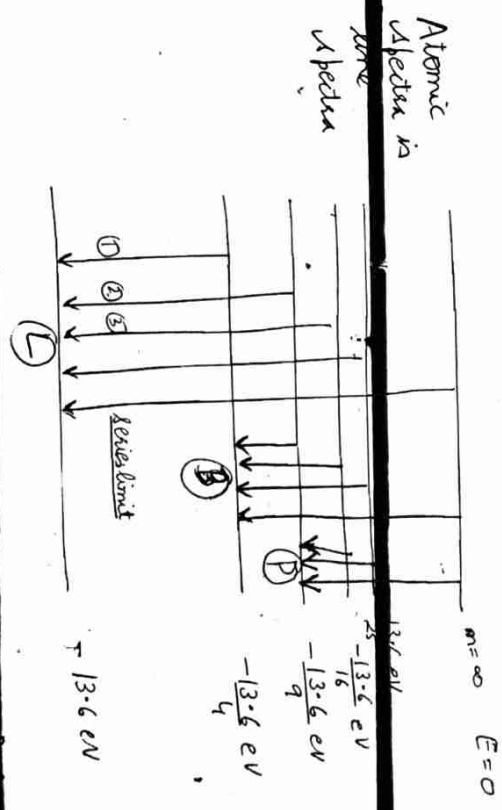
$$E_n = -13.6 \left(\frac{Z^2}{n^2} \right)$$

Remember, $E_n = -(k \cdot E)_n$

$$1 \leq n < \infty$$

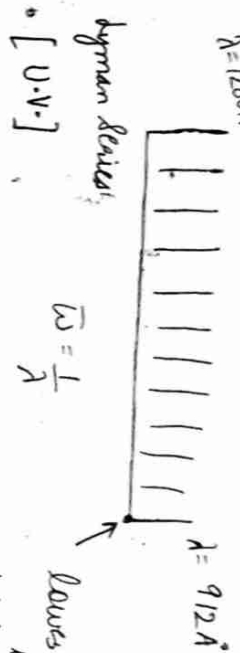
For e- to move out of atom, $k \cdot E > P \cdot E$

$$\text{① } n = \infty \quad \underline{E = 0} \quad \frac{1}{2} m v^2 = \frac{Z e^2}{4\pi\epsilon_0 r}$$



$n=1$ to $n=1$	: Lyman series	UV Region
$n=1$ to $n=2$	: Balmer series	Visible Region 3647 Å - 6563 Å
$n=2$ to $n=3$	: Paschen series	IR
$n=3$ to $n=4$	: Brackett series	IR
$n=4$ to $n=5$	: Pfund series	IR

$$\lambda = 1200 \text{ \AA}$$



$$\bar{\omega} = \frac{1}{\lambda}$$

Lowest Wavelength
Highest Wavelength

$$\frac{1}{\lambda} = R \left[\frac{1}{n_2^2} - \frac{1}{n_1^2} \right]$$

$$\frac{1}{\lambda_{\min}} = R \left[\frac{1}{1^2} - \frac{1}{\infty} \right] = R \Rightarrow \lambda = \frac{1}{R} = \frac{1}{1.09 \times 10^7} = 912 \text{ \AA}$$

$$\frac{1}{\lambda_{\max}} = R \left[\frac{1}{1^2} - \frac{1}{2^2} \right] = \frac{3R}{4} \Rightarrow \lambda_{\max} = \frac{4}{3R} = \frac{4}{3 \times 1.09 \times 10^7} = 1200 \text{ \AA}$$

note that we are always interested in this series $\therefore$ λ (n=3 to n=2) λ (n=4 to n=3) refer to this series

Balmer Series, $\frac{1}{\lambda_{\min}} = R \left[\frac{1}{2^2} - \frac{1}{\infty} \right] = \frac{R}{4}$

$$\Rightarrow \lambda_{\min} = \frac{4}{R} = 4000 \text{ \AA}$$

$$\frac{1}{\lambda_{\max}} = R \left[\frac{1}{2^2} - \frac{1}{3^2} \right] = \frac{5R}{36} \Rightarrow \lambda_{\max} = \frac{36}{5R} = 7000 \text{ \AA}$$

$$\Rightarrow \lambda_{\max} = \frac{36}{5R} = 7000 \text{ \AA}$$

If I see from very sensitive spectrometer, I can see few lines in between the lines also, called fine structure.

Bohr's Model Drawbacks (couplings)

1) Not valid for more than 1 e- atoms.

2) It does not explain fine structure of lines. No logic on reason given for quantization of angular momentum.

3) When atom is kept in external field, it does not follow the behaviour: Zeeman Effect (Magnetic field), Stark Effect (Electric field).

4) Phase orbits do not exist. Quantum Mechanical Probabilities.

5) Note that if I take Hamiltonian as $\left(\frac{p^2}{2m} + V_e \right) \cdot I$ will get same values as Bohr Model. Unless I change Hamiltonian, I won't get fine structure by Quantum Mechanics.

That change is SPIN. (Remember, in scattering we add the term $\propto \frac{1}{2} \vec{S} \cdot \vec{S}$ in Hamiltonian)

Electron spin introduced in 1925 by Goudsmit & Uhlenbeck.

(1) To convert Gyromagnetic Ratio

$$\left(\frac{\text{Magnetic Moment}}{\text{Angular Momentum}} \right) = \frac{\mu}{L} = -\frac{e}{2m}$$

$$\vec{\mu}_L = -\frac{e}{2m} \vec{L}$$

it needed to be introduced

(2) To ~~explain~~ fine structure of atomic spectra

(3) To explain Zeeman Effect

(4) To explain Stern-Gerlach Experiment.

Now $\vec{\mu}_L = -\frac{e}{2m} \vec{L} = -\frac{e}{2m} g_L \vec{L}$ $g_L = 1$

Quantum Mechanically $\vec{\mu}_L = -\frac{e}{2m} \vec{L} = -\frac{e}{2m} g_L \vec{L}$ $g_L = 2$

Orbital $L = 0 \Rightarrow L = 0 \Rightarrow \mu_L = 0$

$\psi_{l,m} = |\psi\rangle = \sqrt{l(l+1)} \hbar = \frac{\sqrt{3}}{2} \hbar$ $\vec{\mu}_L = -\frac{e}{2m} \vec{L}$

but if $\boxed{\uparrow\downarrow}$ $\psi = 0 \Rightarrow \mu_L = 0$ } fully filled orbitals

Orbital $L = 1 \Rightarrow L = \sqrt{2}$

$\vec{J} = \vec{L} + \vec{S}$: LS coupling

$\vec{\mu}_J = \frac{e}{2m} g_J \vec{J}$: g_J : Lande's "g" splitting factor

This is called vector model of atoms for multi electron atom : LS coupling for single-electron atom : spin-orbit coupling

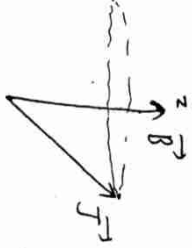
If I keep the atom in magnetic field ($\vec{B}$),

if $\vec{B}$ external $\Rightarrow$ (Zeeman Effect) / (Paschen Back Effect) (weak $\vec{B}$) (strong $\vec{B}$)

if $\vec{B}$ internal $\Rightarrow$ LS coupling or spin-orbit coupling (multi-electron atom) (single-electron atom)

$\vec{T} = \vec{\mu}_J \times \vec{B}$ ($\mu_J \propto \vec{J}$) $\Rightarrow \vec{T} \perp \vec{J}$ ($\Rightarrow$ Precession)

Torque will try to align $\vec{\mu}_J$ along $\vec{B}$. hence $\vec{\mu}_J$ will precess i.e. $\vec{J}$ will precess in direction of $\vec{B}$.



Remember from space quantization that in whatever direction I apply $\vec{B}$, $\vec{J}$ will align at specific angles to that direction i.e. $J_z = m_J \hbar$ since $m_J < j \Rightarrow \vec{J}$ cannot be $\parallel$ to $\vec{B}$

when $\vec{J}$ is precessing, J_z will be const, $J_z = m_J \hbar$

$\frac{d\vec{J}}{dt} = -\frac{e}{2m} g_J (\vec{J} \times \vec{B})$ Larmor Precession $j < m_j < j$ $(j+1)$ orientations

$E = -\vec{\mu}_J \cdot \vec{B}$ Potential Energy of Orientation of Dipole

$= +\frac{e}{2m} g_J \vec{J} \cdot \vec{B}$

$= +\frac{e}{2m} g_J B J_z$

$E = +\frac{e \hbar}{2m} \bullet B \cdot g_J m_j$

Atom Geelach Experiment

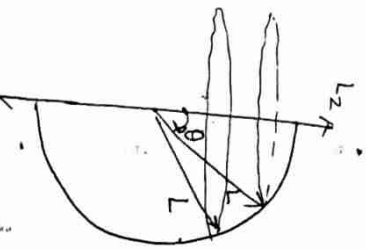
It is "VIVID CONFIRMATION OF VECTOR MODEL OF AN ATOM"

Vector Model :- $\vec{J} = \vec{L} + \vec{S}$

In Bohr Model, S was not there; only $L = n\hbar$

$E = -13.6 \frac{Z^2}{n^2}$ Coming out due to quantization of L

It is demonstrating : ① Orbital Motion or Space Quantization
② Spin Motion governed by



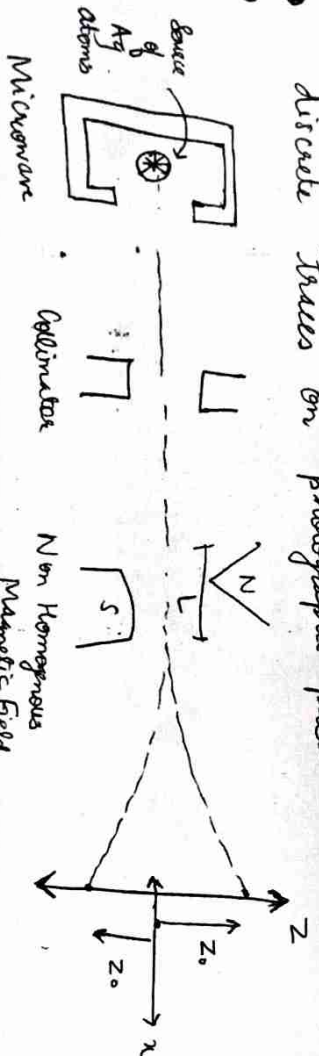
$$S^2 = s(s+1)\hbar^2$$

$$S_z = m_s \hbar$$

$$-s < m_s < s$$

Total angular momentum of an atom results from the combination of spin and orbital angular momenta of its electrons. Since any momentum is a vector quantity, we can represent the total angular momentum as addition of orbital and spin angular momentum vectors.

Beam of neutral silver (Ag) atoms is passed through non-homogeneous magnetic field giving rise to 2 discrete traces on photographic plate



A & M PHYSICS (2)

Ag ($Z=47$) $5s^1$: e^- are responsible for ΔE .

Note that only valence e^- are responsible for jumps.

For fully filled orbitals, $L=0$ and $S=0$ i.e. $J=0$

$$\left. \begin{matrix} n s^2 \\ n p^6 \\ n d^{10} \end{matrix} \right\} \begin{matrix} L=0 \\ S=0 \end{matrix}$$

Note that Energy level is defined by J . Hence they contribute nothing to ΔE .

In spectroscopy, valence electrons are called Optically Active Electrons.

For Ag, $L=0$ (s orbital)

$$\vec{J} = \vec{L} + \vec{S} = \vec{S}$$

$$S_z = m_s \hbar$$

$$\vec{\mu}_J = -\frac{e}{2m} \cdot g_J \vec{J}$$

$$\vec{\mu}_S = -\frac{e}{2m} \vec{S}$$

Whenever it is left in field, Torque acts giving rise to Precession.

$$E = -\vec{\mu}_s \cdot \vec{B} = -(\mu_s)_z B$$

$$(if \vec{B} = B \hat{z})$$

$$(\mu_s)_z = -\frac{e}{2m} g_s m_s \hbar$$

$$(\mu_s)_z = -\frac{e \hbar}{2m} g_s m_s$$

$$\frac{e \hbar}{2m_e} = \mu_B : \text{Bohr Magnetron}$$

$$\mu_B = 9.27 \times 10^{-24} \text{ Joule/Tesla}$$

$$\frac{e \hbar}{2m_p} : \text{Nuclear Bohr Magnetron}$$

✓ $(\mu_s)_z$ can take no of value that m_s can have.
Hence Stern-Gerlach experiment also gives quantization of magnetic moment.

✓ It should not be a charged particle. Otherwise $F = q(\vec{v} \times \vec{E})$ will act and circular path will be obtained, hence no trace on plate. Hence, we took Δ neutral particles.

We want only force in z direction.
Experiment will not succeed in $\vec{B}$ is Homogeneous, because otherwise no net force will act on dipole moment.

$$\vec{F} = (\vec{\mu} \cdot \vec{\nabla}) \vec{B}$$

$$\vec{F} = 0 \text{ for homogeneous field}$$

★ The quantity $\left(\frac{e \hbar}{2m}\right)$ forms a natural unit for the measurement of atomic magnetic dipole moments and is called Bohr Magnetron, denoted by μ_B .

$\vec{B}$ is non homogeneous

$$\vec{B} = B_x \hat{i} + B_y \hat{j} + B_z \hat{k} = B_z \hat{k}$$

$$\text{and } \left(\frac{\partial B_z}{\partial z}\right) \neq 0 ; \frac{\partial B_z}{\partial z} = 0 = \frac{\partial B_z}{\partial y}$$

$$F_z = \mu_z \left(\frac{\partial B_z}{\partial z}\right)$$

$$= \mu_B g_s m_s \left(\frac{\partial B_z}{\partial z}\right)$$

$$\text{distance } x = L$$

$$\text{velocity } v_x = v_x \text{ (const.)}$$

$$t = \left(\frac{L}{v_x}\right)$$

$$v_z \hat{i} = 0$$

$$a_z = \left(\frac{F_z}{m}\right) \Rightarrow z = \frac{1}{2} a_z t^2 = \frac{1}{2} a_z \left(\frac{L}{v_x}\right)^2$$

$$= \frac{1}{2} \frac{\mu_B \left(\frac{\partial B_z}{\partial z}\right) g_s m_s L^2}{m v_x^2}$$

$$\frac{1}{2} m v_x^2 = kT \text{ that can be considered by temp. of oven}$$

$$\text{Normally, it is } \frac{3}{2} kT \quad \left[\begin{array}{l} \text{like } 300 \text{ K: } \frac{3}{2} kT \\ > 300 \text{ K: } \frac{5}{2} kT \end{array} \right]$$

$$\Rightarrow z = \frac{\mu_B \left(\frac{\partial B_z}{\partial z}\right) L^2}{3 kT} \frac{g_s m_s}{2}$$

$$g_{-3} = 2$$

o Cadmium } fully
Mercury } free
oil

Home 2 traces.

fixed $s = \frac{1}{2}$

$$\Rightarrow Z = \pm \frac{1}{2} \left(\frac{\mu_B \left(\frac{\partial B_z}{\partial z} \right) L^2}{3kT} \right) = \pm 2$$

$\sum_j (2m)^{L_j}$
 (1 dependent) $\rightarrow$ (8 applied) $\rightarrow$ (m; dependent)
 4 Quantum numbers
 State, level, sub level, Spectral Term

State: Complete description of $\psi (n, l, m_l, m_s)$
We require all the 4 set of quantum numbers.

Energy level defined by $\vec{J}$ ($= \vec{L} + \vec{S}$)

Rule to 1-77 on
78

more the
more the

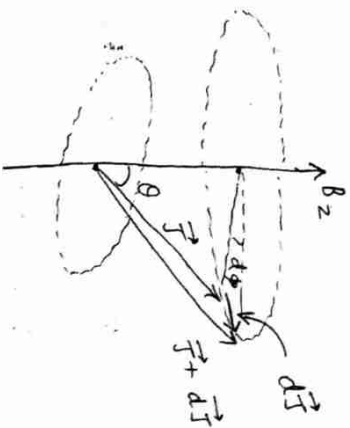
over the energy $\exists$ exists $\Rightarrow \mu_5$ exists
hence in $\exists$ exists $\Rightarrow \mu_5$ exists

general
 $E_S < E_P < E_D$ for

If $\vec{B}$ exists $\Rightarrow$ $\vec{J}$ will precess around $\vec{B}$ called Larmor Precession

line : 2 subpanels (of different levels of course) at 1/24 in

EXTERNAL MAGNETIC FIELD



→ 5
T
→ 5

$$\omega_p = \left(\frac{d\phi}{dt} \right) = \frac{\left(\frac{dJ}{J \sin \theta} \right)}{dt}$$

$$\tau = \frac{d\vec{J}}{dt} = \vec{\mu}_J \times \vec{B} = \mu_J B \sin \theta$$

$$\Rightarrow \omega_p = \frac{\mu_B \sin \theta}{J \sin \theta} = \frac{e}{2m_e} B g \gamma$$

$$\left(\frac{\mu_j}{T} = \frac{e}{T_m} g_j \right)$$

$$W_P = \frac{e}{2m} g \mu_B$$

$\Rightarrow$ cup is independent of n .
all have same cup

$$\vec{E} = -\vec{\mu} \cdot \vec{B} = \frac{e}{2m} \vec{S} \cdot \vec{B}$$

$$E = \mu_B \cdot B \quad \text{J m}^{-1}$$

$$= \frac{e}{2m} \vec{S} \cdot \vec{B} = \mu_B \frac{1}{2} N_{m_j} = (Z+1) \mu_B$$

$$\gamma_p = \frac{e}{4\pi m} B g J$$

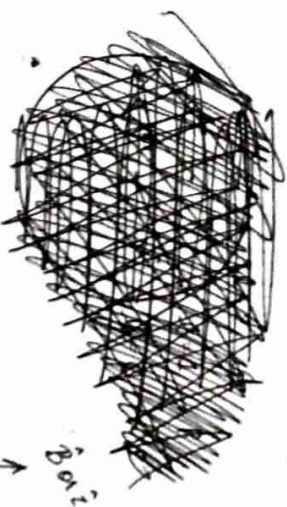
m_j : Magnetic Moment Quantum No.

→ For energy level will be split into $(2l+1)$ energy levels.

sublevel.

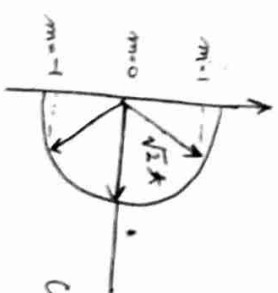
l	0	1	2	3	4	5
Orbital	s	p	d	f	g	h

② $(2l+1)$ gives different orbitals i.e. no. of values of m_l i.e. number of orbitals in the sublevel



Zooming $l=1$

from Top view



Counting only valence electron

$x = \text{Multiplicity} = (2s+1)$ for single e^- atom

Many electron atoms: $2s+1$ ($L > S$)

$2s+1$ always multiplicity =

$2L+1$ ($S > L$)

$$L = 2s(s+1) + 1 = \text{no. of energy levels}$$

eg.

Na: $1s^2 2s^2 2p^6 3s^1$

X $L=0$

$S = \frac{1}{2}$

$J = \frac{1}{2}$

Term Symbol: $3s^2 3p^1$

Multiplicity 3

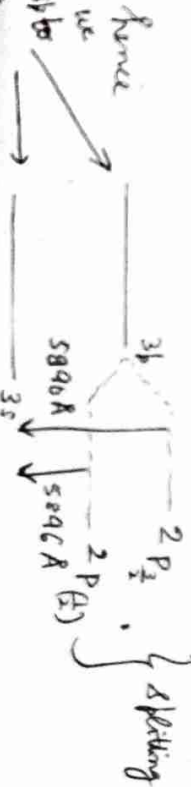
3s

For excited Na atom,

$l=1$ (p)

$S = \frac{1}{2}$

$J = L \pm S = \frac{1}{2}, \frac{3}{2}$



D_1 and D_2 lines

Note that we have more levels from 3s.

$\Delta E = -\frac{5.89 \times 10^{-19}}{m} \left[\frac{1}{n_1^2} - \frac{1}{n_2^2} \right]$

This is called ΔE caused due to spin

③ Term symbol corresponds to 1 energy level i.e. 1 particular value of J

④ When subjected to B ext. each J i.e. each level will break into $(2J+1)$ sublevels

Energy level : Transition between 2 energy levels (Note that Sodium transitions are 2 lines)

Energy sublevel : Component : Transition $1/2$

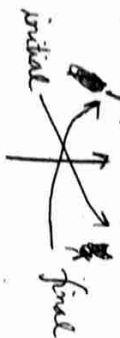
(Components are those that is Zeeman effect on Sodium Transitions)
(Of course not 2 sublevels of same level)

Spectral Term L_j : Multiplets : Transition $1/2$ 2 spectral levels.

But what are allowed transition?

It is governed by selection Rule:

$$P(\text{transition}) = \langle f | A | i \rangle \neq 0$$



Reason for transition

Selection rule

eg $\langle i | \mu_z | f \rangle \neq 0$

or allowed transition $\int \psi_i^* \mu_z \psi_f d\tau > 0$

we get, $\Delta m_j = \pm 1, 0$ for Zeeman effect

When 2 e- are said to be equivalent?

$2p^1 3p^1$: non equivalent

$2p^1 2p^1$: Equivalent

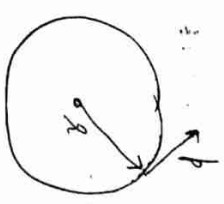
Equivalent means :- n and l quantum numbers

are same.

IT IS APPLICABLE AND OBSERVABLE IN ALL ATOMS.... IT IS MORE DOMINANT IN HEAVIER ATOMS WHERE L and S ARE INDIVIDUALLY COUPLED TO FORM INDIVIDUAL ATOMS AND ONLY THEN COUPLED TOGETHER. THIS COUPLING IS

spin e- produces its own μ_s

NOT STRONG IN LIGHT ATOMS WHERE L 's and S 's are INDIVIDUALLY COUPLED AND THEN COMBINED TO FORM J



Orbiting e- $\Rightarrow$ current loop $\Rightarrow$ creates its own magnetic field

For current loop, $B = \frac{\mu_0}{2\pi} \left(\frac{I}{r} \right)$

$$= \frac{\mu_0 e L}{4\pi m r^3} = \left(\frac{e}{m} \right) \left(\frac{\mu_0}{4\pi} \right) \frac{L}{r^3}$$

$$\vec{B}_{\text{int}} = \frac{e}{m} \frac{\mu_0}{4\pi} \frac{\vec{L}}{r^3}$$

In c-frame of reference, nucleus is rotating with ω . Hence it will experience force due to field created by nucleus.

$\vec{B}_{int} = 0$ only if $\vec{L} = 0$

$\vec{T} = \vec{\mu}_s \times \vec{B}$

$= \mu_s B \sin \theta$

$d\omega = T d\theta = -\vec{\mu}_s \cdot \vec{B} = \frac{e}{2m} \cdot 2\vec{s} \cdot \vec{B}$

$= \frac{e}{2m} 2s_z B$

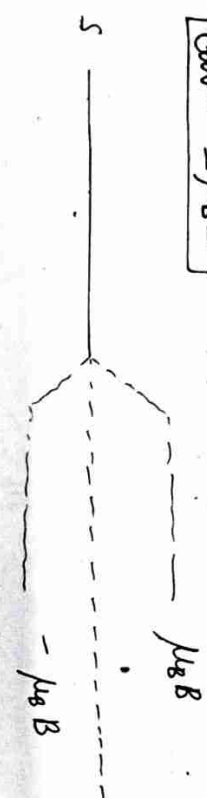
$= \frac{e \hbar}{2m} 2ms \cdot B$

$m_s = \pm \frac{1}{2}$

$d\omega = \pm \left(\frac{e \hbar}{2m} \right) B = \pm \mu_B B$

$dM = \pm \mu_B B$

INTERNAL MAGNETIC FIELD



It will result into fine structure doubling.

This analysis for single valence e^- .

Single e^- atoms μ_s interacting with $\vec{B}_{int}$ to lift the

Its a weak interaction.

$E = \left(\frac{e}{m} \right)^2 \frac{\mu_0}{4\pi} \frac{L}{r^3}$

Self Last Page

$E_{LS} = \left(\frac{\mu_0}{4\pi} \right) \left(\frac{e}{m} \right)^2 \frac{1}{r^3} (\vec{L} \cdot \vec{S}) = \vec{\mu}_s \cdot \vec{B}$

$\Rightarrow \vec{E} = \alpha \frac{1}{r^3} \vec{L} \cdot \vec{S}$

Hence its called spin orbit coupling.

Now E_{LS} is the correction factor in Hamiltonian.

It will now give fine structure, Quantum Mechanically.

$H' = \frac{p^2}{2m} + V(r) + \left[\frac{\alpha}{r^3} (\vec{L} \cdot \vec{S}) \right]$

$H' = H + \frac{\alpha}{r^3} (\vec{L} \cdot \vec{S})$

from, $H\psi = E\psi$
 $E = \frac{-B \cdot 6}{n^2}$

$\langle \frac{1}{r^3} (\vec{L} \cdot \vec{S}) \rangle$

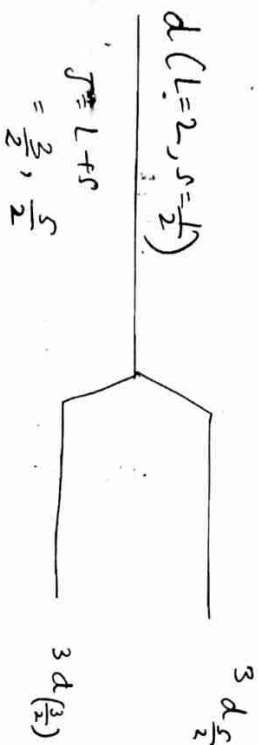
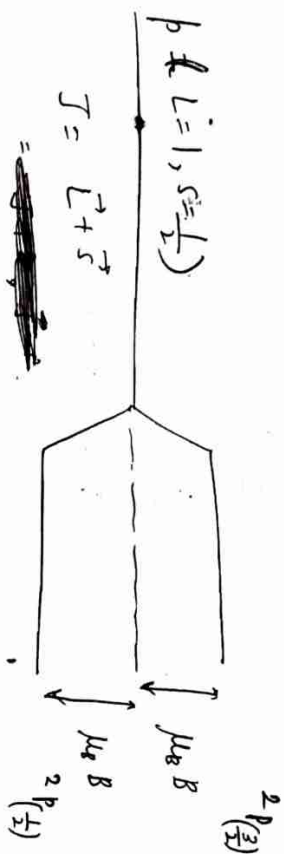
$\vec{J} = \vec{L} + \vec{S}$

$J^2 = L^2 + S^2 + 2\vec{L} \cdot \vec{S}$

$\vec{L} \cdot \vec{S} = \frac{J^2 - L^2 - S^2}{2} = \frac{j(j+1) - l(l+1) - s(s+1)}{2} \frac{\hbar^2}{4}$

$$E_{l,s} = \left(\frac{\mu_B}{4\pi} \right) \left(\frac{e}{m} \right)^2 \left(\frac{h}{2} \right) \frac{1}{\hbar^3} L \cdot S(j+1) = L(L+1) - S(S+1)$$

$$= \alpha < \frac{1}{\hbar^3} [j(j+1) - L(L+1) - S(S+1)] >$$

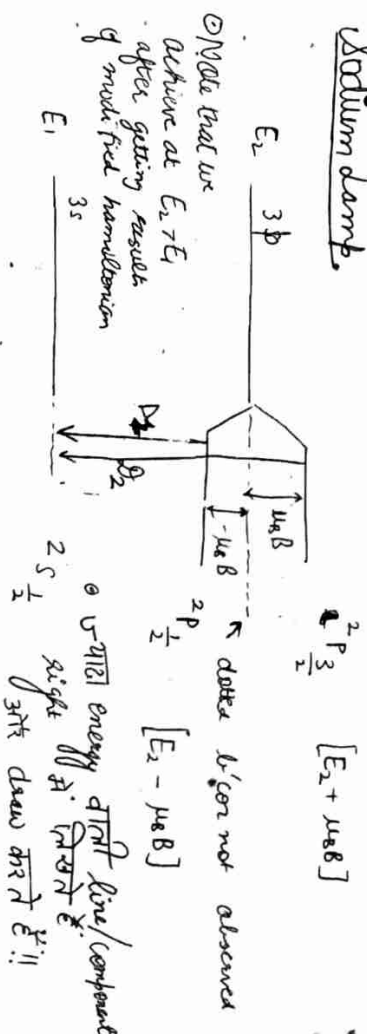


$$= \alpha [j(j+1) - L(L+1) - S(S+1)] < \frac{1}{\hbar^3} >$$

$$H_{atom} < n, l, m | \frac{1}{\hbar^3} | n, l, m >$$

Perfect example for spin orbit coupling are d, and dz lines of sodium lamp.

Sodium lamp



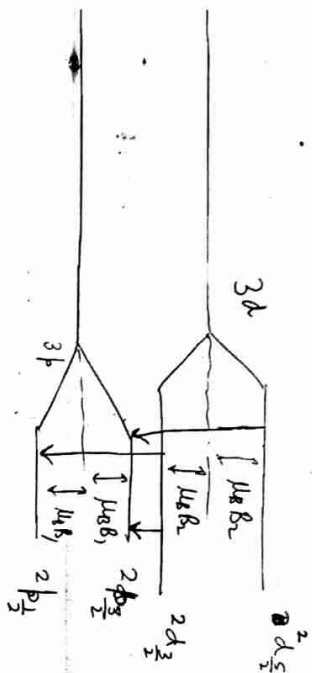
For spin orbit coupling selection rules

$$\lambda_1 = \frac{hc}{\lambda_1} = (E_2 - \mu_B B) - E_1 = (E_2 - E_1) - \mu_B B$$

$$\lambda_2 = \frac{hc}{\lambda_2} = (E_2 - E_1) + \mu_B B$$

$$hc \left[\frac{1}{\lambda_2} - \frac{1}{\lambda_1} \right] = 2\mu_B B$$

But can be calculated which is felt by p electron. No coupling in s electron.



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B_1 and B_2 are not equal in general.

3 spectral lines for $d \rightarrow p$ transitions

$$\left. \begin{aligned} \Delta l &= \pm 1 \\ \Delta j &= 0, \pm 1 \\ \Delta s &= 0 \end{aligned} \right\}$$

Fine Structure of H atom

Dirac's Quantum Mechanical Model

Spin Orbit Coupling

$$\begin{aligned} H_{\text{spin}} &= \frac{h\nu}{4\pi} \left(\frac{e}{m} \right)^2 \frac{h^2}{2} [j(j+1) - l(l+1) - s(s+1)] \\ &= \frac{Z^4 R_\infty^2}{n^3 l(l+1)} \end{aligned}$$

Relativistic Correction in Hamiltonian.

$$H = E - m_0 c^2 + V$$

$$= c^2 (m - m_0) + V$$

$$= m_0 c^2 \left[\left(1 - \frac{v^2}{c^2} \right)^{-\frac{1}{2}} - 1 \right] + V$$

$$= m_0 c^2 \left(1 + \frac{v^2}{2c^2} - \frac{3}{8} \frac{v^4}{c^4} + \dots \right) + V$$

$$= \frac{1}{2} m_0 v^2 - \frac{3}{8} \frac{m_0 v^4}{c^2} + V$$

$$\text{Relativistic Correction} = -\frac{3}{8} \frac{p^4}{m_0^3 c^2}$$

$$\langle H' \rangle = -\frac{3}{8} \frac{p^4}{m_0^3 c^2} \quad \langle p^4 \rangle$$

Solution of $H\psi = E\psi$ gives Bohr's n^{th} level. The departure from these values is given by the above 2 corrections. It comes out to be:

$$\Delta E = -\frac{Z^4 R_\infty^2}{n^3} \left[\frac{1}{j+\frac{1}{2}} - \frac{3}{4n} \right]$$

$$\Delta E = -\frac{5.84 Z^4}{n^3} \left[\frac{1}{j+\frac{1}{2}} - \frac{3}{4n} \right] \text{ cm}^{-1}$$

Dirac's Quantum Mechanical Model

Also see Bohr Model. This is responsible for $E_d > E_p > E_s$!!

eg. $n=3$

$n=3^*$ Bohr Energy

3d

3p

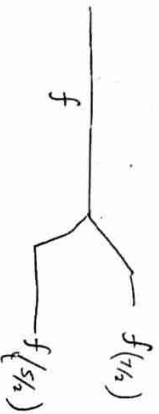
3s

$\Rightarrow$ for $l=0, 1, 2$

$\Delta E = -\frac{3}{8} \frac{p^4}{m_0^3 c^2}$

Hence Energy depends on n value; then on a finer level it depends on l values; then on a further finer level it depends on j values.

is values have no spin orbit coupling



spin orbit coupling only for single e- atoms.

spin orbit coupling Rules of selection:

$$\begin{cases} \Delta l = \pm 1 \\ \Delta j = 0, \pm 1 \\ \Delta s = 0 \end{cases} \quad \left(j=0 \text{ is not allowed} \right)$$

$$\Delta T = - \frac{R \alpha^2 Z^4}{n^3} \left[\frac{1}{j+1/2} - \frac{3}{4n} \right]$$

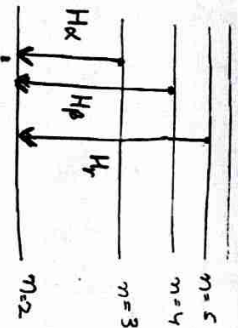
Departure from Bohr's n^{th} level

$$= - \frac{5.84}{n^3} Z^4 \left[\frac{1}{j+1/2} - \frac{3}{4n} \right]$$

This is Dirac's Quantum Mechanical Model to explain fine spectrum of lines.

H_α , H_β and H_γ lines come in the **BAHMER series** of H spectrum

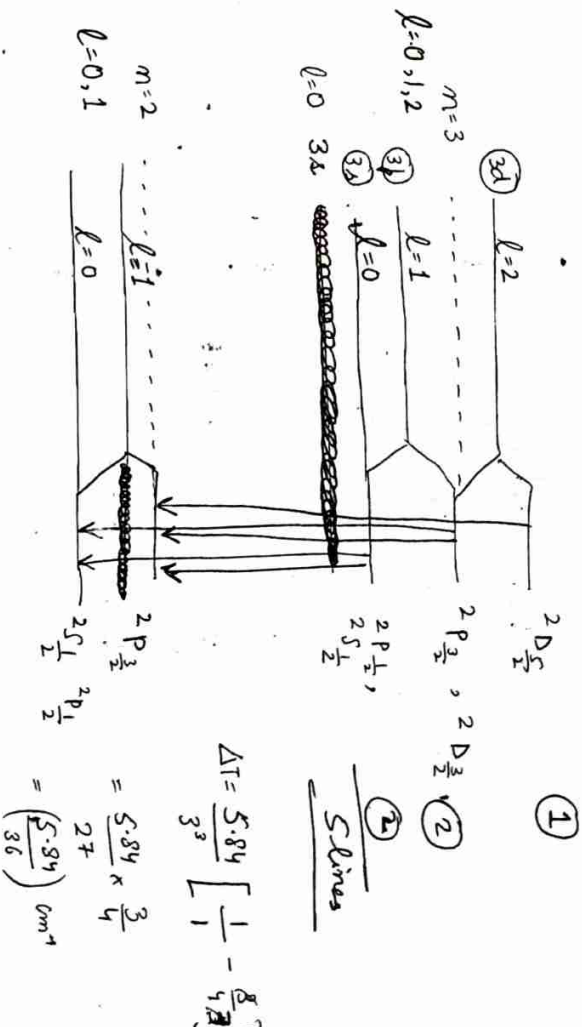
H_α : $n=3$ to $n=2$
 H_β : $n=4$ to $n=2$
 H_γ : $n=5$ to $n=2$



$$\frac{1}{\lambda_{H_\alpha}} = R \left[\frac{1}{2^2} - \frac{1}{3^2} \right] \Rightarrow \lambda_{H_\alpha} = \left(\frac{36}{5R} \right)$$

But when I look at H_α using high resolution spectroscope, I can see 7 lines, not one.

Similarly H_β shows 12 lines that are not explained by Bohr's Model.



According to Dirac's Quantum Mechanical, we can explain 5 lines as ΔT only depends on j and n .

Hence, $2S_{1/2}$ & $2P_{1/2}$ have same ΔT .

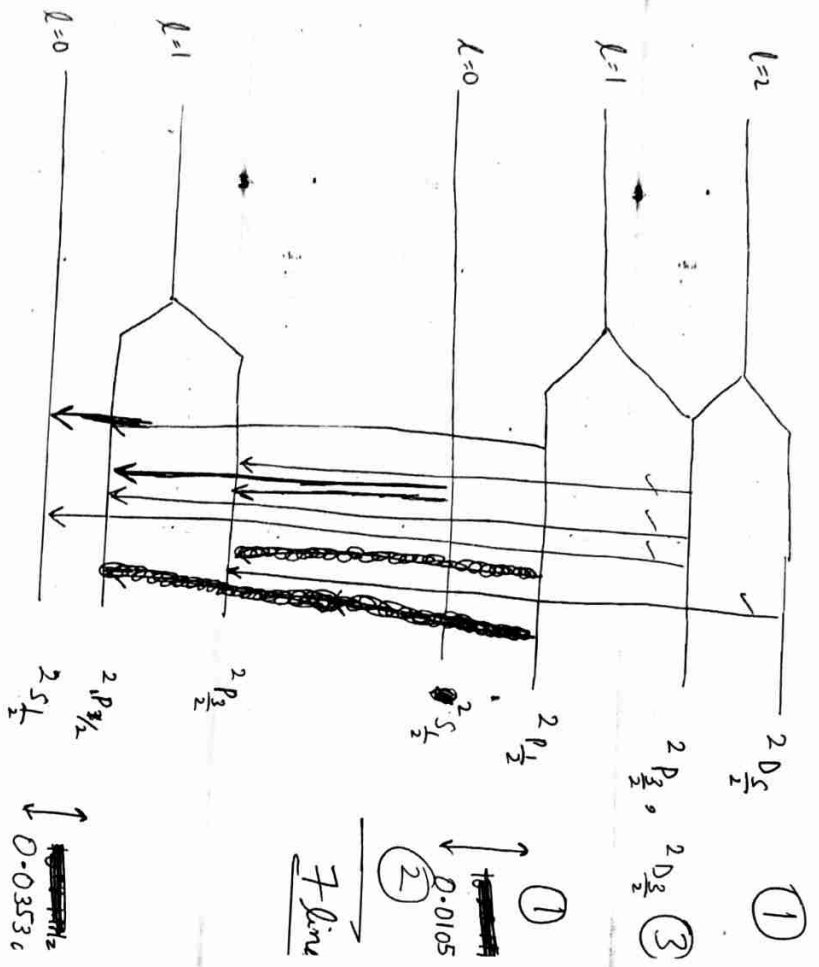
Hence a contradiction?? This was explained by Lamb & Retherford.

Its called LAMB SHIFT.

If says $2S_{1/2}$ and $2P_{1/2}$ are non-degenerate for low l . It does not change energy levels of $2P_{3/2}$ and $3D_{3/2}$.

Now the diagram becomes

Note that $AL \neq 0$ it should be $\neq$



Lamb shift is observed in H like atoms i.e. single e- atoms.

For drawing spectra, EM radiation falls on atom. It interacts with e- in vacuum, and exerts force. Due to this force, interaction of e- with nucleus is weakened. Resulting into removal of degeneracy of lower l values.

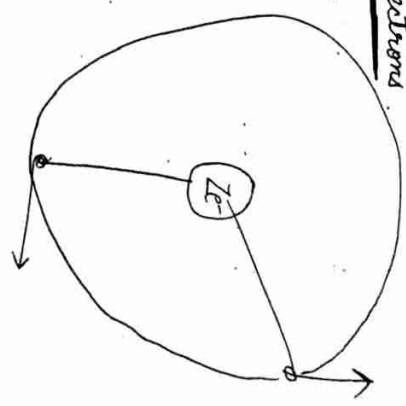
Note that in Lamb shift, the level of $2P_{1/2}$ energy is actually lower than $2S_{1/2}$ energy.

IS and JJ coupling (for many e- atoms)

Phenomenon of many e- atoms. Many e^- means

Many valency electrons

Note that upto now, we were focusing on H-like atoms by the formulae for S-O interaction was devised for single electron atoms



$$\vec{L}_{int} = \vec{L}_1 + \vec{L}_2$$

It's called orbital-orbital interaction: electrostatic interaction

$$H = \frac{p_1^2}{2m} + \frac{p_2^2}{2m} + V(r_1, r_2) \quad \text{Unperturbed Hamiltonian}$$

$L_{max} = |L_1 + L_2|$: lowest energy (due to dmax)
 $L_{min} = |L_1 - L_2|$: highest energy (due to dmin)
 all $\vec{L}$ in same direction $\Rightarrow N+N+N \Rightarrow$ low energy
 $\vec{L} \uparrow$ and other $\vec{L} \downarrow \Rightarrow N+S \Rightarrow$ more energy

$\vec{S}_{spin} = \vec{S}_1 + \vec{S}_2$ It's called spin spin interaction

$$S_{min} = S_1 - S_2 = 0 \quad \text{Max. Energy}$$

$$S_{max} = S_1 + S_2 \quad \text{Min. Energy}$$

Spin-orbit Interaction * Hence more 'operator' implies less energy. $\therefore$ order is $1, 2, 3$ are perturbations in t

✓ lighter Atom: C $1s^2 2s^2 2p^2$: spin orbit interaction is small
 ✓ heavier Atom: spin orbit interaction is high

LS coupling: also called Russell-Saunders coupling

① For lighter atom, where spin-orbit interaction is lowest, spin orbit coupling dominates over other interactions.

$\vec{S} = \vec{S}_1 + \vec{S}_2$
 $|S-S_1| \leq S \leq |S_1+S_2|$

$\vec{L} = \vec{L}_1 + \vec{L}_2$
 $|L_1-L_2| \leq L \leq |L_1+L_2|$

$\vec{L} + \vec{S} = \vec{J}$

$E = \alpha [L \cdot S]$ (LS coupling)

$|L-S| \leq J \leq |L+S|$

Individual e-'s angular momenta couple and form whole angular momentum of atom. It's called J coupling.

2p 3p : Terms due to LS coupling

2p: $L_1=1, S_1=\frac{1}{2}$

3p: $L_2=1, S_2=\frac{1}{2}$

70-100% 0.25 1.00 0.25 1.00 0.25 1.00

$S=0, L=1$
 $\vec{S} \uparrow, \vec{S} \downarrow$
 singlet triplet

BOSON

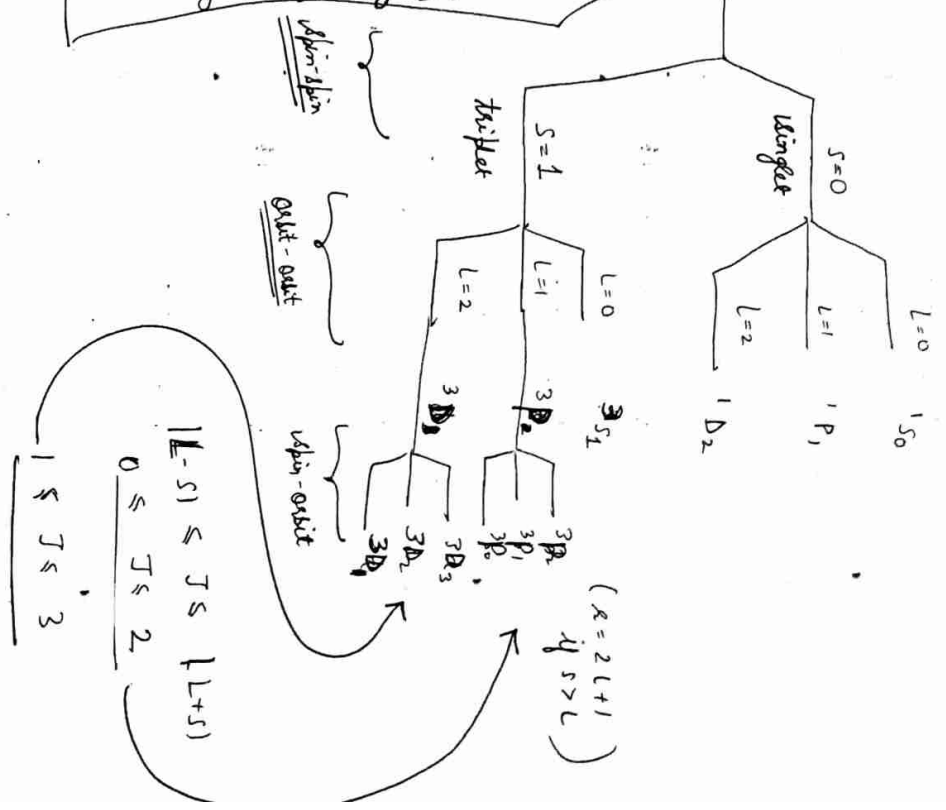
Note that S has value 0 in singlet e- atoms.

$S=\frac{1}{2}$: Fermi

$|S-S_1| \leq J \leq |S_1+S_2|$
 $|L-L_1| \leq L \leq |L_1+L_2|$

$L = l_1 + l_2$
 $0 \leq L \leq 2$
 $|l_1-l_2| \leq L \leq |l_1+l_2|$
 $0 \leq L \leq 2$

(*) Note that total number of possible states of (2p 3p) are $6 \times 6 = 36$. Apply Zeeman Effect and Count.... Final states will be 36 & all will be non-degenerate. For LS coupling, every state has degeneracy = $(2J+1)$.



* In final spin-orbit coupling, order is according to l . Note that here $l=1$ or $l=2$ has no meaning w.r.t. subshell l . It just means addition of m_l 's.

(*) Note that spin-orbit interaction studied for He was a special case of general LS coupling.

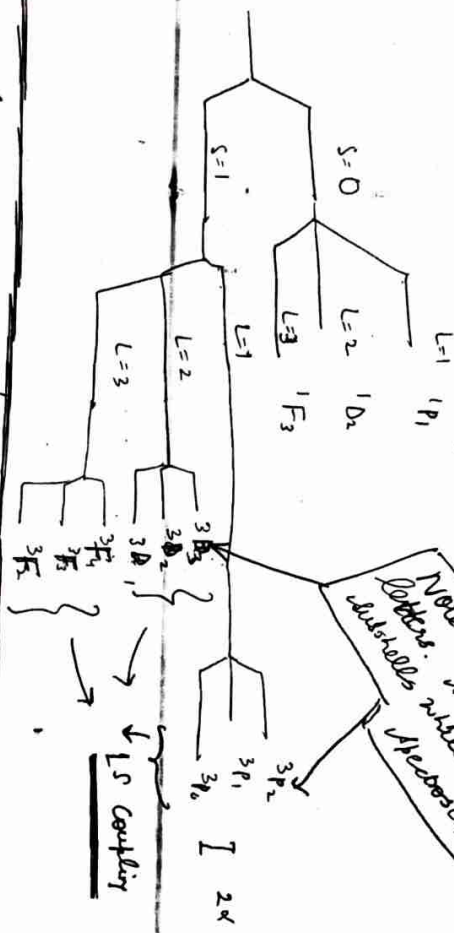
pd

$l_1 = 1 \quad l_2 = 2$

$L = 1, 2, 3$

$s_1 = \frac{1}{2} \quad s_2 = \frac{1}{2}$

$S = 0, 1$



Note that capital letters for subshells while lowercase letters for spectroscopic notations.

Completed subshells
No contribution to L or S
Only contribution from $3p^2$ electrons.

LANDE'S INTERVAL RULE

$E = \alpha L \cdot S$

$= \alpha [J(J+1) - L(L+1) - S(S+1)]$

The relative spacing of the fine structure levels within a multiplet is governed by Lande Interval Rule.

Remain same for successive terms

$E(J+1) - E(J) = \alpha [(J+1)(J+2) - J(J+1)]$

$= 2\alpha [J+1]$

i.e.

$\Delta E \propto (J+1)$

L-S coupling selection rules

$\Delta L = 0, \pm 1$
 $\Delta J = 0, \pm 1$
 $\Delta S = 0$

$\Delta L = 0$ allowed
 $\Delta J = 0$ allowed
 $\Delta S = 0$ allowed

Energy interval on adjoining levels J and $J+1$ of fine-structure multiplets proportional to $(J+1)$. That is to the large of the 2 J values involved

Spectral terms of equivalent electrons due to LS coupling

p^2 : m and l are same for both m_l and m_s will be different to follow Pauli's exclusion principle.

$\vec{S} = s_1 + s_2 = 0, 1$

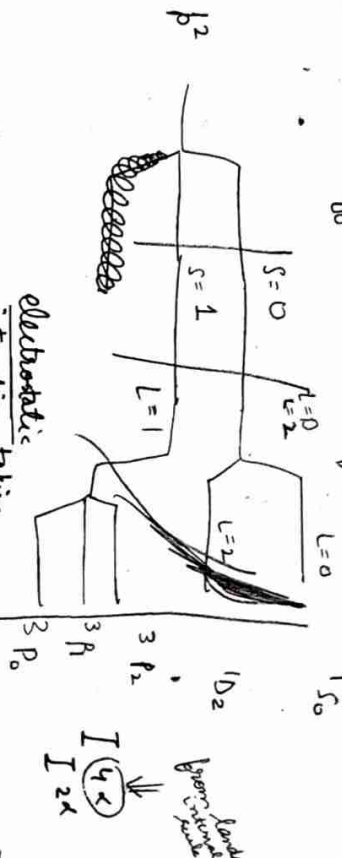
$S=0$: m_s values for both e^- are different

$m_s = (\pm \frac{1}{2})$

$S=1$: m_s values same

$L = l_1 + l_2 = 0, 1, 2$

Along with $S=1$, only that values of L will go that will have different values of m_l .



Multiplet terms of p^2

$p^2 \Leftrightarrow p^4$
 $d^1 \Leftrightarrow d^9$
 $(nd)^2 \Leftrightarrow (nd)^{n-2}$
no. of e in subshells

electronic interaction taking into account Pauli Exclusion Principle
 $L = l_1 + l_2$

$\vec{J} = \vec{L} + \vec{S}$

LS coupling

Total number of possibilities are

$6 \times 6 = 36$

For $2e^-$ in p subshell

Can be seen from $5V$

A&M Physics (4)

J-J coupling

For heavier atoms, for which the spin-orbit (magnetic) interaction term in Hamiltonian predominates over the residual electrostatic interaction and the spin-spin correlation

It occurs for heavier elements when spin-orbit coupling dominates other types of interaction

eg. 4p5d

$$m_l = 4, l_1 = 1, S_1 = \frac{1}{2} \Rightarrow T_1 = \frac{1}{2}, \frac{3}{2}$$

(10-5) to (10+5)

J variation

We cannot write finally common integral J terms as even same J terms are distinct

$$n_0 = 5, l_2 = 2, S_2 = \frac{1}{2} \Rightarrow T_2 = \frac{3}{2}, \frac{5}{2}$$

Each of the four levels is further split by residual electrostatic interaction and spin-spin correlation into a no. of integral J levels.

Combinations

$$\left(\frac{1}{2}, \frac{5}{2}\right)$$

$$\left(\frac{3}{2}, \frac{3}{2}\right)$$

$$\left(\frac{3}{2}, \frac{5}{2}\right)$$

$$J = 0, 1, 2, 3$$

$$J = 1, 2, 3, 4$$

Selection rule for J-J coupling:

$$D \Delta J = 0, \pm 1$$

$$J = 0 \leftrightarrow J = 0$$

And it's NO rule for

ΔS ΔL as they are no longer good quantum numbers

$$|J_1 - J_2| \leq J \leq (J_1 + J_2)$$

After Carbon, atoms typically show relatively seldom. In fact, there is gradual shift from L-S coupling for lighter atoms towards J-J coupling for heavier atoms.

Zemman Effect

Behaviour of atom in external weak/intermediate magnetic field.

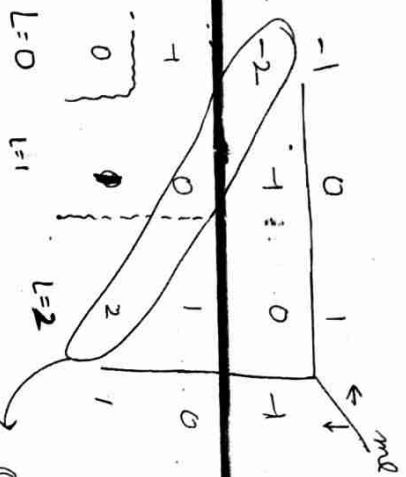
$$E = -\vec{\mu} \cdot \vec{B} = \omega_p J_z$$

If field is external ⇒ Zeeman Effect (spin-orbit interaction)

$$l_1 = 1, l_2 = 1$$

Beit's

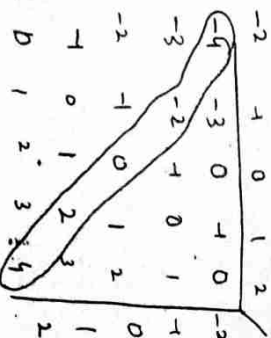
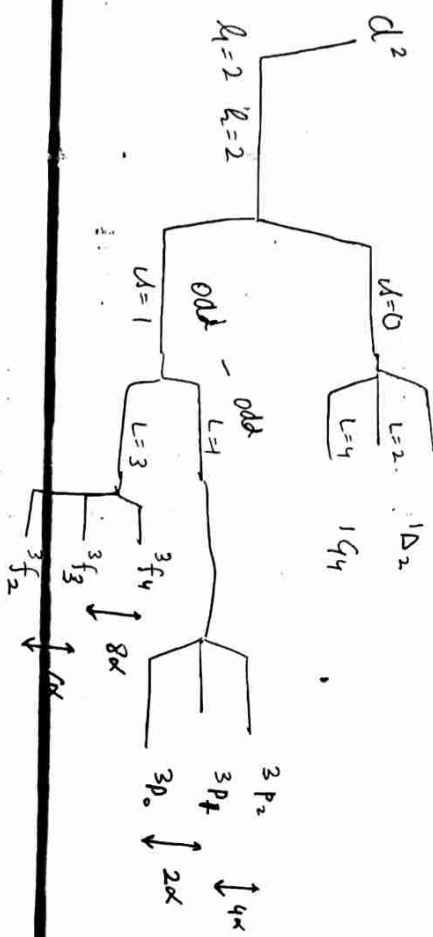
same



Obtained when both have same mJ values. Will not go along with J=1 where mJ have same values.

even-even

1s



Note that in pd or 2p 1s coupling, we could associate all l values with every S. But here we cannot. If we try to associate l=1 and 3 with s=0, in the above example, e- with same quantum states will result.

same quantum states will result.

Angular Velocity of LAMBERT PRECESSION

$$\omega_B = \frac{e}{2m} g B \checkmark$$

$$\Rightarrow E = \frac{e}{2m} g B m_j \hbar$$

$$E = \mu_B g B m_j$$

Note that we have taken vector model i.e. combined L and S as total J.

Note that if field is strong, its called Paschen Back Effect.

Perturbation $\Delta E = \mu_B g B m_j$

Selection Rule comes out to be:

$$\Delta m_j = 0, \pm 1$$

For complete answer, have to mention selection rule.

π component σ components

linearly polarized

Note that spectroscopic notation represented levels. Now, in Zeeman effect, one energy level is split into $(2J+1)$ sublevels.

Transition b/w them will lead to different components.

There are 4 types of Zeeman Effect:

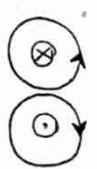
Transverse Zeeman Effect:

Perpendicular $\uparrow$ spectra is viewed perpendicular to direction of field

It will give plane polarized π and σ components

$$\Delta m = 0, \pm 1$$

Longitudinal Zeeman Effect: $\uparrow$ spectra is viewed parallel to direction of applied magnetic field.



It will give circularly polarized components.

RCP and LCP $\Delta m_j = \pm 1$

Normal Zeeman Effect Non Spin Effect: i.e. occurs in atoms where $S=0$ i.e. cannot occur in single e^- atoms.

Atoms should have even electrons (necessary but not sufficient condition). In 1896, Peter Zeeman observed it in Zn and Cd.

It could be transverse or longitudinal depending upon angle of viewing. It is rare.

Anomalous Zeeman Effect: could be both but typically S is non zero.

It is more common and most general case. We will explain this in detail.

Quantum Mechanical Explanation

Both precess about $\vec{B}$

$$\vec{L} \Rightarrow \mu_L = -\frac{e}{2m} \vec{L}$$

$$\vec{S} \Rightarrow \mu_S = -\frac{e}{2m} 2\vec{S}$$

$$\vec{\mu} = \vec{\mu}_L + \vec{\mu}_S = -\frac{e}{2m} (\vec{L} + 2\vec{S})$$

We know $\vec{L}$ and $\vec{S}$ couple to form $\vec{J}$ and $\vec{J}$ precesses about $\vec{B}$. But in strong field, this coupling breaks down and the 2 components $\vec{L}$ & $\vec{S}$ individually precess about $\vec{B}$.

$$\vec{\mu} = -\frac{e}{2m} [(\vec{L} \times \vec{B}) + (2\vec{S} \times \vec{B})]$$

Both will individually precess about $\vec{B}$



$$U = -\vec{\mu} \cdot \vec{B} = \frac{e}{2m} (\vec{L} \cdot \vec{B} + 2\vec{S} \cdot \vec{B})$$

$$= \frac{e\hbar}{2m} (m_l \cos \theta + 2m_s \cos \theta)$$

This is Paschen Back Effect

But Zeeman Effect has:

We are interacting in $\vec{\mu} \cdot \vec{J}$ because in weak fields, $\vec{\mu}_L$ and $\vec{\mu}_S$ are individual non-existent, and therefore, their combination i.e. $\vec{\mu}$ exists.

$$\vec{\mu} = \frac{e}{2m} (\vec{L} + 2\vec{S})$$

L-S coupling: a weak field effect

Note that $\vec{\mu}$ is not parallel to $\vec{J}$ but we are interested in $\vec{\mu}_J$ i.e. component of $\vec{\mu}$ along $\vec{J}$...

In strong field, coupling will break down and $\vec{L}$ and $2\vec{S}$ will individually precess

$\vec{J} = \vec{L} + \vec{S}$: Vector model of atom

μ_J model is appropriate in Zeeman Effect in weak field.

$$\vec{\mu}_J = (\vec{\mu} \cdot \vec{J}) \frac{\vec{J}}{J^2}$$

$$\vec{L} = \frac{\vec{\mu}_J \times \vec{B}}{\mu_J \sin \theta} = \mu_J B \sin \theta$$

$$dU = L d\theta = \mu_J B \sin \theta d\theta$$

$$U = -\vec{\mu}_J \cdot \vec{B} = -(\mu_J)_z B = \mu_B B g m_j$$

$$\vec{\mu}_J = (\vec{\mu} \cdot \vec{J}) \frac{\vec{J}}{J^2} = \frac{-e}{2m} \left[\frac{(\vec{J} + \vec{S}) \cdot \vec{J}}{J^2} \right] \vec{J}$$

$$= \frac{-e}{2m} \left[\frac{J^2 + (\vec{S} \cdot \vec{J})}{J^2} \right] \vec{J}$$

$$= \frac{-e}{2m} \left[1 + \frac{(\vec{S} \cdot \vec{J})}{J^2} \right] \vec{J}$$

$$\begin{aligned} \vec{L} &= \vec{J} - \vec{S} \\ \vec{L} \cdot \vec{L} &= (\vec{J} - \vec{S}) \cdot (\vec{J} - \vec{S}) \\ \Rightarrow \vec{S} \cdot \vec{J} &= \frac{J^2 + S^2 - L^2}{2} \end{aligned}$$

due to L-S coupling

Note that $\vec{\mu}$ is not antiparallel to $\vec{J}$, but rather we are interested in only antiparallel component of $\vec{\mu}$ i.e. $\vec{\mu}_J$ i.e. the component along $\vec{J}$.

Now all are quantized

$$\Rightarrow J^2 = j(j+1) \hbar^2$$

$$S^2 = s(s+1) \hbar^2$$

$$L^2 = l(l+1) \hbar^2$$

Refer to derivation

P-205, 206

$$= \frac{-e}{2m} \left[1 + \frac{j(j+1) + s(s+1) - l(l+1)}{2j(j+1)} \right] \vec{J}$$

$$\vec{\mu}_J = \frac{-e}{2m} g_j \vec{J}$$

$$g_j = \frac{1 + \frac{j(j+1) + s(s+1) - l(l+1)}{2j(j+1)}}{2j(j+1)}$$

* The importance of the g factor is that it directly gives the

$$\Rightarrow \vec{\mu}_L = -\frac{e}{2m} g_L \vec{L}$$

$$\vec{\mu}_S = -\frac{e}{2m} g_S \vec{S}$$

Now, $\vec{J} = \vec{L} + \vec{S}$

$$\Rightarrow \vec{\mu}_J = -\frac{e}{2m} g_J \vec{J}$$

$$g_J = \frac{1}{2} \left[\frac{J(J+1) + S(S+1) - L(L+1)}{J(J+1)} \right]$$

$$4D_{\frac{1}{2}} \Rightarrow g = 0$$

for purely orbital motion, $S=0$ & $J=L$

$$\Rightarrow g_J = 1 + \frac{0}{L(L+1)} = 1$$

for purely spin motion, $L=0$ & $J=S$

$$g_J = 1 + \frac{2S(S+1)}{2S(S+1)} = 2$$

If $\vec{J}$ precesses $\Rightarrow \vec{\mu}_J$ will precess as $\vec{\mu}_J$ is antiparallel to $\vec{J}$.

$\vec{J}$ precesses about $\vec{B} \Rightarrow \vec{\mu}_J$ precessing about $\vec{B}$

and

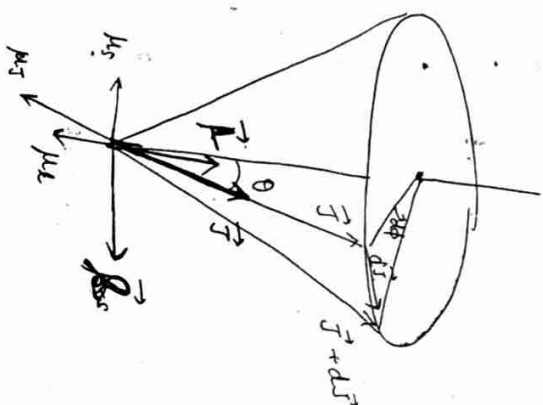
$$J_z = m_j \hbar$$

$$E = \mu_B J_z$$

$$= \frac{e}{2m} g B J_z$$

$$= \left(\frac{e\hbar}{2m} \right) g_J m_j$$

$$E = \mu_B B (g_J m_j)$$



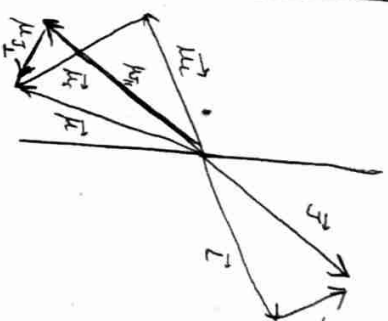
$$E = -\vec{\mu}_J \cdot \vec{B} = \frac{e}{2m} g_J B J_z$$

$$= \frac{e\hbar}{2m} g_J B m_j$$

$$E = \mu_B B (g_J m_j)$$

Normal Zeeman Effect

we assume $\vec{\mu}_L$ and $\vec{\mu}_S$ are not $\Delta E = \vec{\mu}_J \cdot \vec{B}$. But we have taken so. In.



$\vec{\mu} \neq \vec{J}$ because

$$\vec{\mu} = \frac{e}{2m} [\vec{L} + 2\vec{S}]$$

$$\vec{J} = \vec{L} + \vec{S}$$

But, $\vec{L}, \vec{S}, \vec{\mu}_L, \vec{\mu}_S$ and therefore $\vec{\mu}$ all precess about $\vec{J}$.

($\vec{J}$ in turn precesses about $\vec{B}$)

$\Rightarrow \vec{\mu} = \vec{\mu}_{J\parallel} + \vec{\mu}_{J\perp}$
The $\vec{\mu}_{J\perp}$ averages to zero due to the precession motion
 $\Rightarrow \Delta E = \vec{\mu}_{J\parallel} \cdot \vec{B}$

Normal Zeeman Effect

$$E = \mu_B B \quad g = 1$$

$$\Delta E = \mu_B B \quad \Delta m_l$$

$$\Delta m_l = 0, \pm 1$$

(if $s=0$)
i.e. $g = g_L = 1$

Normal Zeeman Effect

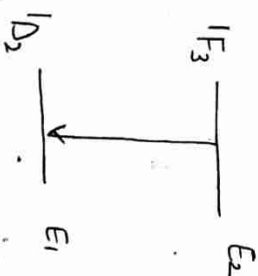
3 components

Longitudinal Normal Zeeman Effect: 2 circularly polarized components

Anomalous Zeeman Effect: 1 line is split into several components

Draw Zeeman pattern of $1F_3 \rightarrow 1D_2$

Now in presence of B , it will split into $2J+1$ sublevels.



$$(2L+1) = 0 \Rightarrow L=0$$

Hence Normal Zeeman Pattern

$$\Rightarrow J=L=3 \text{ for } 1F_3$$

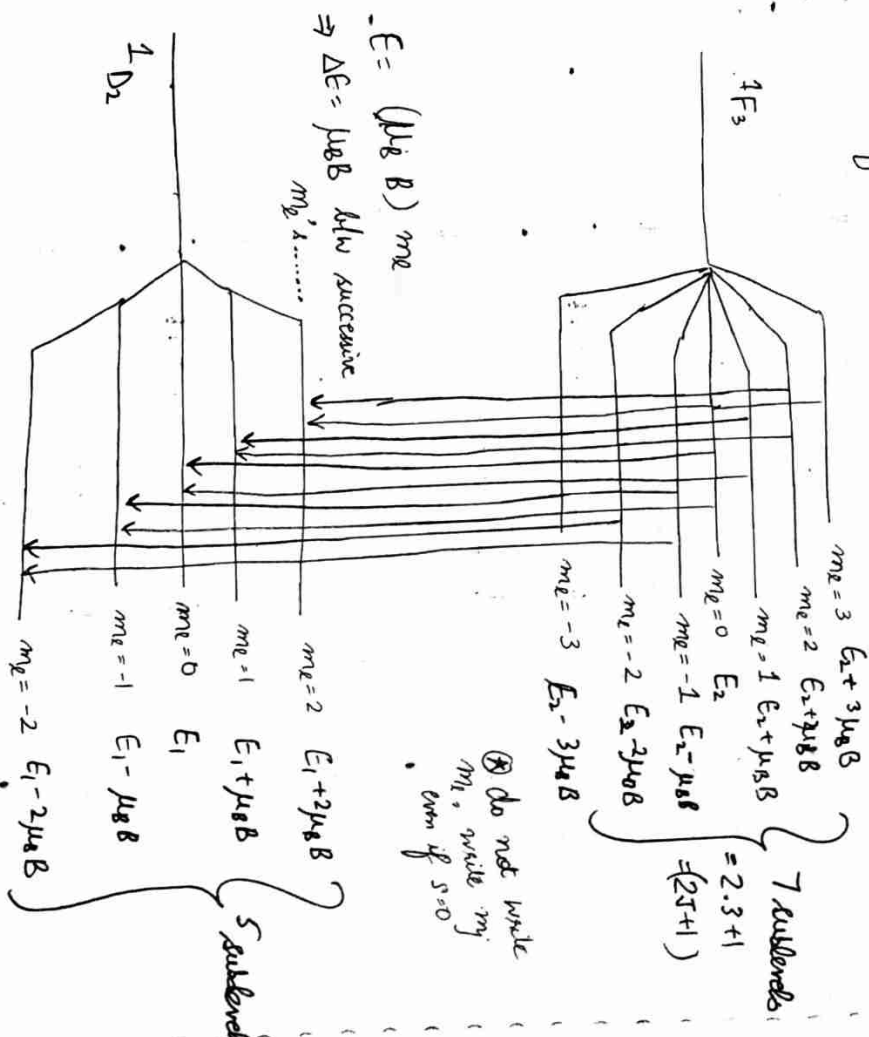
$$J=L=2 \text{ for } 1D_2$$

π Component: E parallel to External Magnetic Field

σ Component: E perpendicular to External Magnetic Field

Intensity ratios are derived from Correspondence principle.
i.e. σ Component together have same intensity as π Component.

Energy level will split into $2J+1$ levels in presence of external field.



$$\Delta m_l = 0, \pm 1$$

Note that $m_l = 0 \rightarrow m_l = 0$ is allowed

For all red lines, $\Delta E = (E_2 - E_1)$, hence only 1 line in spectrum. It corresponds to $\Delta m_l = 0$

Green: $m_l - m_l = -1$: It has minimum energy.

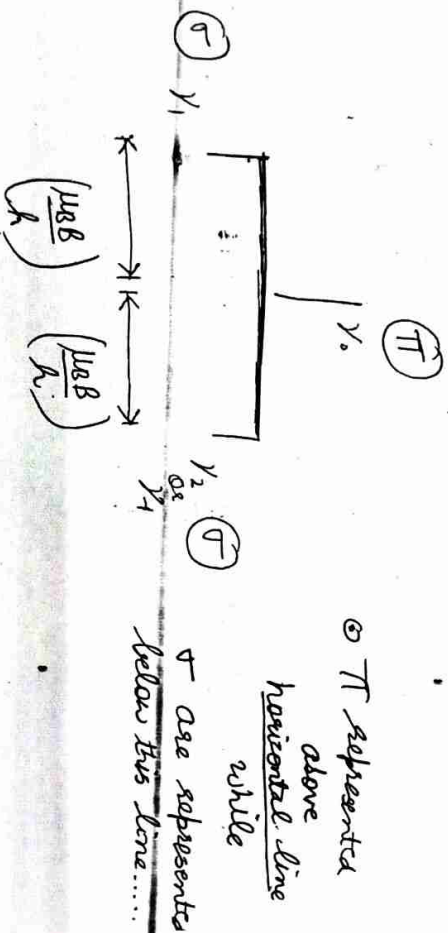
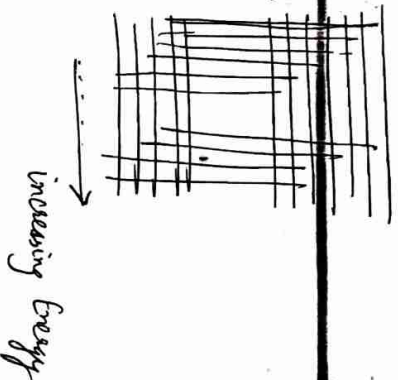


① In exam ~~also~~ also more energy lines on right and less energy lines on left.

$$\gamma_0 = \frac{\Delta E}{h} = \left(\frac{E_2 - E_1}{h} \right)$$

$$\gamma_1 = \frac{\Delta E}{h} = \frac{E_2 - E_1}{h} + \frac{\mu_B B}{h}$$

$$\gamma_2 = \frac{\Delta E}{h} = \frac{E_2 - E_1}{h} - \frac{\mu_B B}{h}$$



② π represents above horizontal line while

σ represents below this line...

In longitudinal, γ_0 will be missing

$$\text{Zeeman shift} = \frac{\mu_B B}{h} = \left(\frac{eB}{4\pi m} \right)$$

③ Note that irrespective of no. of transitions, there are 3 components in Zeeman effect, where $S=0$.

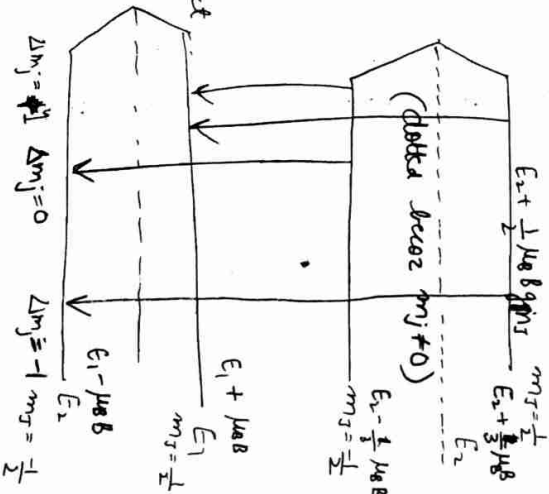
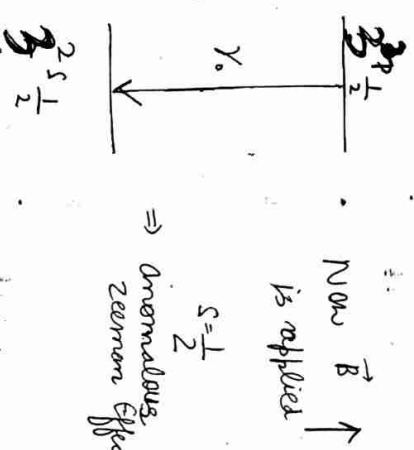
We can calculate B from the Zeeman shift. Anomalous Zeeman Pattern

Q Zeeman Effect of Sodium lines D_1 and D_2

D_1 : 4 components D_2 : 6 components

$$^2P_{1/2} \rightarrow ^2S_{1/2} \quad ^2P_{3/2} \rightarrow ^2S_{1/2}$$

D_1 line



- Now note that

$$E = -\mu_B B (g m_J)$$

We have to take g into account for both levels

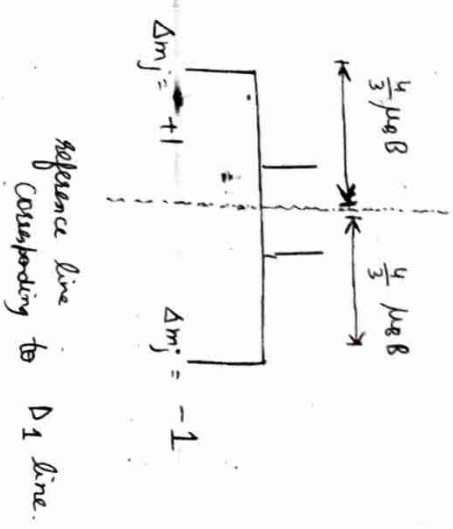
Hence 2 different components for $\Delta m_J = 0$

$$g(^2S_{1/2}) = 2$$

$$g(^2P_{1/2}) = \frac{1}{4} \cdot \frac{3}{2} + \frac{1}{2} \cdot \frac{3}{2} - 1 \cdot 2 = 1 - \frac{1}{2} = \frac{1}{2}$$

$$= 1 + \frac{3}{4} + \frac{3}{4} - 2 = 1 - \frac{1}{2} = \frac{1}{2}$$

$$\gamma_1 = \gamma_0 - \frac{4}{3} \mu_B$$



D₁: $\Sigma \rightarrow \Sigma$ line
D₂: $\Sigma \rightarrow \Sigma$ line

★

$$\gamma_1 = \gamma_0 - \frac{4}{3} \frac{\mu_B}{h} \quad \sigma \quad \Delta m = +1$$

$$\gamma_2 = \gamma_0 - \frac{2}{3} \left(\frac{\mu_B}{h} \right) \quad \pi \quad \Delta m = 0$$

$$\gamma_3 = \gamma_0 + \frac{2}{3} \left(\frac{\mu_B}{h} \right) \quad \sigma \quad \Delta m = -1$$

$$\gamma_4 = \gamma_0 + \frac{4}{3} \left(\frac{\mu_B}{h} \right) \quad \sigma \quad \Delta m = -1$$

★

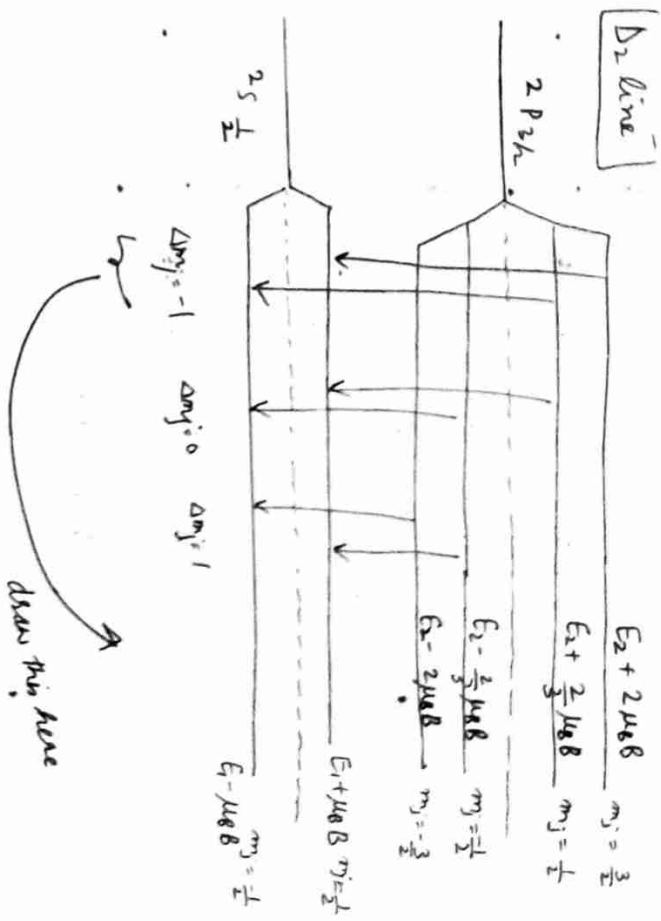
selection rule for Zeeman:

$$\Delta m = 0, \pm 1$$

$$M_j = 0 \quad \leftarrow \rightarrow \quad M_j = 0 \quad \text{if} \quad \Delta J = 0$$

eg. $3P_1 \rightarrow 3S_1$ transition

D₂ line



$$g(2P_{3/2}) = 1 + \frac{\frac{3}{2} \cdot \frac{5}{2} + \frac{1}{2} \cdot \frac{3}{2} - 1 \cdot 2}{\frac{15}{4}} = 1 + \frac{10}{15} = 1 + \frac{2}{3} = \left(\frac{4}{3} \right)$$

$$1 + \frac{\frac{15}{4} + \frac{3}{4} - \frac{8}{4}}{\frac{15}{4}} = 1 + \frac{10}{15} = 1 + \frac{2}{3} = \left(\frac{4}{3} \right)$$

★ Note that selection rules do not absolutely prohibit transitions that violate the rule, but only make such transitions very unlikely.

★ Spin has no classical analogue. In a purely quantum context and any spatial motion is accounted as a part of orbital angular momentum. If e^- is considered as a charged body rotating about its own axis, then our magnitude of momentum (angular) would demand the equivalent velocity of turning electron to be much greater than c . i.e. $v_{eq} \gg c$ which is not possible.

* Spectroscopy is that branch of Physics which deals w. the observation and interpretation of radiation emitted and absorbed by atoms and molecules, thereby, throwing light on their structure.

* Rydberg - Ritz Combination Principle says that the wave number of spectral lines can be expressed by the difference of spectroscopic terms in such a way that other differences of these terms also give the wave numbers of lines in the same spectrum.

* Allowed transitions are governed by Laporte's rule which says that in an electric dipole transition, the parity of the state of the atom must change.

Proof Transition Rate for x-component of dipole moment = $\langle \psi_f | A | \psi_i \rangle$

$$= \int_{-\infty}^{\infty} \psi_f^* \left(\sum_j e_j x_j \right) \psi_i dx$$

Let ψ_i be even
Now if $\sum e_j x_j$ is odd function

$\Rightarrow$ if ψ_f is even $\Rightarrow$ integrand = 0

$\Rightarrow \psi_f$ is odd

• And vice versa.

Hence Parity will undergo change.

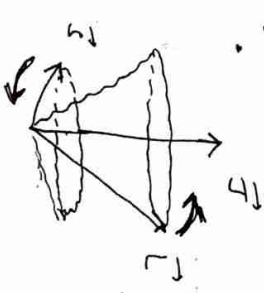
* $\vec{L}$ can never be aligned exactly parallel to $\vec{B}$ as pre1 is always less than $\sqrt{L(L+1)}$

* It has been concluded from experimental data that the gyromagnetic ratio for electron spin is twice the corresponding ratio for orbital angular momentum i.e. $\frac{\mu_s}{\mu_L} = 2 \left(\frac{m_s}{m_L} \right) \Rightarrow g_s = 2 g_L$ ("e" is 1)

* Spin Orbit Interaction

Angular Momenta of atomic electron, $\vec{L}$ and $\vec{S}$ interact magnetically, which is known as spin orbit interaction.

They exert torques on each other. These internal torques do not change the magnitudes of the vectors $\vec{L}$ and $\vec{S}$ but cause them to precess about their result $\vec{J}$. If atom is in free space and no external torque acts, then total angular momentum $\vec{J}$ remains conserved.



$$J^2 = L^2 + S^2 + 2(\vec{L} \cdot \vec{S})$$

Spin Orbit Interaction, together with relativistic correction, explains the Hydrogen Fine structure.

Line Shift Phenomenon is also observed in fine structure

* Energy levels expressed in wave numbers are called ter

* The "state" of an atom is the condition of motion of all the electrons. It is specified by listing four quantum numbers of each electron.

* Energy level: A collection of states having the same energy, in the absence of external magnetic or electric field constitutes an 'energy level'.

* Energy sublevel: An external field splits one energy level into several sublevels, each characterized by one or more magnetic quantum numbers.

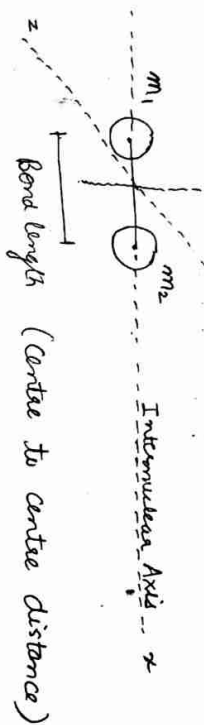
* Multiplet is a collection of spectral lines or transition between two terms, eg 6 components

Molecular Physics (1)

24/02/2012

ROTATIONAL SPECTRA

• We study diatomic molecules and generalize the results.



Asymmetric Molecules : $I_{xy} = I_{yz}$

Asymmetric Molecules : $I_{xy} \neq I_{zz}$

Spherical Top : $I_{xx} = I_{yy} = I_{zz}$

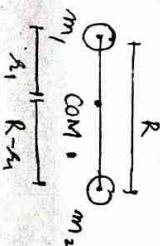
Rigid rotator molecules : 'r' (Bond length) is fixed i.e. no stretching along internuclear axis.
 1 about internuclear axis is negligible.
 Rotation is possible only along line passing through Centre of Mass, and $\perp$ to internuclear axis.

Rotating Cores

Now we know,

$m_1, R_1 = m_2 (R - R_1)$

$R_1 = \left(\frac{m_2 R}{m_1 + m_2} \right) \Rightarrow R - R_1 = \left(\frac{m_1 R}{m_1 + m_2} \right)$



$\bar{O} = U'_e + G(v) + F(v) - [U'_e + G(v) + F(v)]$
 Remember the notations

Thank God, let's fix

For rigid rotator,

$I_{CM} = m_1 R_1^2 + m_2 (R - R_1)^2 = \frac{m_1 m_2 R^2}{(m_1 + m_2)} + \frac{m_2 m_1^2}{(m_1 + m_2)}$

$= \left(\frac{m_1 m_2}{m_1 + m_2} \right) R^2$

$I_{CM} = \mu R^2$ R: Bond length

$J_{CM} = \mu R^2 \omega = I_{CM} \omega$

$E = \frac{J^2}{2I} = \frac{(I_{CM} \omega)^2}{2 I_{CM}} = \frac{1}{2} I_{CM} \omega^2$

• No matter due to what reason, J is created, it will quantum mechanically take values corresponding to

$\sqrt{j(j+1)} \hbar$ ($j=0, 1, 2, \dots$)

$E_j = \frac{J^2}{2I_{CM}} = \frac{[j(j+1)\hbar]^2}{2I_{CM}} = \left(\frac{\hbar^2}{8\pi^2 I_{CM}} \right) j(j+1)$ Zero Energy

We usually define Energy in wave numbers as J_{CM} Numbers i.e.

$F(J) = \frac{E(J)}{hc} = \frac{h}{8\pi^2 I_{CM} c} j(j+1)$

"Born - Oppenheimer" Approximation has been used, to solve J values quantum mechanically for rigid rotator

Now $F(J)$ can have values 0, 2B, 6B, 12B, 20B
hence discrete energy spectrum.

Now due to what property of molecule rotation occurs??
Note every molecule rotates. H_2 , N_2 , O_2 molecules i.e. (homonuclear) diatomic molecules do not give rotational or vibrational spectra or rotational-vibration spectra.

Only heteronuclear molecules possess rotation.
Heteronuclear molecules possess dipole moment. e.g. HCl , HBr , CO

Presence of permanent dipole moment is reason for rotation.
When it interacts with incident source, it interacts with $\vec{E}$ and torque $= (\vec{p} \times \vec{E})$ acts.

For rigid rotator molecule, selection rules of permanent dipole moment, give: $\rightarrow$ This is known by the study of matrix elements of dipole transition

Rigid Rotator Molecules: $\Delta J = \pm 1$
Non Rigid Rotator: $\Delta J = \pm 1$ (more)

For (Non Rigid Rotator)

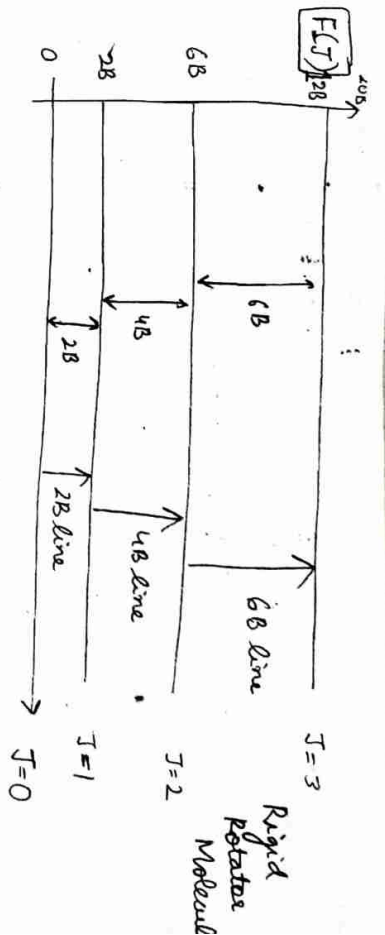
$$\frac{J_y^2}{2I_y} + \frac{J_z^2}{2I_z} \rightarrow$$

$$E = j(j+1)B - D[j(j+1)]^2$$

$\uparrow$ centrifugal distortion coefficient [CDC]
only this term in rigid rotator

Order of B $\approx 10 \text{ cm}^{-1}$
Order of D $\approx \frac{1}{1000} \text{ cm}^{-1}$

Centrifugal distortion effect for non rigid rotator is



$\rightarrow$ Rotational levels are increasing gaps
Order of B is 0.001 eV

Rotational spectra are typically observed in absorption and in emission spectra. In emission spectra, it will come if initially, there is sufficient population in excited level.

$\circ$ Remember more $|J|$ means more energy

$$\Delta J = +1 \text{ (absorption)}, -1 \text{ (emission)}$$

Depending upon energy of transition oscillator, the various jumps can occur. [Microwave source]

All frequencies of transition oscillator are not easily available.

$$\because \Delta J = \pm 1 \Rightarrow \Delta E = j(j+1)B - j(j-1)B = j(2) = 2jB$$

$$\Rightarrow \Delta E = 2jB$$

$$\Rightarrow \Delta(\text{spectral line}) = 2B$$

uniform gaps in rotational spectra of diatomic rigid rotator molecules.

→ If gap is not uniform ⇒ ~~non rigid rotator~~ ~~non rigid rotator~~ molecule

If uniform gap ⇒ rigid rotator (nature of molecule)

$$\Rightarrow B = \frac{h}{8\pi^2 c I_{cm}}$$

$$\Rightarrow I_{cm} = \frac{h}{8\pi^2 c B} \quad (\text{Moment of Inertia})$$

$$\Rightarrow \mu R^2 = I_{cm} \quad (\text{Bond length})$$

$$\Rightarrow R = \sqrt{\frac{I_{cm}}{\mu}}$$

there 3 things were calculated from seeing gap.

$$H^1 Cl^{35.5} \quad H^2 Cl^{35.5}$$

$$\mu = \frac{1 \times 35.5}{36.5} \quad \mu = \frac{2 \times 35.5}{37.5}$$

$$\text{If } \mu \uparrow \Rightarrow B \downarrow$$

⇒ from shift in B, we can obtain information about isotope. This is called isotope effect.

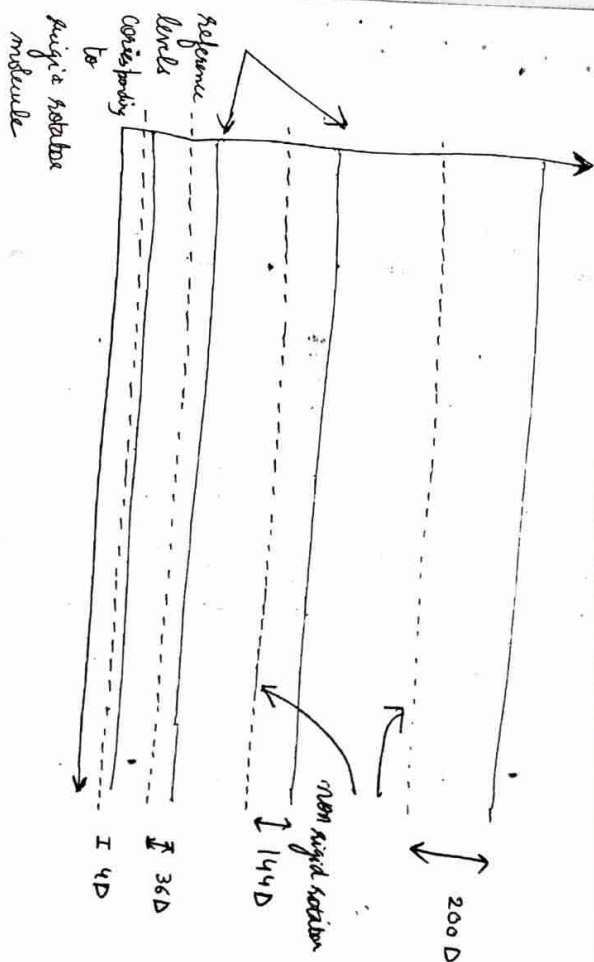
⇒ 4th information we get is Isotope Effect from gap.

For ~~rigid rotator~~ (Non Rigid Rotator)

$$F(J) = B J(J+1) - D(J+1)^2(J+1)^2$$

$$F(0) = 0 ; F(1) = 2B - 4D ; F(2) = 6B - 36D ;$$

$$F(3) = 12B - 144D ; F(4) = 20B - 400D$$



$F(0) = 0$	$\Delta J = 1$	$\Delta J = 2$
$F(1) = 2B - 4D$	$2B - 4D$	$6B - 36D$
$F(2) = 6B - 36D$	$4B - 32D$	$10B - 140D$
$F(3) = 12B - 144D$	$6B - 108D$	$14B - 364D$
$F(4) = 20B - 400D$	$8B - 256D$	

gap is non uniform

non rigid rotator molecule

If 3 lines are given ⇒

$$2B - 4D : \text{known}$$

$$4B - 32D : \text{known}$$

$$x_1$$

$$x_2$$

$$x_3$$

Q8) CO : $J=0 \rightarrow J=1$

$F(J) = B J(J+1)$
 $F(0) = 0$
 $F(1) = 2B$
 $2Bc = 1 \text{ Hz}$

For the curve that has no minima, it corresponds to unstable molecule in which there is repulsion between the 2 nuclei at all separations.

VIBRATIONAL SPECTRA

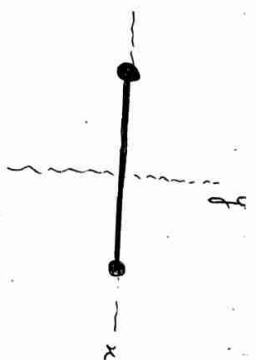
In non homogeneous electric field, permanent dipole moment experiences

$\vec{F} = (\vec{p} \cdot \vec{\nabla}) \vec{E}$

$\Rightarrow$ Oscillatory force due to oscillating $\vec{E}$.
 $\Rightarrow$ vibration

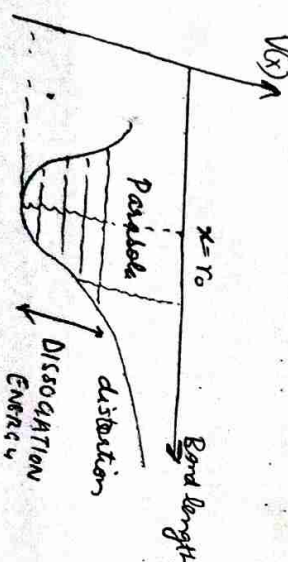
Molecule behaves like simple harmonic oscillator.

~~For rigid rotator molecule, not vibration along internuclear axis. Only vibrational motion along direction perpendicular to internuclear axis passing through centre.~~



$F_y = -ky$
 $V = \frac{1}{2}ky^2$ (Parabolic Potential)

But molecules may not be simple harmonic oscillator

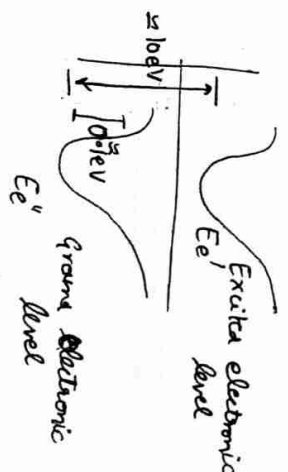


Potential for diatomic molecule
 Combination of Parabolic & Aberratic curves.

$V(x) = \frac{1}{2}Fx^2 - \frac{1}{3}gx^3$
 $x = r - r_0$

Dissociation Energy: Amount of energy required to dissociate the bonds.

Note that we are not changing electronic levels. The parabolic curve drawn is for a particular electronic configuration.



E'' : double prime for ground state electron
 E' : single prime for excited electron
 Assume the primes like basis of

In vibrational spectra, we are not interested in excitation of electronic levels, we are not providing that much energy.

Harmonic Oscillator

In lower levels, $V(x) = Fx^2$

At higher levels: $V(x) = -gx^3$ [ignoring it for simplicity analysis]

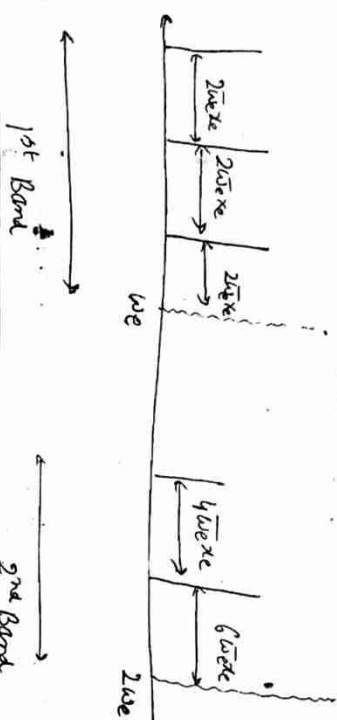
$V = \frac{1}{2}kx^2$ Quantum

$E(v) = (v + \frac{1}{2}) \hbar \omega_e$

wavenumber $\tilde{\omega} = \frac{1}{2\pi c} \sqrt{\frac{k}{m}}$
 $\Rightarrow \tilde{\omega} = \frac{1}{2\pi c} \sqrt{\frac{k}{m}}$

$\tilde{\omega} = \frac{1}{2\pi c} \sqrt{\frac{k}{m}}$
 $\tilde{\omega} = \left(\frac{2\pi}{c}\right)$

- Atomic spectra: line spectra
Molecular spectra: Band spectra



Classical frequency for a harmonic oscillator is: $\nu_{\text{osc}} = \frac{1}{2\pi} \sqrt{\frac{k}{\mu}}$. For anharmonic oscillator, this frequency decreases as v (quantum no.) increases.

Vibration spectra occurs in IR region.....

★ Born-Oppenheimer Approximation

Quantum Mechanical Hamiltonian operator for a molecule, containing terms for electronic & nuclear motion is very complex. In order to solve Schrödinger's equation, Born-Oppenheimer approximation is used, according to which:

(i) first, due to slow moving massive nuclei, the wave equation for electronic motion alone is solved, in which nuclear coordinates appear simply as parameters.

(ii) then, the wave equation for motion of nuclei alone is solved in which electronic eigenvalues of electronic motion occurs as part of potential energy function of nuclear motion.

(iii) complete wave function = $\psi_e \psi_n$

★ Some Characteristics:

① Electronic spectra in emission as well as absorption. Vibrational Rotational & Rotational in absorption.

② Electronic spectra band has a sharp edge, called

Ground structure of vibrational rotational

③ Each vibrational rotational band consists of an intense band called Fundamental Band, which is accompanied by a few weak bands.

④ Electronic: homonuclear as well as heteronuclear. Vibrational rot, rotational: heteronuclear only.

★ R_e : equilibrium

② $R < R_e$: more nuclear repulsion
③ $R > R_e$: ~~more~~ electronic attraction

★ Zero Point energy for rotational spectra = 0
for vibrational spectra = $\frac{1}{2} \bar{\nu}_0$ $\nearrow$ increases in both cases

★ Molecular Spectra

All the spectral lines arising from transitions between rotational levels J' and J'' associated with a given pair of vibrational levels v' and v'' of a given pair of electronic levels form a band.

Thus for a given band, J' and J'' vary from line to line while v' and v'' are constants.

All the bands which arise from transitions between vibrational levels of a given pair of electronic levels constitute a band system, the values of v' and v'' varying from band to band.

Thus, a single electronic transition gives rise to a band system. Each line in each band of the system arises due to a change in all the three energies E_e, E_v & E_r .

$$\bar{E}_0 = \bar{E}_e + \bar{E}_v + \bar{E}_r$$

For a given band, $\bar{E}_e$ and $\bar{E}_v$ are constants, while $\bar{E}_r$ changes from line to line. Position in band where $\bar{E}_r = 0$ is called Band Origin, its wave number is $(\bar{\nu}_e + \bar{\nu}_v)$.

For a band system similar, $\bar{E}_e$ is const, while $\bar{E}_v$ varies from band to band. Position in the system where $\bar{E}_v = 0$ is called System Origin its wave number is $\bar{\nu}_e$.

For vibrational motion,

$\Delta v = +1$ Band :- Most intense line is $v=0$ to $v=1$

i.e. $\bar{\omega}_1 = \bar{\omega}_e (1-2x_e)$

$\Delta v = +2$ Band

At room temperature, usually $v''=0$ is seen. But for heavier molecules like I_2 , $\bar{\omega}_e$ is less $\Rightarrow v''=0,1,2$ can be seen at room temperature

Population is governed by Maxwell Boltzmann Statistics.

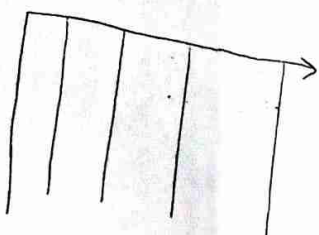
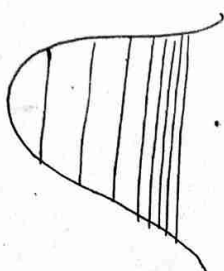
$$N_1 = N_0 e^{-\left(\frac{\Delta E}{kT}\right)}$$

ΔE : Energy $3/k$

$v=0$ to $v=1$ transition

In certain liquid molecules, where rotation is prohibited due to their molecular structure, only then pure vibration occurs. Otherwise normally, vibration & rotation both occur if energy excitation equivalent to vibrational levels is exact when the molecules.

We can also say with 0.1 eV gap, many 0.001 eV gaps of energy can be included. Hence within 1 vibration energy gap, there can be many rotational energy gaps.



In vibration,

gap $\downarrow$

Non-rotation gap $\uparrow$ with increase in quantum number J

with increase in quantum number J as difference in spectra of

Vibration - Rotation Spectra

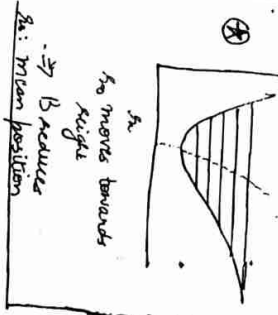
Molecules having permanent dipole moment interact in Electric Field and undergoes vibration as well as rotational motion. Combined effect is

$$G(v, J) = G(v) + F(J)$$

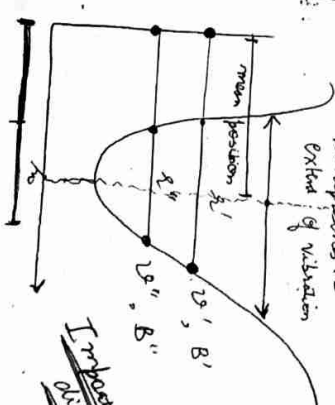
$$= \left[\left(v + \frac{1}{2} \right) \bar{\omega}_e - \left(v + \frac{1}{2} \right)^2 \bar{\omega}_e x_e \right] + B J(J+1)$$

B: Rotational constant $= \left(\frac{h}{8\pi^2 c \mu r^2} \right)$ in vibrational level v

The infrared spectra of a molecule like HCl consists of intense band known as fundamental band, accompanied by no. of weak bands called overtone at wave numbers approx. double, triple, etc. of fundamental band.



$\Rightarrow$ No. moves towards right
 $\Rightarrow$ B reduces
 $\Rightarrow$ mean position



Also the mean No. moves towards right with increase energy levels

intermolecular separation, x

In increasing vibrational state, intermolecular distance increases. Always difference is not very significant

Now

$$r' > r''$$

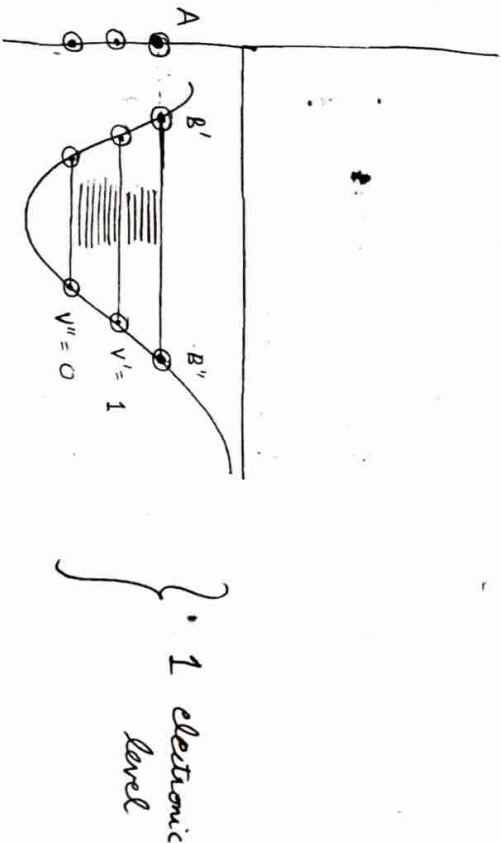
$$B' < B''$$

$$B' - B'' < 0$$

$\Rightarrow B' - B''$ is small negative quantity.

Molecule is excited with every excited molecular state vibrations

Now spectral line will come when $\Delta G(v, J)$ is non zero.



Note that rotational energy starts from zero. i.e. zero. Point Energy = 0. But in vibrational motion, zero point energy $\neq 0$. Many rotational energy are indicated in this level.

Using ground state of vibrational - rotational level

$$G(v'', J'') = \left(v'' + \frac{1}{2}\right) \bar{\omega}_e - \left(v'' + \frac{1}{2}\right)^2 \bar{\omega}_e x_e + B'' J''(J''+1)$$

$$G(v', J') = \left(v' + \frac{1}{2}\right) \bar{\omega}_e - \left(v' + \frac{1}{2}\right)^2 \bar{\omega}_e x_e + B' J'(J'+1)$$

Excited state

- For same electronic level, $\bar{\omega}_e, x_e$ is same
- For same vibrational level, B is same

$$\bar{\omega}_{v', v''} = \Delta G(v', J') - G(v'', J'')$$

$$\bar{\omega}_{v', v''} = (v' - v'') \bar{\omega}_e - (v' - v'')(v' + v'' + 1) \bar{\omega}_e x_e + B' J'(J'+1) - B'' J''(J''+1)$$

Rigid Rotator Diatomic Molecule

$$\Delta v = v' - v'' = 1$$

$$\Delta J = \pm 1$$

(absorption)
(-1 for emission)

For Non-rigid rotator Molecule

$$\Delta v = \pm 1, \pm 2$$

$$\Delta J = \pm 1$$

Note that $\Delta J = 0$ is not observed !!

$$\text{For } \Delta v = 1$$

* We normally study the spectra for normal absorption case $\Delta v = 1$

$$\bar{\omega}_{v', v''} = \bar{\omega}_e - 2(v'' + 1) \bar{\omega}_e x_e + B' J'(J'+1) - B'' J''(J''+1)$$

$$\text{Specific case } v'' = 0 \text{ to } v' = 1 : \bar{\omega}_{v', v''} = \bar{\omega}_e - 2 \bar{\omega}_e x_e + B' J'(J'+1) - B'' J''(J''+1)$$

Band

vibrational line

Every line of vibrational spectra will show a fine structure and will be a band due to different values of

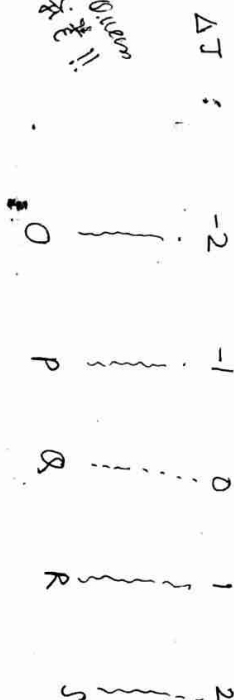
Now,

$$\bar{\omega}_{v,s} = \underbrace{\bar{\omega}_e - 2\bar{\omega}_e x_e}_{\bar{\omega}_v : \text{const. for a particular transition}} + B'J'(J'+1) - B''J''(J''+1)$$

ΔG_v will give $\bar{\omega}_v$

The fine lines will come from ΔG_v .

Now, considering $\Delta G(J)$ for



$\Delta J = 0$: Q branch

$\Delta J = 1, -1$: R branch, P branch

$\Delta J = 2, -2$: S branch, O branch

not observed in diatomic vibrational spectra. Occurs in electronic spectra.

hence in vibrational band, only P and R branch are observed to be seen.

R branch

$$\Delta J = +1$$

$$J' - J'' = 1$$

$\Rightarrow$

$$\bar{\omega}_{v,s}(R \text{ branch})$$

$$= \bar{\omega}_v + B'(J''+1)(J''+2) - B''J''(J''+1)$$

$$= \bar{\omega}_v + (J''+1) [(B' - B'') J'' + (2B' - B'')]$$

$$= \bar{\omega}_v + 1 [2B' - B'']$$

Put $J''=0$

Put $|\Delta J| = -1$ for P branch, $[J' - J'' = -1]$

where $J'' = 1, 2, 3, 4, \dots$

$$\bar{\omega}_{v,s}(P \text{ branch}) = \bar{\omega}_v + B'(J''-2)(J''-1) - B''J''(J''+1)$$

$$\bar{\omega}_P = \bar{\omega}_v - (B' + B'') J'' + (B' - B'') J''^2$$

$$\bar{\omega}_R = \bar{\omega}_v + (B' - B'') J'' + (B' + B'') J''^2 + 2B''$$

Note that $\bar{\omega}_P$ and $\bar{\omega}_R$ are bands (for different values of J'' , different lines) Mathematically both of them can be combined into a single equation.

Put

$$\bar{\omega}_s = \bar{\omega}_v + (B' + B'') m + (B' - B'') m^2$$

where m : parameter

if $m = -J''$: J'' starting from +1

we get P branch

if $m = (J''+1)$: we get R branch

Note that $m \neq 0$ i.e. $|m| > 0$

It's called

For P branch, replacing J values (1, 2, 3, 4, ...)

$$\bar{\omega}_P(1) = \bar{\omega}_v - 2B''$$

$$\bar{\omega}_P(2) = \bar{\omega}_v - 2(B' + B'') + 4(B' - B'')$$

$$= \bar{\omega}_v + 2B' - 6B''$$

$$\bar{\omega}_P(3) = \bar{\omega}_v - 3(B' + B'') + 9(B' - B'')$$

$$= \bar{\omega}_v + 6B' - 12B''$$

Refer Fig. 11 on P-313

For R values, $m = J'' + 1$ J : starts from 0

For R series

$$\bar{\omega}_R(0) = \bar{\omega}_V + 2B'$$

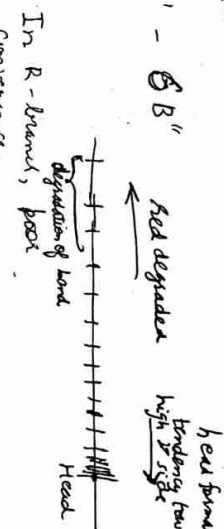
$$\bar{\omega}_R(1) = \bar{\omega}_V + 2(B' + B'') + 4(B' - B'')$$

$$= \bar{\omega}_V + 6B' - 2B''$$

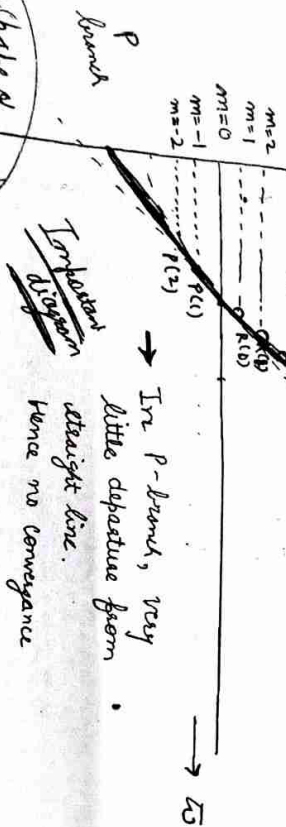
$$\bar{\omega}_R(2) = \bar{\omega}_V + 3(B' + B'') + 9(B' - B'')$$

$$= \bar{\omega}_V + 12B' - 6B''$$

$\bar{\omega}_V$ vs m



Head formation
Tendency is observe towards higher frequency side i.e. violet side hence Red is degraded.



shape of the curve should be like. Fig 12, P 314

if $B' - B'' > 0 \Rightarrow$ Violet Degraded Parabolae: formed in electronic spectra
But $B' - B'' < 0 \Rightarrow$ Red Degraded Parabolae: formed in electronic spectra

For violet of parabolae

$$\frac{d\bar{\omega}}{dm} = 0 \Rightarrow (B' + B'') + 2m(B' - B'') = 0$$

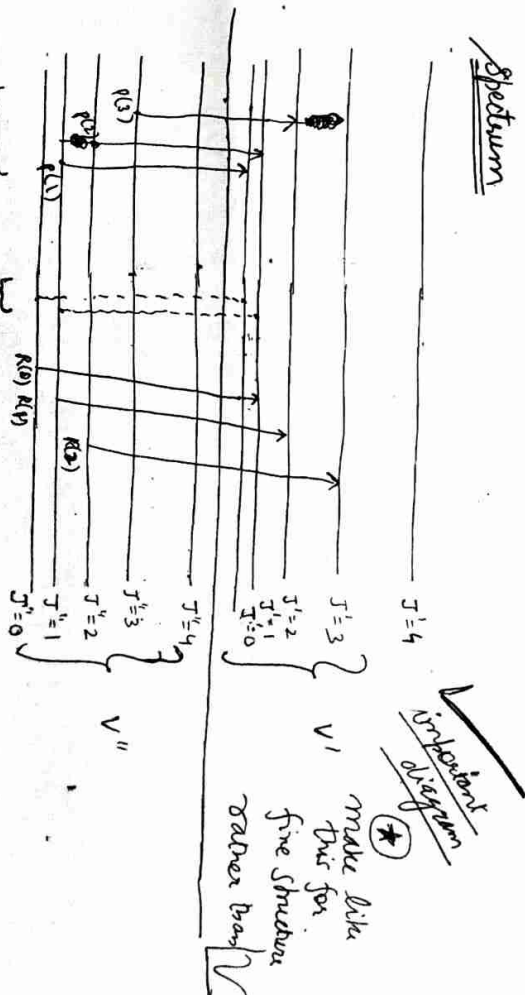
$$m = - \frac{(B' + B'')}{2(B' - B'')}$$

① Vibration Rotation spectra shows poor tendency of head formation towards higher wave number side violet side (is tendency)

② [Only bands degraded to only 1 side are formed, i.e. Red degraded only]

Reason for both is $K' > K''$ but difference is small $B' < B''$
 $(B' - B'')$ is small negative quantity

Spectrum



Important diagram

P Branch $\Delta J = -1$
Q branch $\Delta J = 0$
R branch $\Delta J = 1$

Given 3 lines of infrared spectrum, we can have info about:

- Types of Bond
- I_{cm}
- Isotopic Effect

⑧ $\lambda_{ir} = 14.7 \mu m$

$$\frac{1}{\lambda} = \frac{1}{2\pi c} \sqrt{\frac{k}{\mu}}$$

$$\mu = \frac{m_1 m_2}{m_1 + m_2} = \checkmark$$

$\Rightarrow$ K: Calculated ✓

$J=0 \rightarrow J=1$ Rotational $\Delta E = 2B \times c = \frac{2h}{8\pi^2 I}$

$$\nu = \frac{1}{4\pi^2 I}$$

⊛ Practical difficulty to get emission spectra. B coz in order to observe it, we need sufficient of particles in higher energy state. Its difficult to raise them to such low level. Raman Oscillator not very good.

$$20.68 (J+1)$$

$$E(J+1) - E(J) = B(J+1)(J+2) - B J(J+1)$$

$$= 2B(J+1)$$

$$2B = 20.68 \quad B = \frac{h}{8\pi^2 c I_{cm}}$$

Hence I_{cm} obtained.

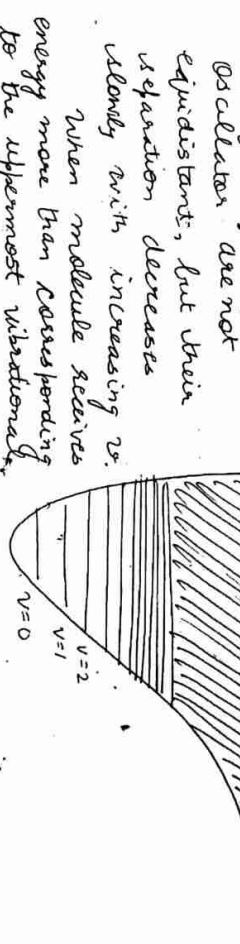
$$K: k_1, k_2 = \mu_1 \omega_1^2; \mu_2 \omega_2^2; \mu_3 \omega_3^2$$

$$H = \left(\frac{J^2}{2I} \right) = \left[\frac{J_{x^2}}{2I_{xx}} + \frac{J_{y^2}}{2I_{yy}} + \frac{J_{z^2}}{2I_{zz}} \right]$$

⊛ Spacings between vibrational levels are considerably larger than the spacing between rotational level of a molecule, infra larger than kT at room temperature.

Hence most of the molecules in a sample exist in $v=0$ state, with ~~only~~ only their zero point energy. Excitations are heavy molecules like I_2 that have small $\omega_e \Rightarrow$ they can exist @ $v=0, v=1, v=2$ at room temperature! $v=1$ or $v=0$ at RT are still appreciable amount of eff.

⊛ Energy levels of anharmonic oscillator are not equidistant, but their separation decreases slowly with increasing v .



When molecule receives energy more than corresponding to the uppermost vibrational level, it dissociates into atoms and excess energy appears as ungratified kinetic energy of these atoms. Hence a continuum joins the uppermost level.

It is similar to ungratified state beyond series limit in H atom spectrum which follows Rydberg path (as shown).



⊛ There are about 50 rotational levels associated with a single vibrational levels for HCl molecule.

⊛ B_v can be assumed to be $B_v = B_e - \alpha \left(v + \frac{1}{2} \right)$

where $B_e = \left(\frac{h}{8\pi^2 \mu r_e^2 c} \right)$

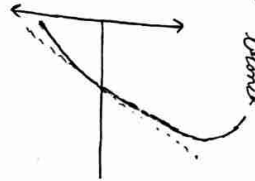
For vibrational-rotational motion,

$$\bar{\omega}_R = \bar{\omega}_v + (B' - B'')m + (B' - B'')m^2$$

$$m = \pm 1, \pm 2, \pm 3, \dots$$

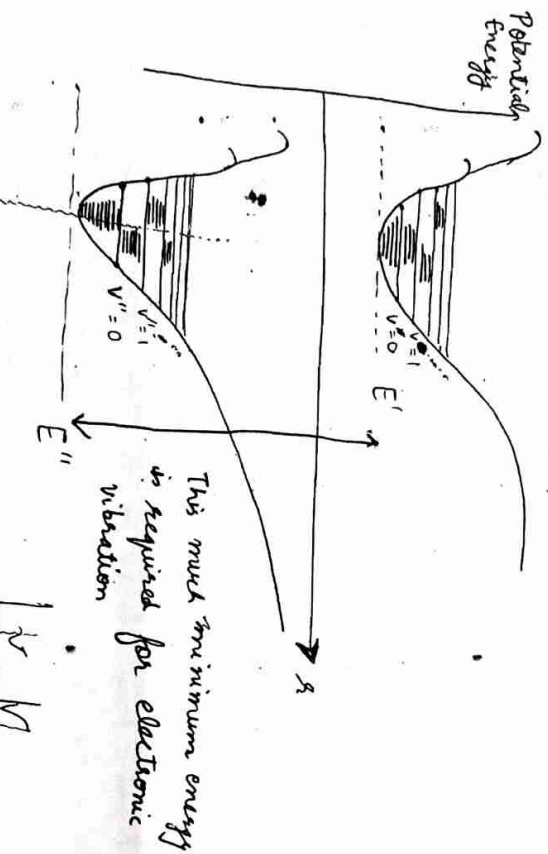
Positive: R branch
Negative: P branch

- The spectrum formation has
- poor tendency for head formation
- red-degraded character



Electronic spectra

No condition like possession of dipole moment to all molecules give electronic spectra. It is of order of several eV.



In electronic motion, bond length may be altered.

There are 4 allowed transitions depending upon m value corresponding to l_z about internuclear axis.

Selection rules

$$\Delta m_l = 0, \pm 1, \pm 2, \pm 3$$

Transition $\Sigma \quad \Pi \quad \Delta \quad \Phi$

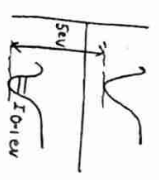
In Σ transition, $\Delta m_l = 0$. It is 0 for rigid rotator molecules. [Electronic spectra can be obtained in emission molecules.]

- * Also unlike vibrational or rotational level [It is always higher in lower levels]
- * They have absorption spectra because population is higher in lower levels

Again note that for rigid rotator molecule, only $\Delta m_l = 0$ is allowed.

$$E = E_{el} + E(v) + E(J)$$

eg. if $\Rightarrow$ 5 eV $\Rightarrow$ 5.1 eV $\Rightarrow$ 5.150 eV
provided E_{el} changes $\Rightarrow$ new E_{el} as well as E_v changes $\Rightarrow$ new E_v as well as E_J changes $\Rightarrow$ new E_J as well as E changes $\Rightarrow$ new E , new v , new J .



(Note that quanta are so small st such energy level) (continuum is possible)

$$\Rightarrow \bar{\omega} = \frac{\Delta E(e)}{hc} + \frac{\Delta E(v)}{hc} + \frac{\Delta E(J)}{hc}$$

$$= \Delta T(e) + \Delta G(v) + \Delta F(J)$$

$$\bar{\omega} = \bar{\omega}_e + \Delta G(v) + \Delta F(J)$$

(note that it is different from the one earlier)

Whenever energy is given in units of eV (typically from white source), ~~electronic~~ levels of molecule change due to change of electronic level of individual atoms.

In electronic spectra, Δv is unrestricted. When Δv is ~~unrestricted~~ ^{not considered} ~~considered~~, it is called fine structure.

Note that intensity of transitions are governed by Population rules defined by Maxwell Boltzmann statistics.

Gross / Coarse structure of Electronic Band Spectrum

$$\bar{D} = \bar{\omega}_e + G(v') - G(v'') \quad (2 \text{ diagrams})$$

In Gross structure, we neglect ΔT for time being.

$\Delta v =$ unrestricted (Restriction is only the no. of vibrational levels in single electronic level)

$$G(v) = \left(v + \frac{1}{2}\right) \bar{\omega}_v - \left(v + \frac{1}{2}\right)^2 \bar{\omega}_v x_v$$

$$\bar{\omega}_v = \frac{1}{2\pi c} \sqrt{\frac{k}{\mu}} \quad \text{where } \bar{\omega} = \sqrt{\frac{k}{m}} \Rightarrow \bar{\omega} = \frac{1}{\sigma} = \frac{c}{\sigma} = \left(\frac{2\pi c}{\lambda}\right)$$

$$= \frac{\omega}{2\pi c} = \frac{1}{2\pi c} \sqrt{\frac{k}{\mu}}$$

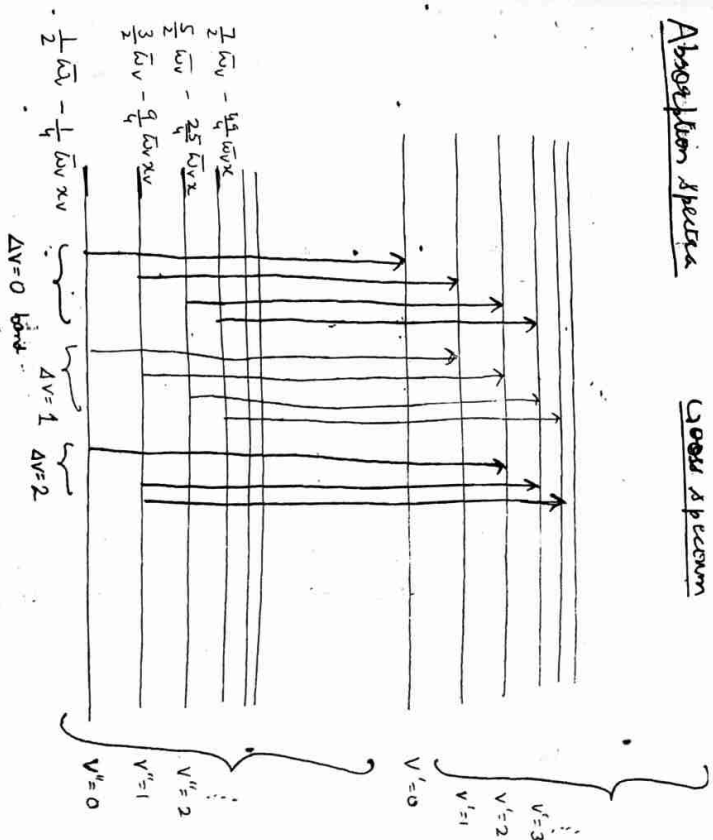
In ground state, $G(v'') = \left(v'' + \frac{1}{2}\right) \bar{\omega}_v - \left(v'' + \frac{1}{2}\right)^2 \bar{\omega}_v x_v$

In excited state, $G(v') = \left(v' + \frac{1}{2}\right) \bar{\omega}_v - \left(v' + \frac{1}{2}\right)^2 \bar{\omega}_v x_v$

Electronic energy is the energy which the molecule possess if nuclei were fixed, & consists of
 $\rightarrow$ K.E. & P.E. of extranuclear electrons
 $\rightarrow$ P.E. of repulsion between nuclei.

Absorption Spectra

Gross Spectrum



Note that separation are not of const. width $\Rightarrow$ Band of lines

Similarly, we can draw from $\Delta v = -1, -2, \dots$

Note that it is Absorption Spectra. We can draw Emission spectra just by drawing opposite lines given that enough population is there in excited state.

$$\bar{\omega}_{v'} = E_e + G(v') - E_g - G(v'')$$

$$= [T(v') - T(v'')] + [G(v') - G(v'')]$$

$$= \bar{\omega}_e + \left(v' + \frac{1}{2}\right) \bar{\omega}_v - \left(v' + \frac{1}{2}\right)^2 \bar{\omega}_v x_v - \left(v'' + \frac{1}{2}\right) \bar{\omega}_v + \left(v'' + \frac{1}{2}\right)^2 \bar{\omega}_v x_v$$

Note that $\bar{\omega}_v' \neq \bar{\omega}_v$
 (different electronic levels have different)

$\Delta v=1$
 $\Delta v=2$

All are bands having distinct wave numbers for all lines. These bands are called PROGRESSIONS.

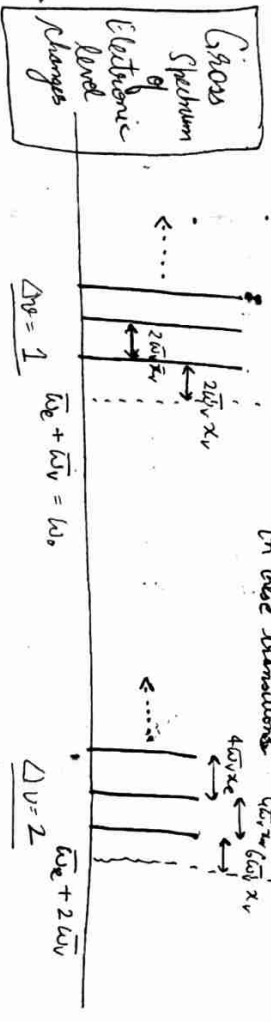
Progression are Band systems

Band or Progression

$v=0 \rightarrow v=1 \Rightarrow \omega = \bar{\omega}_e + \frac{3}{2} \bar{\omega}_v - \frac{9}{4} \bar{\omega}_v x_v - \frac{1}{2} \bar{\omega}_v' + \frac{1}{4} \bar{\omega}_v' x_v$
 $= \bar{\omega}_e + \bar{\omega}_v - 2 \bar{\omega}_v x_v$
 $v=1 \rightarrow v=2 \Rightarrow \omega = \bar{\omega}_e + \bar{\omega}_v - 4 \bar{\omega}_v x_v$
 $v=2 \rightarrow v=3 \Rightarrow \omega = \bar{\omega}_e + \bar{\omega}_v - 6 \bar{\omega}_v x_v$

All total $v''=0$
 $v'=1, 2$
 Progression
 for $v''=0$

Let us say $(\bar{\omega}_e + \bar{\omega}_v) = \text{reference level } \omega_0$



→ Note that subsequent height of lines decrease due to lower population in these transitions.

Δv=2 Band or Progression

$v=0 \rightarrow v=2 \Rightarrow \omega = \bar{\omega}_e + 2 \bar{\omega}_v - 6 \bar{\omega}_v x_v$
 $v=1 \rightarrow v=3 \Rightarrow \omega = \bar{\omega}_e + 2 \bar{\omega}_v - 10 \bar{\omega}_v x_v$

Note that $\omega_e' \neq \omega_e''$



Now note that every line of every band will show further fine structure due to different rotational levels.

$\Delta E = \Delta E_e + \Delta G(v) + \Delta F(J)$
 $= \bar{\omega}_e + \bar{\omega}_v + \Delta F(J)$
 $= \omega_0 + \Delta F(J)$

Selection Rule

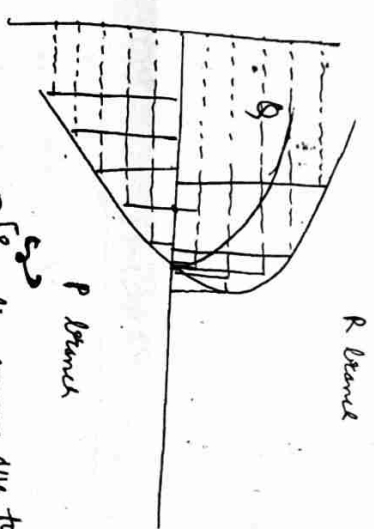
$\Delta J = \pm 1$ for rigid rotator molecule
 $\Delta J = 0, \pm 1$ for symmetric molecule
 $\Delta J = \pm 2, \pm 1$ exceptional cases in certain molecules that are heavy & non rigid rotator.

These transitions are called Branches. Bands are

$\Delta J = +1 : R \text{ branch}$
 $\Delta J = -1 : P \text{ branch}$

Note that here $x' > x''$ (Possible case)
 $\Rightarrow (B' - B'') < 0$: Negative quantity [Not small]

Hence good FORTRETT PARABOLA is formed.
 $\omega = \bar{\omega}_0 + (B' + B'') m + (B' - B'') m^2$



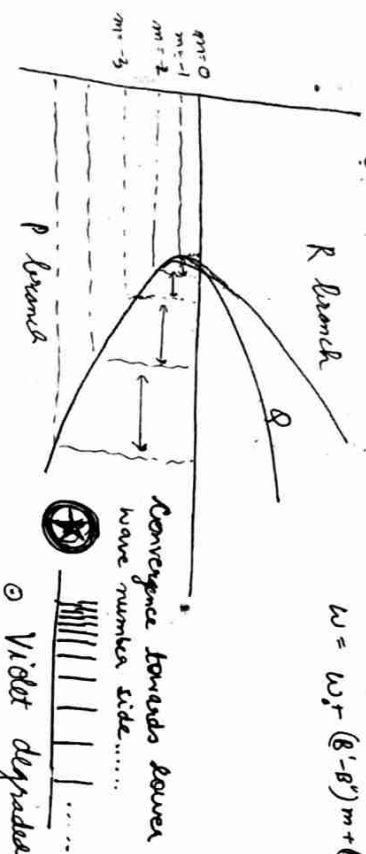
These lines converge due to step 4 of parallel curve

- Good tendency for band formation.
- Red degraded
- Converge towards higher wave number side

$B' - B'' > 0$: Positive Quantity

$\Delta : mJ''$

$\omega = \omega_0 + (B' - B'')m + B''m^2$



Vertical in P branch
Convergence in towards lower wave number or red side, hence violet degraded.

Similarly, we can do for Q, as well as S branches.

$\bar{\omega} = \bar{\omega}_0 + B'J'(J'+1) - B''J''(J''+1)$

$\bar{\omega} = -\bar{\omega}_0 + (B' - B'')J'(J'+1)$ [Q branch]

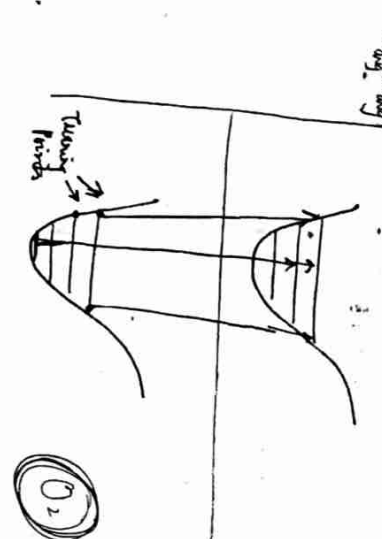
Again $(B' - B'')$ can be negative [symmetric molecules] or positive

FRANK-CONDON PRINCIPLE

'Most favoured transitions' as 'Most Probable Transition' in electronic spectra are given by this principle.

It is the 'secondary' criteria. Primary criteria is the Population according to Maxwell Boltzmann curve.

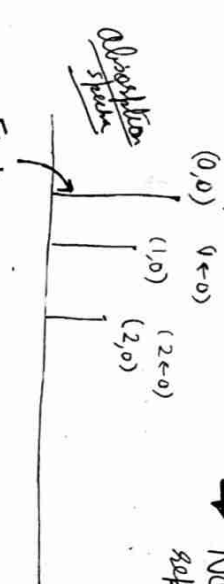
All transitions must be represented by vertical lines.



During transitions, internuclear distances are const. Electrons are so small and swift that in their time of transition, heavy nuclei remain at the same positions.

For ground state transitions, must occur from middle of level. Any other transitions occur from turning points. '0_0' is more at ends for higher state.

Note that transitions according to F.C. Principle are most probable and most intense. Other transitions are of course allowed but are less probable.

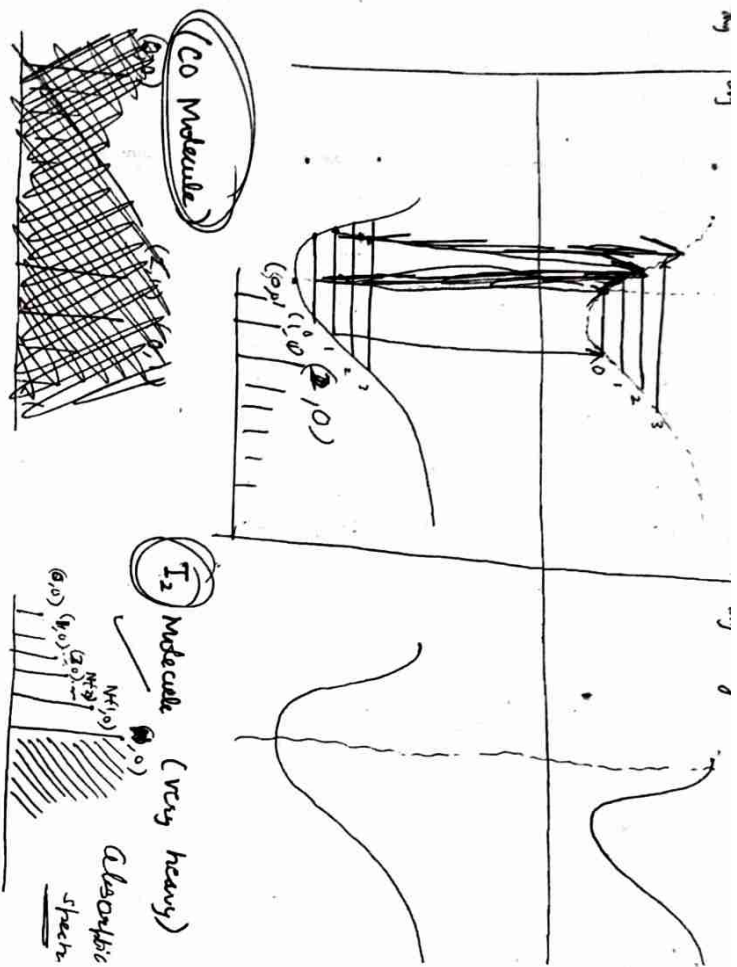


Note how the transitions are represented (x, y)

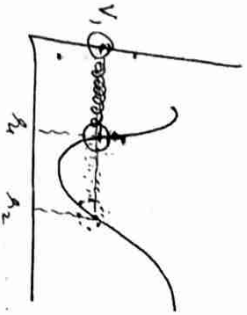
(x ← y) : Absorption spectra
(x → y) : Emission spectra
Vertical lines from turning points of v=1 cut at v'=1, 2 respectively.

⇒ (1,1) & (2,1) have equal intensity !!





→ Note that in vibrational energy state graphs, bond length is shown by a component of points on curve



Bond length r_1 and r_2 have same Potential Energy.

→ Order of gap of spectral lines :- $\approx 20 \text{ cm}^{-1}$: rotational
 $\approx 1000 \text{ cm}^{-1}$: vibrational
 $\approx 50,000 \text{ cm}^{-1}$: electronic

(*) Homonuclear molecules (H_2, O_2, N_2) which give neither rotational nor vibrational spectra b'coz they do not have permanent dipole moments. give electronic spectra b'coz the instantaneous dipole moments change during the redistribution of electronic charge which accompanies electronic transition.

(*) Same Δv , different $\Delta J \Rightarrow$ Branches : which is a Band
 $\Delta J = 1$ Branch R
 $\Delta J = -1$ Branch P

while

$\Delta v = \text{const}$, $v' = \text{const}$ or $v'' = \text{const}$ Progression which are
 $(v' \text{ and } v'' \text{ varying})$ $(v'' \text{ varying})$ $(v' \text{ varying})$
 [This is also called a dispersion] [v'' progression] [v' progression] Bond systems

जो vary कर रहा है, उसको Progression
 i.e. जो Progress कर रहा है !!

Molecular Physics (4)

28/02/12

Raman Spectra

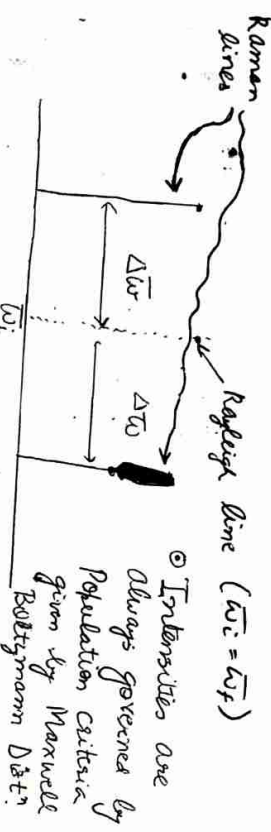
① In solution, generally specimen is taken in vapour state.

In Raman spectroscopy, we can take in any phase.

② Here, permanent dipole moment is not required. Induced Dipole Moments are reason for Raman Spectra.

$\gamma_i \rightarrow$ Specimen $\rightarrow \gamma_f$

If γ_f compares higher as well as lower freq waves $\Rightarrow$ Raman Scattering



It is a characteristic of molecule. It typically lies in infrared region.

Intensity $\propto$ intensity STOKES

$\omega_i + \Delta\omega$: lower intensity ANTI-STOKES

Typical freq. in Photo Electric Effect : UV

If freq. is higher, say γ -ray $\Rightarrow$ Electron is removed as we as the photon comes out with reduced frequency.

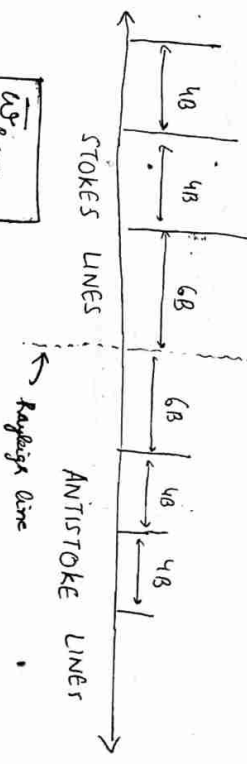
It's called Compton Effect.

✓ But in Raman Spectra, the unusual thing is that some photons comes out with higher energy, though intensity is less.

✓ In Compton Effect, interaction with atoms (particularly e^-). In Raman Effect, interaction with molecules. Atomic Phenomenon vs Molecular Phenomenon.

Also $\Delta\omega$ lies in infrared region $\Rightarrow$ vibrational energy levels.

Fine Structure of Raman Spectra



$$\omega_{\text{Raman}} = \omega_i \pm 4B \left(j + \frac{3}{2} \right) \quad j = 0, 1, 2, 3, \dots$$

Note that Stokes lines are not confirmation of Raman Effect. Only once Anti-Stokes are obtained, we are confirmed of Raman Effect. But Anti-Stokes lines are low intensity lines, hence difficult to observe. When lasers came, then it became possible to detect them. hence, we call it Laser Raman Spectroscopy.

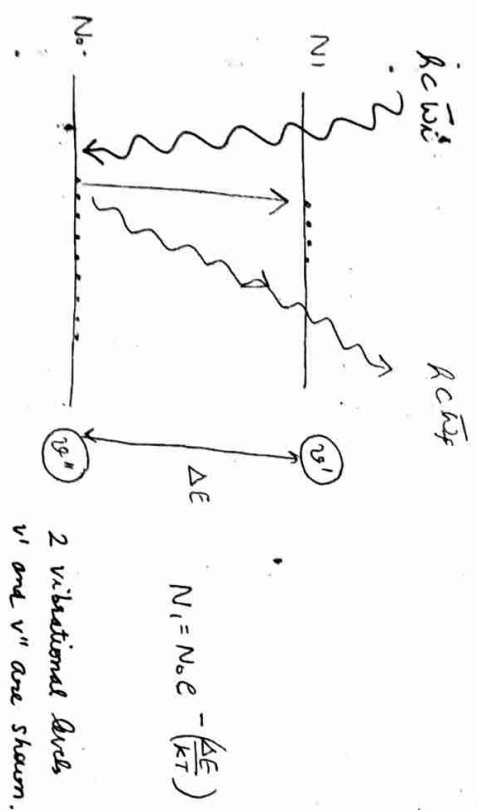
Quantum Mechanical Explanation

$\Delta v = \pm 1$. Vibrational Raman Spectra

$\Delta J = \pm 2$. Rotational Raman Spectra

For vibrational $\omega_f = \omega_i \pm [G(v+1) - G(v)]$

$$\Rightarrow \Delta\omega_{\text{Raman}} = \pm \omega \left[v + \frac{3}{2} - v - \frac{1}{2} \right] = \pm \omega$$



Let $\Delta E = E(v') - E(v'') = \underline{h\omega_s}$

Visible light is incident upon v'' i.e. $\omega_i > \omega_s$

If elastic collision, no energy transfer

$h\omega_i + E(v'') = E(v') + h\omega_s$: Rayleigh scattering
 $\Rightarrow \omega_i = \omega_s$

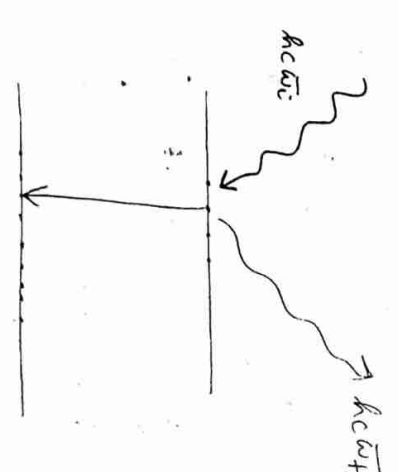
If inelastic collision

$h\omega_i + E(v'') = E(v') + h\omega_s$: Stokes line (Raman scatter)
 $\Rightarrow \omega_s = \omega_i + \left(\frac{E(v') - E(v'')}{h} \right)$

Raman shift : characteristic of molecule

High intensity laser maximum population in ground state. Incident & scattered light has nearly same energy.

$h\omega_i + E(v'') = E(v') + h\omega_s$



$\Rightarrow \omega_s = \omega_i + \left(\frac{E(v') - E(v'')}{h} \right)$: Anti Stokes line

Lower intensity line due to lower population in excited level

→ Inelastic collision b/w photon & molecules is, therefore, responsible for Raman Effect.

o Note that NO NEW PHOTON IS SCATTERED and NO PHOTON IS COMPLETELY ABSORBED. (Compton)

Selection rule for Fine structure

$\Delta v = \pm 1$ [0 for Rayleigh]
 $\Delta J = 0, \pm 2$ [0 for Rayleigh]

Selection rule is due to induced dipole moment. Hence occurs in all molecules irrespective of possession of permanent dipole moment.

For vibrational Raman spectrum,

$\omega_s = \omega_i \pm \left[G(v+1) - G(v) \right]$
 $\omega_s = \omega_i \pm \Delta \omega_{\text{Raman}} = \omega_s$

For fine structure, or rotational elements, we are

$\bar{\omega}_{\text{Raman}} = \bar{\omega}_i \pm [F(J+2) - F(J)]$ **Note** that we are not talking about fine lines within coarse vibration lines. i.e. a. talking about only fine lines on left

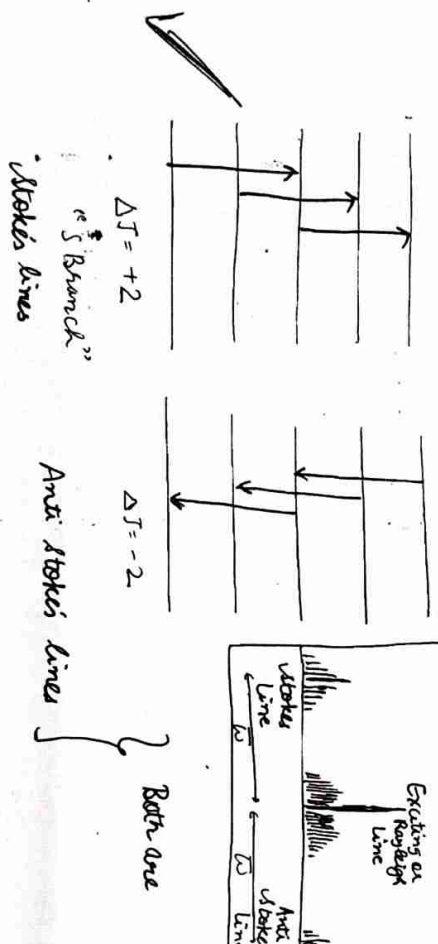
where $F(J) = B J(J+1)$

$\Rightarrow \bar{\omega}_{\text{Raman}} = \bar{\omega}_i \pm B [J(J+2) - J(J+1)]$ lines on left of Rayleigh line in far-IR end spectra.

$\bar{\omega}_{\text{Raman}} = \bar{\omega}_i \pm 4B \left(J + \frac{3}{2} \right)$

$\Rightarrow \bar{\omega}_{\text{Raman}} = \bar{\omega}_i \pm 6B, \bar{\omega}_i \pm 10B, \bar{\omega}_i \pm 14B, \dots$

From given lines, we can calculate $B \Rightarrow I_{\text{cm}} \Rightarrow \text{Bond length}$



Homonuclear diatomic molecules do not give vibrational rotational spectra but give Raman spectra. Called Infrared Inactive but Raman Active.

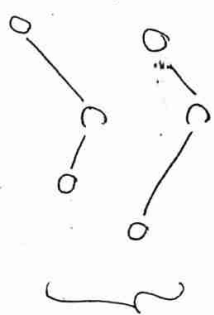
For CO_2 molecule $\text{O}=\text{C}=\text{O}$ asymmetric linear molecule.

linear mode

Permanent Dipole Moment $\Rightarrow$ only Raman spectra



Bond Mode (symmetric or asymmetric)



No centre of symmetry $\Rightarrow$ Dipole Moment

No centre of symmetry $\Rightarrow$ Both active

Centre of symmetry $\Rightarrow$ Only Raman Active

These apart from I_{cm} , Bond length , type of symmetry is also known from spectra.

→ Measuring Anc was initial source of light. It has drawbacks:

1. It extended source of light $\Rightarrow$ large quantity of sample is required, leading to wastage of precious material.

2. Some of the light of incident rays enter spectrometer tube. It will mask weak Raman lines.

3. Total energy = E_{hv} incident $\Rightarrow$ whole incident photon

may be caused by excitation of materials with UV light. Fluorescent will mask Raman's weak lines. [Note: Raman's lines are weak and difficult to see.]

Analysis of coloured salt solutions was difficult due to limited frequency available in mercury arc. Lasers are available over a wide range of frequencies.

All these drawbacks were removed after the use of LASER. Hence before 1966, Raman spectroscopy was finding little use in molecule structure determination.

As an IR Laser (inert) are used. They can be focused on small area with wide power range. Hence normal density grating can be used to record spectra. Also they have wide frequency range.

$$\lambda = \frac{1}{514.5 \text{ nm}}$$

21 Fine structure cannot be resolved otherwise

$$\frac{h^2}{8\pi^2 I} = B = 1.43 \times 10^{-5}$$

$$I_0 = 8.4 \times 10^{12} \text{ Hz}$$

$$E(j) = \frac{I^2}{2I} = \frac{j(j+1) \frac{h^2}{8\pi^2 I}}{8\pi^2 I}$$

$$E(v=1) - E(v=0) = h\nu_0$$

$$\text{Equating the ①, } j(j+1) = \frac{6.62 \times 8.4 \times 10^0}{1.6 \times 10^{-13}}$$

* Vibrational Raman spectra gives exactly with wave no. of main vibrational rotational spectra in near infra-red spectrum while

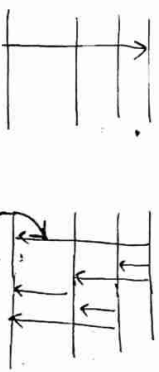
Rotational Raman spectra vibration = 2 [separation of lines in microwave spectrum]

* Induction or Polarization by incident light cause dipole moment in homonuclear molecules $\Rightarrow$ rotation. n.v. r.v. a...

* Luminescence (2114)

Phosphorescence

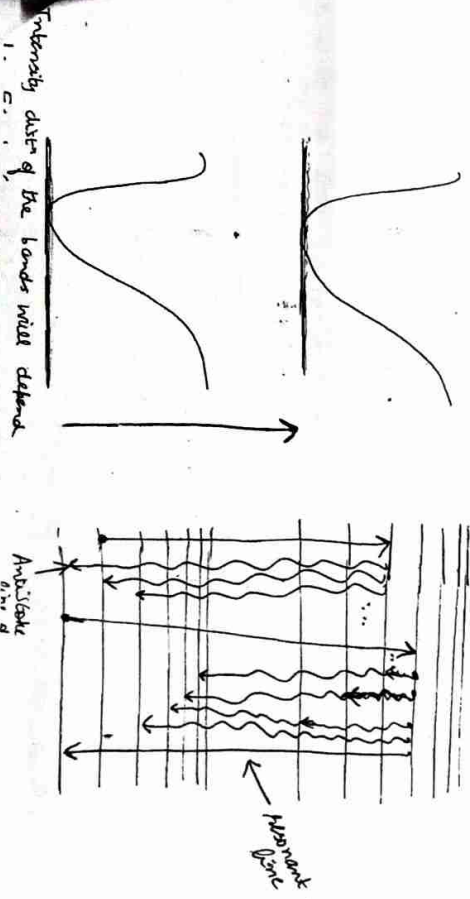
When a molecule is incident upon by a "suitable frequency" i.e. one of the absorption frequencies, molecule may be raised to any excited level (rot/vib/electronic).



→ this frequency is absorbed
→ all these frequencies will be emitted

Now, molecule has option to come down instantaneously. It's called Fluorescence. The γ released are less than incident.

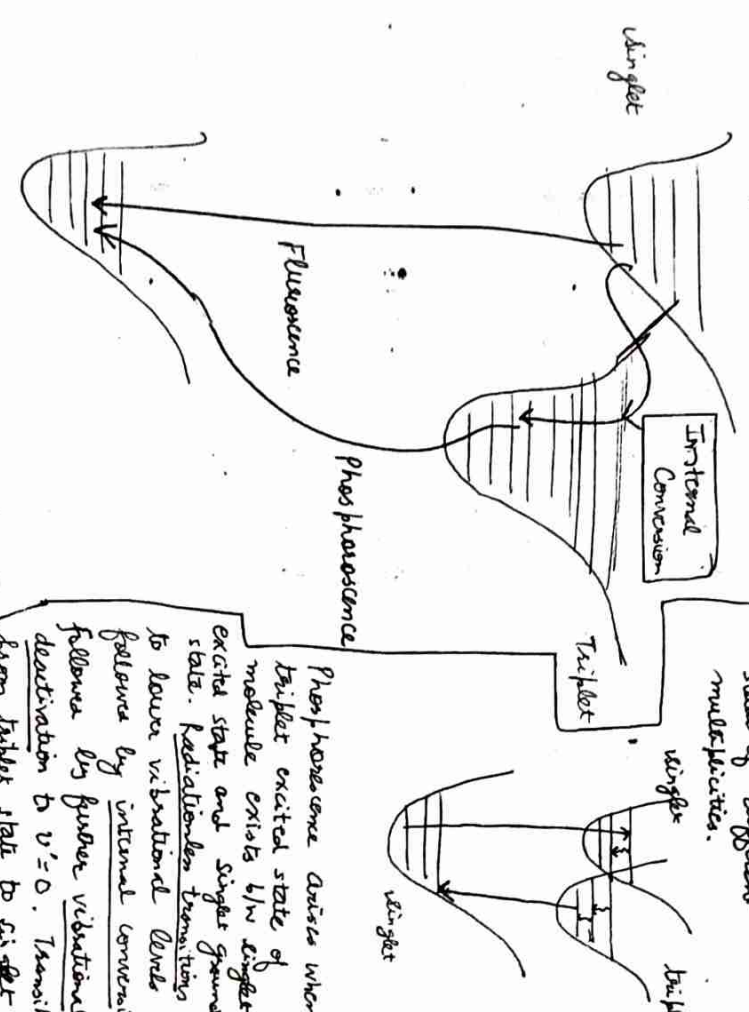
In Phosphorescence, energy is emitted even after source of excitation is removed. It is delayed emission of radiation.



These 'I' molecules are raised to higher energy. Incident photon is absorbed.

All frequencies emitted are less than that of resonant &

Singlet, $\Delta = 0$ $\lambda = 1$ Triplet, $\lambda = 1$ $\lambda = 3$



Delayed emission causes due to transitions that connect electronic states of different multiplicities.

Phosphorescence arises when triplet excited state of molecule exists b/w triplet excited state and singlet ground state. Radiations transition to lower vibrational levels follows by internal conversion followed by further vibrational deactivation to $v=0$. Transition from triplet state to singlet ground state are responsible for phosphorescent emission.

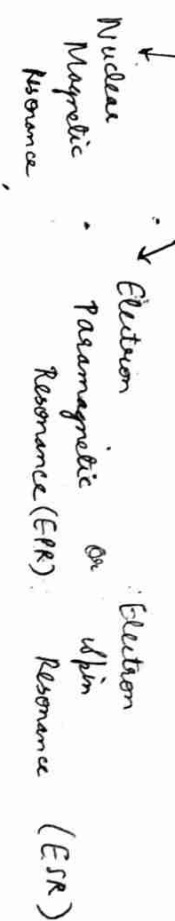
Remember selection rule, $\Delta \lambda = 0$
 $\sigma \rightarrow \sigma$ transition: same multiplicity.
 It has high probability $\Rightarrow$ shorter time.

Phosphorescence occurs in violation of selection rule
 $\Rightarrow$ low probability $\Rightarrow$ longer time
 $\Rightarrow$ delayed emission.

Molecular Physics (5)

NMR and EPR/ESR

NMR and EPR are magnetic resonance techniques



Resonance is matching of frequencies of source and system.

Precession frequency of nucleus is matched with f of oscillating B. Angular Momentum precesses about field. Energy levels are separated by ΔE . ΔE is very small $\approx 10^{-8}$ eV. Remember, we mentioned it in spectroscopy.

$h\nu = 10^{-8} \times 10^{-19}$
 $\gamma \approx 10^6$
 $\gamma = \frac{hc}{E}$ in Radio Frequency Region

$\frac{\Delta E_{\text{electron}}}{\Delta E_{\text{spin}}} \approx 10^{-3} \text{ eV} : \text{Microwave Region}$

It is also called EPR because all paramagnetic salts have $\mu_{\text{spin}} \neq 0$, i.e. have unpaired $e^- \Rightarrow$ Spin Angular Momentum $\neq 0 \Rightarrow$ has magnetic precession $\Rightarrow \Delta E_{\text{electron spin}}$

We know, $\vec{\mu}_L = -\frac{e}{2m} g_L \vec{L}$ $g_L = 1$
 $\vec{\mu}_S = -\frac{e}{2m} g_S \vec{S}$ $g_S = 2$

$\vec{\mu}_J = -\frac{e}{2m} g_J \vec{J}$ $g_J = 1 + \frac{J(J+1) + S(S+1) - L(L+1)}{2J(J+1)}$
 $\vec{\mu}_J$ is component of $\vec{\mu}$ along $\vec{J}$. We are interested in

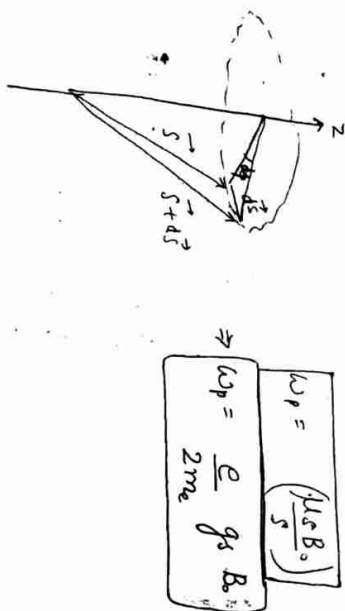
Note that prerequisite of ESR is $S \neq 0$.

Atom with unpaired e- i.e. $S \neq 0$ is kept in steady magnet field (const B_0).

$$\vec{T} = -\mu_B \vec{S} \times \vec{B}_0$$

$\Rightarrow \vec{S}$ will rotate about direction of $B_0 \Rightarrow$ e- will precess

$$\vec{T} = \frac{d\vec{S}}{dt} \Rightarrow d\vec{S} \perp \vec{S} \Rightarrow \text{Precession}$$



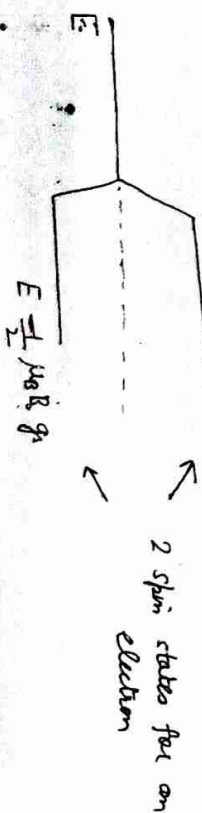
Precisely same derivation for Nuclear Magnetic Moment

Now Δ electron spin will occur due to potential of I and Magnetic field.

$$E = \omega_p S_z = \frac{e}{2m_e} g B_0 m_s \hbar = \mu_B g m_s$$

$$\text{For } e^- : m_s = \pm \frac{1}{2}$$

$$E = \pm \left(\frac{\mu_B g}{2} \right)$$



$$\Delta E = \mu_B g \hbar \gamma = h \nu$$

⚡ This γ has no role in NMR/ESR

$\rightarrow$ Here by absorbing frequency γ or emitting frequency γ , e- can flip between its spin states.

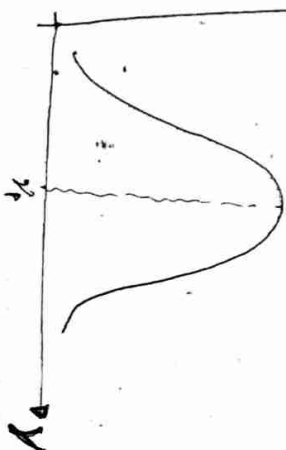
Now I apply another oscillating field,

$$\vec{B}_1 = B_1 \sin(2\pi \nu t)$$

from H_{system} Oscillator.

This varying field will cause magnetic resonance when γ of B_1 becomes equal to γ precession, then maximum absorption of energy is observed.

Energy absorbed by electron



$\gamma_0 = \gamma_{\text{precession}} = \gamma_2$ for resonance

I am applying fields, let for creating energy separation $\Delta E = \mu_B$ 2nd is applied for creating resonance with the natural frequency γ_0 is 1st for

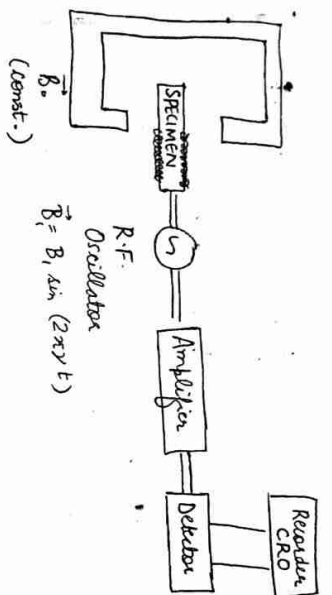
If @ $\gamma = \gamma_0$, I get maximum energy absorption

$$\gamma_0 = \frac{e}{4\pi m_e} g B_{\text{fixed}}$$

$\Rightarrow g$ is obtained

γ value lie in Microwave Region....

$$\Delta E \approx 10^{-3} \text{ eV}$$

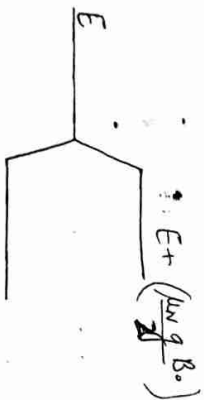


Typical schematic diagram of NMR spectroscopy

(For EPR, change the oscillator to hyperfine oscillation)

Ang. Momentum $|\vec{I}| = \sqrt{I(I+1)} \hbar$

$$\vec{\mu}_{\text{magnetic}} = \frac{e}{2m_p} g \vec{I}$$



$$\omega_p = \frac{e}{2m_p} g B_0$$

$$\Delta E = \mu_p I_z = \left(\frac{e\hbar}{2m_p} \right) g B_0 m_I$$

Nuclear Beta Magnetron

$$= \frac{\mu_N g B_0 m_I}{2}$$

which can be $\pm \mu_N g B_0$
(if only single proton)

Similarly,
NMR frequency $\nu = \frac{\omega_p}{2\pi} = \frac{e}{4\pi m_p} g B_0$

③ In chemical analysis and biological imaging, NMR and EPR are greatly used as low energy levels of microwave and Radio wave, hence no potential damage to living tissues.

MRI = NMR used in Medical Diagnosis

→ NMR is a technique in spectroscopy where RF fields induce transitions b/w nuclear energy levels causing absorption of energy wisely maximum energy absorption occurs at resonant level.

(B) $\vec{J} \Rightarrow \vec{\mu}_S \Rightarrow \vec{I} \text{ in } B \Rightarrow \tau \perp S \Rightarrow dS \perp S$

⇒ Precision

(22) $\gamma_p = \frac{e}{4\pi m_p} g_p B_0$

$$= \frac{1.6 \times 10^{-19}}{4\pi \times 1.6 \times 10^{-24}} \times g_p \times 1.5$$

$$\gamma_p = \frac{\Delta E}{\hbar} = \left(\frac{2\mu_B B}{\hbar} \right) = \frac{2e\hbar}{2m} g_I m_I B$$

(*) $\mu_p = \frac{e}{2m} g \vec{I}$
(*) $(\mu_p)_z = \frac{e}{2m_p} g \cdot I_z = \frac{e\hbar}{2m_p} g m_I$

$2.793 \mu_N = \mu_N g m_I$
 $m_I = \left(\frac{1}{2} \right)$ (no value of $m_I = \frac{1}{2}$)

$\Rightarrow g = 2 \times 2.793 = 5.586$

In water NMR occurs due to Proton
 $\gamma_{\text{proton}} = 5.586$

(Hence $\mu_H = \frac{e}{2m} \times 5.586$
 $= \underline{\underline{\mu_N \cdot 8.793}}$)

Chemical shift

$\Rightarrow$ due to shielding : $B = B(1-\sigma)$

Magnetic field encountered by bare nucleus due to shielding effect of outer electrons

$$\Rightarrow \gamma_P = \frac{e}{4\pi m_p} g_P B \cdot (1-\sigma)$$

We take 2 nucleus A and B

$$B_A = B_0 (1-\sigma_A)$$

$$B_B = B_0 (1-\sigma_B)$$

$$\Rightarrow B_A - B_B = -B_0 (\sigma_A - \sigma_B)$$

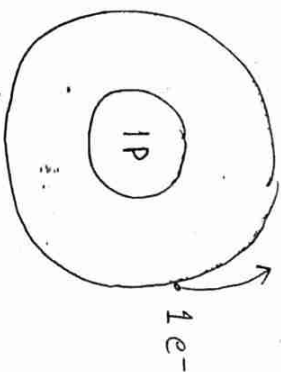
$$\Rightarrow (\sigma_A - \sigma_B) = -\left(\frac{B_A - B_B}{B_0}\right)$$

B_0 : applied
 B_A, B_B : fields at which resonance was observed.

Its important in chemical analysis.

Importance of Neutral H atoms in Astronomy

In start of Universe, abundance of H (neutral) Nucleus Fusion to form He atoms and then further elements were formed.



Neutral H atom

Till now, we have neglected motion of Nucleus.

If I use ultra-high resolution spectroscopy, I see hyperfine lines far more than what explained by

$$\left[H = \frac{p^2}{2m} - \frac{e^2}{4\pi\epsilon_0 r} + \alpha (\vec{L} \cdot \vec{I}) \right] + [\text{Lamb shift}]$$

It is due to motion of nucleus.

$$\vec{J} = \vec{L} + \vec{S} \quad \dots \dots \text{Fine structure}$$

Hyperfine structure

$$\vec{I} = \sqrt{I(I+1)} \hbar$$

$$\Rightarrow \vec{F} = \vec{L} + \vec{S} + \vec{I}$$

$$= (\vec{J} + \vec{I})$$

Total Ang. Momentum of atom

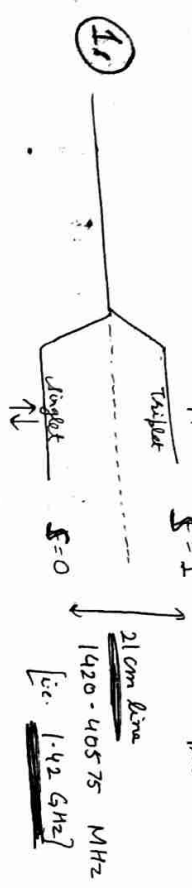
$$\Rightarrow |\vec{J}-\vec{I}| < \vec{F} < |\vec{J}+\vec{I}|$$

$\Delta F: 0, \pm 1$

Now, For $1s, 2s: l=0$
[lowest states]

Hyperfine structure of Neutral Hydrogen Atom

$\Rightarrow \vec{S} = \vec{S}_e + \vec{S}_p$
(electron) (proton)
 $S = 0, 1$



Now selection rule: $\Delta S = 0$

It lies in ^{MICROWAVE} Frequency region

Transition Probability = 2.9×10^{-15} per second

$\Rightarrow$ Transition Time $\approx 10^{14}$ second $\approx$ 10¹⁷ years

10^7 seconds $\approx$ 1 year

Hence, importance only in astronomy where such Time Periods are important.

Due to interatomic collisions, Transition time is reduced a little amount.

- It can provide clear picture of matter power spectrum in period of matter power spectrum in period after recombination,
- by mapping red-shifted 21 cm radiation
- It can provide picture of how universe was reionized,

As neutral H atoms which were ionized due to reionization from quasi-stellar sources which were appear as holes in 21 cm background.

21 cm line

Hydrogen in our galaxy has been mapped by the observation of 21 cm wavelength line of hydrogen gas. At 1420 MHz, this radiation can penetrate the dust clouds and giving us a more complete map of H than that of the stars themselves since their visible light is not able to penetrate the dust clouds. (ionizing)

The 1420 MHz radiation comes from the transition between the 2 levels of the $1s$ ground state, slightly split due to interaction between electron spin and nuclear spin. This splitting is known as hyperfine structure. This splitting of $1s$ ground state is extremely small compared to ground state energy of -13.6 eV (about $\frac{2}{10^6}$ ratio). 2 states come from the fact that both the e^- and proton have ($\frac{1}{2}$) spin, so there are 2 possible states, spin parallel (slightly high energy) and spin antiparallel (slightly low energy) ^{less tightly bound}

NMR

When the nuclear magnetic moment associated with nuclear spin is placed in an external magnetic field, the different spin states are given different magnetic potential energies. In the presence of such static magnetic field which produces a small amount of spin polarization, an RF signal of known frequency can induce a transition

Between spin states.

This "spin flip" places some of the spins, in their higher energy states. If the R-F. signal is then switched off, the relaxation of the spins back of the lower state produces a measurable amount of RF signal @ the resonant frequency associated with the spin flip. This process is called Nuclear Magnetic Resonance (NMR)

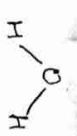
It is used to study chemical structure i.e. composition of different compounds & polymers. Also chemical shifts measurements in high resolution NMR spectroscopy is major source of chemical information

Laser Raman Spectroscopy and Molecular Structure

Raman effect is an important tool to determine molecular structure information of diatomic and polyatomic molecules
→ The vibrational and rotational Raman spectra enable us to determine ω and $B \Rightarrow$ to determine force constant k , bond length, I or for diatomic molecules.

→ For polyatomic molecules, Raman spectra and IR data (vibrational) give information regarding shape (linear or bent) and symmetry of the molecules. According to 'Mutual Exclusion Principle', for molecules with a centre of symmetry the frequencies of IR spectra are not observed in Raman spectra and vice versa. This enables us to deduce structure of molecules.

eg. Molecules of CO_2 and CS_2 are found to have strong IR bands that are not observed in Raman spectra
 $\Rightarrow$ Centre of symmetry $\Rightarrow O=C=O \quad S=C=S$

Molecules H_2O show IR bands i.e. centre of symmetry
Raman spectra $\Rightarrow$  (no centre of symmetry)

Raman Spectroscopy technique used to study vibrational, rotational and other low frequency modes of a system. It relies on inelastic scattering, as Raman scattering of monochromatic light from a laser in the visible, near infrared, or near UV range. The shift in energy of photons gives information about vibrational modes in the system. IR spectroscopy yields similar but complementary results.

The main difficulty of Raman scattering is the separation of weak inelastically scattered light from the intense Rayleigh scattered light. With the advent of lasers and notch filters, it has become an important instrument of spectroscopic analysis.

Uses of Raman Spectroscopy:

(i) determination of molecular structure
(ii) Raman gas Analyzers use real time monitoring of anaesthetic and respiratory gas mixtures during surgery.

(iii) In solid state physics, Raman scattering is used to characterise materials, and thereby Temperature measurement and finding crystallographic orientation.
(iv) Non invasive way of monitoring of biological tissue; and historical documents to determine last rays of preservation.