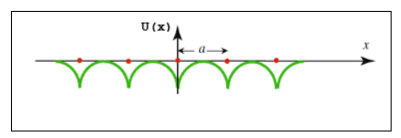
#### Chapter 8: The nearly free electron model and the tight binding model

The electronic structure of solids is usually described by two different approaches, the nearly free electron model and the tight binding model. Their primary differences are as follows:

#### 1- Nearly Free Electron Model (NFEM)

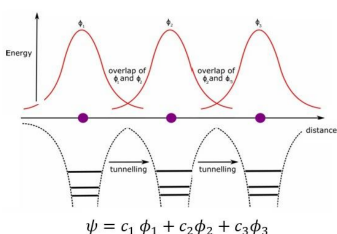
This model considers that electrons in a solid are nearly free, which means they move in a weak periodic potential caused by ions in the crystal lattice. The potential is treated as a small perturbation to the free electron gas,



and the electrons are delocalized over the whole crystal. The model is particularly useful for metals and simple metals where the potential is relatively weak. The mathematical approach is based on solving Schrödinger equation for a free electron gas with a weak periodic potential treated as a perturbation, resulting in forming nearly parabolic energy bands with small deviations due to the weak periodic potential. In conclusion, the model focuses on the delocalized nature of electrons and the influence of weak periodic potential on their motion.

### 2- Tight Binding Model (TBM)

This model assumes that electrons are tightly bound to their respective atoms, and the wavefunctions of neighboring atoms overlap only slightly. This model is more appropriate for describing insulators and semiconductors where electrons are more localized. The mathematical approach is based on the idea that wavefunction is constructed as a linear combination of atomic orbitals (LCAO) from neighboring atoms. The overlap between LCAOs leads to the formation of energy bands that can have non-parabolic shapes, reflecting the atomic orbitals from which they originated.



# (1) Nearly free electron model:

again, this model assumes that electrons in a solid are nearly free, meaning they move in a weak periodic Potential What can be treated as a small Perturbation so H= Ho + U(r); Ho: unperturbed Hamiltonian

$$=-\frac{k^2D^2}{2m}+U(r)$$
 with  $H_0|k\rangle = \mathcal{E}_k^{(0)}|k\rangle$ 
plane waves and

weak compared to the kinetic energy term (-ti202), such bhat Who potential U(F) can be treated as a parturbation on top of the completely solved problem of free electron (Ho = - \frac{\tau^2}{2m}).

plane waves and  $\mathcal{E}_{k}^{(b)} = \frac{k^{2}k^{2}}{2m} \quad j|_{k} = \frac{1}{\sqrt{V}} e^{\frac{1}{\sqrt{K}} \frac{2}{V}}$ IR> form a complete set of orbbogond strates < k/k) = 8, k'

- in this chapter we are using the braket notation to represent wavefunctions in reciprocal space (1k2). I'm coordinate representation, we have  $i(\vec{R}+\vec{R})\cdot\vec{r}$   $(\vec{r}|k) = V_{R}(\vec{r}') = \sum_{k} V(\vec{R}+\vec{R}) e$ 

projection wavefunction coefficients not wave function of IR) on (F) | refunction | ref

from scabbering bheory, the periodic lobential W(F) can scatter electron from state R -> R? and the matrix elements of the Potantial can be written as < R' |U|R') = FT[ U], fourier transform of U(r') evaluated at R-R', transition
amplitude = { D, if R-R' + IT, reciprocal labbice
from initial
state 1k) to
Lind state (k') so scabbering occurs only if the difference between inition and final warrectors is a reciprocal lartice vector. UK can be evaluated using  $L_{\kappa} = \frac{1}{\sqrt{2}} \left( \frac{d^{3}r^{3}}{U(r^{2})} e^{-\frac{1}{2} \left( \frac{r^{2}}{R} \right)^{2}} \right)$ se courit coll Now according to Perturbation bleavy, ble zerobh, first, and Second corrections to energy

Ex in the state IR> is givan by  $\begin{aligned} \mathcal{E}_{R}^{(0)} &= \mathcal{E}_{R}^{(0)} \\ &+ \langle R^{(1)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} | L | R^{(2)} \rangle \\ &+ \langle R^{(2)} | L | R^{(2)} |$ 

the first-order correction is the expectation value (average) of Whe Potential Calculated using the unperturbed states, i'm (RILIK). for a specific k as Shown i'n figure, where bhere is no degenercy at k, we can apply the non-degenerate Perburbation Cheory. let us see how every at Point A is affected when switching on ble weak periodic Potential U(r). None wear periodic potential U(r).

ab point A, we have  $|\vec{k}+\vec{k}\rangle = \frac{1}{\sqrt{3}}e^{-i\vec{k}\cdot\vec{r}}$ , Now for 1D  $|\vec{k}-\vec{k}| = \frac{2\pi}{\alpha}n$ ,  $|\vec{k}-\vec{k}| = 0$ K= 27 n, N=0 => K=0  $\langle \vec{R} | \mu(\vec{r}) | \vec{R} \rangle = \langle \vec{R} | \mu(x) | \vec{R} \rangle = \frac{1}{a} \left( e^{-i\vec{R}\cdot\vec{r}} \mu(\vec{r}) e^{-i\vec{R}\cdot\vec{r}} \right)$  $= \frac{1}{a} \int_{0}^{a} d^{3} \vec{r} \, L(\vec{r}) = \frac{1}{a} \int_{0}^{a} dx \, L(x) = constant$   $= \frac{1}{a} \int_{0}^{a} d^{3} \vec{r} \, L(\vec{r}) = \frac{1}{a} \int_{0}^{a} dx \, L(x) = constant$   $= \frac{1}{a} \int_{0}^{a} d^{3} \vec{r} \, L(x) = constant$   $= \frac{1}{a} \int_{0}^{a} dx \, L(x) = constant$ simply ab Point B, we have  $|\mathbf{k} - \mathbf{r}| = \frac{1}{\sqrt{n}} e^{-i\mathbf{k}\cdot\mathbf{r}} =$ so, simply, the 1st order correction Exp is just

a constant bhat represents the average potential. its effect is just to shift all energy levels by same amount 40. Note that in some cases Ho is either zero or constant. for example, of U(x) a CS 2xx or U(x) a sin zxx, 6 hour < k | L(x) | k > = Zero. on the other hand if L(x) x cos = x, bha < R / U(x)/k> = 1/2. Now let us find the second order correction of Ex of the skate Ik) at Point A. Now we consider that the bar state 1k) (n=0) is affected by only the closest states | R-R) (n=-1) and | R+R) (n=+1) as shown i  $\frac{1}{2} = \sum_{k' \neq k'} \frac{|\langle k' | L | k \rangle|^{2}}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\langle k' | L | k \rangle|^{2}}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\langle k' | L | k \rangle|^{2}}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} - \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\langle k' | L | k \rangle|^{2}}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\langle k' | L | k \rangle|^{2}}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} - \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} - \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} = \frac{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|}{|\xi_{k'}^{(u)} - \xi_{k'}^{(u)}|} + \frac{|\xi_{k'}^{(u)} - \xi_$  $\xi_{R}^{(z)} = \sum_{k' \neq k} \frac{|\langle k' | L | k \rangle|^{2}}{\xi_{R}^{(u)} - \xi_{R'}^{(u)}}$  $= \frac{|U_{K}|^{2}}{\frac{\xi^{(u)}}{k} - \xi^{(u)}_{R+K}} + \frac{|U_{-K}|^{2}}{\xi^{(u)}_{R} - \xi^{(u)}_{R-K}}; \quad \text{where} \quad \langle k'|U|k \rangle - \frac{U_{K'-k}}{k'-k}$   $\text{when} \quad \xi^{(u)}_{R} = \frac{k^{2}k^{2}}{2m}, \quad \xi^{(u)}_{R+K} = \frac{k^{2}}{2m} |\vec{k} + \vec{k}|^{2}; \quad \xi^{(u)}_{R-K} = \frac{k^{2}}{2m} |\vec{k} - \vec{k}|^{2}$ Now since, the Potential is Very weak, than IUKI2 is very small w.r.t/. bhe energy differences in the denominators,

1'.e  $|U_{K}|^{2} = |U_{-K}|^{2} < |\xi_{R}^{(0)} - \xi_{R+K}^{(0)}|$  or  $|\xi_{R}^{(0)} - \xi_{R-K}^{(0)}|$ , so Ex is so small. In conclusion, far away from the edges of Who 1st B.Z (k=0, + 7/a), energy states are not affected by the Peniodic Potential.

Now let us see the effect of the Periodic Potential at Points very close or at zoncedge i.e at R= ± T/a. a ccording to Laue condition (R'- R = K or equivalently R-R'= IT), and in order for <k'|U|k> to be non-zero, we must have R'-R=F,50

R'= R+K =>  $\vec{R}' = (\vec{R} + \vec{R})^2 = (\vec{R} + \vec{R}) \cdot (k + \vec{R}) = k^2 + |\vec{R}|^2 + 2\vec{R} \cdot \vec{R}$  $k^2 = k^2 + k^2 + 2k^2 + k^2$ , but for elastic scattering  $k^2 = k^2 + 2k^2 + k^2$ 

Nohan  $|\vec{R}+\vec{R}'|^2 = (\vec{R}+\vec{R}) \cdot (\vec{R}+\vec{R}') = \vec{R}^2 + (\vec{R}^2 + \vec{R}^2 + \vec{$ 

=)  $\xi_{k+k}^{(u)} = \frac{1}{2m} |\vec{k} + \vec{k}|^2 = \frac{1}{2m} = \xi_k^{(u)}$  >>  $\xi_k^{(u)}$  and  $\xi_{k+k}^{(u)}$ 

are degenerate at zonc edge, and hance Ex

=>  $\mathcal{E}_{R-K}^{(0)} = \frac{h^2}{2m} |\vec{k}|^2 = \frac{h^2 k^2}{2m} = \mathcal{E}_{k}^{(0)}$  and  $\mathcal{E}_{R}^{(2)} = \frac{1}{2m} \frac{|L|R|^2}{2m} = \frac{1}{2m} \frac{|L|R|^2}{2$ 

Now when  $\xi_{R}^{(r)} \approx \xi_{R+17}^{(r)}$ , the Non-Leg Pert bheory will Note work any more. However, this case can be treated by the Legenerate Per. theory. So at Zone boundary, we have Ek = ER+18 and bhis happens when |R = |R+17 |. to fix bhis problem, we diagonalize It in the space of the stabes blad cause the degeneray. bhe  $R = \pm \frac{\pi}{a}$  are symmetric,  $|k+k|^2 = \frac{1}{n} =$ hen we have a 2x2 matrix for H < k| H|k> = < k| Ho|k> + < k| U|k> = En + L. < R+F 14/ R+F) = < R+F1 HO1 R+F7> + < R+F1 11/ R+F7> = ER+F7 + 10  $\langle \vec{R} + |\vec{r}| + |\vec{R}\rangle = \langle \vec{R} + |\vec{R}| + |\vec{R}\rangle + \langle \vec{R} + |\vec{R}'| + |\vec{R}\rangle$   $= \langle \vec{R} + |\vec{R}'| + |\vec{R}\rangle + |\vec{R}'| + |$ 

NON ( R+ 13 / 12) = 0 as they are orthogonal, so < k+17 1 H/k> = Ur and lines, < RIH R+17>= (R+17111) = UK So  $H = \begin{pmatrix} \xi_R + \mu_0 \end{pmatrix}$   $\mu_R$   $\mu_R$  = ( \( \xi\_{h} + \Li\_{0} \) \\ \( \xi\_{h} + \Li\_  $= \left( \begin{array}{ccc} \mathcal{E}_{R}^{(0)} + \mathcal{U}_{\bullet} & \mathcal{U}_{K}^{\dagger} \\ \mathcal{U}_{K} & \mathcal{E}_{R}^{(0)} + \mathcal{U}_{\bullet} \end{array} \right) \left( \begin{array}{c} |R\rangle \\ |R+K\rangle \end{array} \right) = \mathcal{E} \left( \begin{array}{c} |R\rangle \\ |R+K\rangle \end{array} \right)$  $= \frac{1}{2} \left| \frac{\xi_{k}^{(0)} + \mu_{0} - \xi}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^{(0)} + \mu_{0} - \xi} \right|^{2} = \frac{1}{2} \left| \frac{\mu_{k}}{\xi_{k}^$ => E = E(0) + LIO + | LIK zerobh first selond correbibus so due to ble fariblit weak Potential, the two states 12) and 18+47> refle each others; i.e 12) is Pushed down E = = Ex + Wo - I WITI, and the state 1 R+17) is rughed up with E= Ex+10+1UEI, 50 Charling agap of size DE= 2/UK/

similarly ab k=+x/a, mixing
1R> with 1R-17> would give bhe same result. 2 MH [ Find the 1st and 2nd -3 Example 1: corrections to energy due to the Constant Potential L(x)=Wo ; wo>o  $\mathcal{E}_{R} = \langle R | U(x) | R \rangle = \langle R | W_{0} | R \rangle = W_{0} \langle R | R \rangle = W_{0}$ ER = { | < k'| U(x) | k > | 2 k' + k - (6) NOW < K'| U(x) k> = wo < k'|k> = wo & k, k'  $=> \mathcal{E}_{R}^{(r)} = W_{0} \sum_{k' \neq k} \frac{\delta_{R,k'}}{\xi_{R}^{(o)} - \xi_{R'}^{(o)}} = D \text{ as } k' \neq k$ so the constant potential uniformly shifts all energy levels by we at first order. There is no 2nd order correction be cause the constant potential does not mix different R states. Hence, No energy gaps are created by bhis constant Potential. the energy for any state (k) is  $E_{R} = E_{R}^{(1)} + E_{R} = \frac{k^{2}k^{2}}{2m} + W_{0}$ ab R=0 => ER=Wo and  $k = \pm T/a =$   $E_R = \frac{k^2}{2m} (\pm \frac{\pi}{2})^2 + w_0 = \frac{k^2 \pi^2}{2ma_1^2}$ 

Example 2: consider she periodic potential W(x) = 2Vo cos (27x) of 1D solid with labbic spacing Find Ex, Ex, and bhe energy gap created at R=± T/a. Now let 1k>= \frac{1}{\ta}e^{ikx} => \langle k| = \frac{1}{\ta}e^{-ikx}, so En = < RIU(x) | h> = 1 ( = ihx 2 Vo cus 27 x e ihx dx =  $\frac{2 \text{Vo}}{a} \int_{0}^{\infty} dx \cos(\frac{2\pi}{a}x) = 0$ , as we integrate the cos over ibs Period  $\frac{2\pi}{2\eta_{a}} = 9$  $\xi_{k}^{(2)} = \sum_{k' \neq k} \frac{|\langle k' | U(x) | k \rangle|^{2}}{|\langle k' | L_{k'} \rangle|^{2}}; \text{ when } |k' \rangle = \frac{1}{\sqrt{a}} e^{ik'x}$ Now (k'| L(x)|h) = 200 (e-ik'x as 23x e ikx dx = 2 \( \frac{2 \quad \qu =  $\frac{V_0}{a} \left( e^{-ikx} \left[ e^{i(R+\frac{23}{a})x} + e^{i(R-\frac{23}{a})x} \right] dx$  $= \frac{V_0}{a} \left[ \int_{0}^{a} e^{-i(R' - (R + \frac{2\pi}{a}))X} + \int_{0}^{a} e^{-i(R' - (R - \frac{2\pi}{a}))X} \right]$ - vo [a 8k', R+23 + a8k', R-23] only two terms contribute from k'=k+23 and k'=k-23

$$= \frac{|V_0|^2}{\xi_{h}^{(0)} - \xi_{h+2\pi}^{(0)}} + \frac{|V_0|^2}{\xi_{h}^{(0)$$

Now 
$$\langle R|H|R \rangle = \langle R|H_0+U(x)|R \rangle$$

$$= \langle R|H_0|R \rangle + \langle R|U|R \rangle$$

$$= \langle R|H_0|R \rangle + \langle R|U|R \rangle$$

$$= \langle R|K|R \rangle = \langle R|K \rangle$$
and
$$\langle R|H|R \rangle = \langle R|H_0+U|R \rangle + \langle R|K|U|R \rangle$$

$$= \langle R|H_0|R \rangle + \langle R|K|U|R \rangle$$

$$= \langle R|R \rangle + \langle R|R \rangle + \langle R|R \rangle + \langle R|R \rangle$$

$$= \langle R|H_0|R \rangle + \langle R|R \rangle + \langle R|R \rangle$$

$$= \langle R|R \rangle + \langle R|R \rangle + \langle R|R \rangle$$

$$= \langle R|H_0|R \rangle + \langle R|R \rangle + \langle R|R \rangle$$

$$= \langle R|R \rangle + \langle R|R \rangle + \langle R|R \rangle$$

$$= \langle R|R \rangle + \langle R|R \rangle + \langle R|R \rangle$$

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$$= \langle R|R \rangle + \langle R|R \rangle$$

$$= \langle R|R \rangle + \langle R|R \rangle +$$

$$| \frac{1}{a} | \frac{$$

$$=> E^{\dagger} = E_{R}^{(6)} + V_{0}, \quad E^{\dagger} = E_{R}^{(6)} - V_{0}$$

$$PE = E^{\dagger} - E^{\dagger} = 2V_{0} \equiv \text{band gap}$$

Let us find ble eigenstates: ble eigenvolue equabion reads  $H\left(\frac{1k}{|k-h|}\right) = \mathcal{E}\left(\frac{1k}{|k-h|}\right) \Rightarrow \left(H-\mathcal{E}\right)\left(\frac{1k}{|k-h|}\right) = 0$ y called eigenvector  $= \left( \begin{array}{ccc} \xi_{\mathsf{N}}^{(0)} - \xi & V_{\mathsf{O}} \\ V_{\mathsf{O}} & \xi_{\mathsf{N}}^{(0)} - \xi \end{array} \right) \left( \begin{array}{c} |h\rangle \\ |h-|\tau\rangle \end{array} \right) = \left( \begin{array}{c} 0 \\ 0 \end{array} \right)$  $for \ \mathcal{E}_{+} = \mathcal{E}_{R}^{(0)} + Vo = \begin{cases} \mathcal{E}_{R}^{(0)} - \mathcal{E}_{R}^{(0)} - V_{0} & V_{0} \\ V_{0} & \mathcal{E}_{R}^{(0)} - \mathcal{E}_{R}^{(0)} - V_{0} \end{cases} \begin{pmatrix} |R|^{+} \\ |R-|T|^{+} \end{pmatrix} = \begin{pmatrix} 0 \\ 0 \end{pmatrix}$  $= \left( \begin{array}{ccc} -V_0 & V_0 \\ V_0 & -V_0 \end{array} \right) \left( \begin{array}{c} |R\rangle^+ \\ |R-R\rangle^+ \end{array} \right) = \left( \begin{array}{c} 0 \\ 0 \end{array} \right) = \left( \begin{array}{c} -V_0 \\ 0 \end{array} \right) = \left( \begin{array}{c}$ => bobh arbitrary, =>  $|k-H>^+=|k>^+$  $= \left( \begin{array}{c} |k\rangle^{+} \\ |k\rangle^{-}| \end{array} \right) = \left( \begin{array}{c} | \\ | \end{array} \right)$ Now the eigenstate can be written in terms of the eigenvectors as  $|\Psi\rangle = \frac{1}{|\pi|} |k+|\pi\rangle e^{i(R+|\pi|).r^{2}} = \frac{1}{|\pi|} |k+|\pi\rangle e^{i(R+|\pi|).r^{2}} = \frac{1}{|\pi|} |k+|\pi\rangle e^{i(R+|\pi|).r^{2}}$ =  $\frac{1}{(k+\frac{2\pi}{a}n)} \times e^{i(k+\frac{2\pi}{a}n)} \times e^{i(k+\frac{2\pi}{a}n)} = \frac{1}{(k+\frac{2\pi}{a}n)} \times e^{i(k+\frac{2\pi}{a}n)} \times$ ab k = T/a, we have -iT/ax  $|Y\rangle = |k\rangle e^{iT/a} + |k-t\rangle e^{iT/a}$ , So for  $\xi_{+}$ , we have  $|V\rangle^{\dagger} = |h\rangle^{\dagger} e^{i\frac{\pi}{4}x} + |k-\pi\rangle e = e + e = 2\cos\frac{\pi}{4}x$ 

let us normalize | 4>+ using < y+| 4+>=1 let 14> = 2A as Ix; A is normalization constant =>  $< V^{\dagger} | V^{\dagger} > = 1 => HA^{2} \int cos^{2} (x) dx = 1 => 2A^{2} L = 1$ =>  $A = \sqrt{21}$  $\begin{vmatrix} \delta \\ \frac{1}{2} \end{vmatrix} \Rightarrow A = \sqrt{\frac{1}{2}}$  $= |Y|^{t} = \frac{2}{\sqrt{2L}} \cos \frac{\pi}{2} x = \sqrt{\frac{2}{L}} \cos \frac{\pi}{2} x$ similarly for E = Ex -Vo, one finds 14>= 1&>e 17/4x + 1h-K> e = e -e -e  $|x-y| = -2i \sin \frac{\pi}{4}x$   $|x-y| = -2i \sin \frac{\pi}{4}x$  $\langle Y^{\dagger}|Y'\rangle = \frac{2L'}{L} \left( \cos \frac{\pi}{2} x \sin \frac{\pi}{2} x dx = \frac{-2ia}{2\pi L} \left[ \cos \left(\frac{\pi L}{a}\right) - 1 \right) \right)$  $= -\frac{iq}{\pi L} \left[ \left( \cos N\pi \right)^2 - 1 \right] = -\frac{iq}{\pi L} \left[ \left( \cos (N\pi) \right)^2 - 1 \right]; N is in begin$ but L=Na

Tightly Bound dectrons and The tight Binding modes
in general, then are several unabhoods of band structure
Calculations, such as Parturbation uncolond tight-binding
wellhod, Pseudo Potaniel mebled; and obliers. in the last section,
, we discussed the Perturbabish method, where we assumed
that electrons are delocalized and move avound we discusse
that electrons are delocalized and move around the sample under a periodic potombial. in this section, we discusse under a periodic potombial. In this method is also called the bight-binding method. This method is also called
linear compination of about orbitals ( LCAO).
hon, we will solve ble singe particle schrodinger equation
for the states in a crystal by expanding Block states in term
1 1 is a white the
1 a chank of all will
here we consider a 1D bight-Binding wave function) of specing a porbital (atomic wave function)
$\int \int \int \int \int da \rightarrow a$
- moderations  Im> L=Na   Note to the site of the moderations  Im> Divine notations
1 / January St Per
aromic orbitals are localized around about sites
atomic orbitals are localized around about my with $\langle \vec{r}   m \rangle = \phi(\vec{r} - \vec{R}m)$ ; $\vec{R}m$ position of nucleus $m$ with $\langle \vec{r}   n \rangle = \phi(\vec{r} - \vec{R}m)$ ; $\vec{R}n$
wir on black bluese orbitals are hardly overlap, so

 $\langle m|n\rangle = \int f(\vec{r} - \vec{R_m}) \phi(\vec{r} - \vec{R_n}) d^2r \approx 0$  almost zero for  $m \neq n$ and for m=n, (m/m)= \[ |\phi(\varphi-\varR\_m)|^2 d'v = 1 So (m/n) = 8mn, i.e bhebe orbitals are orbhonon - Now for an isolabed atom (labeled m), the atomic ham: Itonian Itab is Itab = IT + Vm = \frac{P^2}{2m} + Vm Hab Im) = \xi\_{ab} Im)

(K+Vm) Im) = \xi\_{ab} Im) clearly, a single atomic orbital does not solven Sabit by Block's bheorem. However, if we make ... a linear combination of abomic orbitals, then the regulbant Tight binding wavefunction does satisfy Block's theor. let us check first blu wormalization just symbols I= < 4nk2(r) | 4nk2(r) > = ) 4nk luk d'r? = 1 ZZ ( -ik. R \* (-R) e ik. R') d'r

where R, R' sums over all translation vectors 0 -> N-1 or 1->N Now using  $\int d^{\dagger}_{m}(\vec{r}-\vec{R}) d_{n}(\vec{r}-\vec{R}') d^{\prime}_{r} = 1, \text{ when } m=n \text{ and } \vec{R}=\vec{R}'$  = 0, otherwise $\frac{1}{N} \sum_{R} \sum_{R} e^{iR^{2} \cdot (R^{2} - R^{2})} \delta_{R,R}$  $=\frac{1}{N}\sum_{R}e^{0}=\frac{1}{N}\sum_{R}I=\frac{1}{N}\left(\sum_{R}I\right)=\frac{N}{N}=I$ So eq h (1) is correctly nor modified. Let us also check black eq (1) does satisfy Black's toheoven => Ynk(r+R') = \frac{1}{\nu R} \in e^{i\vec{R}.\vec{R}} \pha\_n(r-\vec{R}+\vec{R}') = \frac{1}{\nu R} \in e^{i\vec{R}.\vec{R}} \pha\_n(r-\vec{R}'-\vec{R}')  $= \frac{1}{\sqrt{N}} \sum_{R''} e^{i\vec{R} \cdot (R' + R'')} \phi_{n}(\vec{r} - \vec{R}'') = e^{i\vec{R} \cdot \vec{R}'} \sum_{R''} e^{i\vec{R} \cdot \vec{R}'} \phi_{n}(\vec{r} - \vec{R}'')$ y R" is dummy 10b R"→R = e i k. R' Ynk (r');

Now again, let us consider one dimensional labbice with one atom/cell and let us first have one eletron in the chain. i.e, then is one dectron localized on atom (m), and let us imagine bhat it can hop between these mulit. 11> p) (r-R)

L=Na So bhe total hamiltonian of the electron is the total potential felt by the electron on site |m> that is produced by all other atoms including the

= tr + \(\frac{1}{2}\); just abbrévialion i't is usefull first to consider, the electron interacts with a single muchies (m) and isolable blus from blu sum, so

1'e (+ + Vm)1m> = Eab 1m); bhis is for an isolabed about

Now relate that the condition (m/n) = 8mm is good only when nuclie an four from each other. For simplicity we will go with this assumption. Now Record What

et (1) can be wribben into Dirae Notation as

(1) can be written into pirae ik. R ikna

(N) = Cn (n) ; Cn = In e = In e in 1)

(N) # of about or lastice

N! # of about or lastice

N! # of about or last its

Now schoolinger eg n reads H 14>= & 14> i'n mabrix form and using 14>= & cm 1m>, we set EH Cm/m = E E Cm/m ; mulbiply by <n/r

Folial bhe motory

M (n) H/m > Cm = E E Cm <n/m > elements Hum

Sum | E Hnm Cm = E Cn | ---- (2) Now weed to find bhe materix elements Hom. Now the fall hamiltonian. H= IT+Vm + ZV; where I removed the about on from the sum ; now take bhe  $H|m\rangle = (|T+V_m|)|m\rangle + \sum_{j\neq m} V_j |m\rangle$   $\sum_{k=1}^{\infty} |m\rangle$ inner product with <nl Eat Im> <n11+1m> = Eab <n1m> + <n1 \( \frac{\xeta}{J+m} \) (n) | | | = Eab Sum + (n) \( \frac{\xi}{\text{th}} \) = \( \text{vilm} \) = \( \text{curry} \) \( \text{other undike} \) \( \text{vilm} \) = \( \text{vilm} \) = \( \text{other bin of all nuclike} \) \( \text{other undike} \) \ + <n/ \( \frac{5}{1} \rightarrow \frac{1}{2} \rightarr

and (n/ { Vilm > = } Vo, --- as indicated before 1-k; In+m Hopping, it represe the transition amplitude for an electron to hop from initial - Now, to a good approximation state Im> to final state We can assume nearest 1 ns Kircis } hegabire. This is neighbor hopping si'c O; obherwise due to that when if | n-m> 1 => No holfing \ allows get close } the Potential barrier is reduced in 5° (n/ ½ V)/m) = { Vo; if n=m± 1) order to minimize ive it is hard for an dectoron to hopp back and o, other with energy and make the z system stabl forbh between sites black of m are for away. m+1 t ( Sn, m+1 + Sn, m-1) -- (3) So Ham = (Eab + Vo) Sum hopp one she p huppone to left let us assume while where step to rijht i's N mielie, so It is NXN mabrix, bha  $\varepsilon_o$  is called on – site energy  $H = \begin{pmatrix} z_0 & -t & 0 & 0 & --- \\ -t & z_0 & -t & 0 & 0 & --- \\ 0 & -t & z_0 & -t & 0 \\ 0 & -t & z_0 & -t & z_0 \\ 1 & 0 & 0 & -t & z_0 \end{pmatrix}$ 

Now Plussing -9 " (3) into (2) and using ch= i kan we se E Hum Cm = E Cu or Cn= u e ikan 5 constant of £ [ ε δ δ mm - t (δ n, m+1 + δ n, m-1)] cm = ε ch normalization Eu Cn - E ( Cn-1 + Cn+1) = ECn  $\frac{1}{\sqrt{N}} \xi_{0} e^{ikqn} - t \left( \frac{1}{\sqrt{N}} e^{ikq(n-i)} + \frac{1}{\sqrt{N}} e^{ikq(n+i)} \right) = \xi \frac{1}{\sqrt{N}} e^{ikqn}$ Everikan teikan (e-ika teika) = Ee ikan € - 2 t cos (ka) = € E= Eo - 26 Ces(kay) dispersion relation Note that there is one energy eigenvalue for each h, so there is only one band. Let us plat &-& vs k

4-4 E-Es = -26 cs (Ray in John 15th Bit band widle h = 26 - (26) ー リセ for large enough 1D lattice bandwidth: the difference in energy between the highest and lowest allowed levels within a given band

Weaking) Sharbing with a Single abon, we set zero band wielth, Now by adding more about to 6h labbice, 6h bandwidth Jets -> sinde abom with sharpe Nugars; only 5 Eo-2161 -> atomic spaling

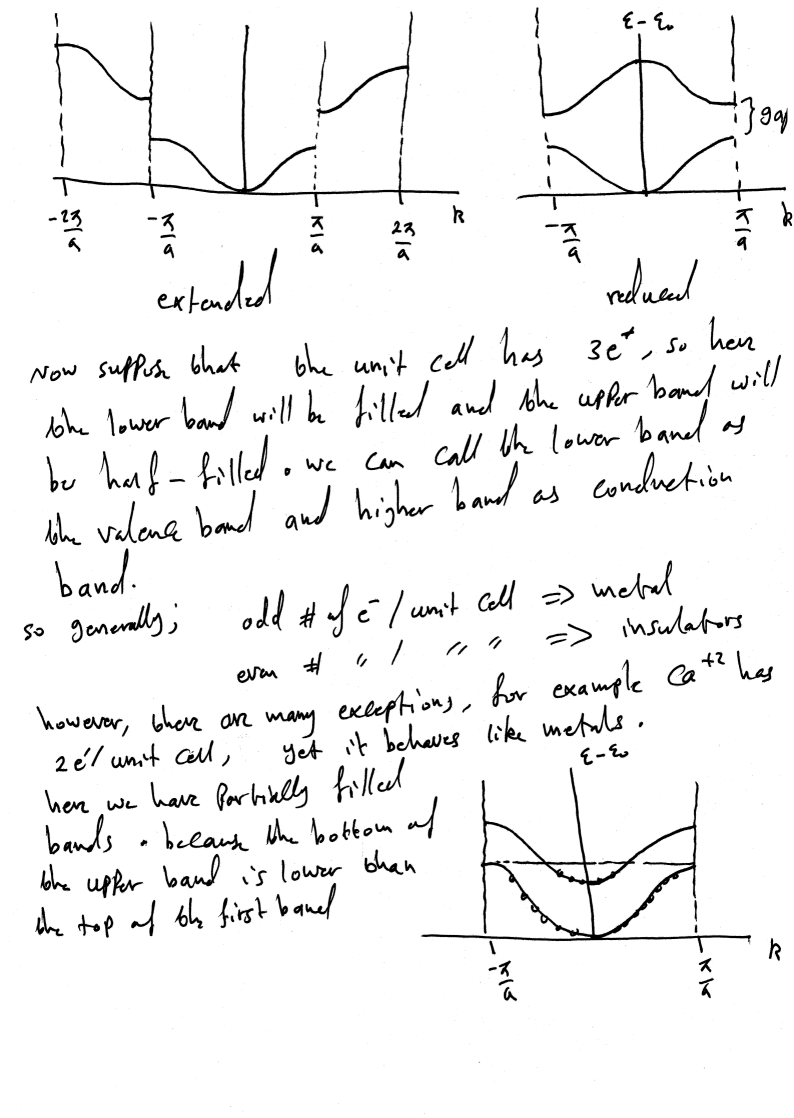
let us look at the shape of the band near R=0. Now using Cosx=1-x2, for xcc1, we set Cus (kg) = 1- (kg)/2 5. E= E\_0-26 (1- (ka)2) = E\_0-2b+ka2k2 so it is quadratic in R i.e EXR2 (parabolic) so the electron near the bottom of the band moves like free electron with an effetive mass mt defined as  $\frac{k^2 k^2}{2m^4} = (a^2 k^2) \implies m^4 = \frac{k^2}{26a^2}$ Now each Mowed 1k) state can have 2 electrons with opposite spins. so having one electron/abom means, there are N deetrous what ab T=0 fill only half of the band. so the fermi surface here is composed of two points Recall that the # of available R = + 1 states in the band is N, which can accommodate up to 2N electrons of the atoms are divalent, then When will be 2N elections that can completely fill the band so fermi surface is bounded by R=II

so for monovalent atoms, where there is only one valence electron for each atom, we have a half-filled band wit an empty states available above Ef. Hence electrons can easly be moved across Ex by an applied dectric field and produce accurrent. Where materials are called metals; with heat capacity CXT. - for divident atoms, when then are 2 valence electrons! about, we will have a completely filled band, so there are no empty states available a bour Ef for concluebili These materials are called insulators with zero heat

These materials are called insulators with zero heat

This is theoretical prediction of the TBM based on non-overleaping bands. However, in realty, most divalent solids (like Ca and Mg) are metals due to band overlap

A.L. I - Now So for we assumed that then was one orbitalor atom and hence we got one band. Now bhen are some situations, where there are more than one orbital atom; lite for example having two different atoms /unit cell or having two different orbitals on the same atom ( 15 and 2P for example). - for 2 orbitals / want cell; and without so bing Whis problem, Non spectrum is like as follows when there are two energy bands, separated by a gap



The general expression for bhe tight-binding dispersion uf a monoatomic lablice is given by  $\mathcal{E}(\mathbf{k}) = \mathcal{E}_{0} - \mathcal{E}_{i} \, \mathbf{e}_{i} \, \mathbf{k} \cdot (\mathbf{k}_{i} - \mathbf{k}_{i})$ Now considering only nearest - neighbor hopping yields  $\mathcal{E}(k) = \mathcal{E}_0 - E \mathcal{E}_0 = i \vec{k} \cdot \vec{R}$ ; where  $\vec{R}$  is a vector connecting agiven labbice site to one of its nearest neighbors. - for a squan labbice, R= tai; and R= taj => E(k) = E - t (eikxa - ikxa + eikya)  $= \mathcal{E}_{o} - 2t \left( \operatorname{Ccs} k_{xa} + \operatorname{Ccs} k_{ya} \right)$   $\varepsilon^{\max} \text{ when } \operatorname{cosk}_{x} a = \operatorname{cosk}_{y} a = -1, \ \varepsilon^{\max} = \varepsilon_{o} + 4t$   $\varepsilon^{\min} \text{ when } \operatorname{cosk}_{x} a = \operatorname{cosk}_{y} a = +1, \ \varepsilon^{\min} = \varepsilon_{o} - 4t$ - for a simple - cubic lattice, it yields E(K) = & - 2t (Cos kxa + Cos kya + Cos kza), bandwidth = 12t - in Homework #6, we discuss formation of bound gaps in diatomic chain ABABAB--- and diatomic squar labbile with on-site energies EA and EB, respectively.

I need to go over problem 4 in HW#6 to make sure you understand how the bandgap forms and how to determine its magnitude for the 1D diatomic chain of atoms ABABAB.....

Tight Binding Density of states: we already found that the DOS of all bands for asolid in d-dimensions was  $D(\varepsilon) = \frac{2\ell}{(2\pi)^d} \frac{2\pi}{7(4/2)} \frac{d-1}{|\nabla \varepsilon_R|}$ L= Na for 1D solid of spacing a, & = & -2 t Cos(ka); t>0 So  $D_1(\Sigma) = \frac{2L}{2\pi} \frac{2\sqrt{\pi}}{\sqrt{\pi}} \frac{1}{|\nabla_R \Sigma|} = \frac{2L}{\pi} \frac{1}{|\nabla_R \Sigma|}$ Now  $\nabla_{\mathbf{k}} \mathcal{E}(\mathbf{k}) = \frac{\partial \mathcal{E}}{\partial \mathbf{k}} = +2\mathbf{k} \sin(\mathbf{k}\mathbf{a}) \cdot \mathbf{a} = 2\mathbf{a} \mathbf{t} \sin(\mathbf{k}\mathbf{a})$ =>  $D_1(\epsilon) = \frac{dL}{\pi} \frac{1}{2a + \sin(ka)} = \frac{L}{\pi a + \sqrt{1 - \cos^2(ka)}}$ ; where  $\sin(ka) = \sqrt{1 - \cos^2(ka)}$ Now using coska =  $\frac{\varepsilon_0 - \varepsilon}{2t}$  =>  $P_1(\varepsilon) = \frac{L}{\pi at} \frac{1}{\sqrt{1 - (\varepsilon_0 - \varepsilon)^2}}$ Ht<sup>2</sup> => P1(E) = L 1 Tat \[ 1 - \left(\frac{\\ \ 2b}{2b}\right)^2\] Note that  $D_1(x)$  is singular  $(\infty)$ , when  $\left(\frac{x_0-x_1}{20}\right)^2=1$ => \( \frac{\xi\_{0} - \xi\_{0}}{2\tr} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) => \( \xi\_{0} - \xi\_{0} = \pm 1 \) = \( \xi\_{0} - \xi\_{0} = \xi\_{0} = \pm 1 \) = \( \xi\_{0} - \xi\_{0} = \pm 1 \) = \( \xi\_{0} - \xi\_{0} = \xi\_{0} = 1'-c & = & - 26 and & + = & + 24 are blue singular points when Dos diverges

also Note black PI(E) i's E = Eo, 50 PI(E) Now how many states are there Es-26 between & - 2 b and & + 2 b 2 t cosodo => 0 = 7/2 when 2= 20+26 => sivo = -1 = 2 = [ Th - (-7/2) ] = 2 = X as expected.

Please go over problem 6 in HW#6 to find out how to calculate DOS of 1D diatomic solid with two different atoms per unit cell ABABAB....

## Block and Wannier representations: (optional)

We have seen that a bloch electron is conventionally labeled by an index n and awave vector  $\vec{R}$ , now in the right-binding model, the TB wave function was expande in terms of atomic orbitals  $\phi_n(\vec{r}^2-\vec{R}^2)$  as  $\psi_n(\vec{r}^2-\vec{R}^2) = \frac{1}{\sqrt{N}} \sum_{\vec{R}} e^{i\vec{R}\cdot\vec{R}} \phi_n(\vec{r}^2-\vec{R}^2)$ ; ---(5)

bhore, we have made abad assumption bhab the atomic orbitals, are orbhogonal i'e  $\int \phi_m(\vec{r}-\vec{R}_m)\phi_n(\vec{r}-\vec{R}_m) = 0$ 

However, bhis assumption is satisfactory when atoms are for away from each other, bhat thier orbitals are not overlapsed. However, in 1937, Gregory wannier professel ar a new set of functions bhat are perfetty localized ab abomic sites and orthogonal. These set of functions are called wanhier functions Wn (r), R). so bhob  $V_{nR}(\vec{r}) = \frac{1}{N} \sum_{\vec{R}} e^{i \vec{R} \cdot \vec{R}} w_n(\vec{r}, \vec{R})$ , Note that  $w_n(\vec{r}, \vec{R})$ the right-binding wavefunction is now is blu Fourier fransform of Bloch war function;

Who (P,R) = IN E e YNR (r) -- (6)

Who (P,R) = VN R

NOW using block Chewren for Ynk (12) = eikorite (12), 6ha of (6) reads wn (r) R) = 1 & e-ik. R eik. r Unk (F) = 1 & e & (r-R) Unk (r) bhus, baking into account the periodicity of unk (r2), we See that Wn (rik) defends on the difference r-k, so  $\omega_n(\vec{r},\vec{R}) = \omega_n(\vec{r}-\vec{R}),$  so Ynk(r)= 1 & eikik wn(r-R) ---- (7)  $W_{n}(\vec{r}\cdot\vec{R}) = \frac{1}{NN} \sum_{k} e^{-ik\cdot\vec{R}} V_{nk}(\vec{r}) --- (8)$ The wappier functions form an orbhonormal set, i'e  $\int d^{3}r^{3} \omega_{n}(\vec{r},\vec{k}') \omega_{m}(\vec{r},\vec{k}')$   $= \int d^{3}r^{3} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$   $= \int_{N} \sum_{R} \sum_{R} \frac{1}{N} e \qquad \forall_{n} R(r) \forall_{m} k'(r)$ = 1 \( \frac{1}{R} \). \( \frac{1}{R'} - \frac{1}{R'} \) \( \frac{1}{R'} -= I (\( \Secold \Secold \text{Nnm}\)
= \( \lambda \) \( \lambda \) \( \text{Recold blab} \), who
\( \text{X} \) \( \text{R} \) \( \text{R} \) \( \text{R} \) Recall What, when R = R'  $\sum_{R} 1 = N$ 

let us switch to Divae notations with

(r/R) = Ynk(r) => 1R> = 1 & e ik.R? eq (q) in

Divae not  $\langle \vec{\mathbf{r}} | \vec{\mathbf{r}} \rangle = W_n(\vec{\mathbf{r}} - \vec{\mathbf{r}}) = \langle \vec{\mathbf{r}} | \vec{\mathbf{r}} \rangle = \frac{1}{\sqrt{N}} \sum_{\vec{\mathbf{r}}} e^{-i\vec{\mathbf{r}} \cdot \vec{\mathbf{r}}} \hat{\vec{\mathbf{r}}} \hat{\vec{\mathbf{r}}} = e^{-i\vec{\mathbf{r}} \cdot \vec{\mathbf{r}}} \hat{\vec{\mathbf{r}}} \hat{\vec{\mathbf{$ Note bhat bobh 1k) and 1k) are orbhonormal (k'1k) = 1 2 2 e - i R'.R' e i R'.R' (R' | R)  $= \frac{1}{N} \sum_{R}^{\infty} \sum_{R'} e^{-iR' \cdot R' + iR \cdot R} = \frac{1}{N} \sum_{R'} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R' \cdot R'} = \frac{1}{N} \sum_{R' \in \mathbb{N}} e^{-iR' \cdot R'} = \frac{1}{N}$ and similarly

(R'IR) = \frac{1}{N} \frac{\x}{R'} \frac{\x}{R'-R'} \frac{\x'}{R'-R'} \frac{\x'}{RR'} \frac{\x' Now as mentioned early, blue deetron wave function can be expanded in terms of warnier functions; i've expanded in terms of warnier functions; i've  $\Psi(r) = \sum_{R} C(R) W(\vec{r} - \vec{R})$  or is  $147 = \sum_{R} C(R) |\vec{R}\rangle$ so  $H |\Psi\rangle = \sum_{R} |\Psi\rangle = \sum_{R} |\Psi\rangle = \sum_{R} |\Psi\rangle$  $\mathcal{E}_{R'}(CR')(\overline{R'}) + \mathcal{E}_{R'}(CR')(\overline{R'})$ mulbiply by < R => \( \int \c(R') \) \( \text{HRR'} = \int \c(R) \) \( -- \) (9)

Now using the completnee of wannier functions E ir/ (RI = 1, ble hamiltonian H can be written as  $H = \sum |R\rangle\langle R| H \sum_{R} |R'\rangle\langle R' = \sum_{R} \sum_{R} \langle R|H|R'\rangle|R\rangle\langle$ H = E,R, HRR' 1R> <R'1 a gain the 1D atoms chain can be solved using 6his H.

let us call or label ble wannier functions by 10>= 1R) Now med. 11 TO 1 1 Now using the TB hamiltonian in materix form (eq 3), we get H = 80 \[ 16>(1) + 6 \[ 16>(1-1) + 16>(1-1) Now plugging bhis H into HIV>= E14> and using

14>= \( \frac{2}{2} \) C(16), we get 2. \( \langle => \( \( \( \) \( => 40 CL + K (CL+1 + CL-1) - ECL = 0

 $\frac{\mathcal{E}}{\nabla N} = \frac{1}{2} \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \right) \right) \right)}{2} + \frac{1}{2} \left( \frac{1}{2} + \frac{1}{2} + \frac{1}{2} \left( \frac{1}{2} + \frac$