The band structure is given by the eyenstates 4 (r) with energy Eqn where g is one of the N allowed wave veetors in the 1st BZ consistant with the Born-Von Karmen bouday conditions, and n is the integer valued band index Since each nth BZ also and ins exactly N allowed wave vectors \overline{k} , and since each \overline{k} can be written Uniquely as $\overline{k} = g + \overline{k}$, where \overline{g} is in the 1st BZ and K is the RL vector that I is closest to, we could also represent the state Gn as the state 4, (r) where k is in the oth BZ and k is obtained from if by making the needed translation $k = \overline{q} + \overline{k}$ to pert q noto the non BZ.

Representing the band sourtwe as then is called The "Reduced Zone Scheme", Representing this state by the with I in the nth BZ is called the " Extended zone scheme", More to say about the soon!

Connext - For free electrons the energy eigenstates one specified for by a wave verter to that can take a value anywhere in k-space. For an electron in a periodic potential, we said eigenstates should be specified by a discrete band index n and a crystal momenta of that must lie only in the 1st BZ. But if we take the problem of the periodic potended ad let TS >0, we recorder The fill electron case. So how con these two ways of tabeling eigenstates both be valid? Answer: We can represent even free electron states wany the bird nidex n ad 1st BZ & as granten nubers. For any k, anywhere in k-space, we can always write $\vec{k} = \vec{g} + \vec{K}$ where \vec{K} is the R.L. vector closest to \vec{k} ad of therefore must be in the 1st B.Z. This follows from the equivalence of all points in the R.L. (since it is a Bravais latter) and from the definition of the 1st BZ as the set of nove vectors doson to the origin than to any other R in the R.L.

Then for any h= g+ K in the nothe Brillow Zone write $\mathcal{E}^{\circ}(\overline{k}) = \frac{\pi^2 k^2}{2m} = \frac{\pi^2 (\overline{\xi} + \overline{k})}{2m} = \frac{\mathcal{E}_n(\overline{\xi})}{\pi}$ energy at crystal momentum og mi band index "n" This the band index gives the B.Z. in which the correganding free election wave vector to lies. Ex: in a 1D B.L. of lattice constant a R.L. vectors are $t_n = (\frac{\pi}{a})n$, $n = 0, \pm 1, \pm 2, -...$ Bragg planes at $k = \frac{Kn}{2} = (\frac{T}{a})n$ 4 Bragg planes can translate the section of cure in Ath BZ back into 3/2 free electron E°(h) 1st BZ by subtrach K = K = 0 $K_1 = K_2 = k$ K = K = 1 = K = 1 = 1 $I^{st}BT = 1$ $Z^{rd}BT = 1$ $Z^{rd}BT = 1$ $Z^{rd}BT = 1$ $Z^{rd}BT = 1$ & appropriate K from Te 3rd BZ representing free Mection states 1st BZ by I in 1st BZ. segnents of curve labeled nor n' Tutegers label correspondy sections of the curves in the Came from the nth two equivalent representations BZ of free electrons

Comments.

O Because to entere the matrix equation as a parameter via the terms Extk, we expect that The eigenvalues En (h) are smooth continuous functions of k. (2) Because En(k) à periodic m k, En(k+k)=E(k), must have we support that En (Ti) has a maximin and a minim at vories continuously between then. This is why En (Th) is referred to as on energy band (3) Since $4_{kn}(\vec{r}) = \sum_{k} e^{i(k-k)\cdot \vec{r}} - C_{k-k}$ is no larger a pure plane wave, 4kn 5 NOT an eigen state of momentum. This is reasonable since with the periodic potential turned on the system no longer has translational invariance. However the system does have invariance in translations by B.L. vectors R. This is why 4kn mixes only wavevectors related to k by k'=k-K for Kin R.L. Because of the defference, h is called the "crystel momentum" -it is not the true momentum. In collosions, the

custol the momentum of an electron is perforg "conserved" modulus a R.L. vector K.

D Because 4/ Rn (F) and En (F) are periodic in k-space, the states k ad k+K are not physically different. Neverthe less it is sometics convenient to represent states by a k that is not in the 1st BZ. This is most often done when one wants to make closer analogy to fill electrons. One therefore has three annonly used schemes for represhity band structure. We illustrate these in ID Reduced Zone scheme ; restrict to 1st BZ $E_3(k)$ for all energy bands n Ez(k) TEILL Tak Repeated tone scheme : show all values of the with En (Th) periodec in Th $\mathcal{E}_{s}(k)$ (E2(4) TEICH Ø 古 -21 -II a a ett BIT

Extended Zone scheme : show energy bad n for using k vectors in nth BZ. ϵ_{2i} $\frac{1}{2\pi} - \frac{1}{2\pi} + \frac{1}{\pi} = \begin{bmatrix} 0 & \pi & 2\pi & \pi \\ \pi & \pi & \pi & \pi \end{bmatrix} = \begin{bmatrix} \pi & \pi & \pi \\ \pi & \pi & \pi \\ \pi & \pi & \pi \end{bmatrix}$ Note: However one represents the bad structure, each band contains only N allo allowed values of k consistant with the Born - von Korman baunday conditions. => each band can contain 2N électron states (2 ber spin Tart) N= # tors in crystal (= # cons q we have a single BL and not a tattice with a basis) (5) If one computers the expectation of velocity in a Block stat electron state, one finds $\overline{U}_{n}(\vec{k}) = \langle \Psi_{nk} | \frac{\pi}{2} \overline{V} | \Psi_{nk} \rangle = \int d^{3}r \ \Psi_{nk}(\vec{r}) \ \pi \overline{V}_{nk}(\vec{r})$ proof follows. Marin point is that the eigenstates of an electron in a periodec potential, although they are not legenstates of momentum, nevertheless have a

well defined, generally finite, value for their average velocity. Therefore, in spite of the interaction of the electron with the periodic potential, the electron in its eigenstate moves with a steady velocity even if no net force (for example from an electric field) is applied to it. This is contrast to Ducke's classical perfore in which collesions between The election of the cons served to reendomize any initial electron velocity on the collision time scale I . The reason Drude's puture does not hold is because of the wave nature of the electron and its coherent scattering of the cons,

$$\frac{A \text{ non perturbative proof for } \overline{V_{F}} = \frac{12E}{R + 2E}$$

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$$\frac{12E}{R}$$

$$\frac{12E}{R} = 12E \text{ Bloch function is known to be of the form [ignore book inder]} = \frac{1}{V_{E}(\vec{x})} = -e^{i\vec{k}\cdot\vec{x}} = U_{E}(\vec{x})$$

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.

Since
$$H_{-\overline{K}}$$
 is Hermitian,

$$\int_{\Omega} \frac{\partial U_{\overline{K}}}{\partial \overline{F}} H_{-\overline{K}} U_{\overline{K}}^{*} d^{3}\overline{x} = \left[\int_{\Omega} U_{\overline{K}} H_{-\overline{K}} \frac{\partial U_{\overline{K}}}{\partial \overline{F}} d^{3}\overline{x}\right]^{*}$$

$$= \int_{\Omega} U_{\overline{K}}^{*} H_{\overline{K}} \frac{\partial U_{\overline{K}}}{\partial \overline{F}} d^{3}\overline{x} \qquad (7)$$

$$J_{huo}, (3), (6) \text{ and } (7) \text{ together imply}$$

$$\int_{\Omega} \left[\frac{\partial U_{\overline{K}}}{\partial \overline{F}} H_{\overline{K}} U_{\overline{K}}^{*} + U_{\overline{K}}^{*} H_{\overline{K}} \frac{\partial U_{\overline{K}}}{\partial \overline{F}}\right] d^{3}\overline{x} = 0 \qquad (8)$$
However, (1) and (3) directly imply

$$\mathcal{E}_{\overline{K}} = \int_{\Omega} U_{\overline{K}}^{*} H_{\overline{K}} U_{\overline{K}} d^{3}\overline{x} \qquad (9)$$
Differentiate eq. (7) with respect to \overline{K} and exploit (8) we have

$$\frac{\partial \mathcal{E}_{\overline{K}}}{\partial \overline{K}} = \int_{\Omega} U_{\overline{K}}^{*} \frac{\partial H_{\overline{K}}}{\partial \overline{K}} U_{\overline{K}} d^{3}\overline{x}$$

$$= \frac{\hbar^{2}}{M} \int_{\Omega} U_{\overline{K}}^{*} (\frac{1}{t} \nabla + \overline{K}) U_{\overline{K}} d^{3}\overline{x} = -\overline{V}_{\overline{K}}$$
or
$$\frac{1}{K} \frac{\partial \mathcal{E}_{\overline{K}}}{\partial \overline{K}} = \frac{1}{M} \int_{\Omega} \Psi_{\overline{K}}^{*} (\frac{\pi}{t} \nabla) \Psi_{\overline{K}} d^{3}\overline{x} = -\overline{V}_{\overline{K}}$$

and a second second

Density of states g(E) = # single electron states per unit energy, per unit and, at energy E gn(E) is density of states in band n $g(\varepsilon) = \sum_{n} g_n(\varepsilon)$ $g_n(\varepsilon) = \int \frac{d^3k}{4\pi^3} \delta(\varepsilon_n(k) - \varepsilon) \leftarrow includes factor$ $<math>\overline{k} \stackrel{\text{in}}{=} \frac{1}{B_{\tau}^2} \delta(\varepsilon_n(k) - \varepsilon) \leftarrow includes factor$ Alternatively $g_n(\varepsilon)d\varepsilon = \int \frac{d^3k}{4\pi^3} x \begin{cases} 1 & \varepsilon f \\ 0 & o \\ \end{pmatrix} \leq \varepsilon + d\varepsilon$ Let Sn(E) be the surface in K-space of constant energy E, and let Sk [k? be the perpendicular distance between Sn(E) ad Sn(E+dE) $S_n(E)/\int S_n(E+dE)$ I he with computer the shaded whene suface area x wedth $g_n(\varepsilon)d\varepsilon = \int \frac{dS}{4T^3} Sk(k)$ = voline S, (E)

But we have $\varepsilon + d\varepsilon = \varepsilon + |\overline{\nabla}_{k} \varepsilon_{n}| Sk(\overline{k})$ since \$\$ En is mormal to surface \$n(E) Si $\Rightarrow d\varepsilon = |\vec{\nabla}_{k}\varepsilon_{h}| sk(\vec{k})$ $\Rightarrow sk(\vec{k}) = d\varepsilon$ $|\vec{\nabla}_{k}\varepsilon_{h}|$ $g_n(\varepsilon) = \int \frac{dS}{4\pi^3} \frac{1}{\left[\overline{V_k}\varepsilon(\tilde{k})\right]}$ $S_{h}(\mathcal{E})$ Since the band energies E (k) are periodic in The on the RL they are bounded above and below Hence there are to where En (to) is a local max or min. At such values The En(k) vanishes and the integrand in gn dwerges One can show that these donvergences are integrable in 3D, hence Jn(E) stays finite, but dgn will have a dwigence. Such singulanting dE are called van Hove sujularties If there is a van Hove snyulauty on the Ferri surface (ie at a Teo such that $E_n(\overline{k}_0) = E_{\overline{k}})$ this can cause some anomalous looking behavior in quantities flat depend on dg

Energy gaps in the Weak Potential Approximation Assume U is a weak justurbation Consider the effect of the periodec potential U on the fre election state with wavevector to an energy Ex. The potential will mix in plane waves with wave vectors R-R, for K in the R.L., and it will shift the energy eigenvalue from \mathcal{E}_{k}° to a new \mathcal{E} . For weak $\mathcal{U}(\vec{r})$, \mathcal{E} is close to \mathcal{E}_{k}° , Schrödingers equation for the Fourier coefficients ck of the election ergenstate is $(\mathcal{E} - \mathcal{E}_{k-k}) C_{k-k} = \sum_{\substack{k'=k' \\ k' \neq k}} \overline{U}_{k'-k} C_{k-k'}$ (X)____ The right hand side can be no bigger than O(T)(since by normalization we will assume that $\sum_{k} |C_{k} \times |^{2} = 1$, and so each $|C_{k} - \times | \le 1$) > The left hand side can be no bigger than O(V) O suppose k is not near any Brasg plane on an energy scale of O(U), ie. 15%-Ege-k >> U for all K in the R.L. Then it is only $(E - E_{k})$ that can be shall; $(E - E_{k} - k)$ must be bigger than O(U) for all K#0.

> we most have ck-K ~ O(U) for all K # 0 Since we start with the free election state at E, then $c_{\mu} \sim O(1)$. From (+) setting R=0 we have $(\mathcal{E}-\mathcal{E}_{k}^{\circ})C_{k} = \sum U_{k}C_{k-k'}$ $k'\neq 0$ but since $U_{k'} \sim O(U)$, and $e_{k-k'} \sim O(U)$, and grading the reaft have the right hand side is O(U'). Since CK ~ O(1) we must then have $(E - \varepsilon_k) \sim O(U^2)$ So when k is not near any Brogg plane, The scattered waves mix in an amount Ck-k ~ O(V) and the energy shifts $(\mathcal{E} - \mathcal{E}_{\mathbf{k}}) \sim O(\mathcal{U}^{2})$ The periodic potential has relatively little effect on the free election state if U small.