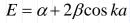
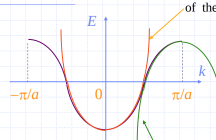
## (10) Correspondence to the free electron model



Free electron model is the bottom parabola of the cos function.



Approximated by  $E = \frac{\pi^2 k^2}{2m}$ 

*m* is different from the value for a real electron.

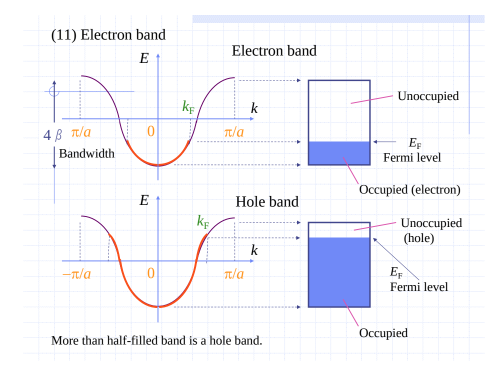
(Effective mass)

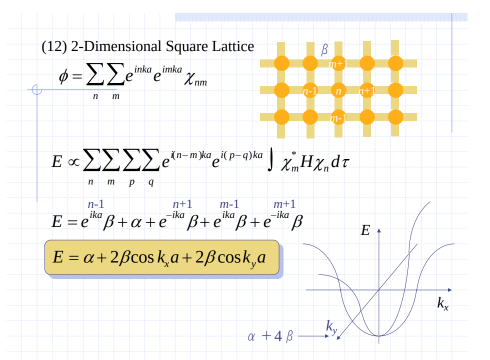
 $4 \beta \text{ large} \rightarrow m \text{ small} \rightarrow \text{ mobile}$ 

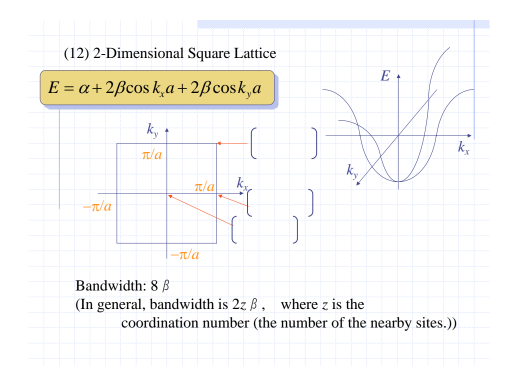
 $4 \beta \text{ small} \rightarrow m \text{ large} \rightarrow \text{not mobile}$ 

Upper half of the band is approximated by free electron with m < 0. Opposite response to electric field,

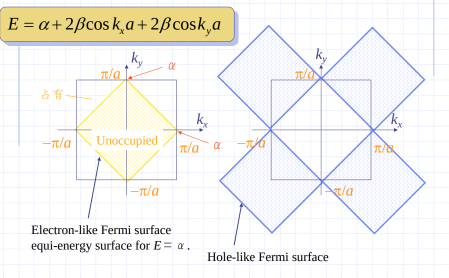
→ Hole with + charge







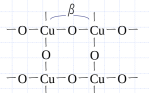
#### (12) 2-Dimensional Square Lattice

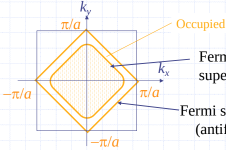


# (12) 2-Dimensional Square Lattice

Cupper oxide high-temperature superconductor

$$(La_{1-x}Sr_x)_2CuO_4$$





Fermi surface of the hole-doped superconductor, (La<sub>1-x</sub>Sr<sub>x</sub>)<sub>2</sub>CuO<sub>4</sub>

Fermi surface of the parent La<sub>2</sub>CuO<sub>4</sub> (antiferromagnetic insulator)

$$\text{La}_2\text{Cu}^{2+}\text{O}_4 \rightarrow \text{Cu}: d^9 \rightarrow d_{x^2-y^2}$$
 band has one electron (Half-filled)

## (13) Tight-bonding method for more than two atoms in a cell

LCAO-MO 
$$\phi = \sum_{i} c_i \chi_i$$

For 2 atoms, i=1, 2



Bloch function is made from  $\chi_i$  for the whole crystal.

$$\chi_i \to \sum e^{inka} \chi_i(n)$$

Crystal orbital is constructed instead of molecular orbital as,

$$\phi = \sum_{i} \sum_{n} c_{i} e^{inka} \chi_{i}(n)$$

Calculate  $E = \frac{\int \phi * H \phi d\tau}{\int \phi * \phi d\tau}$  from this, and similarly to the usual Mo theory,

 $\frac{\partial E}{\partial c_i} = 0$  leads to simultaneous equations of  $c_i$ , and the secular equation is

$$\begin{vmatrix} \alpha_{11}(k) - E & \beta_{12}(k) & \dots \\ \beta_{21}(k) & \alpha_{22}(k) - E & = 0 \end{vmatrix} = 0$$

Since  $\chi_i$  is Bloch function, the matrix elements,  $\alpha$  and  $\beta$  are functions of k:

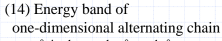
$$\alpha_{ii}(k) = \int (\sum_{i} e^{-imka} \chi_{i}^{*}(m)) H(\sum_{i} e^{inka} \chi_{i}^{*}(n)) d\tau$$

$$= \alpha_{i} + \sum_{i} \beta_{ii}(n) e^{inka} \qquad \beta_{ii}(n) = \int \chi_{i}^{*}(0) H \chi_{i}(n) d\tau$$

$$\beta_{ij}(k) = \int (\sum_{i} e^{-imka} \chi_{i}^{*}(m)) H(\sum_{i} e^{inka} \chi_{j}^{*}(n)) d\tau$$
Nearby atoms
$$= \sum_{i} \beta_{ij}(n) e^{inka} \qquad \text{When interaction } \beta \text{ exists in the } r \text{ direction,}$$
add a term  $\beta e^{ikr}$ .

Nearby atoms

$$\beta_{ii}(n) = \int \chi_i^*(0) H \chi_i(n) d\tau$$



$$\beta$$
 is alternately  $\beta_1$  and  $\beta_2$ .

Two atoms 1 and 2 in a cell.

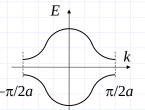
For simplicity, put  $\langle \chi_1 | H | \chi_1 \rangle = \langle \chi_2 | H | \chi_2 \rangle = \alpha = 0$ .  $\langle \chi_1 | H | \chi_2 \rangle = \langle \chi_2 | H | \chi_1 \rangle^* = \beta_2 e^{-ika} + \beta_1 e^{ika}$ 

 $\beta_2$  in the -a direction from 1

 $\beta_1$  in the a direction from 1

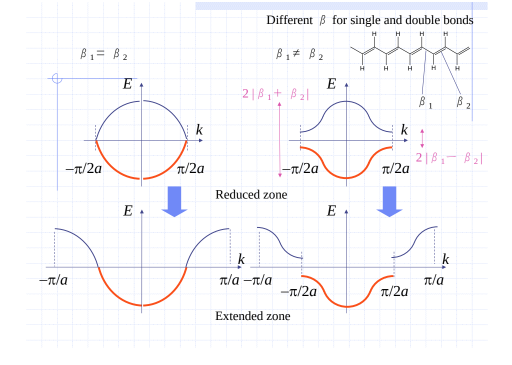
Secular equation is,

$$\begin{vmatrix} -E & \beta_2 e^{-ika} + \beta_1 e^{ika} \\ \beta_2 e^{ika} + \beta_1 e^{-ika} & -E \end{vmatrix} = 0$$

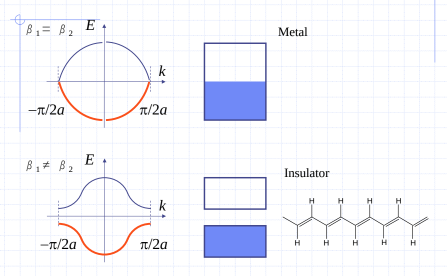


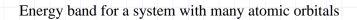
The solution is depicted in the right:

$$1 + \cos 2x = 2\cos^2 x$$



# Peierls Insulator: Long periodicity generates a new energy gap and makes the system insulating.





LCAO-MO from all atomic orbitals in a cell

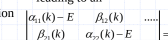
(N, each atom may have more than one.) 
$$\phi = \sum_{i} c_{i} \chi_{i}$$



Make the Bloch function of each  $\chi$ 

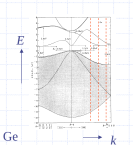
$$\chi_i \to \sum_n e^{inka} \chi_i(n)$$
 leading to an

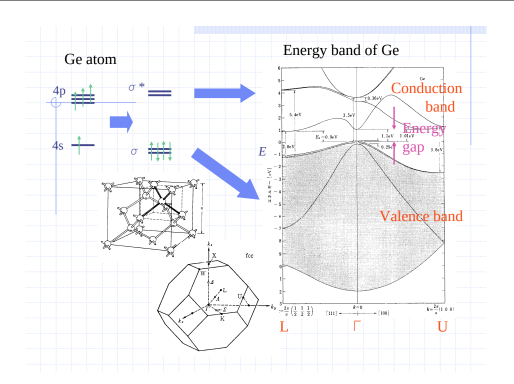
 $N \times N$  secular equation

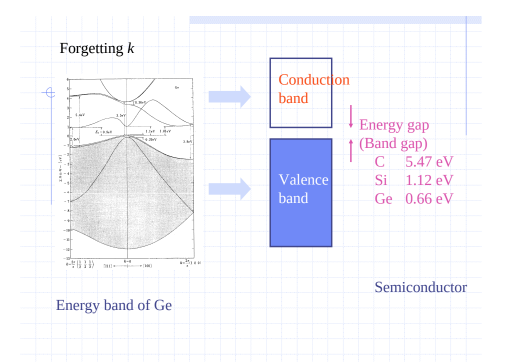


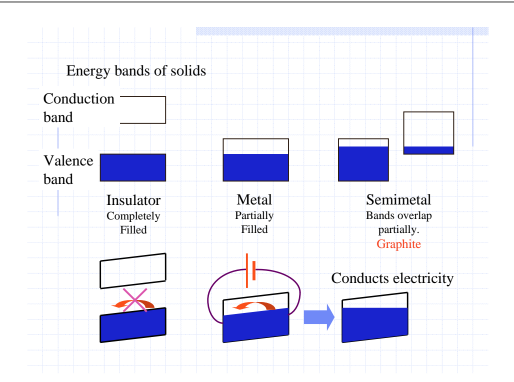
Since each element is a function of k, this secular equation is solved at each k, to give N energy levels.

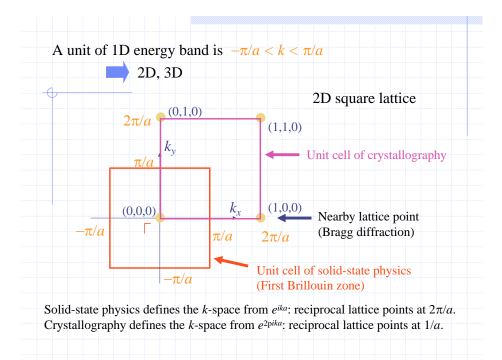
These energy levels for different k are connected to afford continuous energy bands (right).

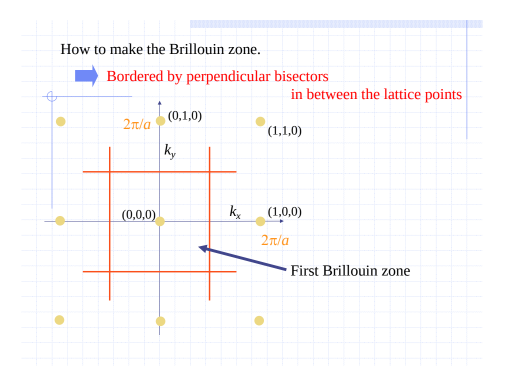


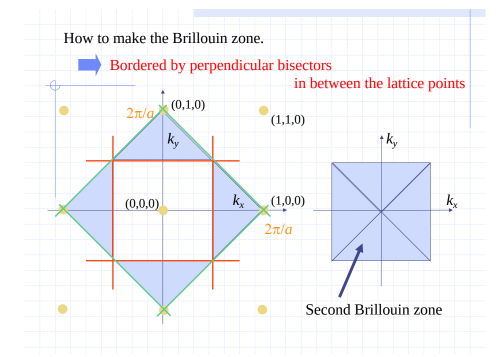


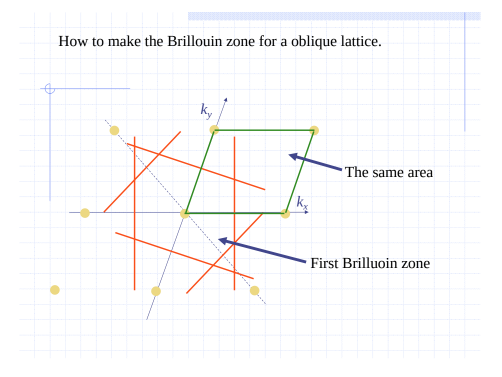


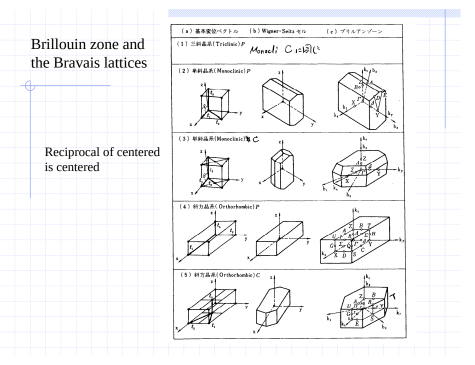








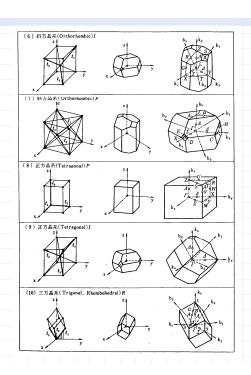




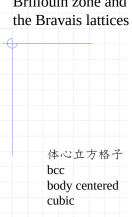
Brillouin zone and the Bravais lattices

Reciprocal of face centered is body centered

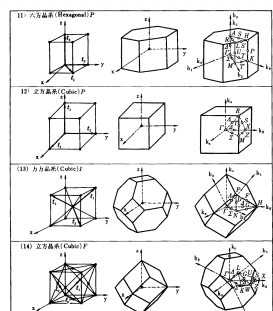
Reciprocal of body centered is face centered



# Brillouin zone and

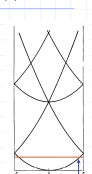


面心立方格子 fcc face centered cubic



### Fermi surface of metals

(1) Alkali metals Li, Na, K

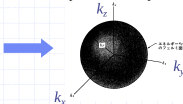


Fermi surface  $\Leftrightarrow$  trajectory of  $k_F \Leftrightarrow$  Surface at  $E=E_F=$ const.

Assuming free electron

$$E = \frac{\hbar^2}{2m} (k_x^2 + k_y^2 + k_z^2) = \text{const.}$$

leads to a sphere in the k-space.



Fermi surface of alkali metals is not deviated from the perfect sphere by 1%.

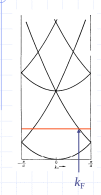
Energy interval in the k-space is  $\Delta k = \frac{2\pi}{Na}$ 

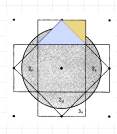
Volume of the Fermi sphere is half of the volume of the 1st Brillouin zone.

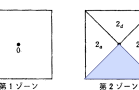
#### Fermi surface of metals

(1) Divalent metals Mg, Al

Volume of the Fermi sphere is the same as the volume of the 1st Brillouin zone.

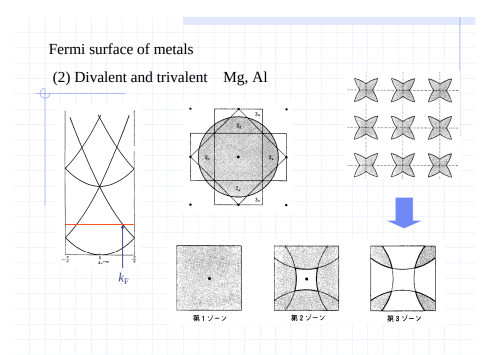


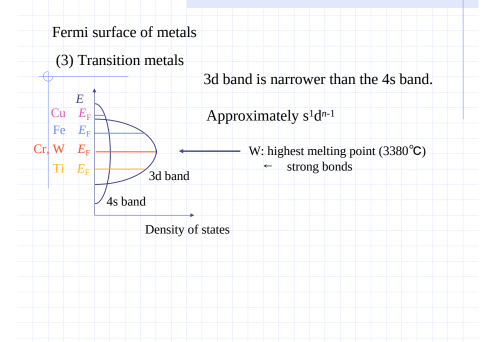


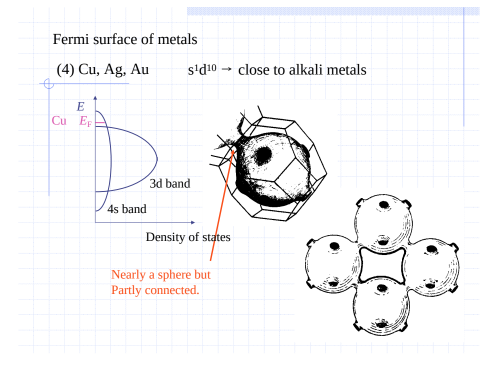


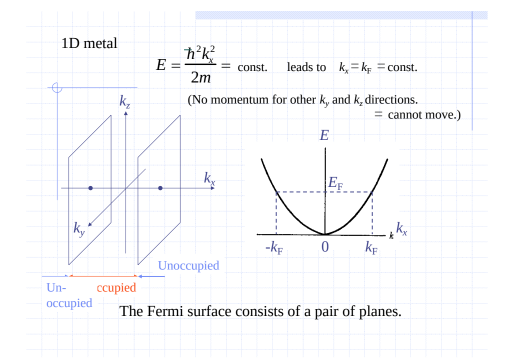


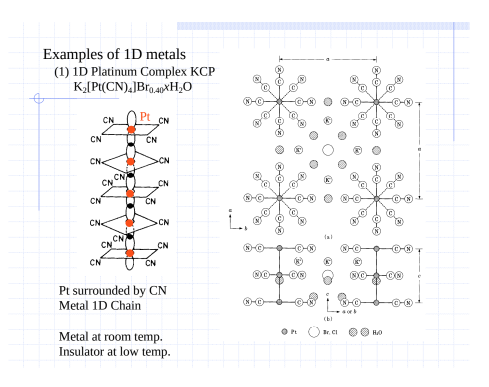
第1 ゾーン

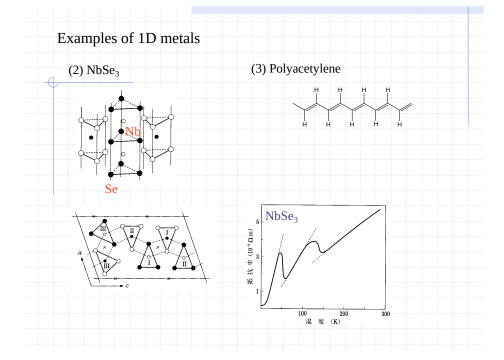


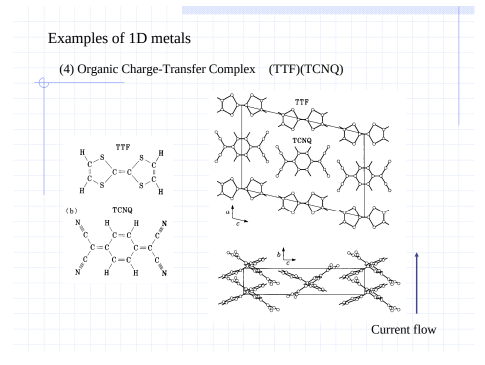


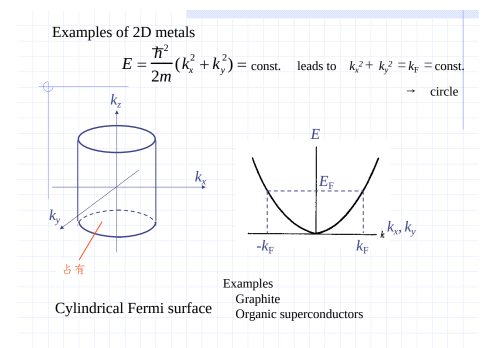


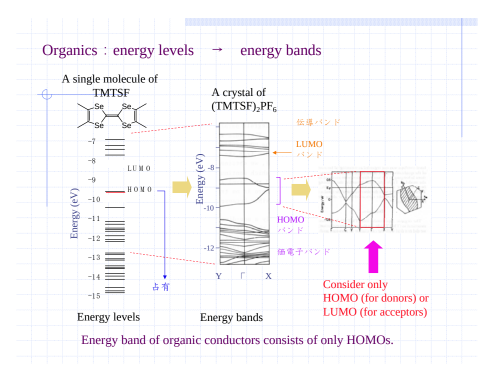


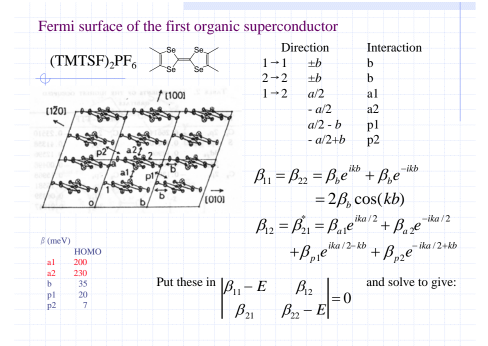


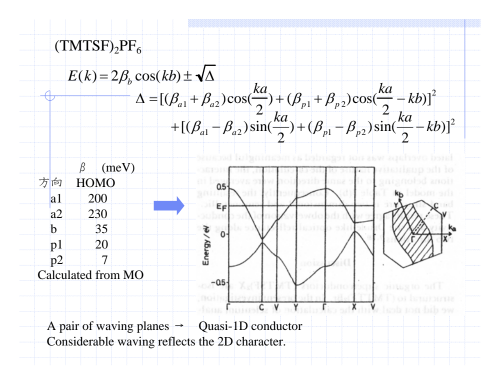


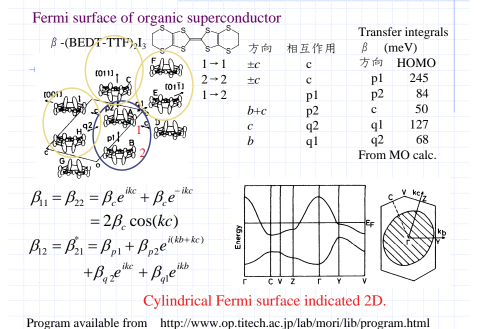










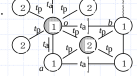


## Excise Band structure of the $\theta$ -phase

This generally found structure is called herringbone structure in organic crystals or alternatively  $\theta$  -phase in organic conductors. Calculate the energy bands of this structure.



A unit cell contains two molecules, numbered 1 and 2. Transfer  $t_a$ , running along a, is between two Molecule 1. Transfer  $t_p$ , running diagonal (a/2, b/2) etc, is between Molecule 1 and 2.



- (1) How many Molecule 1 exist near Molecule 1? From this, obtain the diagonal element  $F_{11}$ .
- (1) How many Molecule 2 exist near Molecule 1? From this, obtain the nondiagonal element  $F_{12}$ .
- (3) Solve the secular equation, and obtain an equation of  $E(k_a, k_b)$  representing the energy band.

$$\cos x + \cos y = 2\cos\frac{x+y}{2}\cos\frac{x-y}{2}$$