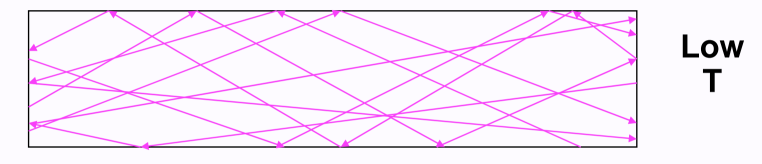
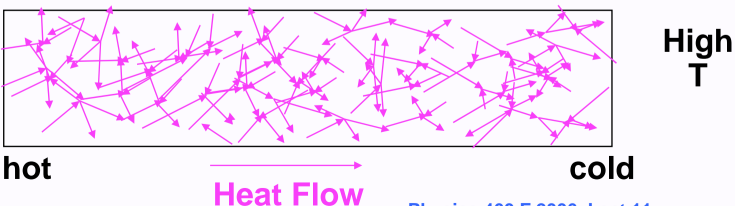
#### Phonons II - Thermal Properties -Continued (Kittel Ch. 5)





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#### Outline

#### Anharmonicity

- Crucial for Thermal expansion and other changes with pressure and temperature
- Gruneisen Constant
- Thermal Heat Transport
   Phonon Heat Conductivity

Low T and High T behaviors

• (Read Kittel Ch 5)

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# • If we expand the energy beyond the harmonic order: $E = E_0 + (1/2) \sum_{ij} \Delta \underline{R}_i \cdot \underline{D}_{ij} \cdot \Delta \underline{R}_j + (1/6) \sum_{ijk} \underline{D}_{ijk} \cdot \Delta \underline{R}_i \Delta \underline{R}_j \Delta \underline{R}_k + \dots \leftarrow \underline{Difficult}$ and Messy

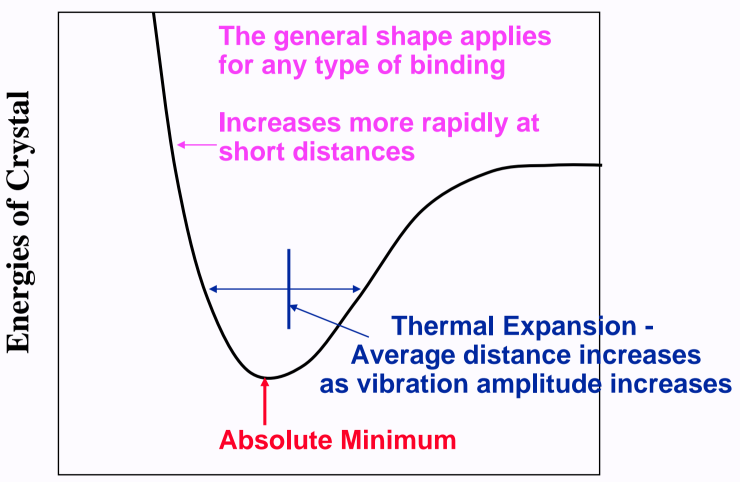
- The problem is fundamentally changed: No longer exactly soluble
- Consequences:

There is thermal expansion and other changes with temperature

The heat capacity is not constant at high T Phonons decay in time

Two phonons can interact to create a third Phonons can establish thermal equilibrium and conduct heat like a "gas" of particles

#### **Thermal expansion - I**



**Distance Between Atoms** 

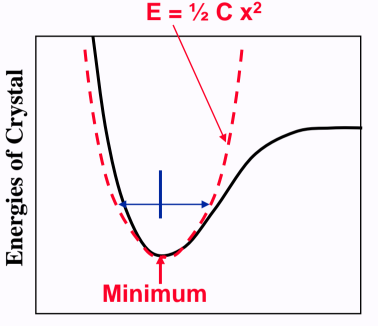
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## **Thermal expansion - II**

There is no thermal expansion In the harmonic approximation!
 Harmonic Approximation

Why? In the harmonic approximation the magnitude of the force is symmetric around the minimum (F = -Cx). Therefore, the vibrations are symmetric  $\Rightarrow$  the average position <x> is always the center, i.e., < x > = 0and

there is no thermal expansion



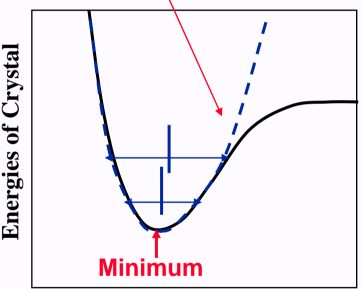
**Distance Between Atoms** 

#### **Thermal expansion - III**

• Anharmonicity causes thermal expansion.

Why? If the potential energy curve is not symmetric, the vibrations are not symmetric  $\Rightarrow$  the average position <x> changes as the magnitude of the vibration increases, i.e., <x> increases and there is thermal expansion

Anharmonic potential energy  $E = \frac{1}{2}C_{x^{2}} + (1/6)A_{x^{3}}$ 



**Distance Between Atoms** 

#### **Gruneisen Constant y**

- Anharmonicity is very complicated because there are all the possible terms involving 3 atoms!
- A simple characterization requires finding a typical measurable quantity.
- This is given by the dimensionless Gruneisen constant  $\gamma$  defined as an average value of the change of vibration frequency per unit dilation:  $\Delta \omega / \omega = \gamma (\Delta V / V) = 3 \gamma (\Delta R / R)$ where L is a characteristic near-neighbor distance.
- Thus  $\gamma = (1/3)(\Delta \omega / \omega)/(\Delta R/R)$
- On the next slide we consider a simple case that shows the relation to anharmonicity

#### **Gruneisen Constant** $\gamma$

 From previous slide: The Gruneisen constant γ defined as an average value of the change of vibration frequency per unit dilation:

 $\Delta \omega / \omega = \gamma (\Delta V / V) = 3 \gamma (\Delta R / R)$ 

where L is a characteristic near-neighbor distance .

- An example is a nearest neighbor interaction with  $E = \frac{1}{2} C \Delta R^2 + (1/6) A \Delta R^3$
- Then the force is given by  $F = -dE/dR = -C(\Delta R) - \frac{1}{2} A (\Delta R)^{2} + \dots$   $= -C [1 + \frac{1}{2} A \Delta R/C] \Delta R$

Since  $\omega^2 \sim C/M \Longrightarrow \Delta \omega / \omega = \frac{1}{2} \Delta C/C = \frac{1}{2} A \Delta R / C$ 

• Thus  $\gamma = (1/3)(\Delta \omega / \omega)/(\Delta R/R) = (1/6) (A/C) \Delta R$ is a measure of anharmonicity

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#### **Other effects of Anharmonicity**

#### Anharmonicity

- Crucial for Thermal expansion and other changes with pressure and temperature
- Gruneisen Constant
- Thermal Heat Transport
   Phonon Heat Conductivity

#### Low T and High T behaviors

• (Read Kittel Ch 5)

**Thermal Energy (Heat) Transport** 

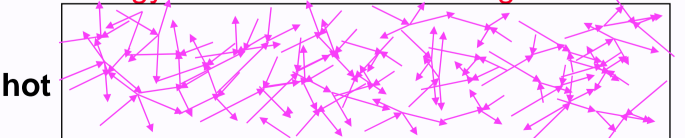
- First consider heat transport in an ordinary gas
- Molecules in the gas have average energy

$$\frac{1}{2} \text{ m v}^2 = 3[\frac{1}{2} \text{ k}_{\text{B}}\text{T}]$$
Factor of 3 in  
and move with average speed v
3 dimensions

- Two limiting cases
  - Ordinary case where molecules scatter from one another very often as they move The molecules diffuse and thermalize with one another This is the usual case for gases like the air in this room
  - Cases where the molecules hit the walls of the container without hitting one another – the molecules transport energy directly from one wall to another

#### **Transport of heat in an ordinary gas**

- Molecules move in all directions and scatter so that they come to local thermal equilibrium in each region.
- How can random motion of molecules cause heat flow in one direction?
- On average, in hotter regions molecules have more kinetic energy. A molecule that moves from a hotter region to a colder region brings energy above the local average. The opposite for a molecule moving from a colder to a hotter region. Either way, there is transport of energy from hotter to colder regions.



Heat Flo

cold

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## **Heat Transport due Molecular Motion**

# Definition: j = heat flow (energy per unit area per unit time ) = - K dT/dx

- If a particle moves from a region with local temperature T to one with local temperature T - ΔT, it supplies excess energy c ΔT, where c = heat capacity per particle. (Note ΔT can be positive or negative).
- On the average for a particle with average thermal speed v in the x direction:

 $\Delta T = (dT/dx) v_x \tau$ , where  $\tau =$  mean time between collisions

• Then  $j = -n v_x c v_x \tau dT/dx = -n c v_x^2 \tau dT/dx$ Density

## Heat Transport due Molecules - II

- This can be simplified in an isotropic case, since averaging over directions gives  $(v_x^2)_{average} = (1/3)v^2$
- This leads to  $j = -(1/3) n c v_x^2 \tau dT/dx$
- Finally we can define the mean free path:

   L = v τ
   and
   C = pa = total basis consolity per unit yell

C = nc = total heat capacity per unit volume

• Then j = -K dT/dx = -(1/3) C v L dT/dxand

K = (1/3) C v L = thermal conductivity

This is a general result for thermal conductivity due to thermal motion of particles - Eq. 42 in Kittel

#### Heat Transport due Molecules - III

- For molecules in a gas we can use classical expressions for C and v, and we can understand some facts about L
- Heat capacity: C = 3Nk<sub>B</sub>
- Average speed: v determined by  $\frac{1}{2}$  Mv<sup>2</sup> = (3/2) k<sub>B</sub>T
- Mean free path L is distance between collisions increases with density temperature independent
- Then

 $K = (1/3) C v L = Nk_B (3k_BT/M)^{1/2} L$ 

• Increases with T for fixed density because v increases

#### Heat Transport due Molecules - IV

- What happens if the mean free path is long compared to the size of the container
  - L ~ dimension of container

and the thermal conductivity is given by K = (1/3) C v L

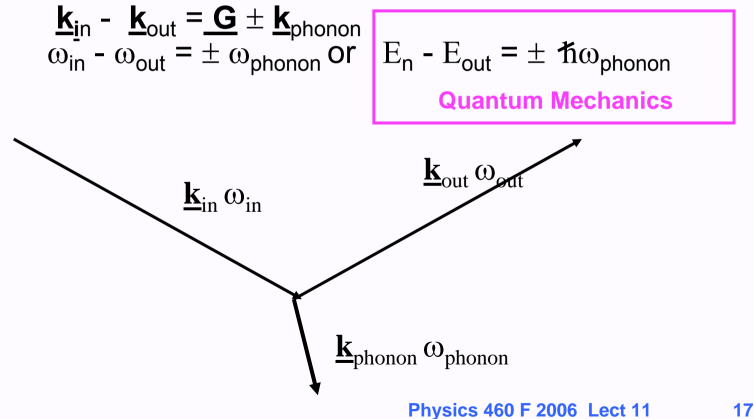
• And the basic idea are still the same

#### What causes heat transport in a solid?

- Motion of the atoms is thermal energy
- But the atoms vibrate around their sites and do not move through the solid
- How does energy moves from hot to cold?
- The amplitude of the vibrational motion is transferred through the solid – hotter atoms transfer energy to colder atoms
- How do we describe this?

#### From Before Inelastic Scattering and Fourier Analysis

- Scattering of neutrons, x-rays, ....
- Inelastic diffraction occurs for



#### **Scattering of Phonons - I**

- The same idea applies to phonons. One phonon can scatter to create two.
- We can say one phonon is destroyed and two are created. This can occur if  $\underline{\mathbf{k}}_{\text{in phonon}} = \underline{\mathbf{k}}_{\text{out phonon 1}} + \underline{\mathbf{k}}_{\text{out phonon 2}} \pm \underline{\mathbf{G}}$  $\omega_{\text{in phonon}} = \omega_{\text{out phonon 1}} + \omega_{\text{out phonon 2}}$  $\underline{\mathbf{K}}_{\text{in phonon}} \boldsymbol{\omega}_{\text{in phonon}}$  $\mathbf{\underline{K}}_{\text{out phonon }1}\omega_{\text{out phonon }1}$ Anharmonic  $\underline{\mathbf{K}}_{\text{out phonon 2}} \omega_{\text{out phonon 2}}$ Interaction

#### **Scattering of Phonons - II**

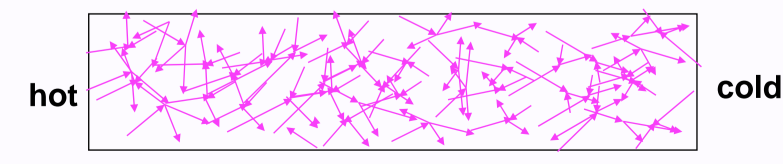
- Also a phonon can absorb (destroy) another phonon and create a different phonon.
- Thus two phonons are destroyed and one is created which can occur if  $\underline{\mathbf{k}}_{\text{out phonon}} = \underline{\mathbf{k}}_{\text{in phonon 1}} + \underline{\mathbf{k}}_{\text{in phonon 2}} \pm \underline{\mathbf{G}}$  $\omega_{\text{out phonon}} = \omega_{\text{in phonon 1}} + \omega_{\text{in phonon 2}}$  $\underline{\mathbf{K}}_{\text{in phonon 1}} \boldsymbol{\omega}_{\text{in phonon 2}}$ Anharmonic  $\underline{\mathbf{K}}_{\text{out phonon}} \boldsymbol{\omega}_{\text{out phonon}}$ Interaction This is proportional to the average occupation of phonons  $\underline{\mathbf{K}}_{\text{in phonon } 2} \boldsymbol{\omega}_{\text{in phonon } 2}$ in state 2 – the probability vanishes if there are no Physics 460 F 2006 Lect 11 phonons to destroy!

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#### **Phonons also act like a gas**

- A phonon is a particle a quantum of vibration
- It carries energy just like a molecule.
- Phonon can come to equilibrium by scattering just like molecules
- Scattering is due to defects and anharmonicity.
- Leads to heat transport just as for molecules

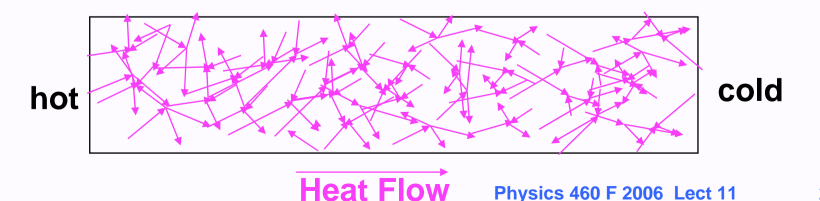
K = (1/3) C v L = thermal conductivity



**Heat Flow** 

#### **Phonons also act like a gas**

- What is different about phonons and gas molecules?
- Speed = v<sub>group</sub> is largest for low energy (v<sub>sound</sub>) and smaller for high energy
- The heat capacity ~  $T^3$  at low T, ~  $3Nk_B$  at high T
- The density of phonons <n> increases with T. Scattering increases ~ <n> since each phonon has scatters from other phonons. Therefore we expect the mean free path to decrease with T



#### **Phonon Heat Transport**

- The same logic applies for phonons as for molecules j = heat flow (energy per unit area per unit time) = - K dT/dx
- If a phonon moves from a region with local temperature T to one with local temperature T - ΔT, it supplies excess energy c ΔT, where c = heat capacity per phonon.
- Then j = (1/3) C v L dT/dx and

K = (1/3) C v L = thermal conductivity

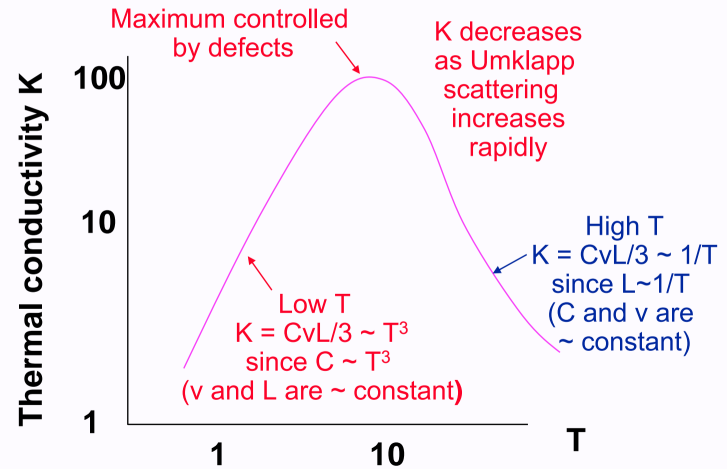
where L = mean free path, C = nc = total heat capacity (a function of temperature)

#### **Phonon Heat Transport - continued**

- What determines mean free path L = v  $\tau$  ?
- At low temperature, the thermal phonons are sound waves that have long mean free paths -L ~ sample size
- At high temperature, phonons scatter from other phonons. The density of other phonons is ~ T, so L ~ 1/T
- At intermediate temperature, phonon scatter from defects and other phonons. The latter decreases very rapidly as T decreases because ONLY Umklapp scattering limits the energy flow.

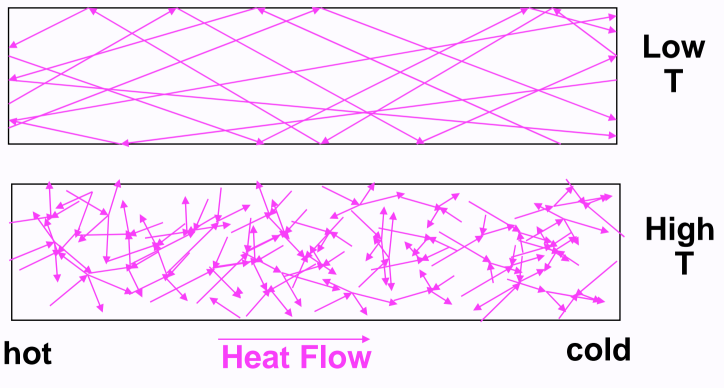
#### **Phonon Heat Transport - continued**

• Behavior in an excellent quality crystal:



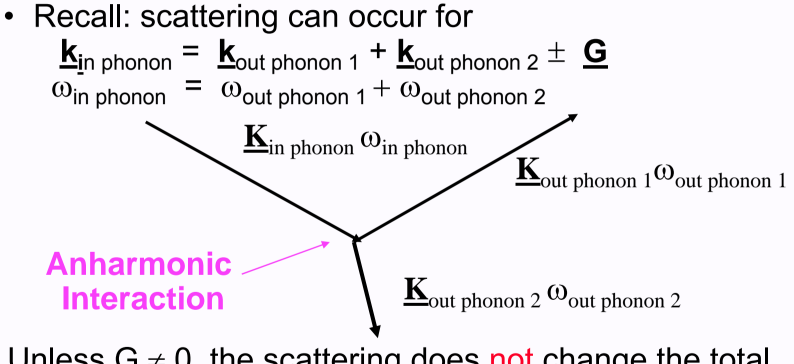
## **Phonon Heat Transport - continued**

- Low T K ~ T<sup>3</sup> increases because density of phonons increases with roughly constant v and L
- High T K ~ 1/T decreases as scattering increases



## **Umklapp Scattering**

• What is Umklapp scattering? Scattering that changes total crystal momentum by a reciprocal lattice vector.



Unless G  $\neq$  0, the scattering does not change the total phonon momentum or energy. Therefore only Umklapp scattering limits the heat flow.

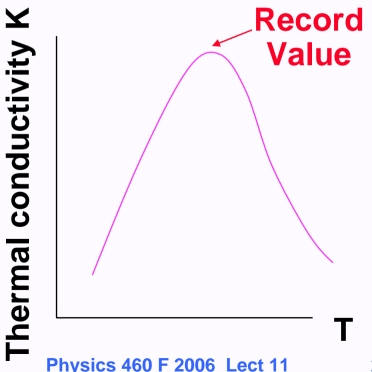
#### **Other effects**

- Maximum value of K affected by all scattering mechanisms
- Impurities scatter phonons
- Isotope effects

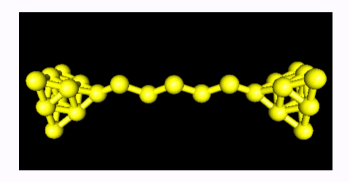
Isotopes do not affect chemical properties, but they do change phonon frequencies and cause scattering of phonons - limits the maximum value of K in high quality crystals

 How did the collapse of the Soviet Union affect thermal conductivity?

- Availability of pure isotopes from Soviet nuclear program has made possible growth of large crystals of pure isotope materials
- New record for heat conductivity for all crystals isotopically pure diamond - better than any metal at room temperature
   New record for heat conductivity for all crystals -Reco Value
- Stimulated new research to create better diamond heat sinks for lasers, etc.



Are there special aspects of thermal conductivity in nanosystems?



Ball and stick model of a 1-atom wide gold "wire" actually made and seen in an electron microscope

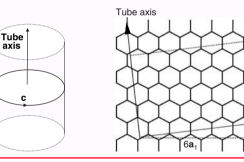
An electrical current leads to very high current densities, heating, ... An electron can "fall" into the well only if it can give up energy to phonons – the rate is reduced because the phonons have only quantized energies

e More later

"Quantum well" for electrons in a semiconductor – used to contain electrons for lasers, electronic devices, ....

Are there special aspects of thermal conductivity in nanosystems?
 Buckyball with

Ball and stick models of a Buckyball and a nanotube



A nanotube is a sheet of graphite rolled up into a long thin tube

Work now to use nanotubes as wires in semiconductor circuits

60 carbon atoms

now made in quantity

31

Perhaps later we can sal a little about electrons

An electrical current in the nanotube leads to very high current densities, heating, ... phonons scatter very little in the tube – good for removing heat --- but the mismatch with other materials makes it hard for the heat to excape – "burn" in the middle Physics 460 F 2006 Lect 11



- Anharmonicty crucial for certain effects Thermal expansion, .... Phonon scattering
- Gruneisen Constant γ
- Heat transport due to thermal motion of particles
  - j = -K dT/dx, K = (1/3) C v L
    - K = thermal conductivity
    - C = heat capacity
    - v = mean velocity

L = mean free path

The same ideas are used later for both thermal and electrical conductivity of electrons!

- Gas of molecules K ~ =  $Nk_B (3k_BT/M)^{1/2} L$
- Phonons act like particles in a gas (We treat heat transport due to electrons later)
- Low T K  $\sim$ T<sup>3</sup> (C  $\sim$ T<sup>3</sup> and v, L  $\sim$  constant)
- K ~ 1/T (C ~  $3Nk_B$ , v ~ constant and L ~ 1/T)

#### Next time

- Start next Part of Course: Electronic Properties of Solids
- Free electron Fermi Gas
- Energy levels in one dimension and three dimensions
- Heat Capacity
- (Read Kittel Ch 6)
- Remember: EXAM MONDAY, OCTOBER 9

#### **Summary of Part I**

• This is an incomplete list of concepts, topics, equations, examples!

- Solids are defined by their "solidity" i.e., to ability to resist shear stress and keep a shape
- A crystal is truly solid (as opposed to a glass which is like a "slow"liquid")

A crystal has a true long range crystalline order

- Crystalline order is defined by the regular positions of the nuclei Crystal Structure = Lattice + Basis
- Bravais lattice of points in real space Examples: line, square, simple cubic, hexagonal, fcc, hcp, bcc
- Primitive Cell

Parallelpiped in 3d Wigner-Seitz cell

• Examples of crystals:

line, square, simple cubic fcc, hcp, bcc, NaCl, ZnS, diamond, graphene

- Diffraction and experimental studies of crystal structures Examples: X-ray, neutron diffraction Appropriate wavelength (k =  $2\pi/\lambda$ )
- Bravais lattice on points in real space and Reciprocal Lattice Reciprocal lattice Brillouin Zone – diffraction occurs only on boundary of BZ
- Diffraction and experimental studies of crystals Bragg Law - easier to describe in terms of reciprocal lattice Elastic scattering : k<sub>in</sub> = k<sub>out</sub> + G Examples:

line, square, simple cubic, hexagonal, fcc, hcp, bcc

• Structure factor – example of diamond Here it is the idea that is important not the details

- Binding of Crystals
   5 basic types
- Simplest examples: pair potentials  $\phi(r)$

Van der Waals. Attractive and Repulsive terms Coulomb interactions in ionic crystals

- For pair potentials, binding is sum of over pairs Count each interaction as ½ for each atom in the pair
- Binding curve
   Minimum

Minimum energy – equilibrium structure

• Bulk modulus

Curvature – B =  $Vd^{2}E/dV^{2}$ 

• Elastic properties

Stress – strain relations - elastic constants  $C_{ij}$ (The ideas that are most important – not the details) Example: cubic crystals –  $C_{11}$ ,  $C_{12}$ ,  $C_{44}$  B=  $(C_{11} + 2C_{12})/3$ Sound waves -  $v_s = (C_{ij}/\rho)^{1/2}$ ,  $C_{ij}$  = appropriate elastic constant,  $\rho$  = mass density

Vibrations of atoms

Harmonic approximation Exact solution for waves in a crystal Dispersion curves  $\omega_{k,m}$  where <u>k</u> is the wavevector in Brillouin Zone and m = 1, ..., 3N

• Counting k states

The number of k states = number of cells in crystal N Example in 1 dimension

Same ideas in 2 and 3 dimensions

Result

Each k state corresponds to a volume of reciprocal space =  $(2\pi)^3/V = V_{BZ}/N$  (2 $\pi/L$  in 1 dimension) (1/N)  $\Sigma$ k inside BZ  $\Rightarrow$  (1/V<sub>BZ</sub>)  $\int_{BZ} dk$ 

Quantization of vibrations
 Ear each vibration with free

For each vibration with frequency  $\omega$ , the quantum is  $\pi \omega$ Phonons act like particles

Observed in inelastic scattering experiments (neutrons, ...)

• Inelastic scattering is a central idea that is used again in transport

• Thermal properties

Fundamental law of probabilities – exp(-E/k<sub>B</sub>T) Planck distribution for phonons

$$= \frac{1}{\exp(\hbar\omega/k_BT) - 1}$$

- Debye and Einstein models
   Debye captures critical features of acoustic modes
- Heat Capacity C
   Low T: C ~ T<sup>3</sup> High T: C ~ constant
- Thermal conductivity K Low T: K ~ T<sup>3</sup> High T: K~ 1/T Maximum as function of T

#### **Summary of Part I Looking Forward**

- What will be important in describing electrons? ٠
- In order to understand:

Metals, insulators, semiconductors, ....

We need:

The structure of the crystal – Lattice, basis, Recip. Lattice Electronic states in in periodic crystal Electrons act like waves/particles The same Brillouin zone as for phonons The same counting of states in the BZ Density of states D(E)

•

## Conductivity – transport of charge The same ideas as for thermal energy $j = \sigma E = \sigma dV/dx$ , $\sigma = (1/3) e^2 v L$ (Ohm's law!)

Scattering - essential for resistance (and for Ohm's law) – is due to phonons, impurities, ...

And we need to understand the crucial differences between electrons ٠ and phonons!

Specific heat - ideas the same but very different result Conductivity – ....