

FYS3410 - Vår 2011 (Kondenserte fasers fysikk)

<http://www.uio.no/studier/emner/matnat/fys/FYS3410/index-eng.xml>

Based on Introduction to Solid State Physics by Kittel

Course content

- **Periodic structures, understanding of diffraction experiment and reciprocal lattice**
- **Imperfections in crystals: diffusion, point defects, dislocations**
- **Crystal vibrations: phonon heat capacity and thermal conductivity**
- **Free electron Fermi gas: density of states, Fermi level, and electrical conductivity**
- **Electrons in periodic potential: energy bands theory classification of metals, semiconductors and insulators**
- **Semiconductors: band gap, effective masses, charge carrier distributions, doping, pn-junctions**
- **Metals: Fermi surfaces, temperature dependence of electrical conductivity**

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FYS3410 lecture schedule and exams: Spring 2011

W/19/1/2011:	Introduction and motivation. Periodicity and lattices	1h
M/24/1/2011:	Index system for crystal planes. Crystal structures	2h
W/26/1/2011:	Reciprocal space, Laue condition and Ewald construction	1h
M/31/1/2011:	Brillouin Zones. Interpretation of a diffraction experiment	2h
W/02/2/2011:	Crystal binding, elastic strain and waves	1h
M/07/2/2011:	Elastic waves in cubic crystals; defects in crystals	2h
W/09/2/2011:	Defects in crystals; case study – vacancies; diffusion	2h
M/14/2/2011:	Crystal vibrations and phonons	2h
W/16/2/2011:	Lattice heat capacity: Dulong-Petit and Einstein models	2h
M/21/2/2011:	Phonon density of states (DOS) and Debye model	2h
W/23/2/2011:	General result for DOS; role of anharmonic interactions	2h
M/28/2/2011:	Thermal conductivity and repetition of crystal vibrations	2h
W/02/3/2011:	no lectures	
M/07/3/2011:	no lectures	
W/09/3/2011:	no lectures	
M/14/3/2011:	Free electron Fermi gas in 1D and 3D – ground state	2h
W/17/3/2011:	Density of states, effect of temperature – FD distribution	1h
M/21/3/2011:	Heat capacity of FEFG	2h
W/23/3/2011:	Repetition	1h
M/28/3/2011:	Mid-term exam	

M/04/4/2011:	Electrical and thermal conductivity in metals	2h
W/06/4/2011:	Bragg reflection of electron waves at the boundary of BZ	2h
M/11/4/2011:	Energy bands, Kronig - Penny model	2h
W/13/4/2011:	Empty lattice approximation; number of orbitals in a band	2h

Påsk uppehåll

W/27/4/2011 no lectures

M/02/5/2011: no lectures

W/04/5/2011: no lectures

M/09/5/2010:	Semiconductors, effective mass method, intrinsic carriers	2h
W/11/4/2010:	Impurity states in semiconductors and carrier statistics	2h
M/16/5/2010:	p-n junctions, Schottky contacts and heterojunctions	2h
W/18/5/2010:	Metals and Fermi surfaces	1h
M/23/5/2010:	Repetition	2h
26-27/5/2010:	Final Exam (sensor:???)	

Lecture 14: Thermal conductivity

- **Repetition: anharmonic crystal interactions – thermal expansion**
- **Phenomenological description of thermal conductivity**
- **Temperature dependence of thermal conductivity as a cause of phonon scattering**
- **Phonon collisions: N and U processes**

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Anharmonic Properties of Solids

Two important physical properties that ONLY occur because of anharmonicity in the potential energy function:

1. Thermal expansion
2. Thermal resistivity (or finite thermal conductivity)

Thermal expansion

In a 1-D lattice where each atom experiences the same potential energy function $U(x)$, we can calculate the average displacement of an atom from its $T=0$ equilibrium position:

$$\langle x \rangle = \frac{\int_{-\infty}^{+\infty} x e^{-U(x)/kT} dx}{\int_{-\infty}^{+\infty} e^{-U(x)/kT} dx}$$

Thermal Expansion in 1-D

Evaluating this for the harmonic potential energy function $U(x) = cx^2$ gives:

$$\langle x \rangle = \frac{\int_{-\infty}^{+\infty} x e^{-cx^2/kT} dx}{\int_{-\infty}^{+\infty} e^{-cx^2/kT} dx}$$

Now examine the numerator carefully...what can you conclude?

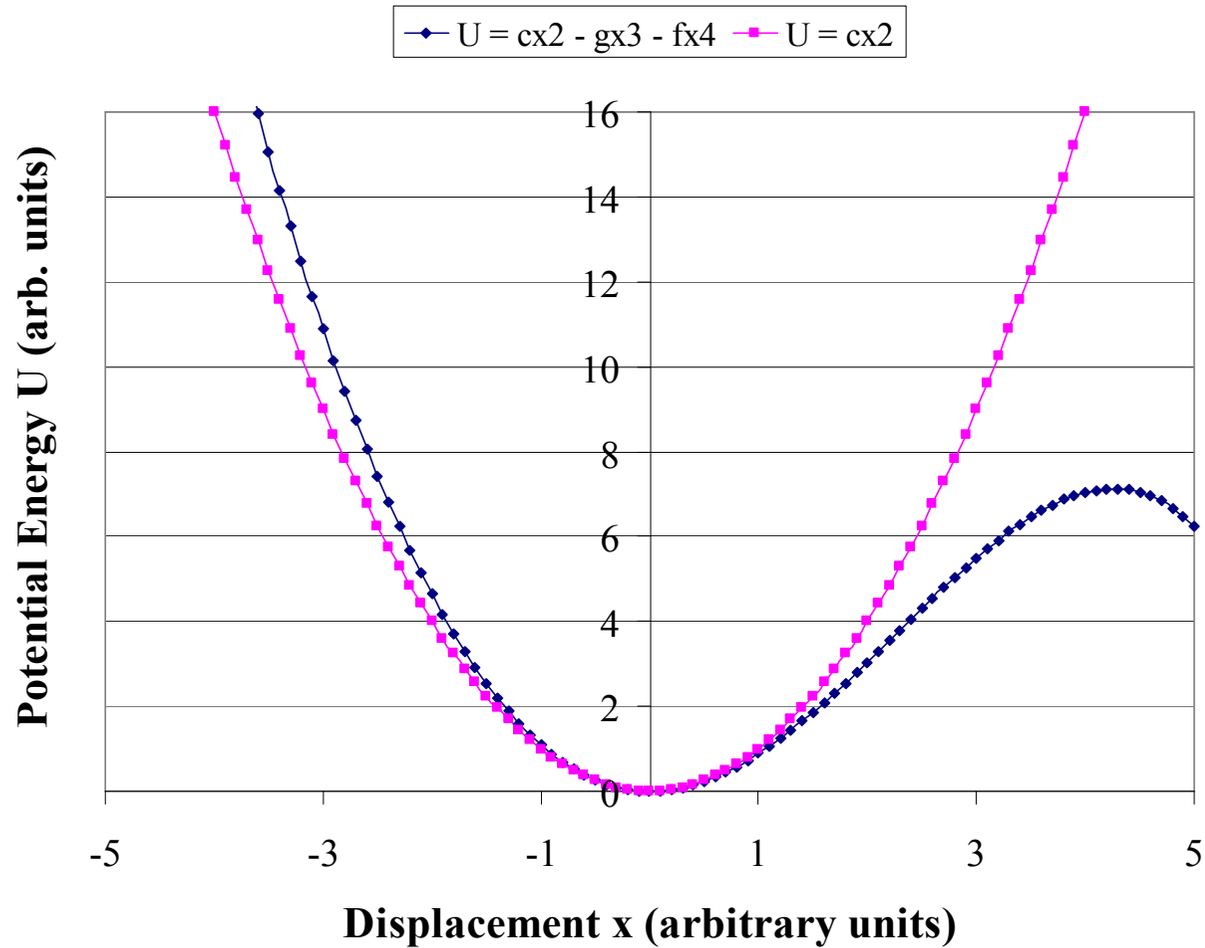
$$\langle x \rangle = 0! \quad \text{independent of } T!$$

Thus any nonzero $\langle x \rangle$ must come from terms in $U(x)$ that go beyond x^2 . For HW you will evaluate the approximate value of $\langle x \rangle$ for the model function

$$U(x) = cx^2 - gx^3 - fx^4 \quad (c, g, f > 0 \quad \text{and} \quad gx^3, fx^4 \ll kT)$$

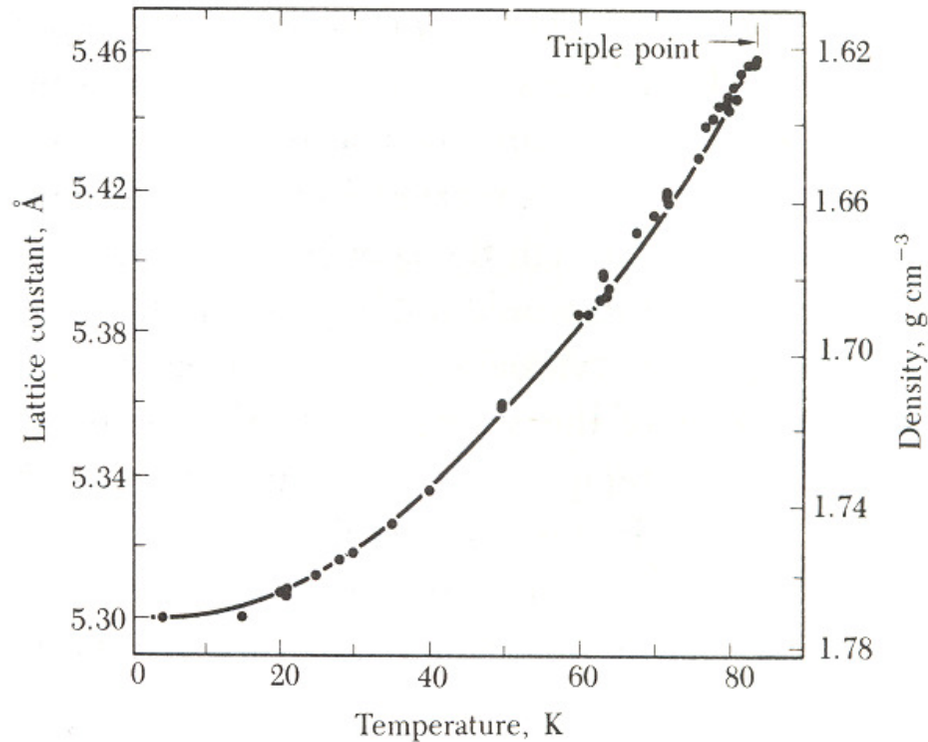
Why this form? On the next slide you can see that this function is a reasonable model for the kind of $U(r)$ we have discussed for molecules and solids.

Potential Energy of Anharmonic Oscillator ($c = 1$ $g = c/10$ $f = c/100$)



Do you know what
form to expect for
 $\langle x \rangle$ based on
experiment?

Lattice Constant of Ar Crystal vs. Temperature



Above about 40 K, we see: $a(T) - a(0) \propto \langle x \rangle \propto T$

Usually we write: $L = L_0(1 + \alpha[T - T_0])$ $\alpha =$ thermal expansion coefficient

Lecture 14: Thermal conductivity

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Phenomenological description of thermal conductivity

When thermal energy propagates through a solid, it is carried by lattice waves or phonons. If the atomic potential energy function is harmonic, lattice waves obey the superposition principle; that is, they can pass through each other without affecting each other. In such a case, propagating lattice waves would never decay, and thermal energy would be carried with no resistance (infinite conductivity!). So...thermal resistance has its origins in an anharmonic potential energy.



Classical definition of thermal conductivity

$$\kappa = \frac{1}{3} C_V v \Lambda$$

Thermal energy flux
(J/m²s)

$$J = -\kappa \frac{dT}{dx}$$

C_V heat capacity per unit volume

v wave velocity

Λ mean free path of scattering
(would be ∞ if no anharmonicity)

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Temperature dependence of thermal conductivity as a cause of phonon scattering

There are three basic mechanisms to consider:

1. Impurities or grain boundaries in polycrystalline sample
 2. Sample boundaries (surfaces)
 3. Other phonons (deviation from harmonic behavior)
- } deviation from perfect crystalline order

To understand the experimental dependence $\kappa(T)$, consider limiting values of C_V and Λ (since v does not vary much with T).

$$C_V \begin{cases} \propto T^3 & \text{low } T \\ 3R & \text{high } T \end{cases} \quad \Lambda \propto \frac{1}{n_{ph}} = e^{\hbar\omega/kT} - 1 \begin{cases} \rightarrow \infty & \text{low } T \\ \frac{\hbar\omega}{kT} & \text{high } T \end{cases}$$

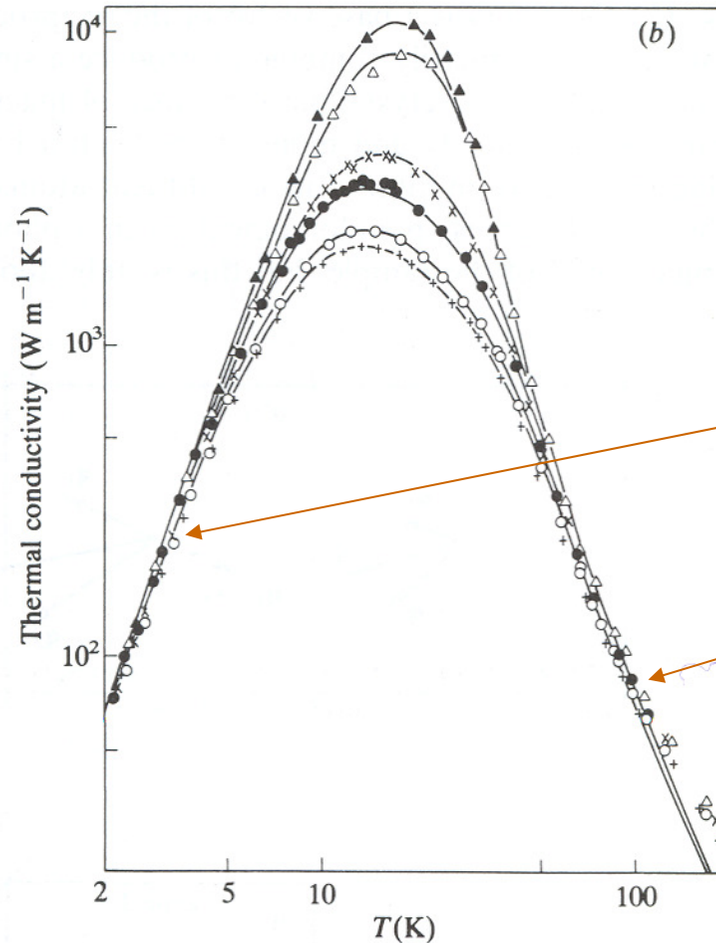
Temperature dependence of thermal conductivity as a cause of phonon scattering

The low and high T limits are summarized in this table:

	C_V	Λ	κ
low T	$\propto T^3$	$n_{\text{ph}} \rightarrow 0$, so $\Lambda \rightarrow \infty$, but then $\Lambda \rightarrow D$ (size)	$\propto T^3$
high T	$3R$	$\propto 1/T$	$\propto 1/T$

How well does this match experimental results?

Temperature dependence of thermal conductivity as a cause of phonon scattering



Experimental $\kappa(T)$

$\propto T^3$

$\propto T^{-1}$?
(not quite)

Figure 5.27 (a) The principal form for the variation of thermal conductivity. (b) Experimental data for LiF crystals containing different amounts of the isotope ${}^6\text{Li}$: \blacktriangle , 0.02% ${}^6\text{Li}$; \triangle , 0.01%; \times , 4.6%; \bullet , 9.4%; \circ , 25.4%; $+$, 50.1%. (After Berman and Brock 1965.)

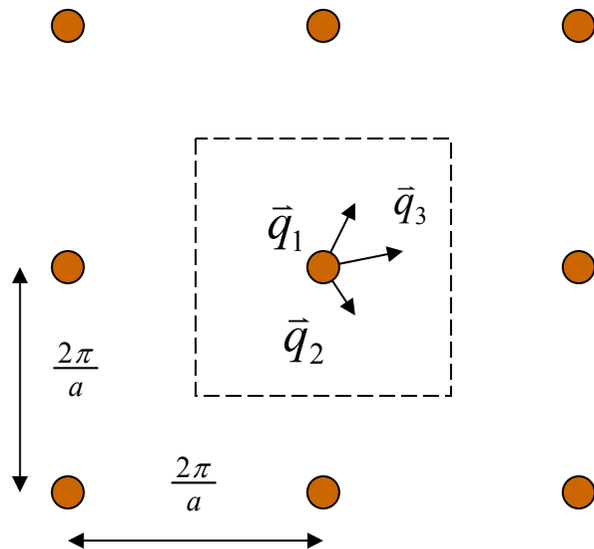
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Phonon collisions: N and U processes

How exactly do phonon collisions limit the flow of heat?

2-D lattice \rightarrow 1st BZ in k-space:



$$\hbar\vec{q}_1 + \hbar\vec{q}_2 = \hbar\vec{q}_3$$

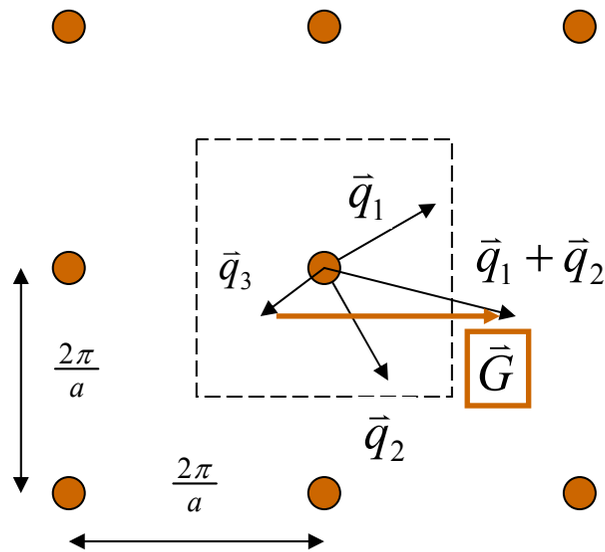
No resistance to heat flow
(N process; phonon momentum conserved)

\rightarrow Predominates at low $T \ll \theta_D$ since ω
and q will be small

Phonon collisions: N and U processes

What if the phonon wavevectors are a bit larger?

2-D lattice \rightarrow 1st BZ in k-space:



Umklapp = “flipping over” of wavevector!

$$\hbar\vec{q}_1 + \hbar\vec{q}_2 = \hbar\vec{q}_3 + \hbar\vec{G}$$

Two phonons combine to give a net phonon with an opposite momentum! This causes resistance to heat flow.

(U process; phonon momentum “lost” in units of $\hbar\vec{G}$.)

\rightarrow More likely at high $T \gg \theta_D$ since ω and q will be larger