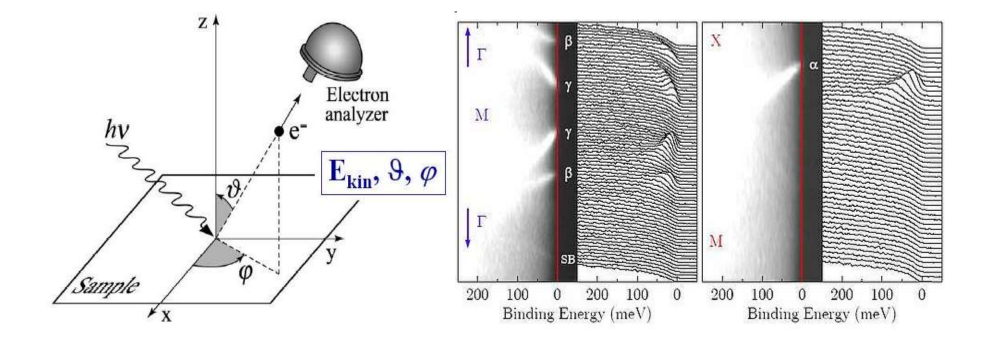
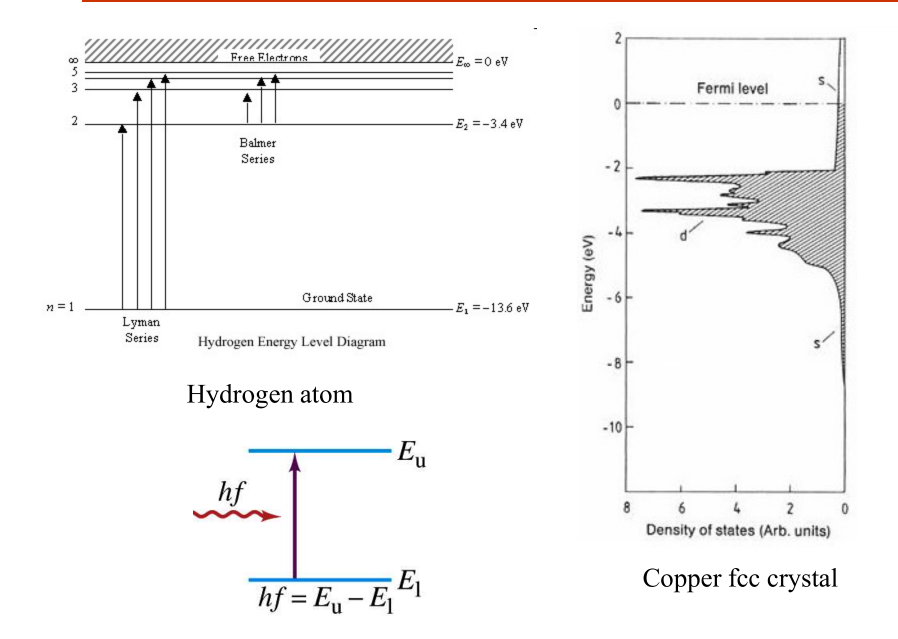


Angle resolved photoemission spectroscopy (ARPES)

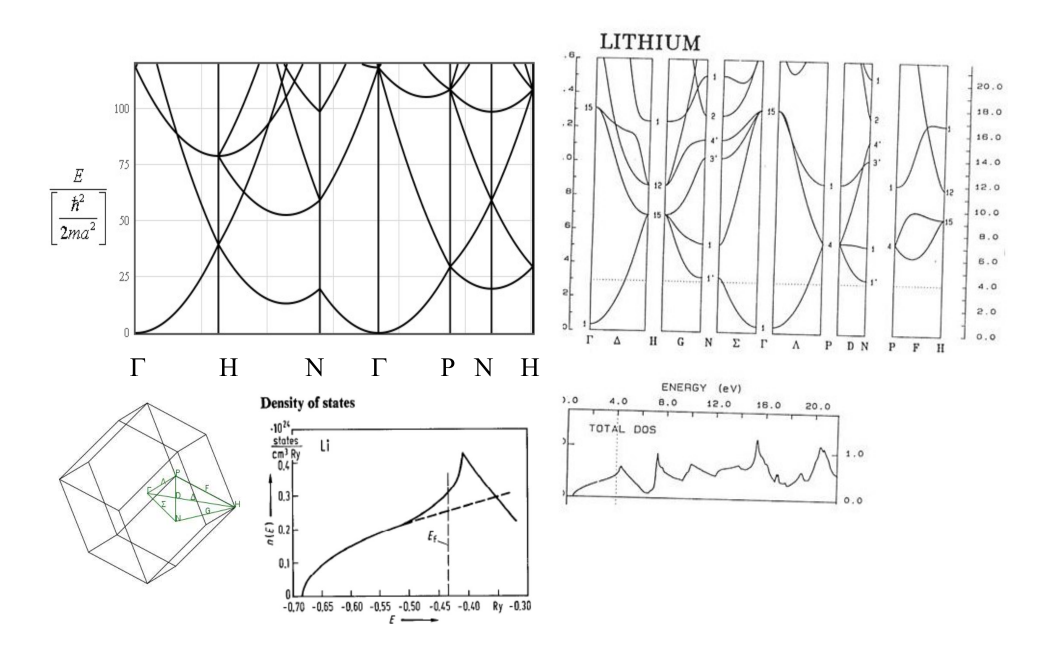


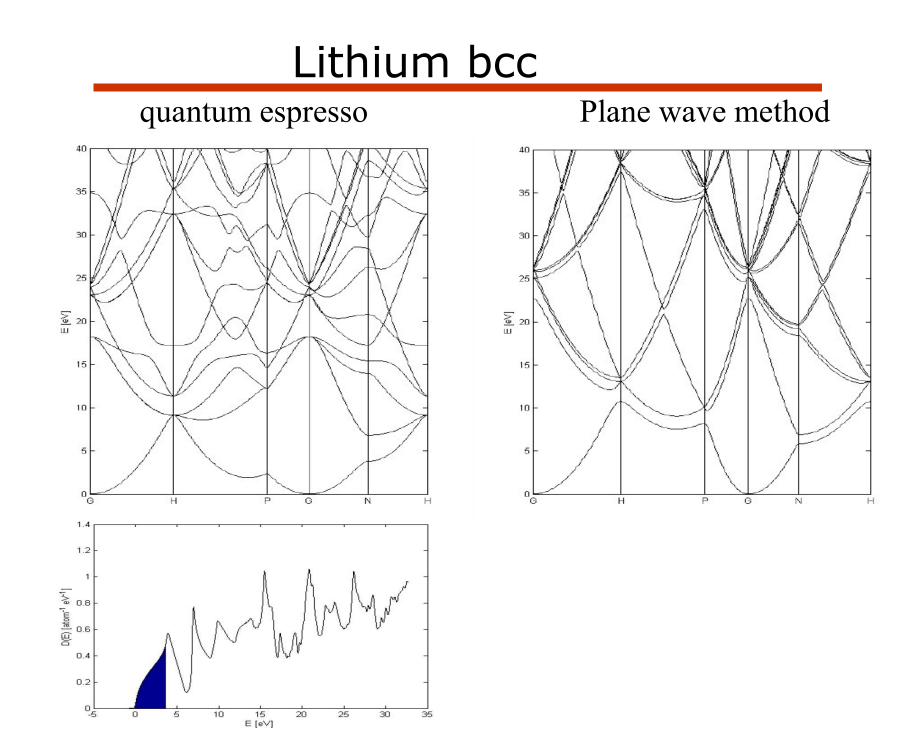
Measure the dispersion relation with angle resolved photoemission

Optical absorption

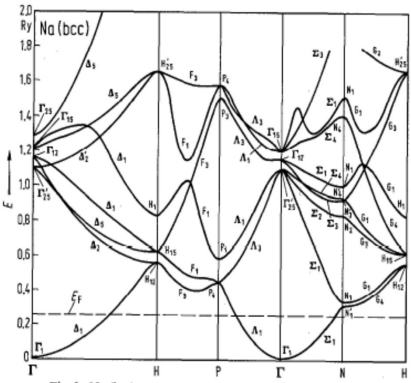


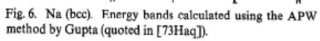
Lithium bcc

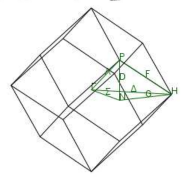


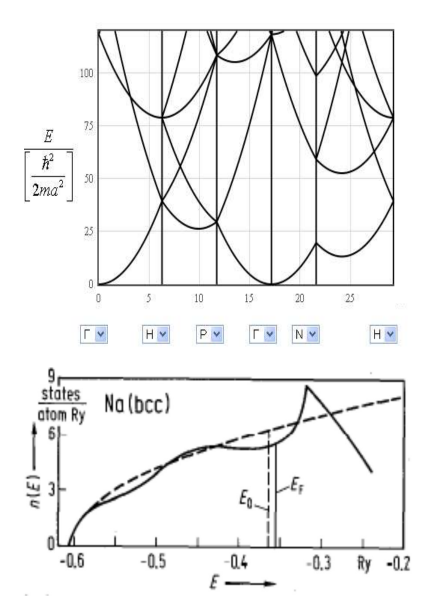


Sodium

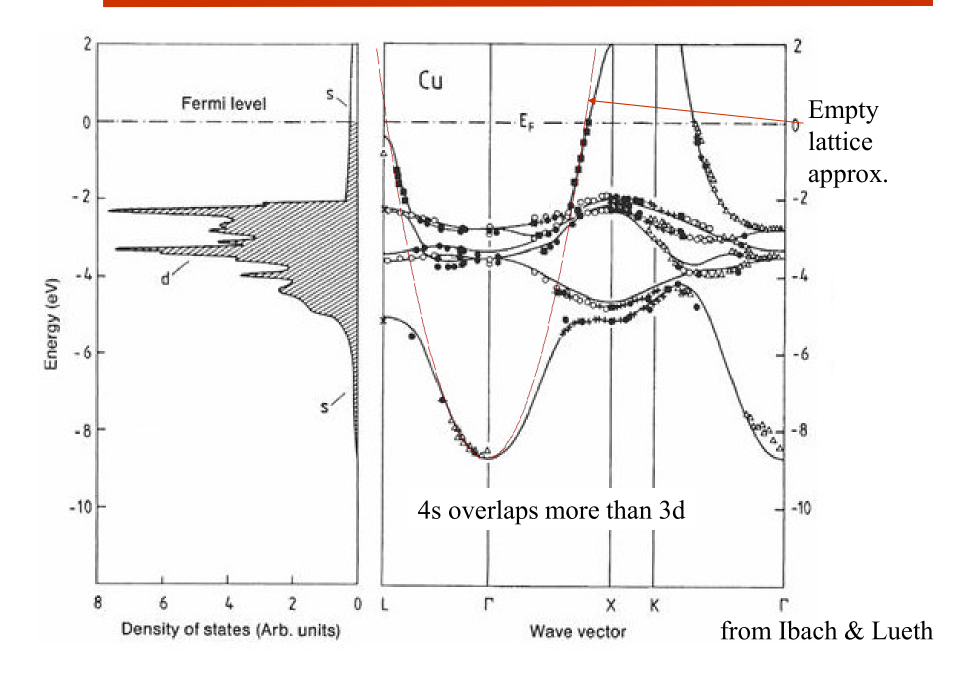




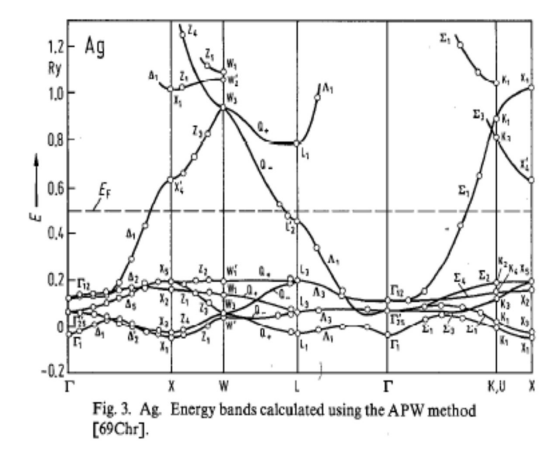




Copper dispersion relation and density of states



Silver



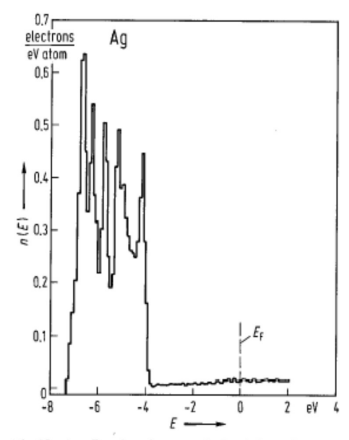
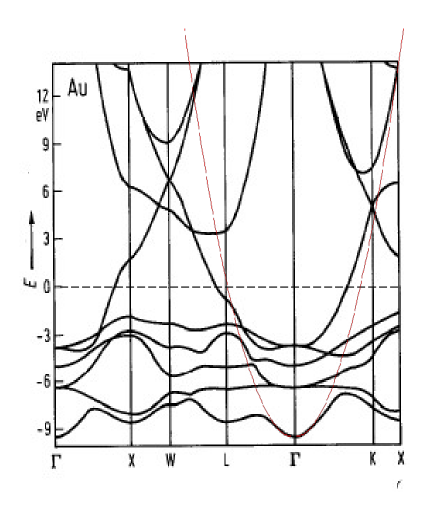


Fig. 15. Ag. Density of states calculated from the energy bands in Fig. 10. Ag [75Fon].

Gold



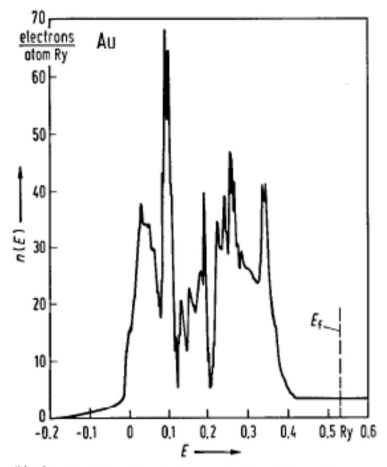
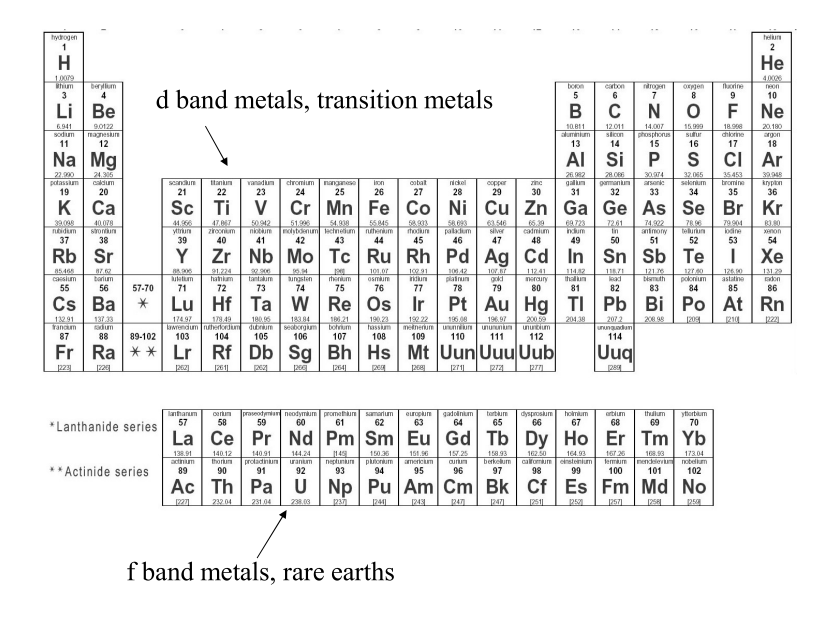
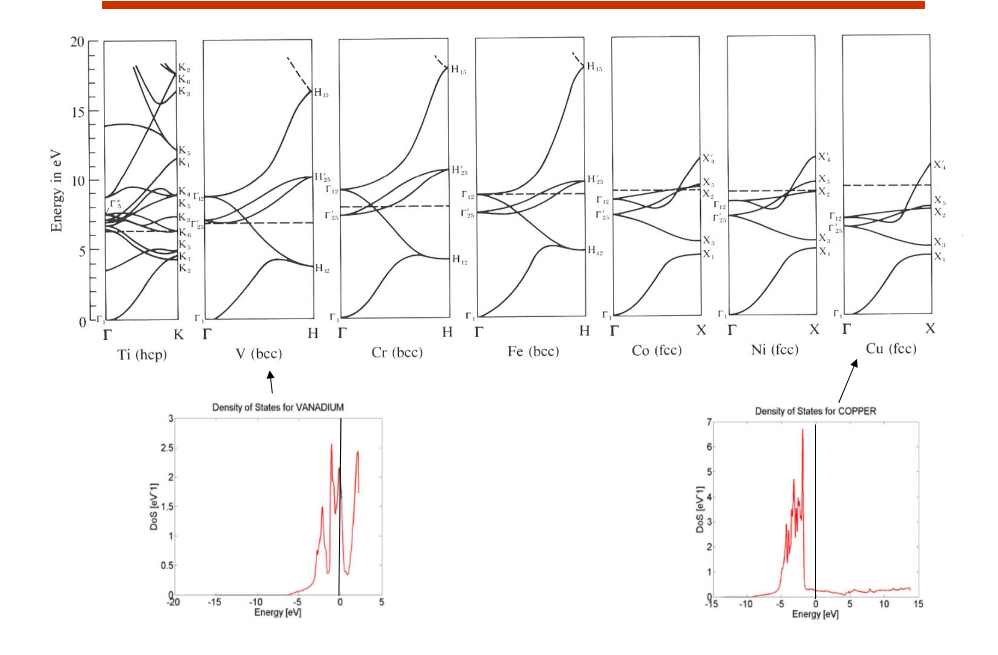


Fig. 9. Au. Density of states calculated from the energy bands in Fig. 4b. Au [71Chr2].



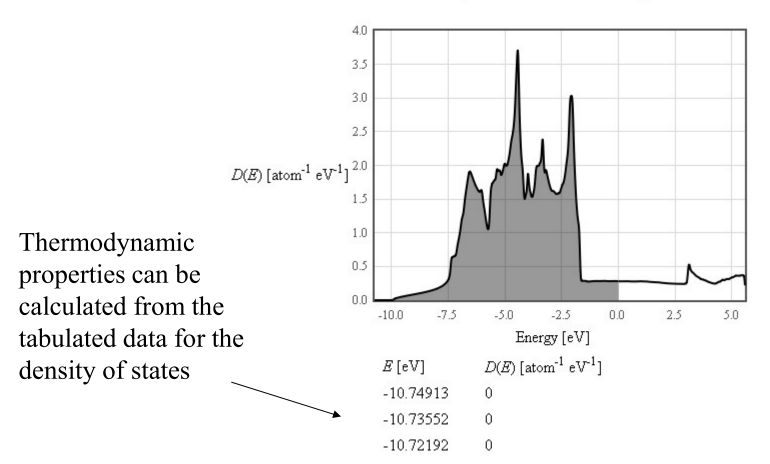
Transition metals



Springer Springer Materials The Landolt-Börnstein Database http://www.springermaterials.com/navigation/bookshelf.html#l_2_106048_ Advanced Search Bookshelf Periodic Table Help For Librarians Feedback Home > Electronic Structure and Transport > Metals: Electron and Phonon States > Band structures and Fermi surfaces of metallic Particles, Nuclei and Atoms elements Molecules and Radicals Band structures and Fermi surfaces of metallic elements Electronic Structure and Transport Introduction i Magnetism Literature survey of calculations and experiments 📆 i Data for Ac...Bi Semiconductivity Data for C...Cu Superconductivity. Data for Dy ... Ir Data for K...Nd Crystallography Data for Ni...Ru Data for Sb ... Ti Thermodynamics Data for TI...Zr References Multiphase Systems Advanced Materials Advanced Technologies Astro- and Geophysics ÷ Inorganic Solid Phases 2 Thermophysical Properties Chemical Safety

Thermodynamic properties of metals

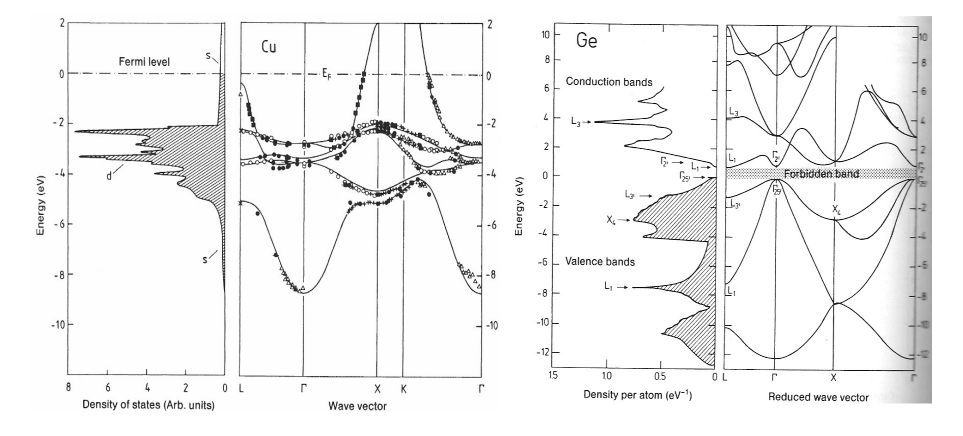
From the band structure measurements, we obtain the electron density of states.



Electron density of states for fcc gold

- o Metals, semimetals, semiconductors, insulators
 - Numerical determination of the thermodynamic properties of metals
 - Chemical potential $\mu(T)$
 - Energy spectral density u(E,T)
 - Internal energy density u(T)
 - Specific heat $c_v(T)$
 - Calculated electron density of states
 - Free electron model in 1-D
 - Free electron model in 2-D
 - Free electron model in 3-D
 - Al fee, Au fee, Cu fee, Na bee, Pt fee, W bee, Si diamond, Fe bee, Ni fee, Co fee, Mn bee, fee, Pd₃Cr, Pd₃Mn, PdCr, PdMn
 - Separable square wave potentials

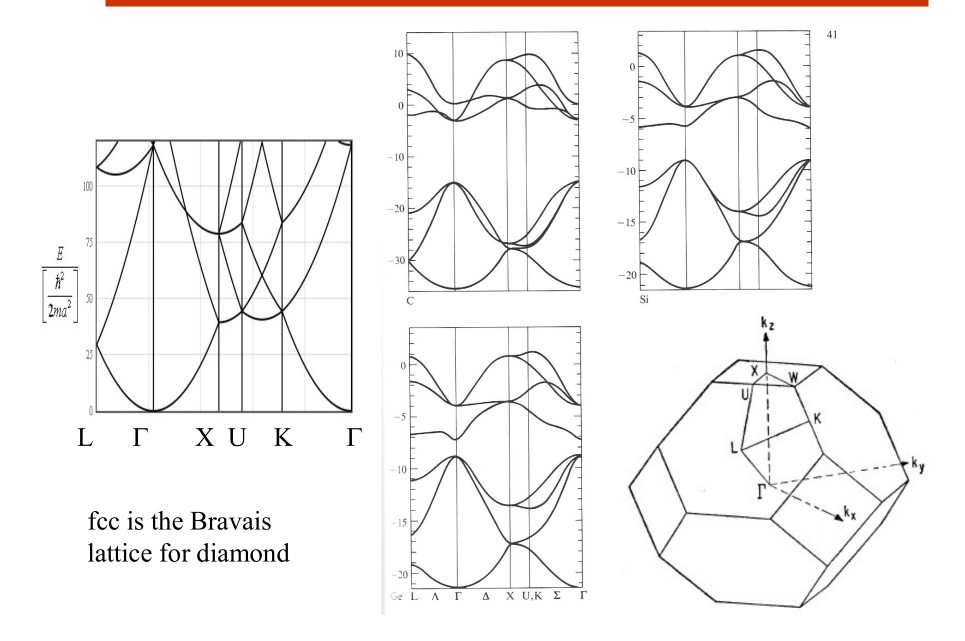
Metals, semiconductors, and insulators



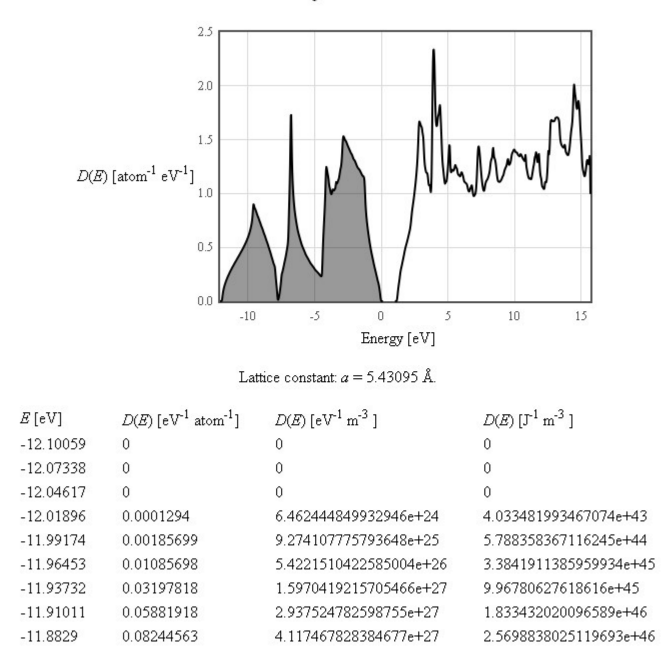
Insulators: band gap > 3 eV

From Ibach & Lueth

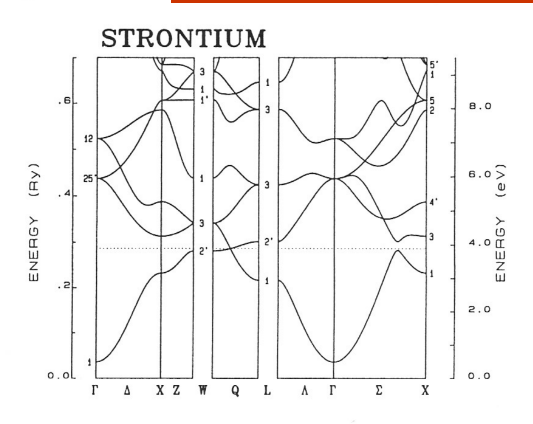
Group IV

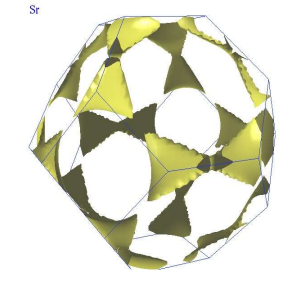


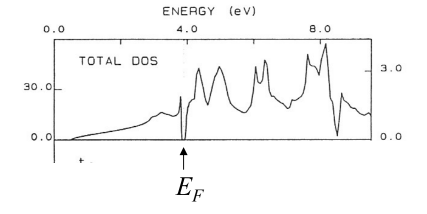
Electron density of states for silicon



Semimetal Strontium







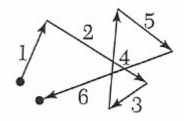
kinetic theory

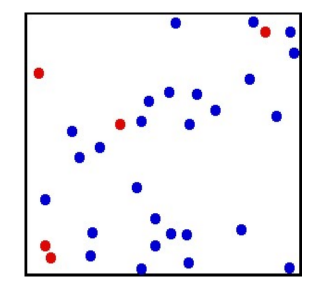
describe electrons as a gas of particles

$$v_F = 10^8 \text{ cm/s.}$$

The average time between scattering events τ_{sc} can be calculated by Fermi's golden rule

mean free path: $l = v_F \tau_{sc} \sim 1 \text{ nm} - 1 \text{ cm}$





Electrons as waves or particles

Scattering of electrons can be thought of as transitions between *k* states or as collisions between particles.

Umklapp scattering of electrons by phonons makes large changes in the momentum of the electrons because of the reciprocal lattice vector **G**.

$$\vec{k}_{el}' = \vec{k}_{el} + \vec{k}_{ph}$$
 \leftarrow phonon emitted