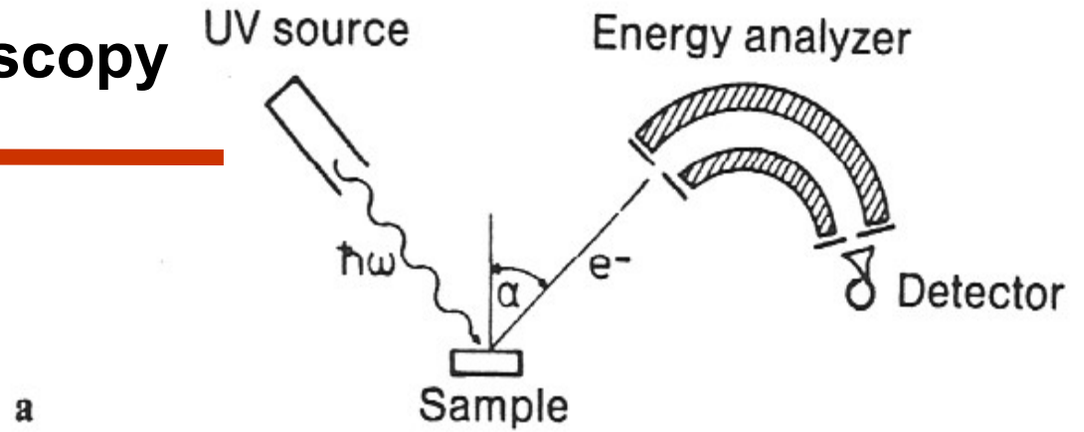


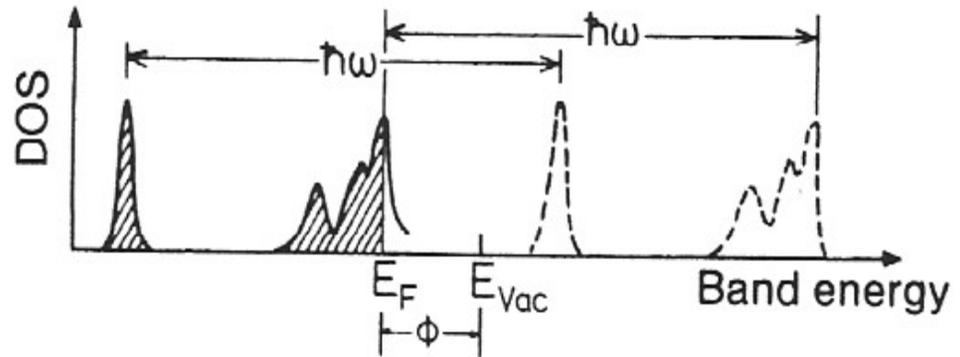
# Photoemission spectroscopy

UPS - Ultraviolet photoemission spectroscopy

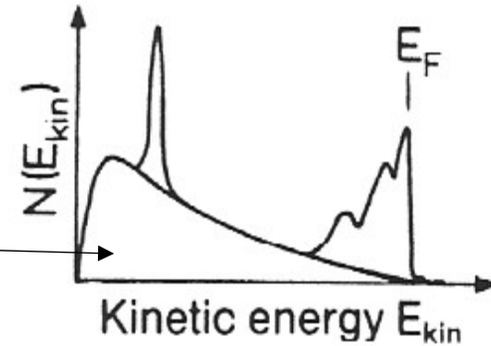
Measure the density of states with photoemission spectroscopy



a



Secondary electrons



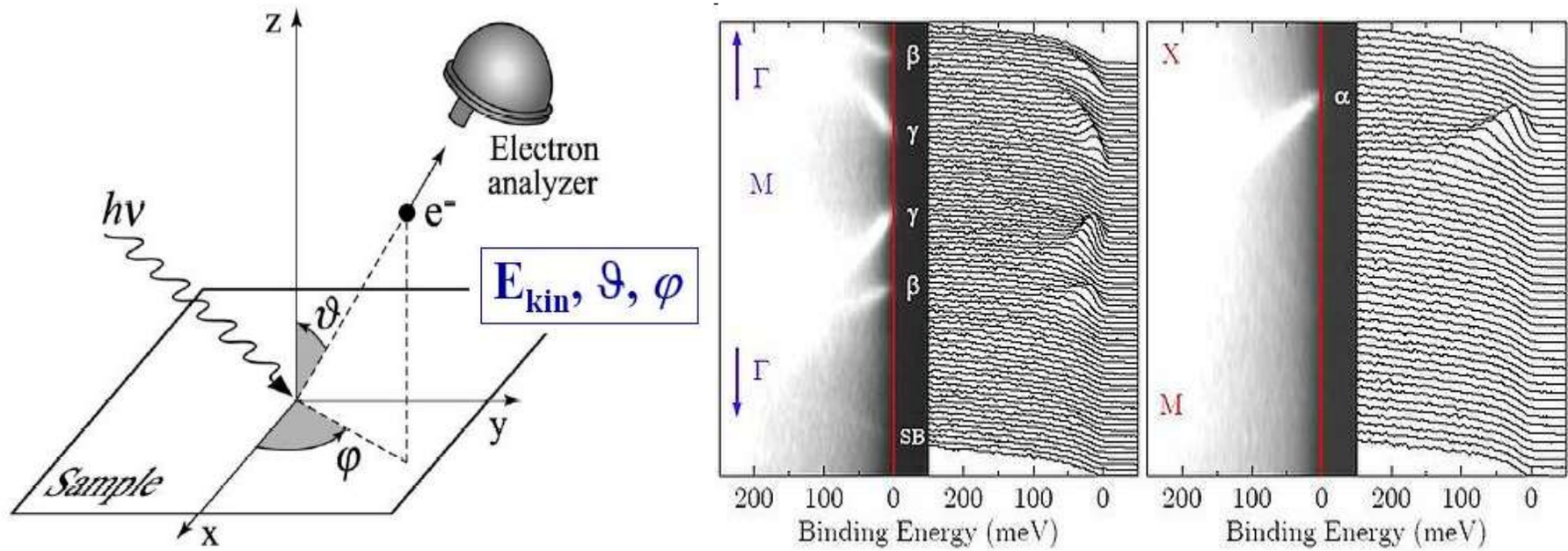
Binding energy

From: Ibach & Lueth

b

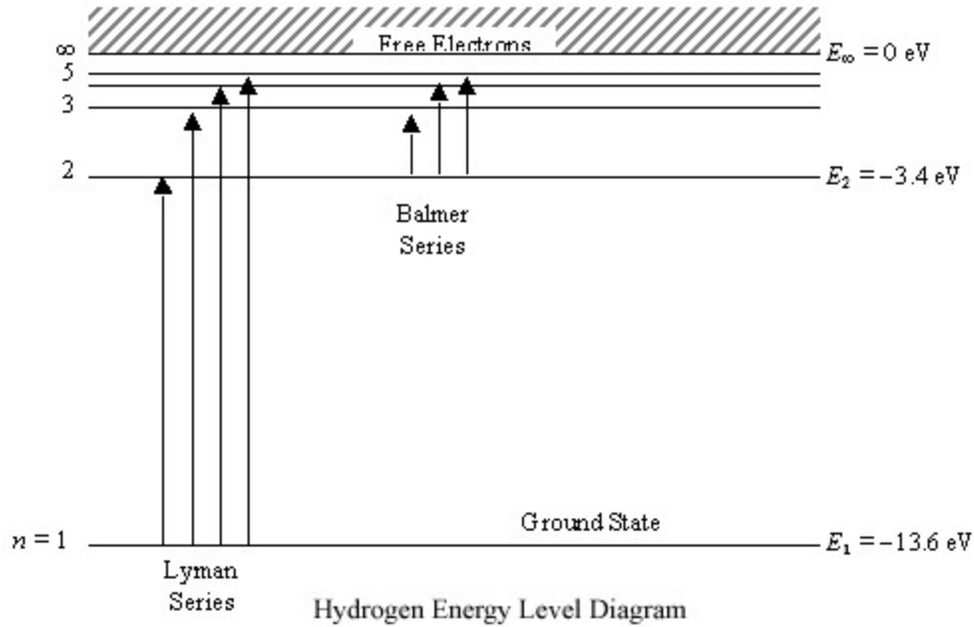
# Angle resolved photoemission spectroscopy (ARPES)

---

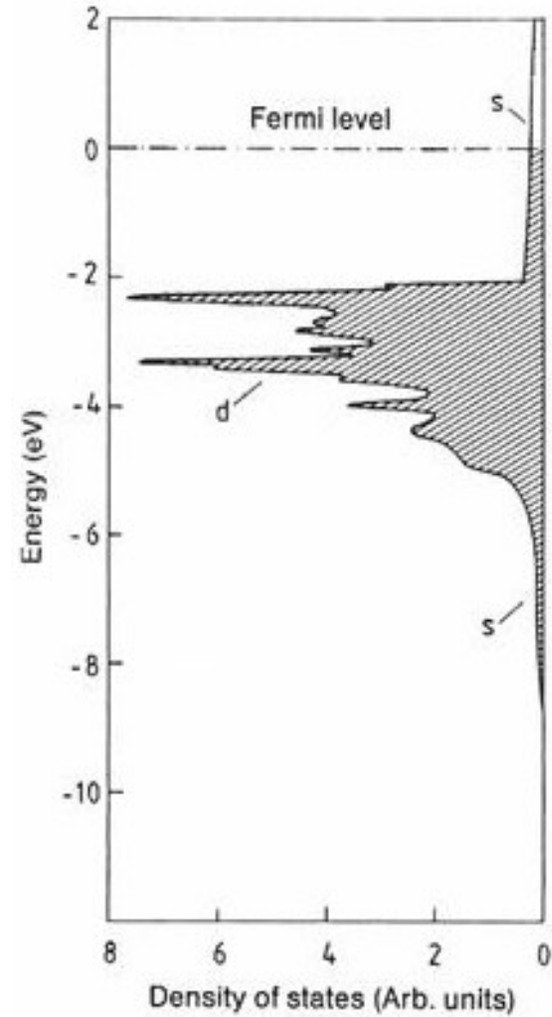
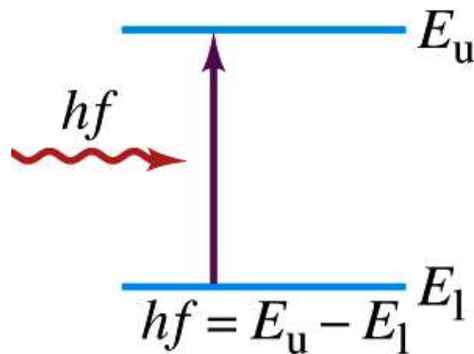


Measure the dispersion relation with angle resolved photoemission

# Optical absorption

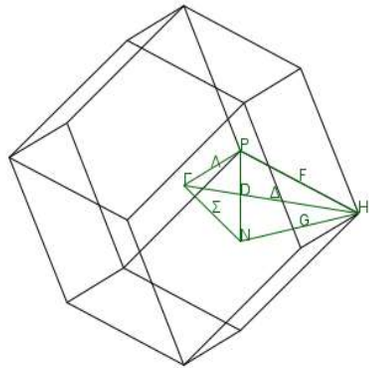
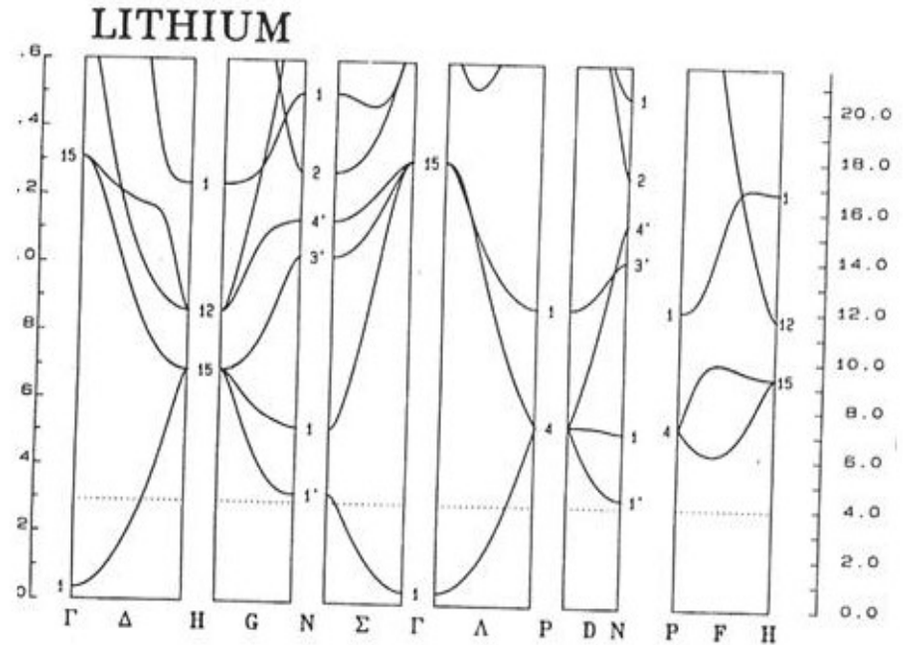
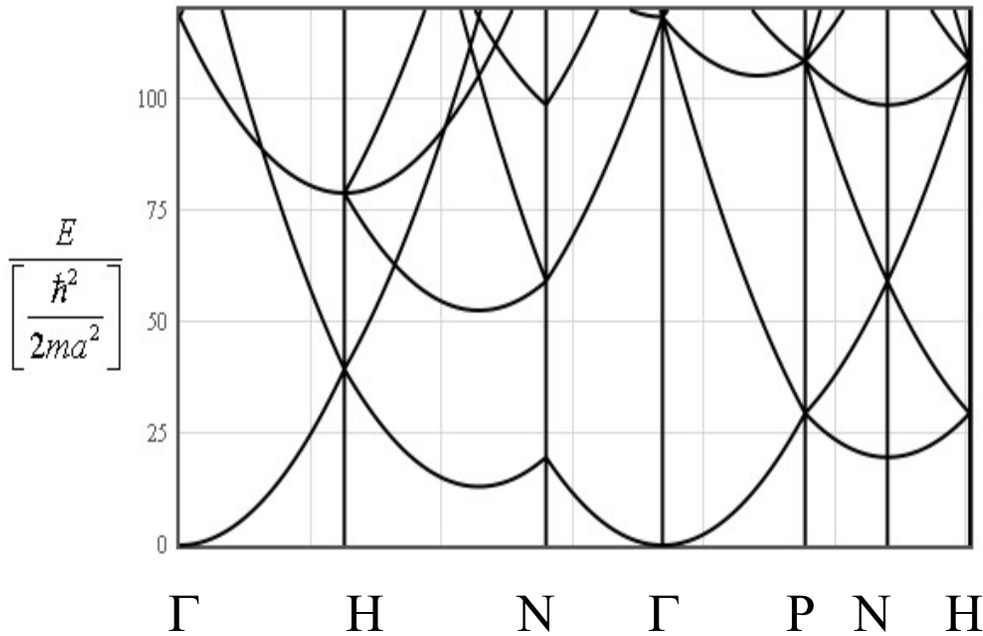


Hydrogen atom

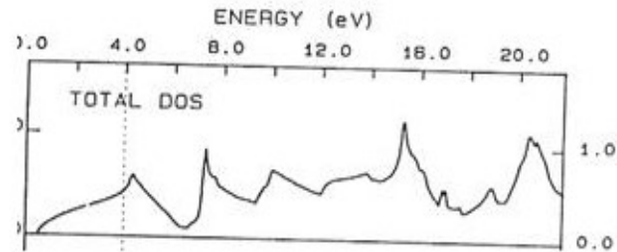
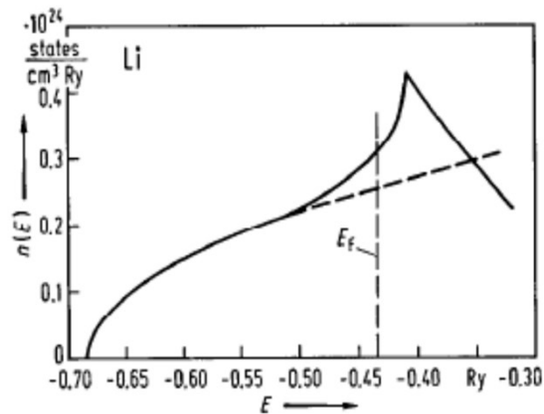


Copper fcc crystal

# Lithium bcc

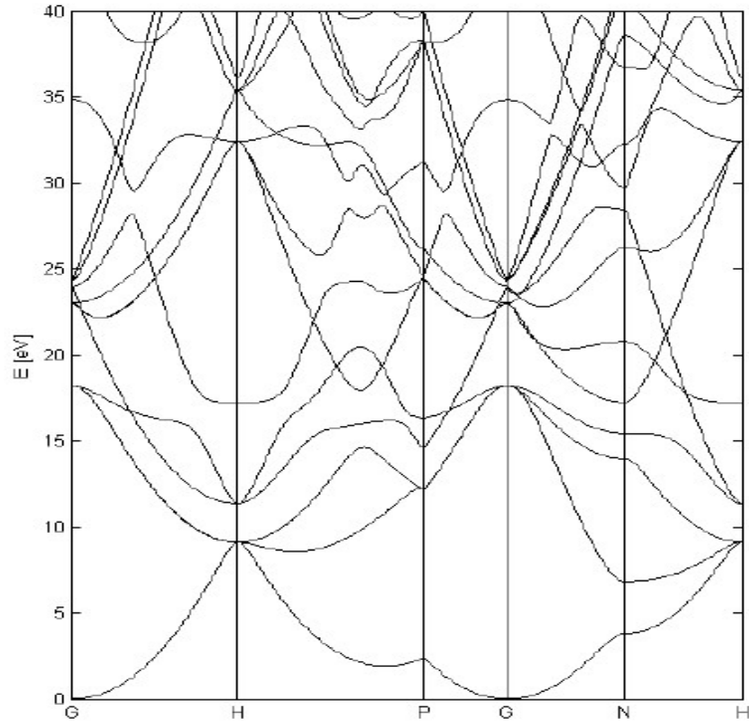


Density of states

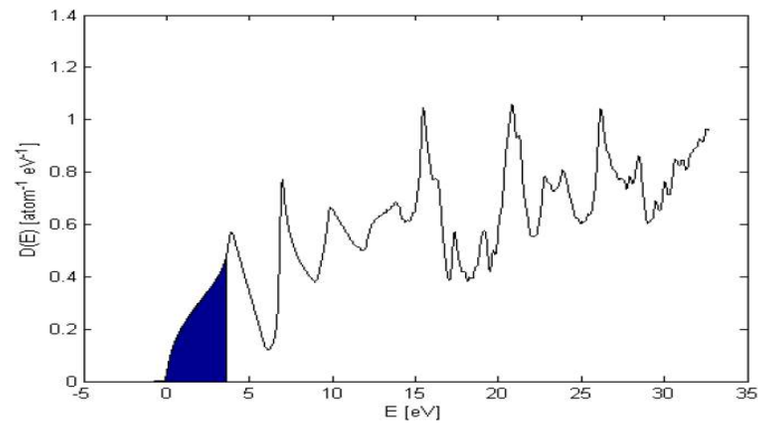
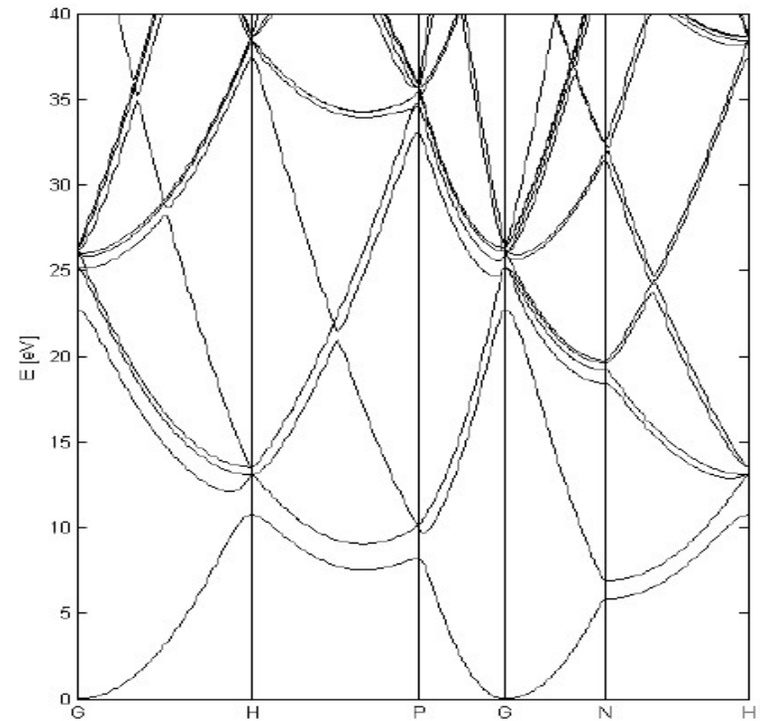


# Lithium bcc

quantum espresso



Plane wave method



# Sodium

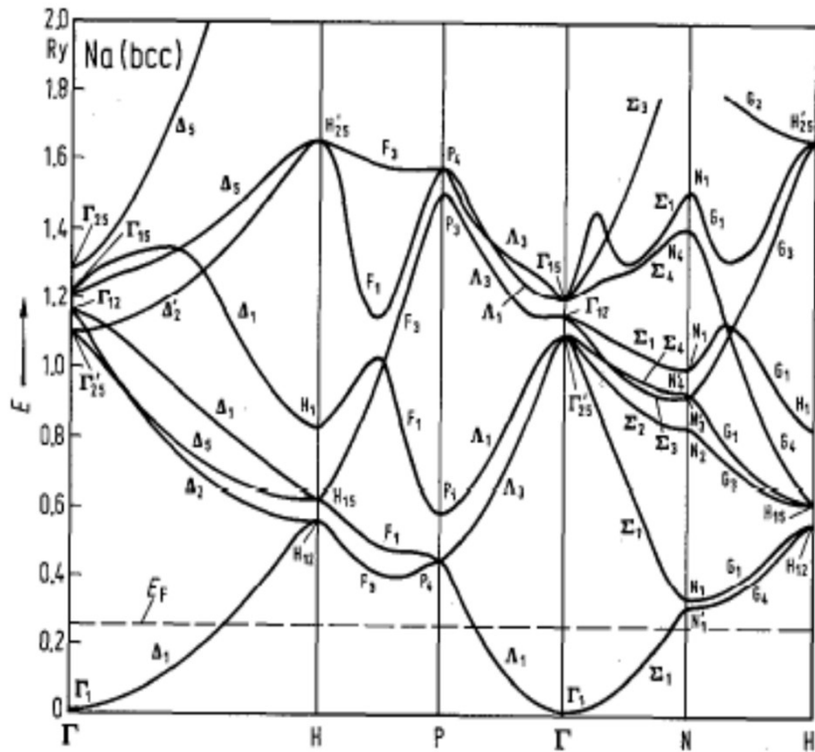
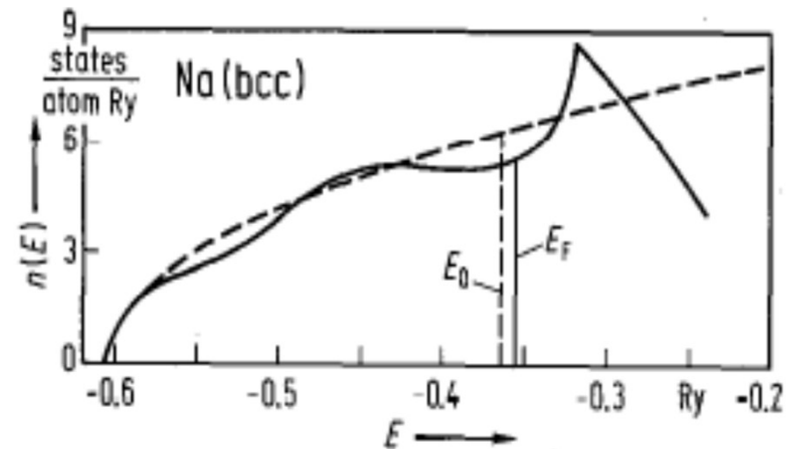
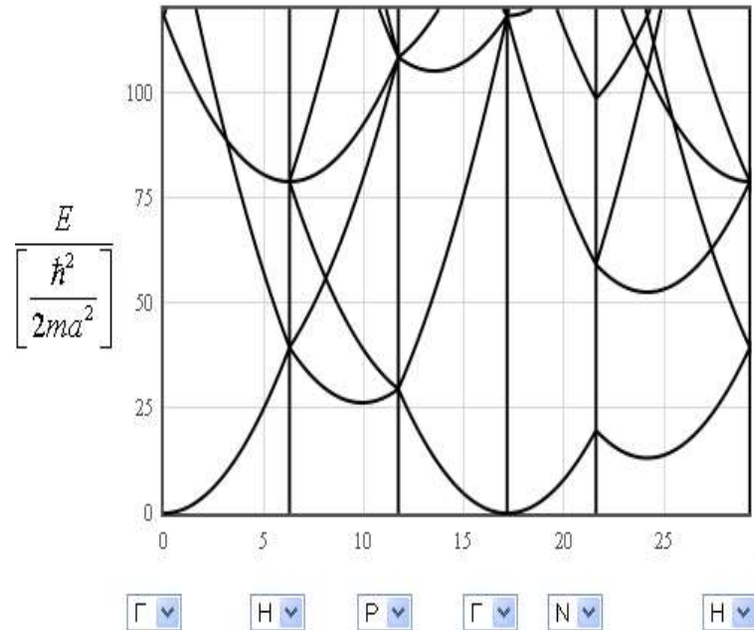
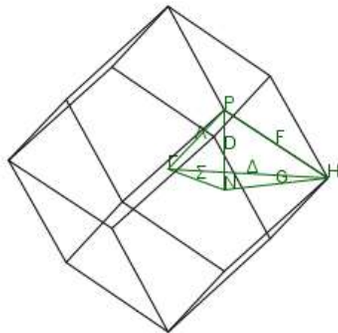
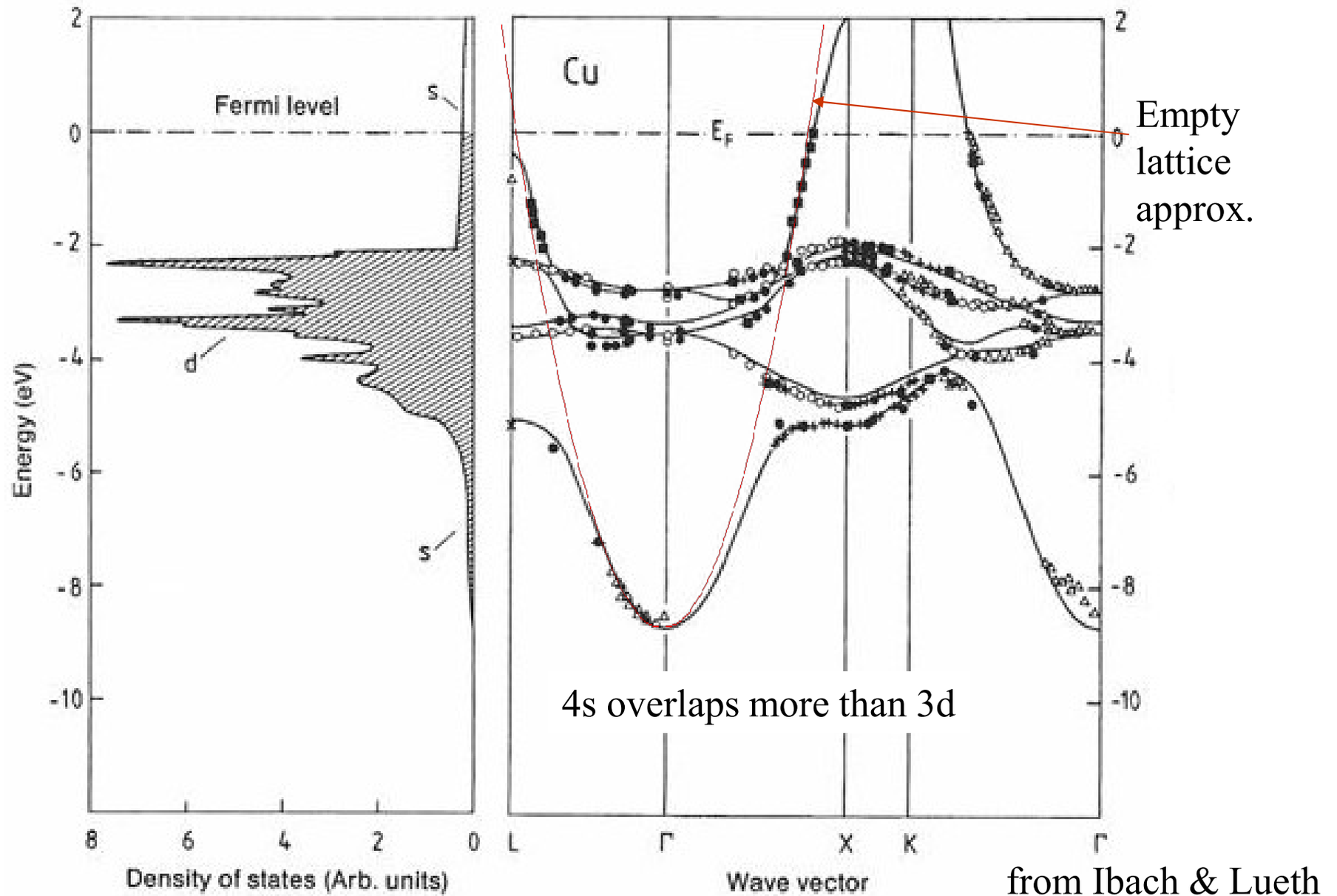


Fig. 6. Na (bcc). Energy bands calculated using the APW method by Gupta (quoted in [73Hag]).



# Copper dispersion relation and density of states



# Silver

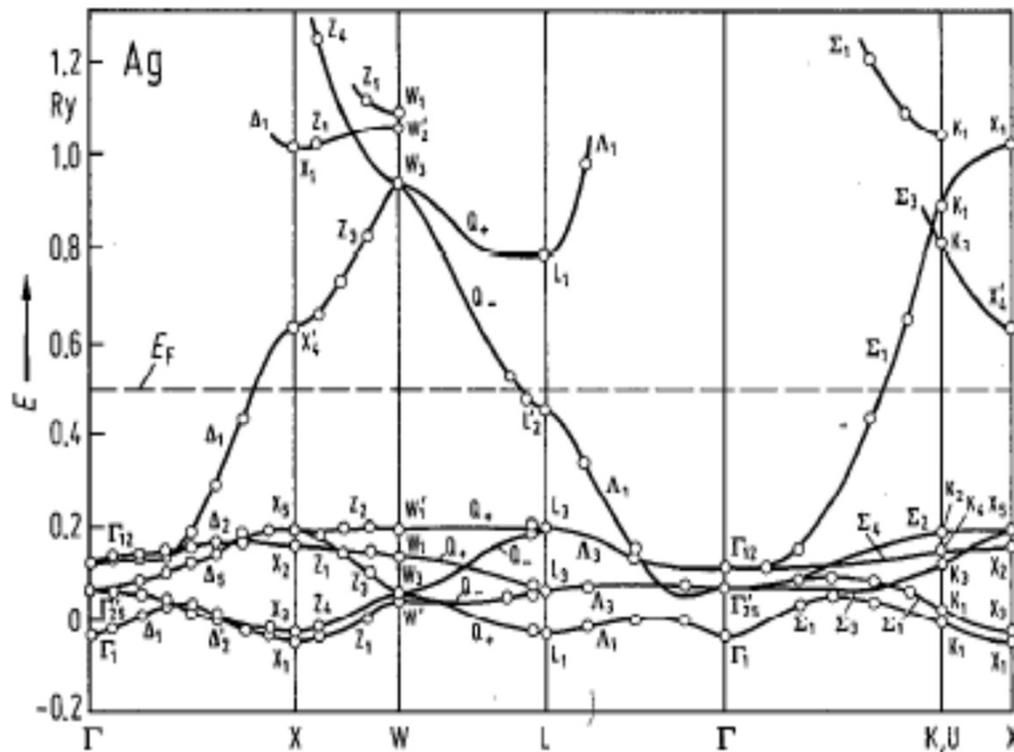


Fig. 3. Ag. Energy bands calculated using the APW method [69Chr].

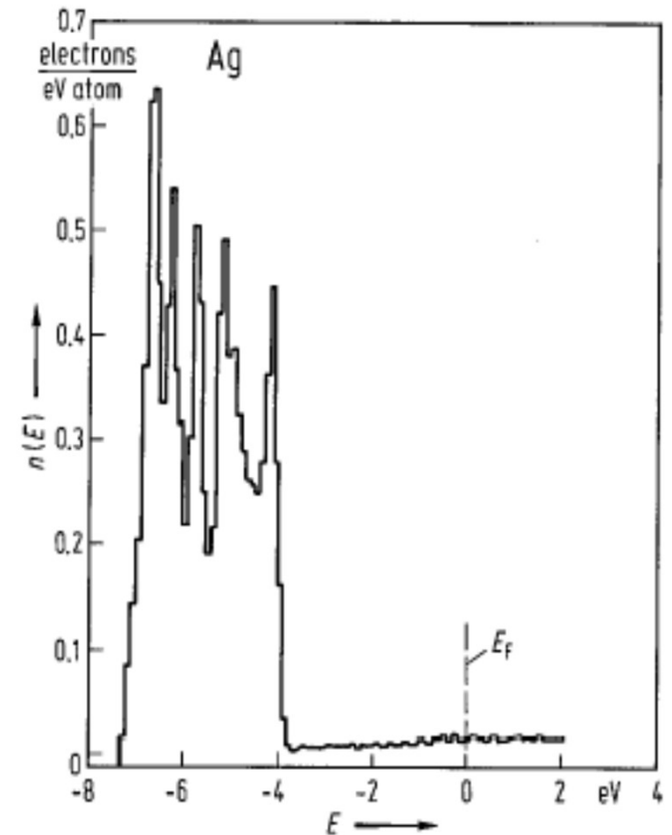


Fig. 15. Ag. Density of states calculated from the energy bands in Fig. 10. Ag [75Fon].

# Gold

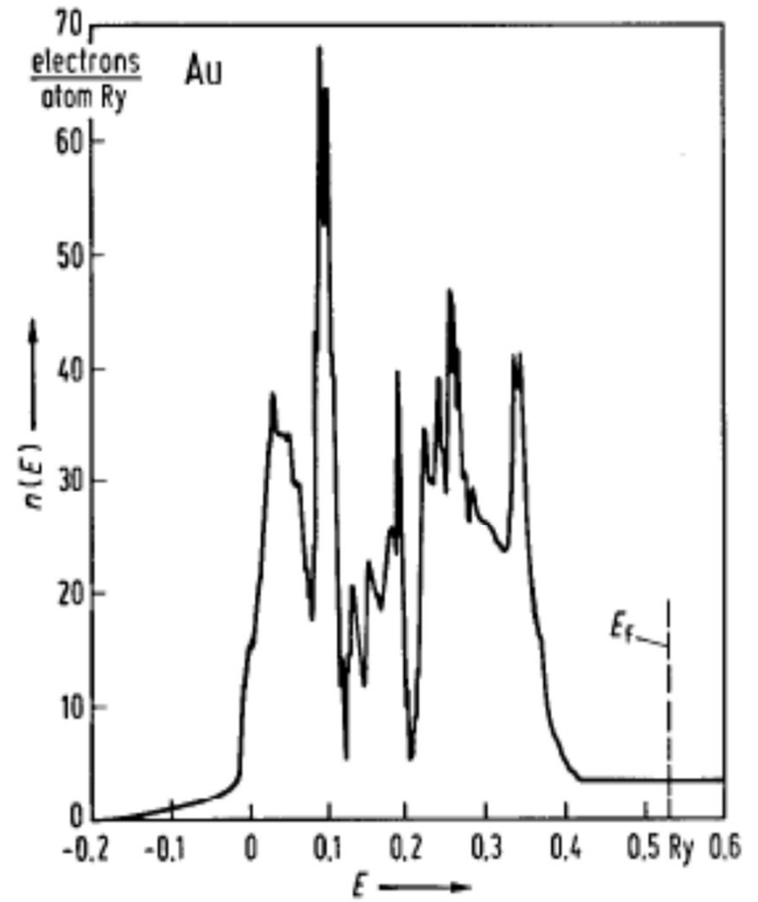
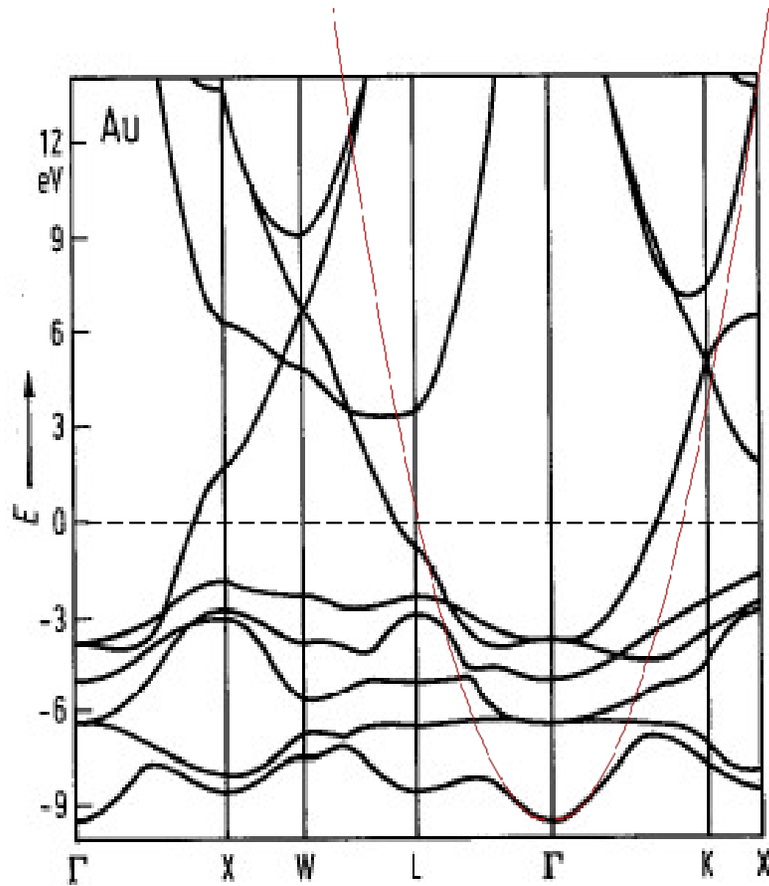


Fig. 9. Au. Density of states calculated from the energy bands in Fig. 4b. Au [71Chr2].

hydrogen 1 <b>H</b> 1.0079																	helium 2 <b>He</b> 4.0026						
lithium 3 <b>Li</b> 6.941	beryllium 4 <b>Be</b> 9.0122																	boron 5 <b>B</b> 10.811	carbon 6 <b>C</b> 12.011	nitrogen 7 <b>N</b> 14.007	oxygen 8 <b>O</b> 15.999	fluorine 9 <b>F</b> 18.998	neon 10 <b>Ne</b> 20.180
sodium 11 <b>Na</b> 22.990	magnesium 12 <b>Mg</b> 24.305																	aluminium 13 <b>Al</b> 26.982	silicon 14 <b>Si</b> 28.086	phosphorus 15 <b>P</b> 30.974	sulfur 16 <b>S</b> 32.065	chlorine 17 <b>Cl</b> 35.453	argon 18 <b>Ar</b> 39.948
potassium 19 <b>K</b> 39.098	calcium 20 <b>Ca</b> 40.078	scandium 21 <b>Sc</b> 44.956	titanium 22 <b>Ti</b> 47.867	vanadium 23 <b>V</b> 50.942	chromium 24 <b>Cr</b> 51.996	manganese 25 <b>Mn</b> 54.938	iron 26 <b>Fe</b> 55.845	cobalt 27 <b>Co</b> 58.933	nickel 28 <b>Ni</b> 58.693	copper 29 <b>Cu</b> 63.546	zinc 30 <b>Zn</b> 65.39	gallium 31 <b>Ga</b> 69.723	germanium 32 <b>Ge</b> 72.61	arsenic 33 <b>As</b> 74.922	selenium 34 <b>Se</b> 78.96	bromine 35 <b>Br</b> 79.904	krypton 36 <b>Kr</b> 83.80						
rubidium 37 <b>Rb</b> 85.468	strontium 38 <b>Sr</b> 87.62	yttrium 39 <b>Y</b> 88.906	zirconium 40 <b>Zr</b> 91.224	niobium 41 <b>Nb</b> 92.906	molybdenum 42 <b>Mo</b> 95.94	technetium 43 <b>Tc</b> [98]	ruthenium 44 <b>Ru</b> 101.07	rhodium 45 <b>Rh</b> 102.91	palladium 46 <b>Pd</b> 106.42	silver 47 <b>Ag</b> 107.87	cadmium 48 <b>Cd</b> 112.41	indium 49 <b>In</b> 114.82	tin 50 <b>Sn</b> 118.71	antimony 51 <b>Sb</b> 121.76	tellurium 52 <b>Te</b> 127.60	iodine 53 <b>I</b> 126.90	xenon 54 <b>Xe</b> 131.29						
caesium 55 <b>Cs</b> 132.91	barium 56 <b>Ba</b> 137.33	57-70 *	lutetium 71 <b>Lu</b> 174.97	hafnium 72 <b>Hf</b> 178.49	tantalum 73 <b>Ta</b> 180.95	tungsten 74 <b>W</b> 183.84	rhenium 75 <b>Re</b> 186.21	osmium 76 <b>Os</b> 190.23	iridium 77 <b>Ir</b> 192.22	platinum 78 <b>Pt</b> 195.08	gold 79 <b>Au</b> 196.97	mercury 80 <b>Hg</b> 200.59	thallium 81 <b>Tl</b> 204.38	lead 82 <b>Pb</b> 207.2	bismuth 83 <b>Bi</b> 208.98	polonium 84 <b>Po</b> [209]	astatine 85 <b>At</b> [210]	radon 86 <b>Rn</b> [222]					
francium 87 <b>Fr</b> [223]	radium 88 <b>Ra</b> [226]	89-102 **	lawrencium 103 <b>Lr</b> [262]	rutherfordium 104 <b>Rf</b> [261]	dubnium 105 <b>Db</b> [262]	seaborgium 106 <b>Sg</b> [266]	bohrium 107 <b>Bh</b> [264]	hassium 108 <b>Hs</b> [269]	meitnerium 109 <b>Mt</b> [268]	ununnium 110 <b>Uun</b> [271]	ununium 111 <b>Uuu</b> [272]	ununium 112 <b>Uub</b> [277]	ununquadium 114 <b>Uuq</b> [289]										

d band metals, transition metals



\* Lanthanide series

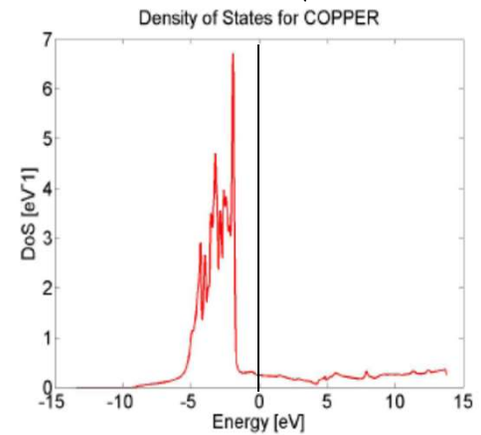
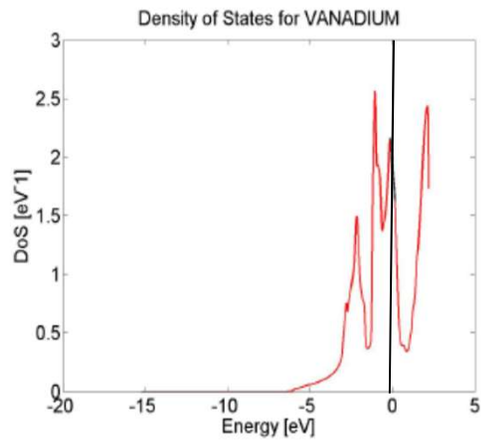
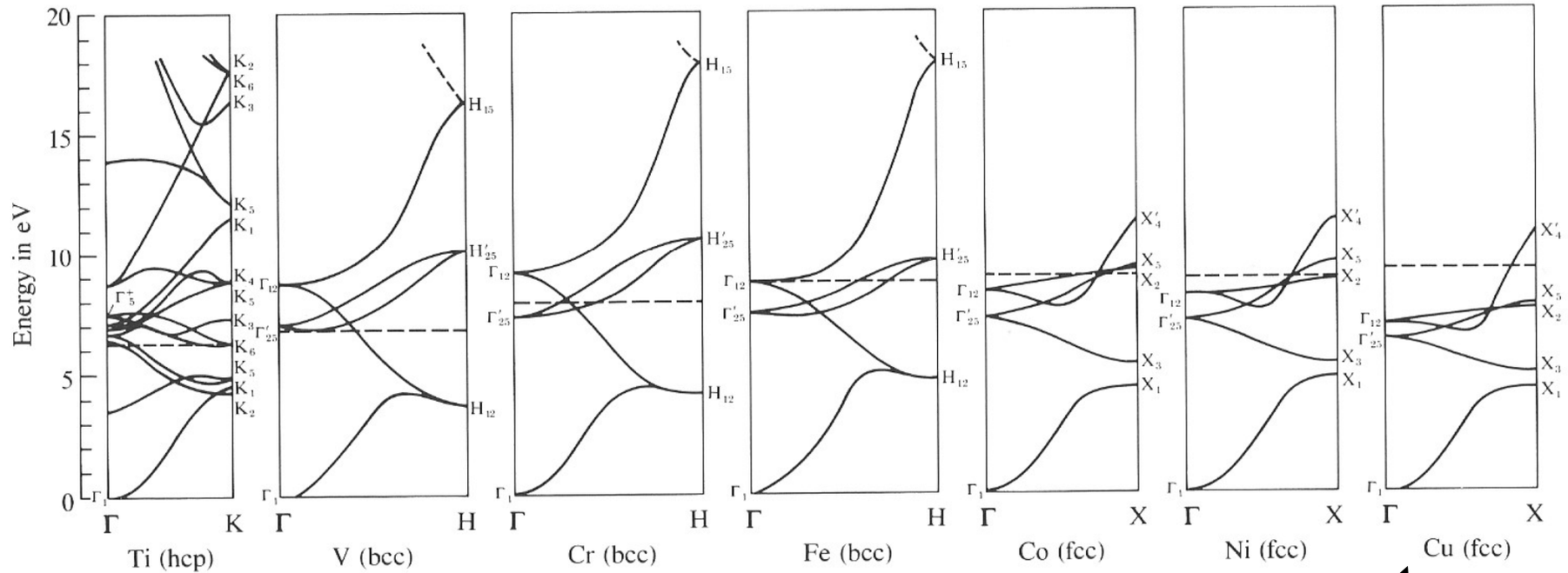
\*\* Actinide series

lanthanum 57 <b>La</b> 138.91	cerium 58 <b>Ce</b> 140.12	praseodymium 59 <b>Pr</b> 140.91	neodymium 60 <b>Nd</b> 144.24	promethium 61 <b>Pm</b> [145]	samarium 62 <b>Sm</b> 150.36	europium 63 <b>Eu</b> 151.96	gadolinium 64 <b>Gd</b> 157.25	terbium 65 <b>Tb</b> 158.93	dysprosium 66 <b>Dy</b> 162.50	holmium 67 <b>Ho</b> 164.93	erbium 68 <b>Er</b> 167.26	thulium 69 <b>Tm</b> 168.93	ytterbium 70 <b>Yb</b> 173.04
actinium 89 <b>Ac</b> [227]	thorium 90 <b>Th</b> 232.04	protactinium 91 <b>Pa</b> 231.04	uranium 92 <b>U</b> 238.03	neptunium 93 <b>Np</b> [237]	plutonium 94 <b>Pu</b> [244]	americium 95 <b>Am</b> [243]	curium 96 <b>Cm</b> [247]	berkelium 97 <b>Bk</b> [247]	californium 98 <b>Cf</b> [251]	einsteinium 99 <b>Es</b> [252]	fermium 100 <b>Fm</b> [257]	mendelevium 101 <b>Md</b> [258]	nobelium 102 <b>No</b> [259]

f band metals, rare earths



# Transition metals



Go Advanced Search

Home

Bookshelf

Periodic Table

Help

For Librarians

Feedback

Particles, Nuclei and Atoms

Molecules and Radicals

Electronic Structure and Transport

Magnetism

Semiconductivity

Superconductivity

Crystallography

Thermodynamics

Multiphase Systems

Advanced Materials

Advanced Technologies

Astro- and Geophysics

Inorganic Solid Phases 

Thermophysical Properties 

Chemical Safety

> Electronic Structure and Transport > Metals: Electron and Phonon States > Band structures and Fermi surfaces of metallic elements

## Band structures and Fermi surfaces of metallic elements

Introduction  

Literature survey of calculations and experiments  

Data for Ac...Bi

Data for C...Cu

Data for Dy...Ir

Data for K...Nd

Data for Ni...Ru

Data for Sb...Ti

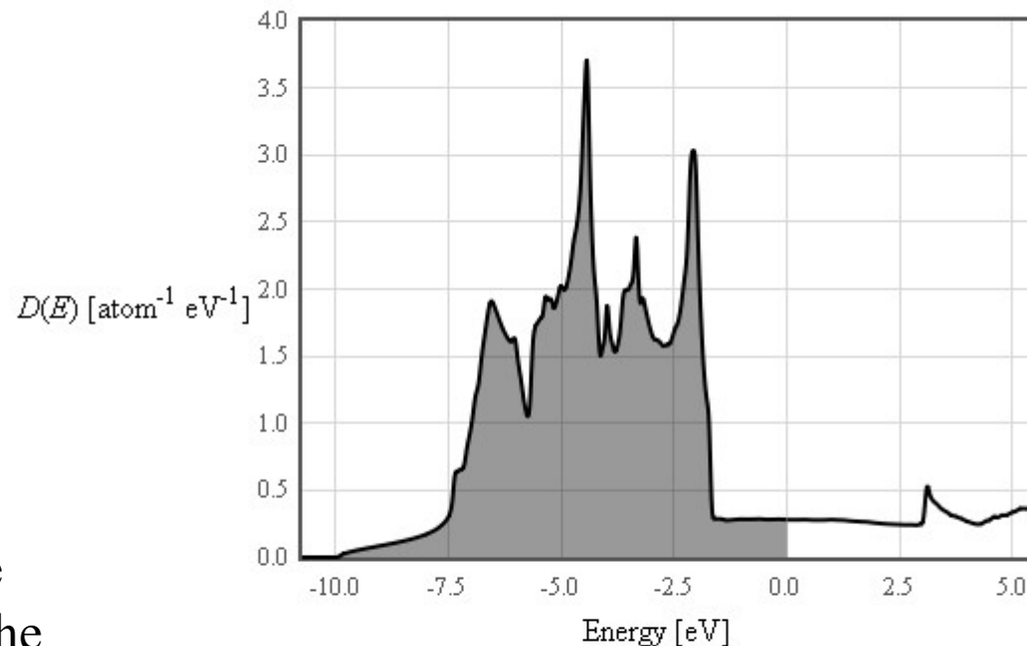
Data for Tl...Zr

References

# Thermodynamic properties of metals

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

Electron density of states for fcc gold

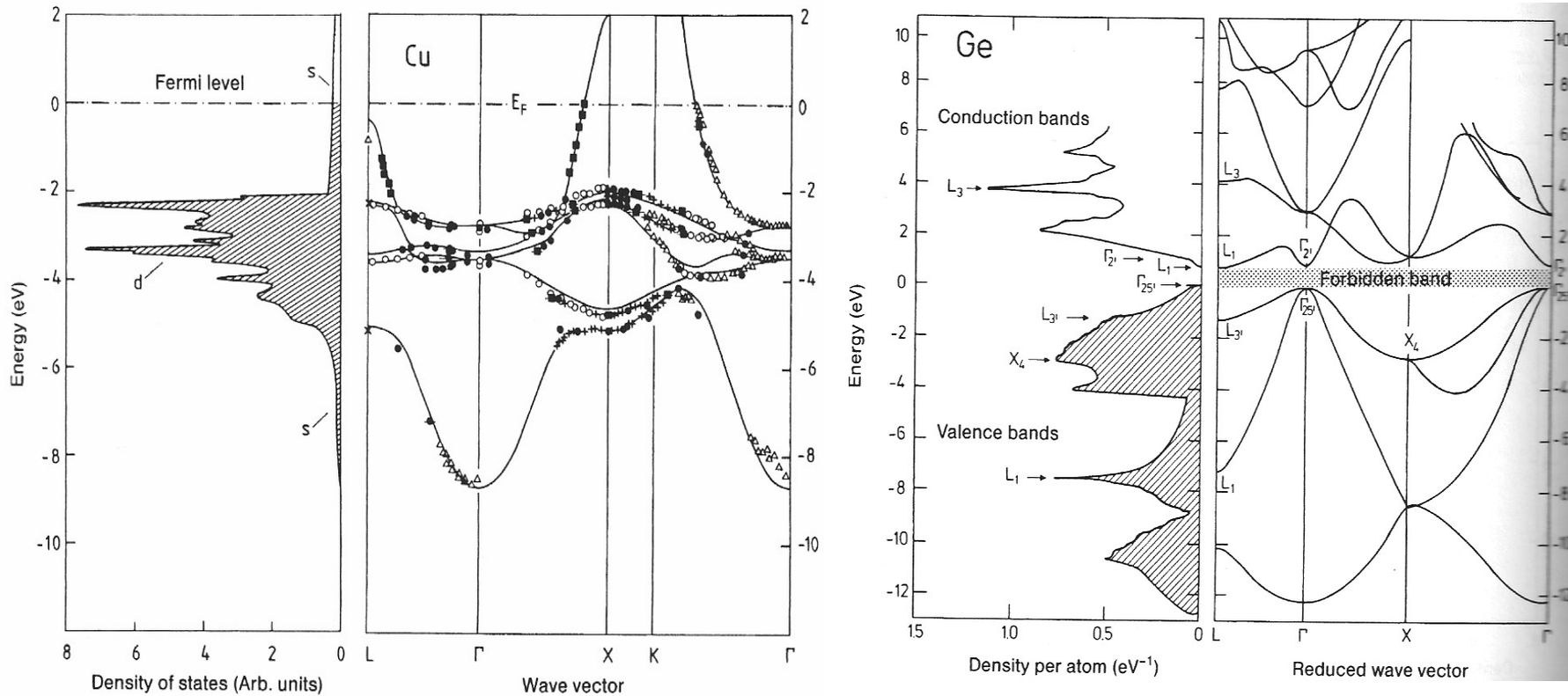


Thermodynamic properties can be calculated from the tabulated data for the density of states

$E$ [eV]	$D(E)$ [ $\text{atom}^{-1} \text{eV}^{-1}$ ]
-10.74913	0
-10.73552	0
-10.72192	0

- 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 fcc, Au fcc, Cu fcc, Na bcc, Pt fcc, W bcc, Si diamond, Fe bcc, Ni fcc, Co fcc, Mn bcc, fcc, Pd<sub>3</sub>Cr, Pd<sub>3</sub>Mn, PdCr, PdMn
    - Separable square wave potentials

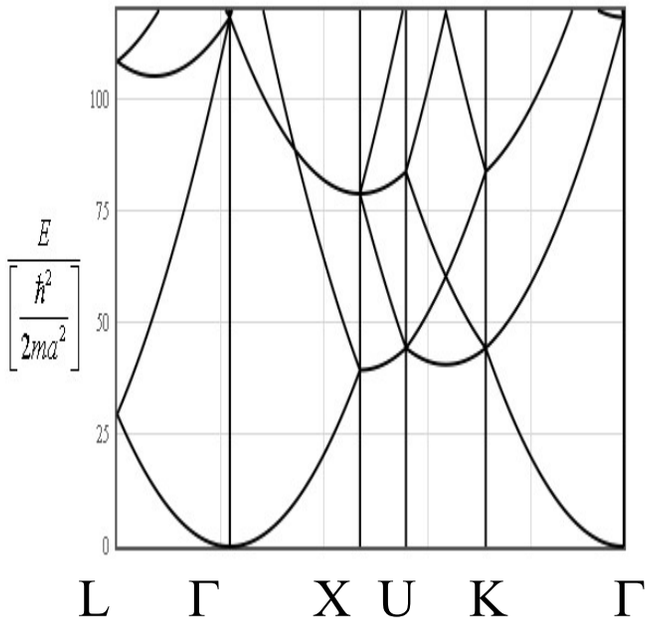
# Metals, semiconductors, and insulators



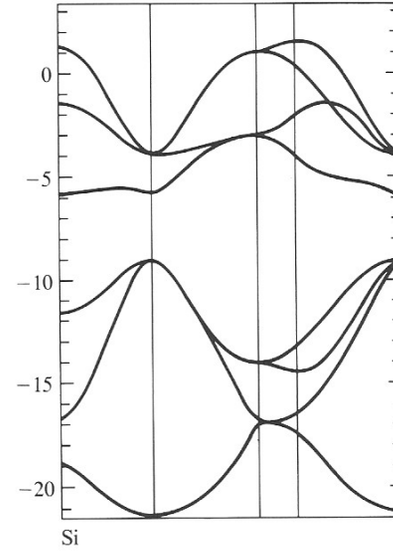
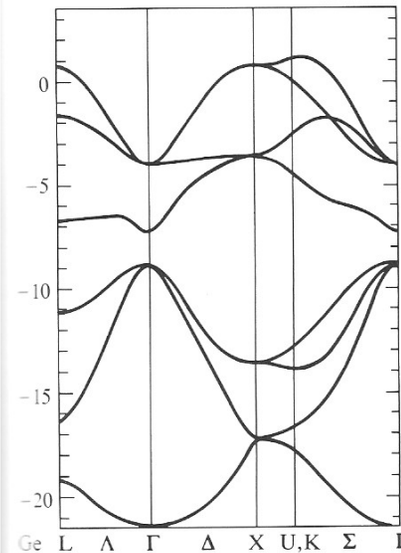
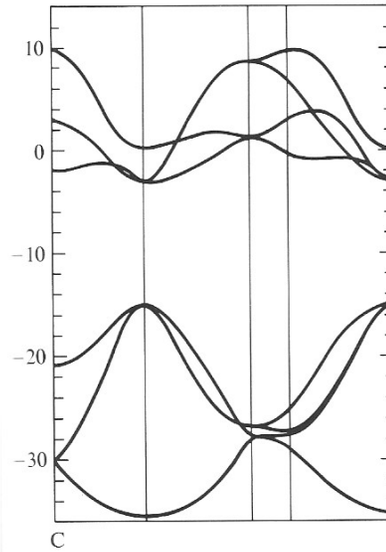
Insulators: band gap  $> 3$  eV

From Ibach & Lueth

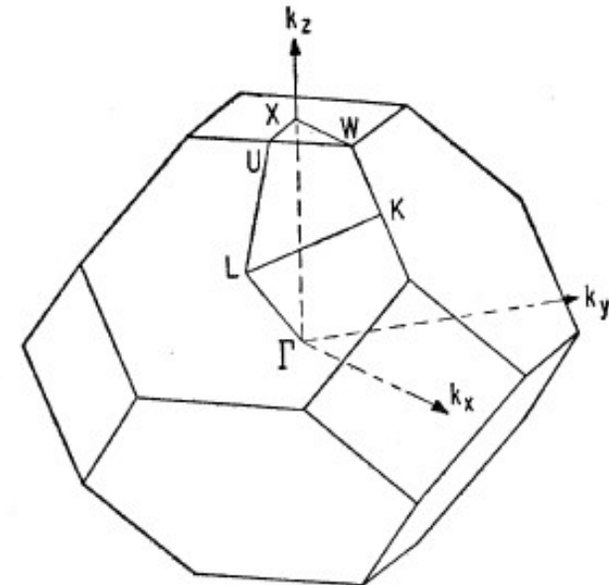
# Group IV



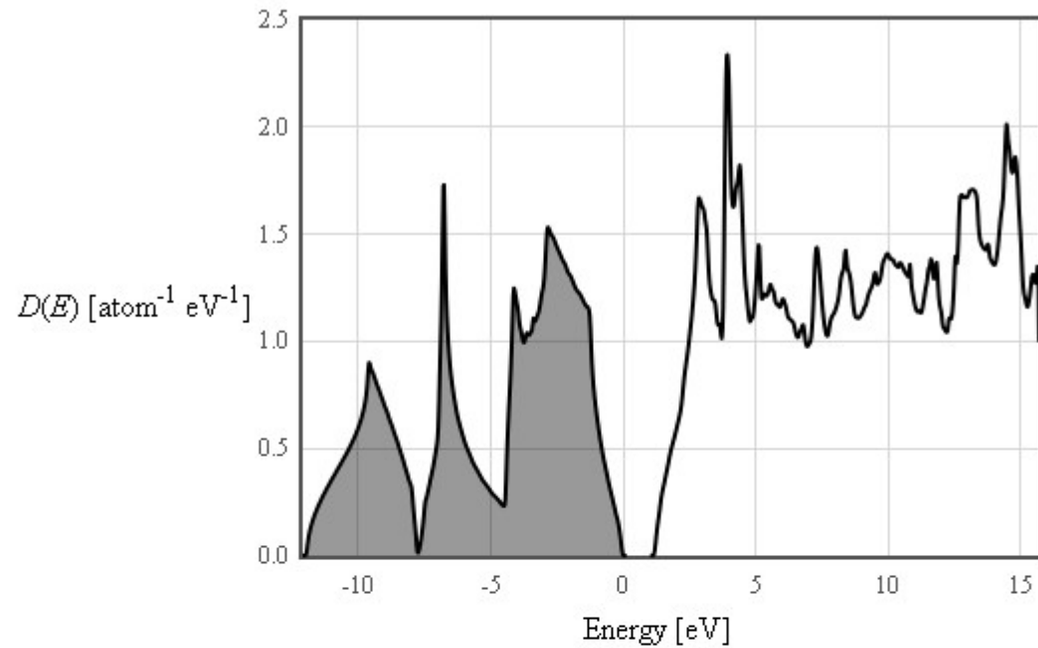
fcc is the Bravais lattice for diamond



41



## Electron density of states for silicon

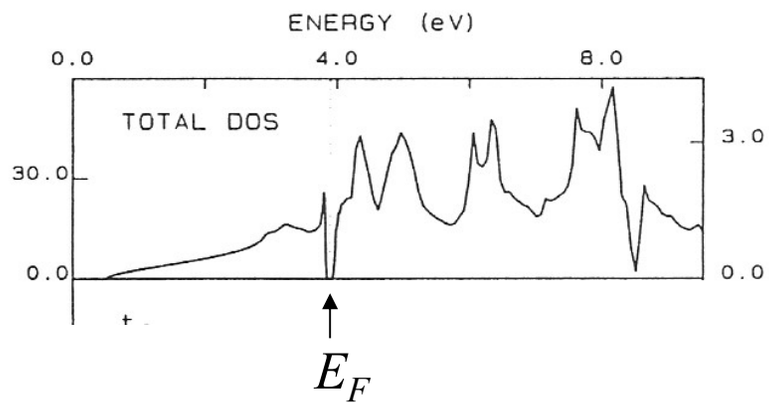
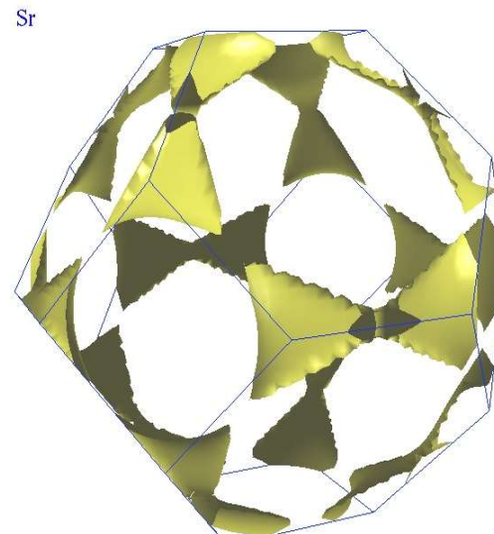
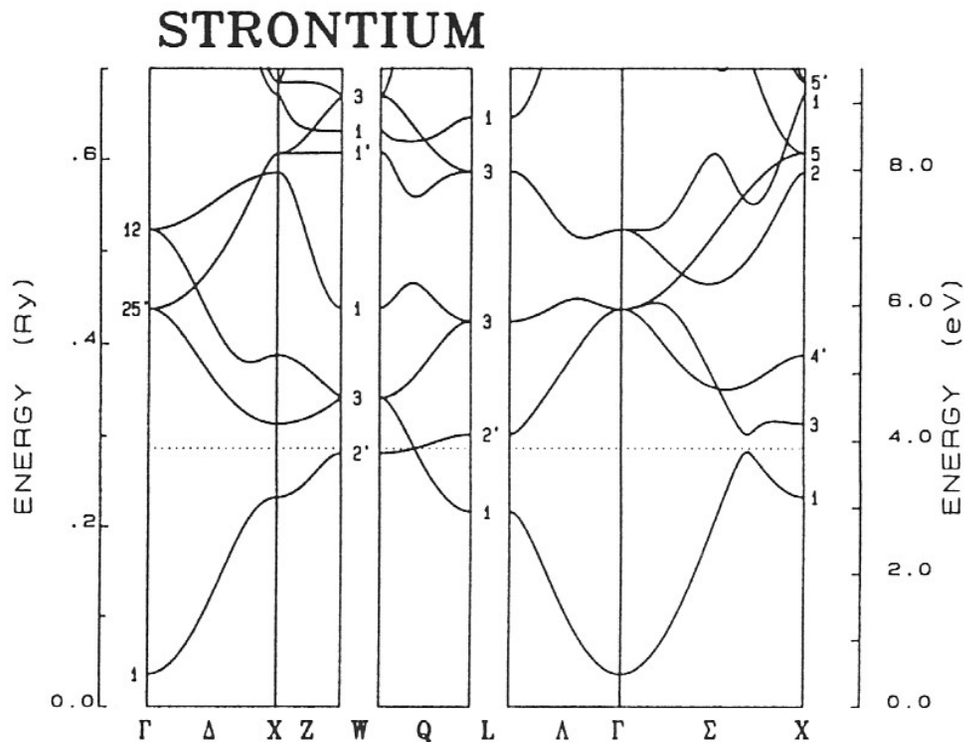


Lattice constant:  $a = 5.43095 \text{ \AA}$ .

$E$ [eV]	$D(E)$ [eV <sup>-1</sup> atom <sup>-1</sup> ]	$D(E)$ [eV <sup>-1</sup> m <sup>-3</sup> ]	$D(E)$ [J <sup>-1</sup> m <sup>-3</sup> ]
-12.10059	0	0	0
-12.07338	0	0	0
-12.04617	0	0	0
-12.01896	0.0001294	6.462444849932946e+24	4.033481993467074e+43
-11.99174	0.00185699	9.274107775793648e+25	5.788358367116245e+44
-11.96453	0.01085698	5.4221510422585004e+26	3.3841911385959934e+45
-11.93732	0.03197818	1.5970419215705466e+27	9.96780627618616e+45
-11.91011	0.05881918	2.937524782598755e+27	1.833432020096589e+46
-11.8829	0.08244563	4.117467828384677e+27	2.5698838025119693e+46

# Semimetal Strontium

00



# kinetic theory

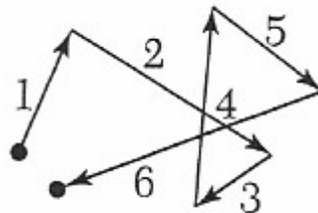
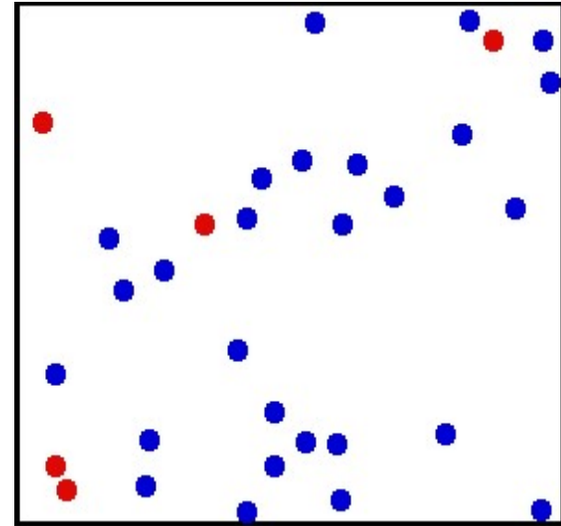
---

describe electrons as a gas of particles

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

The average time between scattering events  $\tau_{sc}$  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  $\mathbf{G}$ .

$$\vec{k}'_{el} = \vec{k}_{el} + \vec{k}_{ph} \quad \longleftarrow \text{phonon emitted}$$

$$\vec{k}'_{el} = \vec{k}_{el} + \vec{k}_{ph} + \vec{G} \quad \longleftarrow \text{Umklapp scattering}$$