Crystal Symmetries and Bindings

Elementary Crystallography

- Solid materials (crystalline, polycrystalline, amorphous)
- Crystallography
- Crystal Lattice
- Crystal Structure
- Types of Lattices
- Unit Cell
- Typical Crystal Structures
 - 14 Bravais Lattices and the Seven Crystal Systems
- Crystal Bindings

Single Crystals

- Single crystals have a periodic atomic structure across its whole volume.
- At long range length scales, each atom is related to every other equivalent atom in the structure by translational or rotational symmetry





Single Crystal

Polycrystalline Solids

- Polycrystalline materials are made up of an aggregate of many small single crystals (also called crystallites or grains).
- Polycrystalline materials have a high degree of order over many atomic or molecular dimensions.
- Grains (domains) are separated by grain boundaries. The atomic order can vary from one domain to the next.
- The grains are usually 100 nm 100 microns in diameter.
- Polycrystals with grains less than 10 nm in diameter are nanocrystalline



Amorphous Solids

- Amorphous (Non-crystalline) Solids are made up of randomly orientated atoms, ions, or molecules that do not form defined patterns or lattice structures.
- Amorphous materials have order only within a few atomic or molecular dimensions.
- Amorphous materials do not have any long-range order, but they have varying degrees of short-range order.
- Examples to amorphous materials include amorphous silicon, plastics, and glasses.
- > Amorphous silicon can be used in solar cells and thin film transistors.



What is a Crystal Lattice

In crystallography, only the geometrical properties of the crystal are of interest, therefore one replaces each atom by a geometrical point located at the equilibrium position of that atom.



Platinum



Platinum surface (scanning tunneling microscope)



Crystal lattice and structure of Platinum

Crystal Lattice

The set of r' defined below for all u_1 , u_2 , and u_3 defines a lattice.

$$\mathbf{r}' = \mathbf{r} + u_1 \mathbf{a}_1 + u_2 \mathbf{a}_2 + u_3 \mathbf{a}_3$$

- An infinite array of points in space.
- Each point has identical surroundings to all others.
- Arrays are arranged in a periodic manner.



Crystal Structure

Crystal structures can be obtained by attaching atoms, groups of atoms or molecules which are called basis (motif) to the lattice sides of the lattice point.



Basis composed of atoms at the positions $\mathbf{r}_j = x_j \mathbf{a}_1 + y_j \mathbf{a}_2 + z_j \mathbf{a}_3$, with *j* = 1,2,...,*s*. Here *x*, *y*, *z* may be selected to have values between 0 and 1. ₇

A two-dimensional Bravais lattice with different choices for the basis



Primitive Cell in 2D

The smallest component of the crystal (group of atoms, ions or molecules), which when stacked together with pure translational repetition reproduces the whole crystal.



The parallelograms 1, 2, and 3 are primitive cells equal in area but the parallelogram 4 is not.

Primitive Cell in 3D

A primitive cell contains only one lattice point and the volume of the parallelepiped is

$$V_c = |\mathbf{a}_1 \cdot \mathbf{a}_2 \times \mathbf{a}_3|$$



x



Primitive Cell

• The primitive cell and, consequently, the entire lattice, is *uniquely* determined by the six lattice constants: a, b, c, α , β and γ .

- Only 1/8 of each lattice point in a unit cell can actually be assigned to that cell.
- Each primitive cell in the figure can be associated with 8 x 1/8 = 1 lattice point.

Five Bravais Lattices in 2D



Typical Crystal Structures

3D – 14 BRAVAIS LATTICES AND SEVEN CRYSTAL TYPES

- Cubic Crystal System (SC, BCC, FCC)
- Hexagonal Crystal System (S)
- Triclinic Crystal System (S)
- Monoclinic Crystal System (S, Base-C)
- Orthorhombic Crystal System (S, Base-C, BC, FC)
- Tetragonal Crystal System (S, BC)
- Trigonal (Rhombohedral) Crystal System (S)





Characteristics of cubic lattices^{*a*}

	Simple	Body-centered	Face-centered
Volume, conventional cell	a^3	a^3	a^3
Lattice points per cell	1	2	4
Volume, primitive cell	a^3	$\frac{1}{2}a^{3}$	$\frac{1}{4}a^3$
Lattice points per unit volume	$1/a^{3}$	$\frac{2}{2}/a^{3}$	$\frac{4}{4}a^{3}$
Number of nearest neighbors	6	8	12
Nearest-neighbor distance	a	$3^{1/2} a/2 = 0.866a$	$a/2^{1/2} = 0.707a$
Number of second neighbors	12	6	6
Second neighbor distance	$2^{1/2}a$	a	a
Packing fraction ^{<i>a</i>}	$rac{1}{6}\pi$	$\frac{1}{8}\pi\sqrt{3}$	$\frac{1}{6}\pi\sqrt{2}$
	=0.524	=0.680	=0.740

^{*a*} The packing fraction is the maximum proportion of the available volume that can be filled with hard spheres.





Index System for Crystal Planes



- Find the intercepts on the axes in terms of the lattice constants
 a1, a2, a3. The axes may be those of a primitive or nonprimitive cell.
- Take the reciprocals of these numbers and then reduce to three integers having the same ratio, usually the smallest three integers. The result, enclosed in parentheses (hkl), is called the index of the plane.

Figure 14 Indices of important planes in a cubic crystal The plane (200) is parallel to (100) and to (100)

Lattice Point Group

By lattice point group we mean the collection of symmetry operations which, applied about a lattice point, carry the lattice into itself.

Crystal lattices can be carried or mapped into themselves by the lattice translations T and by various other symmetry operations. A typical symmetry operation is that of rotation about an axis that passes through a lattice point.

Lattices can be found with rotational symmetry corresponding to rotations by 2π , $2\pi/2$, $2\pi/3$, $2\pi/4$, and $2\pi/6$ radians and by integral multiples of these rotations. The rotation axes are denoted by the symbols 1, 2, 3, 4, and 6.

We can have mirror reflections m about a plane through a lattice point. The inversion operation is composed of a rotation of π followed by reflection in a plane normal to the rotation axis; the total effect is to replace **r** by -**r**.

Symmetry Planes and Axes





Crystal Bindings

Forces hold the atoms together in a solid

- Covalent bonding
- Ionic bonding
- Metallic bonding
- Van der waals bonding
- Hydrogen bonding

All bonding is a consequence of the electrostatic interaction between the nuclei and electrons.

Covalent Bonding

- Covalent bonding takes place between atoms with small differences in electronegativity which are close to each other in the periodic table (*between non-metals and non-metals*).
- The covalent bonding is formed when the atoms share the outer shell electrons (i.e., s and p electrons) rather than by electron transfer.
- > Noble gas electron configuration can be attained.



H_2^+ ion





Each electron in a shared pair is attracted to both nuclei involved in the bond. The approach, electron overlap, and attraction can be visualized as shown in the following figure representing the nuclei and electrons in a hydrogen molecule.



Diamond Structure



- The diamond structure is an example of the directional covalent bonding found in the group IV elements, such as C, Si, Ge.
- ➤ The structure is of the fcc lattice, consisting of the basis of two identical elements (000) and (1/4, 1/4, 1/4).
- Each atom has four nearest neighbours and 12 next nearest neighbours.

Ionic Bonding

- Ionic bonding is the electrostatic force of attraction between positively and negatively charged ions (between non-metals and metals).
- All ionic compounds are crystalline solids at room temperature.
- > NaCl is a typical example of ionic bonding.



Energies of Interactions Between Atoms

The energy of the crystal is lower than that of the free atoms by an amount equal to the energy required to pull the crystal apart into a set of free atoms. This is called the binding (cohesive) energy of the crystal.



Madelung Energy

The main contribution to the binding energy of ionic crystals is electrostatic and is called the **Madelung energy**.

If U_{ij} is the interaction energy between ions *i* and *j*, we define a sum U_i which includes all interactions involving the ion *i*:

$$U_i = \sum_j' U_{ij} , \qquad U_{ij} = \lambda \, \exp(-r_{ij}/\rho) \pm q^2/r_{ij}$$

If we let $r_{ij} \equiv p_{ij}R$, where *R* is the nearest-neighbor separation in the crystal, then

$$U_{ij} = \begin{cases} \lambda \exp(-R/p) - \frac{q^2}{R} & \text{(nearest neighbors)} \\ \pm \frac{1}{p_{ij}} \frac{q^2}{R} & \text{(otherwise).} \end{cases}$$

$$U_{\rm tot} = N U_i = N \left(z \lambda e^{-R/\rho} - \frac{\alpha q^2}{R} \right) \; , \label{eq:Utot}$$

where z is the number of nearest neighbors of any ion and

$$\alpha \equiv \sum_{j}' \frac{(\pm)}{p_{ij}} \equiv$$
 Madelung constant

Sodium Chloride Structure

- Sodium chloride also crystallizes in a cubic lattice, but with a different unit cell.
- Sodium chloride structure consists of equal numbers of sodium and chlorine ions placed at alternate points of a fcc cubic lattice.
- Each ion has six of the other kind of ions as its nearest neighbours.



Metallic elements have only up to the valence electrons in their outer shell.

When losing their electrons they become positive ions.

Electronegative elements tend to acquire additional electrons to become negative ions or anions.

When the Na⁺ and Cl⁻ ions approach each other closely enough so that the orbits of the electron in the ions begin to overlap with each other, then the electron begins to repel each other by virtue of the repulsive electrostatic coulomb force. Of course the closer together the ions are, the greater the repulsive force.

Pauli exclusion principle has an important role in repulsive force. To prevent a violation of the exclusion principle, the potential energy of the system increases very rapidly.

Cesium Chloride Structure

- Cesium chloride also crystallizes in a cubic lattice, but with a different unit cell.
- Cesium chloride structure consists of equal numbers of cesium and chlorine ions placed at alternate points of a simple cubic lattice with Ce⁺ ion at (000) and Cl⁻ at (½, ½, ½).
- Each ion has eight of the other kind of ions as its nearest neighbours.



Figure 18 The cesium chloride crystal structure. The space lattice is simple cubic, and the basis has one Cs^+ ion at 000 and one Cl^- ion at $\frac{1}{2} \frac{1}{2} \frac{1}{2}$.

Comparison of Ionic and Covalent Bonding



Metallic Bonding

- Metallic bonding is found in metal elements. This is the electrostatic force of attraction between positively charged ions and delocalized outer electrons.
- The metallic bond is weaker than the ionic and the covalent bonds.
- A metal may be described as a lowdensity cloud of free electrons.
- Therefore, metals have high electrical and thermal conductivity.





Van der Waals Bonding

- These are weak bonds with a typical strength of 0.2 eV/atom.
- Van Der Waals bonds occur between neutral atoms and molecules.
- Weak forces of attraction result from the natural fluctuations in the electron density of all molecules that cause small temporary dipoles to appear within the molecules.
- It is these temporary dipoles that attract one molecule to another. They are called van der Waals' forces.

- The shape of a molecule influences its ability to form temporary dipoles. Long thin molecules can pack closer to each other than molecules that are more spherical. The bigger the 'surface area' of a molecule, the greater the van der Waal's forces will be and the higher the melting and boiling points of the compound will be.
- ➢ Van der Waal's forces are of the order of 1% of the strength of a covalent bond.



Homonuclear molecules, such as iodine, develop temporary dipoles due to natural fluctuations of electron density within the molecule



Heteronuclear molecules, such as H-Cl have permanent dipoles that attract the opposite pole in other molecules. These forces are due to the electrostatic attraction between the nucleus of one atom and the electrons of the other.



 Van der waals interaction occurs generally between atoms which have noble gas configuration.



Hydrogen Bonding

- A hydrogen atom, having one electron, can be covalently bonded to only one atom. However, the hydrogen atom can involve itself in an additional electrostatic bond with a second atom of highly electronegative character such as fluorine or oxygen. This second bond permits a *hydrogen bond* between two atoms or strucures.
- The strength of hydrogen bonding varies from 0.1 to 0.5 ev/atom.

Hydrogen bonds connect water molecules in ordinary ice. Hydrogen bonding is also very important in proteins and nucleic acids and therefore in life processes.

In the extreme ionic form of the hydrogen bond, the hydrogen atom loses its electron to another atom in the molecule; the bare proton forms the hydrogen bond.



Properties of Different Bonding

Types of Bonding	Ionic	Covalent	Metallic	Van der Waals	Hydrogen
Melting Point	High	Very High	Variable	Low	Low
Hardness	Hard and Brittle	Very hard	Variable	Soft and Brittle	Soft and Brittle
Conductivity	Low	Low	High	Non- conducting	Non- conducting
Examples	NaCl, ZnS	Diamond, Graphite	Au, Cu, Fe	Ne, Ar, Kr	Ice, Organic solids

Exercises

- 1. Draw the primitive unit cell of the fcc lattice and determine the lengths of the primitive lattice vectors a', b', c' (in units of the conventional lattice constant a) and also the angles a', b', c' between the primitive lattice vectors. (Hint: Express the primitive lattice vectors as a linear combination of the lattice vectors a, b, c of the body-centered cubic lattice and use elementary vector algebra.) What distinguishes this unit cell from that of the rhombic Bravais lattice?
- 2. Supposing the atoms to be rigid spheres, what fraction of space is filled by atoms in the primitive cubic, fcc, hcp, bcc, and diamond lattices?
- 3. Take the CsCl structure and assume that the cation radius becomes smaller while the anion radius remains constant. What happens to the binding energy? Show that for small cations the NaCl lattice becomes the preferred structure. For an even smaller cation radius the ZnS lattice has the largest binding energy (Madelung constant A=1.638).