### **Chapter Outline**

### How do atoms arrange themselves to form solids?

- Fundamental concepts and language
- Unit cells
- Crystal structures
  - > Face-centered cubic
  - > Body-centered cubic
  - Hexagonal close-packed
- Close packed crystal structures
- Density computations
- Types of solids

Single crystal

**Polycrystalline** 

Amorphous

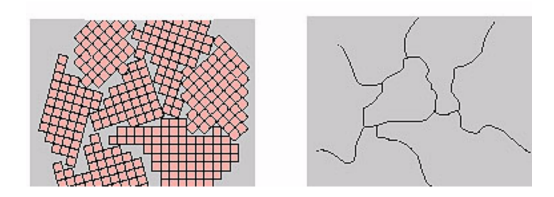
- 3.8–3.10 Crystallography Not Covered / Not Tested
- 3.15 Anisotropy Not Covered / Not Tested
- 3.16 Diffraction Not Covered / Not Tested

### **Types of Solids**

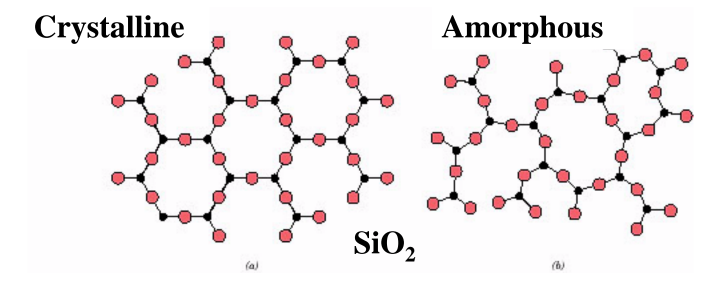
Crystalline material: atoms self-organize in a periodic array

**Single crystal**: atoms are in a repeating or periodic array over the entire extent of the material

Polycrystalline material: comprised of many small crystals or grains

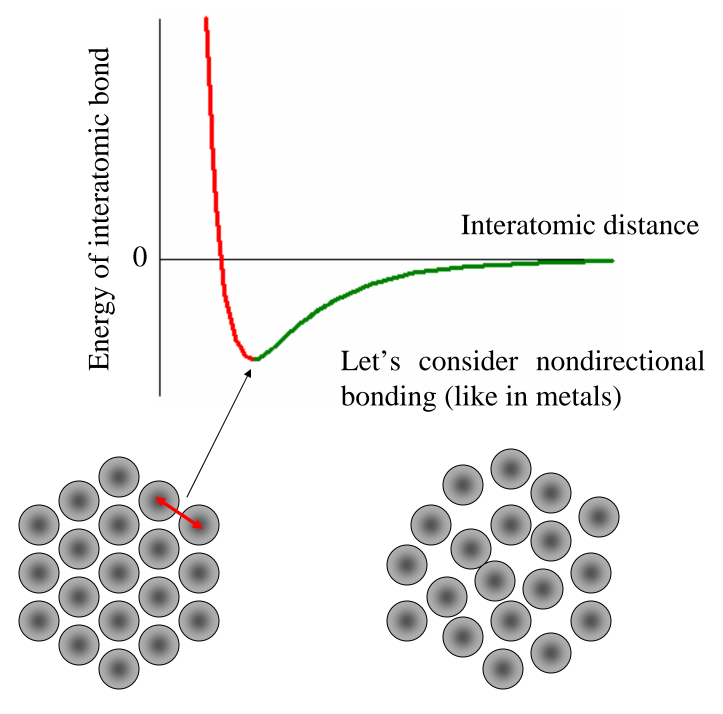


Amorphous: disordered – lack of a systematic atomic arrangement



### **Crystal structures**

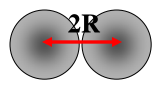
Why do atoms assemble into ordered structures (crystals)?



Energy of the crystal < Energy of the amorphous solid

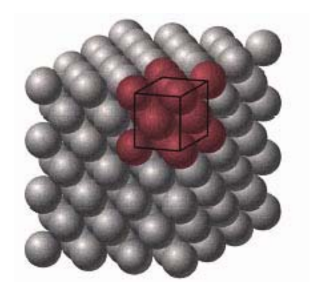
### **Crystal structure**

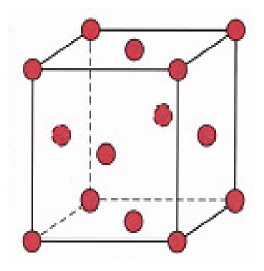
To discuss crystalline structures it is useful to consider atoms as being hard spheres with well-defined radii. In this hard-sphere model, the shortest distance between two like atoms is one diameter of the hard sphere.



- hard-sphere model

We can also consider crystalline structure as a lattice of points at atom/sphere centers.

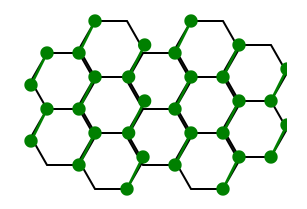




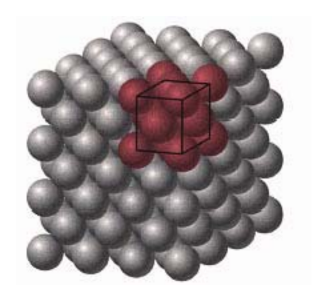
#### **Unit Cell**

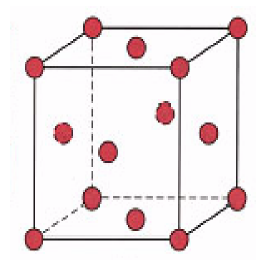
The unit cell is a structural unit or building block that can describe the crystal structure. Repetition of the unit cell generates the entire crystal.

Example: 2D honeycomb net can be represented by translation of two adjacent atoms that form a unit cell for this 2D crystalline structure



Example of 3D crystalline structure:





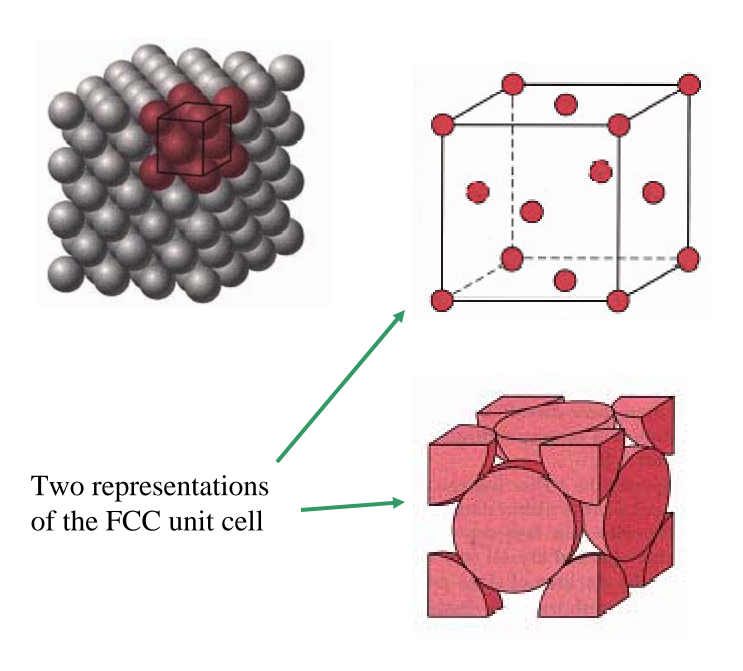
Different choices of unit cells possible, we will consider parallelepiped unit cell with highest level of symmetry

### **Metallic Crystal Structures**

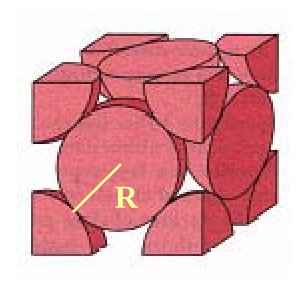
- ➤ Metals are usually (poly)crystalline; although formation of amorphous metals is possible by rapid cooling
- ➤ As we learned in Chapter 2, the atomic bonding in metals is non-directional ⇒ no restriction on numbers or positions of nearest-neighbor atoms ⇒ large number of nearest neighbors and dense atomic packing
- ➤ Atomic (hard sphere) radius, R, defined by ion core radius typically 0.1 0.2 nm
- > The most common types of unit cells are
  - faced-centered cubic (FCC)
  - body-centered cubic (BCC)
  - hexagonal close-packed (HCP).

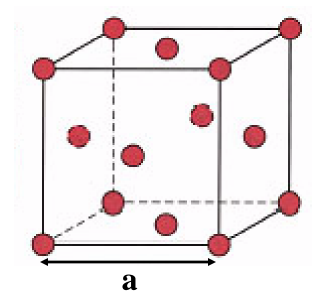
### Face-Centered Cubic (FCC) Crystal Structure (I)

- Atoms are located at each of the corners and on the centers of all the faces of cubic unit cell
- Cu, Al, Ag, Au have this crystal structure



### **Face-Centered Cubic Crystal Structure (II)**

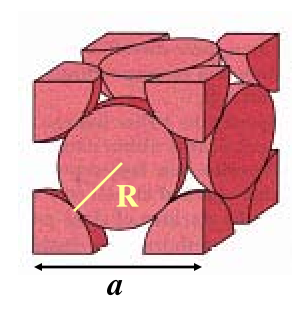




- The hard spheres touch one another across a face diagonal  $\Rightarrow$  the cube edge length,  $a = 2R\sqrt{2}$
- ➤ The coordination number, CN = the number of closest neighbors to which an atom is bonded = number of touching atoms, CN = 12
- Number of atoms per unit cell, n = 4. (For an atom that is shared with m adjacent unit cells, we only count a fraction of the atom, 1/m). In FCC unit cell we have:
  - 6 face atoms shared by two cells:  $6 \times 1/2 = 3$
  - 8 corner atoms shared by eight cells:  $8 \times 1/8 = 1$
- ➤ Atomic packing factor, APF = fraction of volume occupied by hard spheres = (Sum of atomic volumes)/(Volume of cell) = 0.74 (maximum possible)

### **Face-Centered Cubic Crystal Structure (III)**

Let's calculate the atomic packing factor for FCC crystal



$$a = 2R\sqrt{2}$$

**APF** = (Sum of atomic volumes)/(Volume of unit cell)

Volume of 4 hard spheres in the unit cell:  $4 \times \frac{4}{3} \pi R^3$ 

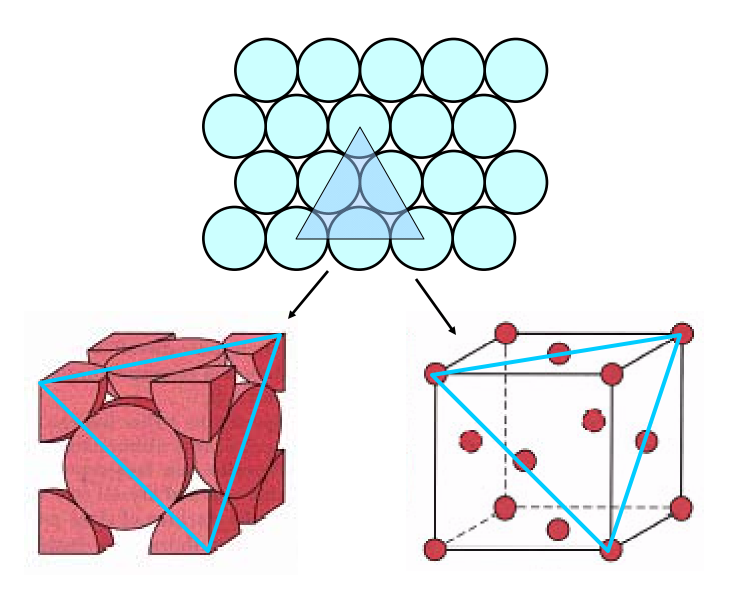
Volume of the unit cell:  $a^3 = 16R^3\sqrt{2}$ 

$$APF = \frac{16}{3} \pi R^3 / 16R^3 \sqrt{2} = \pi / 3\sqrt{2} = 0.74$$

maximum possible packing of hard spheres

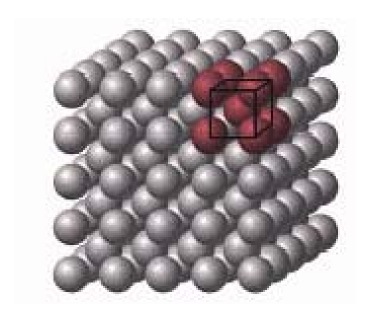
### **Face-Centered Cubic Crystal Structure (IV)**

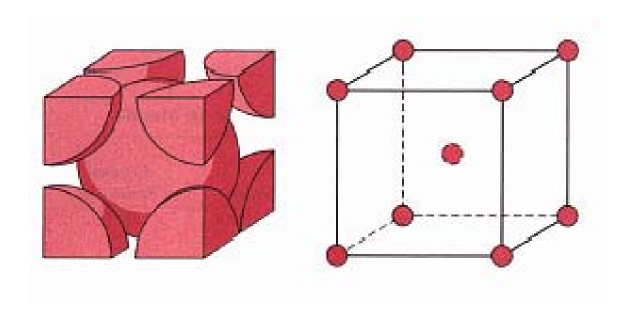
- Corner and face atoms in the unit cell are equivalent
- ➤ FCC crystal has APF of 0.74, the maximum packing for a system equal-sized spheres ⇒ FCC is a **close-packed** structure
- FCC can be represented by a stack of **close-packed** planes (planes with highest density of atoms)



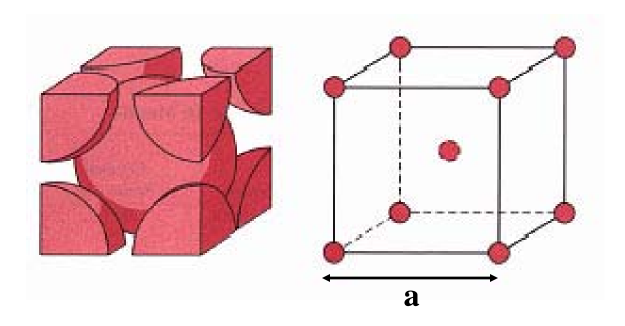
# **Body-Centered Cubic (BCC) Crystal Structure (I)**

Atom at each corner and at center of cubic unit cell Cr,  $\alpha$ -Fe, Mo have this crystal structure





#### **Body-Centered Cubic Crystal Structure (II)**



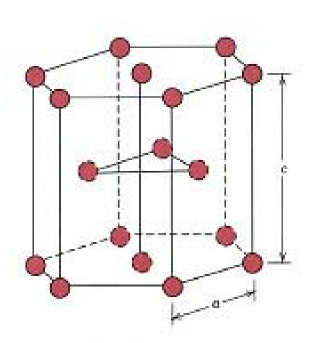
- The hard spheres touch one another along cube diagonal  $\Rightarrow$  the cube edge length,  $a=4R/\sqrt{3}$
- $\triangleright$  The coordination number, CN = 8
- Number of atoms per unit cell, n = 2

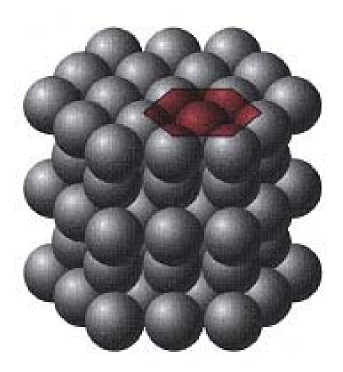
  Center atom (1) shared by no other cells: 1 x 1 = 1

  8 corner atoms shared by eight cells: 8 x 1/8 = 1
- $\triangleright$  Atomic packing factor, APF = 0.68
- Corner and center atoms are equivalent

### Hexagonal Close-Packed Crystal Structure (I)

- ➤ HCP is one more common structure of metallic crystals
- ➤ Six atoms form regular hexagon, surrounding one atom in center. Another plane is situated halfway up unit cell (c-axis), with 3 additional atoms situated at interstices of hexagonal (close-packed) planes
- > Cd, Mg, Zn, Ti have this crystal structure





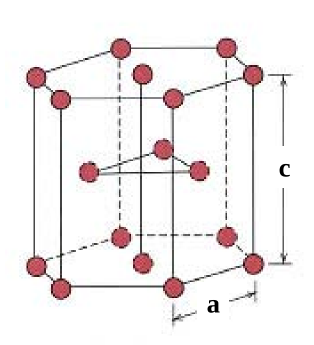
### Hexagonal Close-Packed Crystal Structure (II)

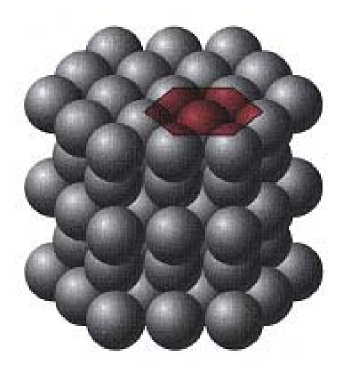
- Unit cell has two lattice parameters a and c. Ideal ratio c/a = 1.633
- $\triangleright$  The coordination number, CN = 12 (same as in FCC)
- Number of atoms per unit cell, n = 6.

  3 mid-plane atoms shared by no other cells: 3 x 1 = 3

  12 hexagonal corner atoms shared by 6 cells: 12 x 1/6 = 2

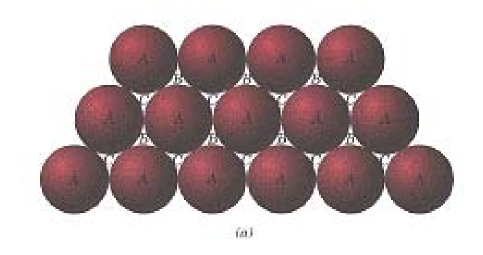
  2 top/bottom plane center atoms shared by 2 cells: 2 x 1/2 = 1
- $\triangleright$  Atomic packing factor, APF = 0.74 (same as in FCC)
- > All atoms are equivalent





# **Close-packed Structures (FCC and HCP)**

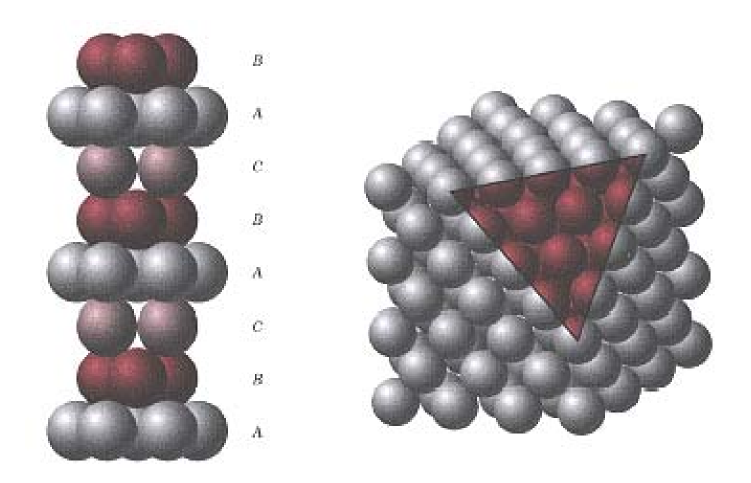
- ➤ Both FCC and HCP crystal structures have atomic packing factors of 0.74 (maximum possible value)
- ➤ Both FCC and HCP crystal structures may be generated by the stacking of close-packed planes
- ➤ The difference between the two structures is in the stacking sequence





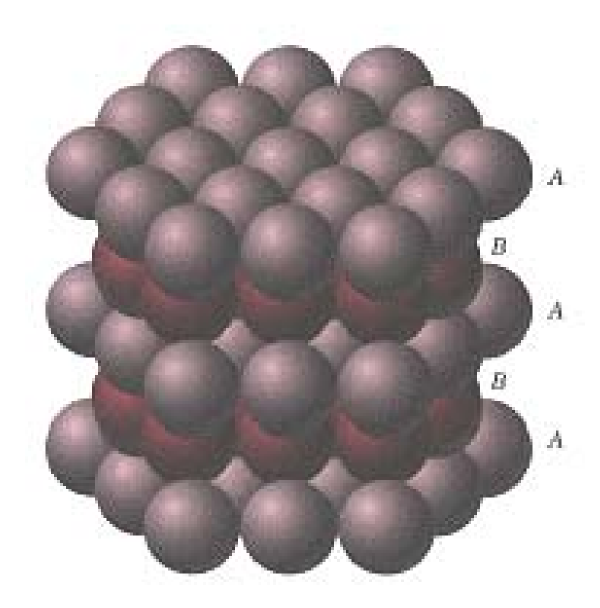
HCP: ABABAB... FCC: ABCABCABC...

FCC: Stacking Sequence ABCABCABC...



Third plane is placed above the "holes" of the first plane not covered by the second plane

# **HCP:** Stacking Sequence ABABAB...



Third plane is placed directly above the first plane of atoms

### **Density Computations**

Since the entire crystal can be generated by the repetition of the unit cell, the density of a crystalline material,  $\rho$  = the density of the unit cell = (atoms in the unit cell, n) × (mass of an atom, M) / (the volume of the cell,  $V_c$ )

Atoms in the unit cell, n = 2 (BCC); 4 (FCC); 6 (HCP)

Mass of an atom, M = Atomic weight, A, in amu (or g/mol) is given in the periodic table. To translate mass from amu to grams we have to divide the atomic weight in amu by the Avogadro number  $N_A = 6.023 \times 10^{23}$  atoms/mol

The volume of the cell,  $V_c = a^3$  (FCC and BCC)  $a = 2R\sqrt{2}$  (FCC);  $a = 4R/\sqrt{3}$  (BCC) where R is the atomic radius

Thus, the formula for the density is:

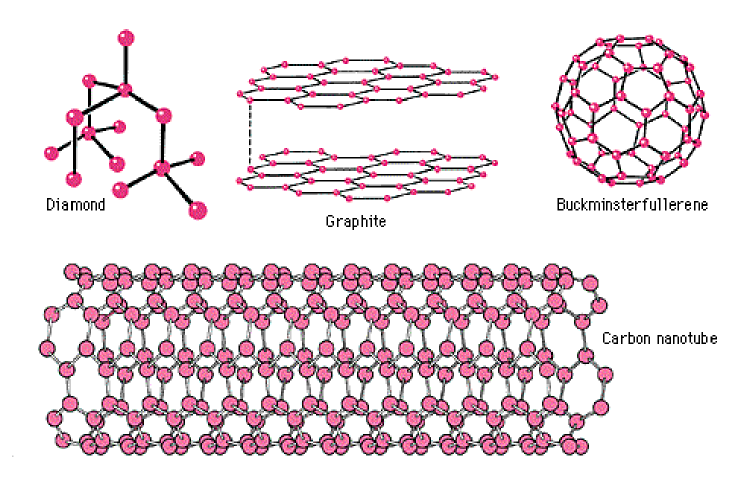
$$\rho = \frac{nA}{V_c N_A}$$

Atomic weight and atomic radius of many elements you can find in the table at the back of the textbook front cover.

#### **Polymorphism and Allotropy**

Some materials may exist in more than one crystal structure, this is called **polymorphism**. If the material is an elemental solid, it is called **allotropy**.

An example of allotropy is carbon, which can exist as diamond, graphite, and amorphous carbon.

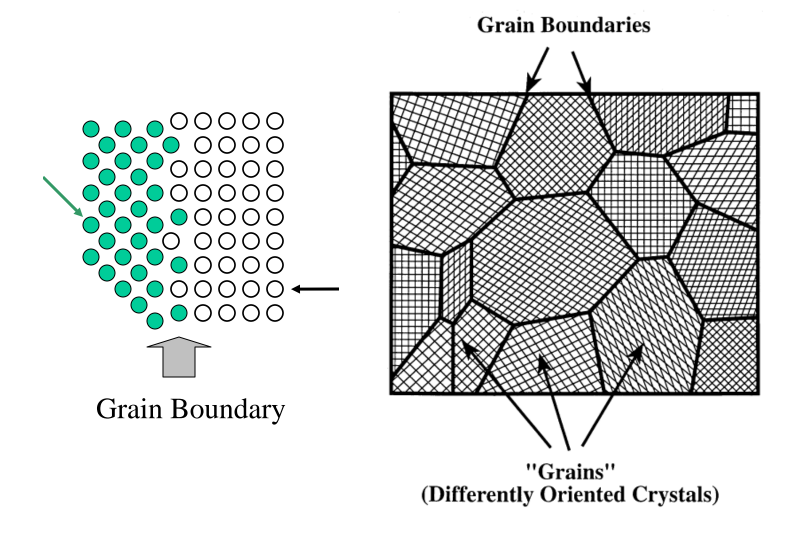


Pure, solid carbon occurs in three crystalline forms – diamond, graphite; and large, hollow fullerenes. Two kinds of fullerenes are shown here: buckminsterfullerene (buckyball) and carbon nanotube.

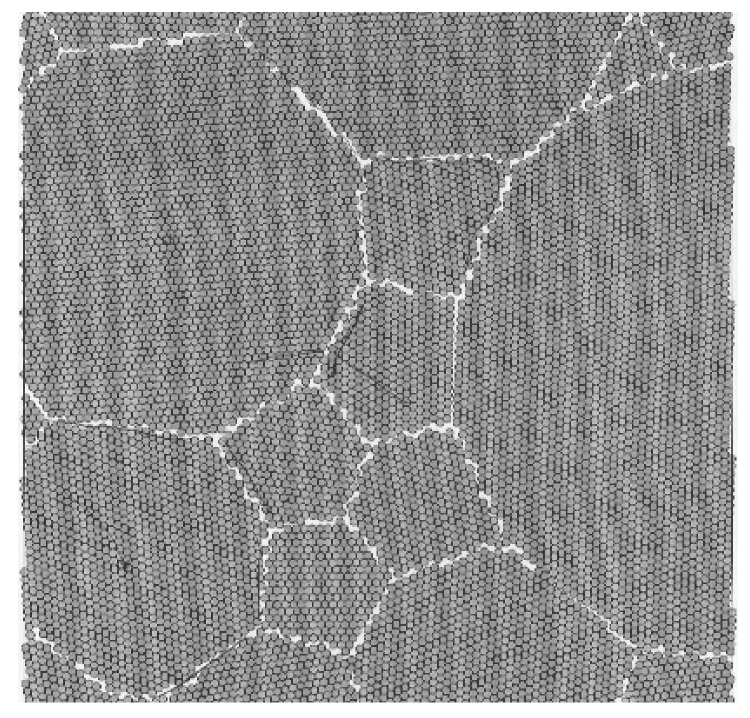
### **Single Crystals and Polycrystalline Materials**

**Single crystal**: atoms are in a repeating or periodic array over the entire extent of the material

Polycrystalline material: comprised of many small crystals or grains. The grains have different crystallographic orientation. There exist atomic mismatch within the regions where grains meet. These regions are called grain boundaries.

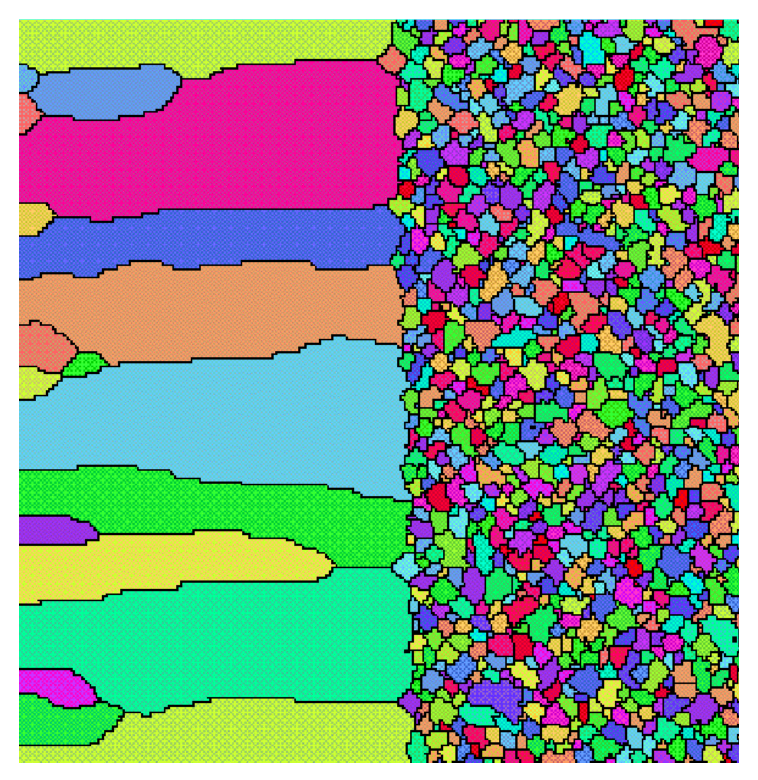


# **Polycrystalline Materials**



Atomistic model of a nanocrystalline solid by Mo Li, JHU

# **Polycrystalline Materials**



Simulation of annealing of a polycrystalline grain structure from http://cmpweb.ameslab.gov/cmsn/microevolproj.html (link is dead)

### **Anisotropy**

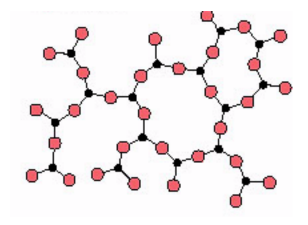
Different directions in a crystal have different packing. For instance, atoms along the edge of FCC unit cell are more separated than along the face diagonal. This causes **anisotropy** in the properties of crystals, for instance, the deformation depends on the direction in which a stress is applied.

In some polycrystalline materials, grain orientations are random, so bulk material properties are **isotropic** 

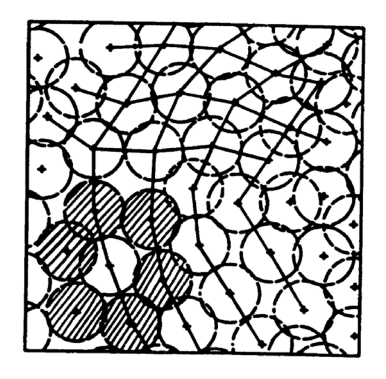
Some polycrystalline materials have grains with preferred orientations (texture), so properties are dominated by those relevant to the texture orientation and the material exhibits anisotropic properties

# **Non-Crystalline (Amorphous) Solids**

In amorphous solids, there is no long-range order. But amorphous does not mean random, in many cases there is some form of short-range order.



Schematic picture of amorphous SiO<sub>2</sub> structure



Amorphous structure from simulations by E. H. Brandt

#### **Summary**

Make sure you understand language and concepts:

- > Allotropy
- > Amorphous
- > Anisotropy
- ➤ Atomic packing factor (APF)
- ➤ Body-centered cubic (BCC)
- Coordination number
- Crystal structure
- > Crystalline
- > Face-centered cubic (FCC)
- > Grain
- Grain boundary
- Hexagonal close-packed (HCP)
- > Isotropic
- > Lattice parameter
- Non-crystalline
- > Polycrystalline
- Polymorphism
- Single crystal
- Unit cell

Homework #1: 2.14, 2.15, 2.20, 3.7, and 3.17

Due date: Monday, September 6.

### Reading for next class:

### **Chapter 4: Imperfections in Solids**

**Point defects** (vacancies, interstitials)

**Dislocations** (edge, screw)

**Grain boundaries** (tilt, twist)

Weight and atomic composition

Optional reading (Parts that are not covered / not tested):

4.9 – 4.10 *Microscopy* 

4.11 Grain size determination