The Structure of Crystalline Solids

ISSUES TO ADDRESS...

- How do atoms assemble into solid structures? (for now, focus on metals)
- How does the density of a material depend on its structure?
- When do material properties vary with the sample (i.e., part) orientation?



Energy and Packing



Dense, ordered packed structures tend to have lower energies.



Materials and Packing

Crystalline materials...

- atoms pack in periodic, 3D arrays
- typical of: -metals

-many ceramics -some polymers

Noncrystalline materials...

- atoms have no periodic packing
- occurs for: -complex structures
 -rapid cooling

"Amorphous" = Noncrystalline



Section 3.3 – Crystal Systems

Unit cell: smallest repetitive volume which contains the complete lattice pattern of a crystal.



- 7 crystal systems
- 14 crystal lattices

a, b, and c are the lattice constants



Fig. 3.4, Callister 7e.

Section 3.4 – Metallic Crystal Structures

 How can we stack metal atoms to minimize empty space?

2-dimensions



Now stack these 2-D layers to make 3-D structures



Metallic Crystal Structures

- Tend to be densely packed.
- Reasons for dense packing:
 - Typically, only one element is present, so all atomic radii are the same.
 - Metallic bonding is not directional.
 - Nearest neighbor distances tend to be small in order to lower bond energy.
 - Electron cloud shields cores from each other
- Have the simplest crystal structures.

We will examine three such structures...



Simple Cubic Structure (SC)

- Rare due to low packing denisty (only Po has this structure)
- Close-packed directions are cube edges.



 Coordination # = 6 (# nearest neighbors)





• APF for a simple cubic structure = 0.52



Body Centered Cubic Structure (BCC)

- Atoms touch each other along cube diagonals.
 - --Note: All atoms are identical; the center atom is shaded differently only for ease of viewing.

ex: Cr, W, Fe (α), Tantalum, Molybdenum

• Coordination # = 8



Callister 7e.

2 atoms/unit cell: 1 center + 8 corners x 1/8



Atomic Packing Factor: BCC

• APF for a body-centered cubic structure = 0.68



Face Centered Cubic Structure (FCC)

• Atoms touch each other along face diagonals.

--Note: All atoms are identical; the face-centered atoms are shaded differently only for ease of viewing.

ex: Al, Cu, Au, Pb, Ni, Pt, Ag

• Coordination # = 12



4 atoms/unit cell: 6 face x 1/2 + 8 corners x 1/8

Atomic Packing Factor: FCC

• APF for a face-centered cubic structure = 0.74



FCC Stacking Sequence

- ABCABC... Stacking Sequence
- 2D Projection

A sites B sites

C sites



• FCC Unit Cell



Hexagonal Close-Packed Structure (HCP)

- ABAB... Stacking Sequence
- 3D Projection



2D Projection
 Top layer
 Middle layer
 Bottom layer

- Coordination # = 12
- APF = 0.74
- *c*/*a* = 1.633

6 atoms/unit cell

ex: Cd, Mg, Ti, Zn



Theoretical Density, p

Density =
$$\rho$$
 = $\frac{\text{Mass of Atoms in Unit Cell}}{\text{Total Volume of Unit Cell}}$

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

where
$$n =$$
 number of atoms/unit cell
 $A =$ atomic weight
 $V_C =$ Volume of unit cell = a^3 for cubic
 $N_A =$ Avogadro's number
= 6.023 x 10²³ atoms/mol





Densities of Material Classes



Crystals as Building Blocks

- Some engineering applications require single crystals: --diamond single
 --turbine blades
 - crystals for abrasives



(Courtesy Martin Deakins, GE Superabrasives, Worthington, OH. Used with permission.) Fig. 8.33(c), *Callister 7e.* (Fig. 8.33(c) courtesy of Pratt and Whitney).

• Properties of crystalline materials often related to crystal structure.

--Ex: Quartz fractures more easily along some crystal planes than others.





Polycrystals

Anisotropic

• Most engineering materials are polycrystals.



Adapted from Fig. K, color inset pages of *Callister 5e*. (Fig. K is courtesy of Paul E. Danielson, Teledyne Wah Chang Albany)

- Nb-Hf-W plate with an electron beam weld.
- Each "grain" is a single crystal.
- If grains are randomly oriented, overall component properties are not directional.
- Grain sizes typ. range from 1 nm to 2 cm (i.e., from a few to millions of atomic layers).

Isotropic

Single vs Polycrystals

- Single Crystals
 - -Properties vary with direction: anisotropic.
 - -Example: the modulus of elasticity (E) in BCC iron:
- Polycrystals
 - -Properties may/may not vary with direction.
 - -If grains are randomly oriented: isotropic. (E_{poly iron} = 210 GPa)
 - -If grains are textured, anisotropic.

E (diagonal) = 273 GPa



Data from Table 3.3, *Callister 7e*. (Source of data is R.W. Hertzberg, *Deformation and Fracture Mechanics of Engineering Materials*, 3rd ed., John Wiley and Sons, 1989.)



Adapted from Fig. 4.14(b), *Callister 7e*. (Fig. 4.14(b) is courtesy of L.C. Smith and C. Brady, the National Bureau of Standards, Washington, DC [now the National Institute of Standards and Technology, Gaithersburg, MD].)



Section 3.6 – Polymorphism

• Two or more distinct crystal structures for the same material (allotropy/polymorphism)

titanium α, β-Ti

carbon diamond, graphite

iron system liquid 1538°C δ-Fe BCC 1394°C **FCC** γ-Fe 912°C BCC α-Fe

Section 3.8 Point Coordinates



Point coordinates for unit cell center are

a/2, b/2, c/2 $\frac{1}{2}\frac{1}{2}\frac{1}{2}$

Point coordinates for unit cell corner are 111

Translation: integer multiple of lattice constants → identical position in another unit cell



Crystallographic Directions



Algorithm

- 1. Vector repositioned (if necessary) to pass through origin.
- 2. Read off projections in terms of unit cell dimensions *a*, *b*, and *c*
- 3. Adjust to smallest integer values
 - 4. Enclose in square brackets, no commas [*uvw*]

ex: 1, 0, ½ => 2, 0, 1 => [201]

-1, 1, 1 => [111] where overbar represents a negative index

families of directions <uvv>



Linear Density

Number of atoms

• Linear Density of Atoms = LD = Unit Hength of direction vector





HCP Crystallographic Directions



Algorithm

- 1. Vector repositioned (if necessary) to pass through origin.
- 2. Read off projections in terms of unit cell dimensions a_1 , a_2 , a_3 , or c

 a_3

2

- 3. Adjust to smallest integer values
- 4. Enclose in square brackets, no commas

 a_2

 a_1

 $-a_3$

[uvtw]

Adapted from Fig. 3.8(a), Callister 7e.

ex: $\frac{1}{2}, \frac{1}{2}, -1, 0$

=> [1120]

dashed red lines indicate projections onto a_1 and a_2 axes

HCP Crystallographic Directions

- Hexagonal Crystals
 - 4 parameter Miller-Bravais lattice coordinates are related to the direction indices (i.e., u'v'w') as follows.







Crystallographic Planes





Crystallographic Planes

- Miller Indices: Reciprocals of the (three) axial intercepts for a plane, cleared of fractions & common multiples. All parallel planes have same Miller indices.
- Algorithm
 - 1. Read off intercepts of plane with axes in terms of *a*, *b*, *c*
 - 2. Take reciprocals of intercepts
 - 3. Reduce to smallest integer values
 - 4. Enclose in parentheses, no commas i.e., (*hkl*)



PRACTICE



Crystallographic Planes

<u>example</u>		а	b	С	C.7
1.	Intercepts	1	1	∞	
2.	Reciprocals	1/1	1/1	1/∞	
		1	1	0	
3.	Reduction	1	1	0	ab
4.	Miller Indices	(110)			X
<u>example</u>		а	b	С	Z
1.	Intercepts	1/2	∞	∞	CZ
2.	Reciprocals	1/1⁄2	1/∞	1/∞	
		2	0	0	
3.	Reduction	2	0	0	
4.	Miller Indices	(100)			a b
					X HOLE MAN

⁺ у

y

Crystallographic Planes



Family of Planes {*hkl*}

Ex: $\{100\} = (100), (010), (001), (\overline{1}00), (0\overline{1}0), (00\overline{1})$

Crystallographic Planes (HCP)

• In hexagonal unit cells the same idea is used



Adapted from Fig. 3.8(a),.



Crystallographic Planes

- We want to examine the atomic packing of crystallographic planes
- Iron foil can be used as a catalyst. The atomic packing of the exposed planes is important.
 - a) Draw (100) and (111) crystallographic planes for Fe.
 - b) Calculate the planar density for each of these planes.



Planar Density of (100) Iron

Solution: At T < 912°C iron has the BCC structure.



Planar Density of (111) Iron





- Diffraction gratings must have spacings comparable to the wavelength of diffracted radiation.
- Can't resolve spacings $< \lambda$
- Spacing is the distance between parallel planes of atoms.

X-Rays to Determine Crystal Structure

• Incoming X-rays diffract from crystal planes.



Measurement of critical angle, θ_c , allows computation of planar spacing, *d*.





Adapted from Fig. 3.20, Callister 5e.



SUMMARY

- Atoms may assemble into crystalline or amorphous structures.
- Common metallic crystal structures are FCC, BCC, and HCP. Coordination number and atomic packing factor are the same for both FCC and HCP crystal structures.
- We can predict the density of a material, provided we know the atomic weight, atomic radius, and crystal geometry (e.g., FCC, BCC, HCP).
- Crystallographic points, directions and planes are specified in terms of indexing schemes. Crystallographic directions and planes are related to atomic linear densities and planar densities.



SUMMARY

- Materials can be single crystals or polycrystalline. Material properties generally vary with single crystal orientation (i.e., they are anisotropic), but are generally non-directional (i.e., they are isotropic) in polycrystals with randomly oriented grains.
- Some materials can have more than one crystal structure. This is referred to as polymorphism (or allotropy).
- X-ray diffraction is used for crystal structure and interplanar spacing determinations.

