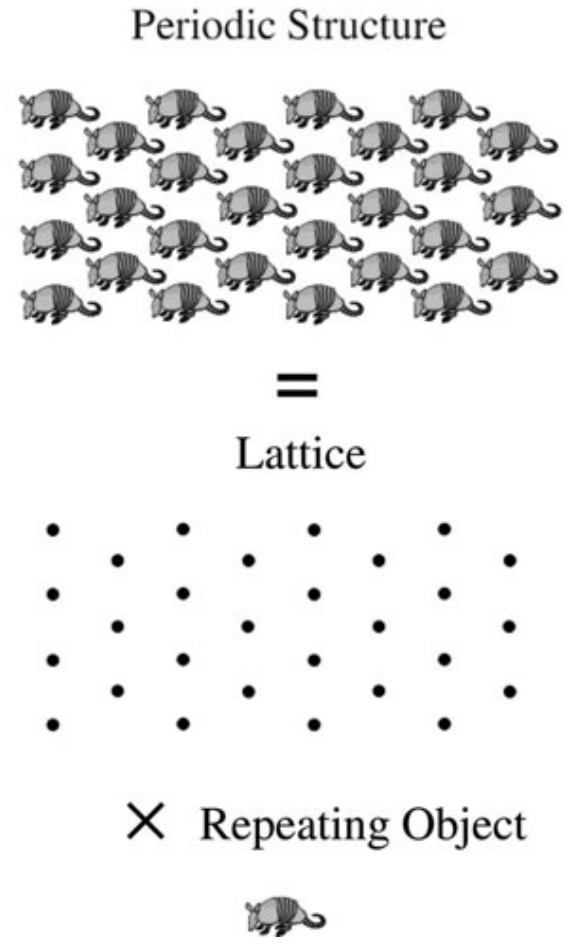
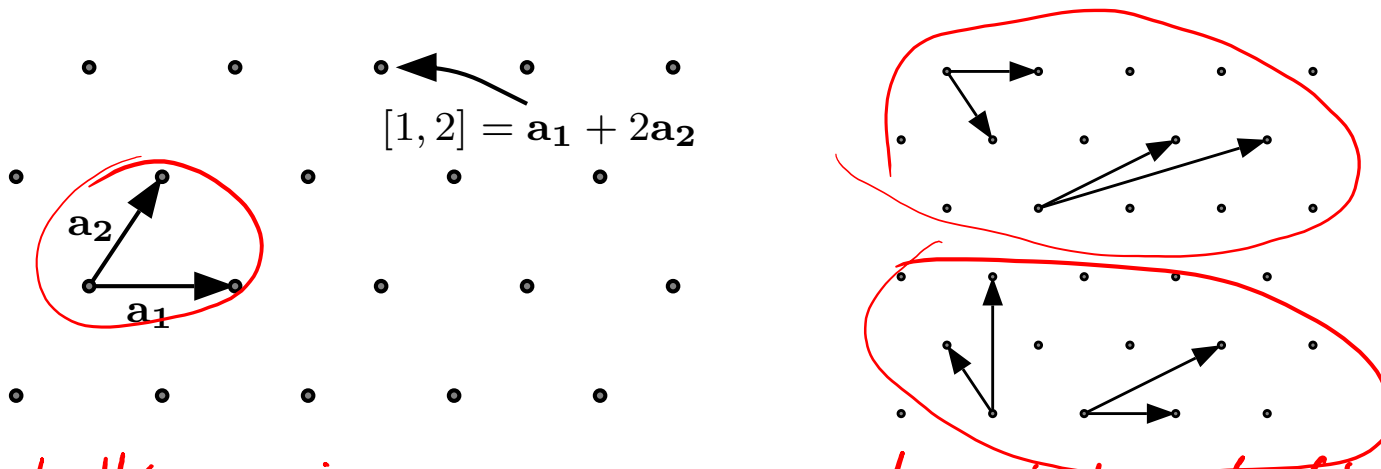


Crystal Structure

Lattices and Unit Cells



1) A lattice is an ∞ set of points defined by integer sums of a set of linear lattice vectors

$$\vec{R} = [h, k] = h_1 \vec{a}_1 + h_2 \vec{a}_2$$

$h = \dots -2, -1, 0, 1, 2, 3 \dots$

$1 \cdot a_1 + 2a_2$
 $[1, 2]$
 $h, k \in \mathbb{Z}$

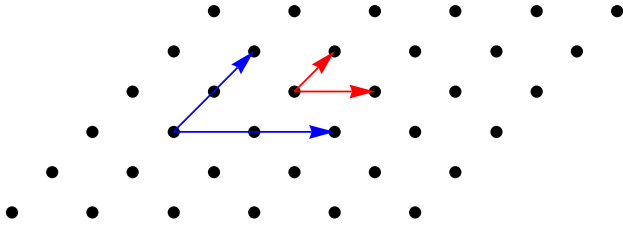
2) A lattice is a set of points where the env. looks the same and eq. to any lattice point.

Primitive lattice vectors

For a 3D lattice, we can find three **primitive lattice vectors (primitive translation vectors)**, such that any translation vector can be written as

$$\vec{t} = n_1 \vec{a}_1 + n_2 \vec{a}_2 + n_3 \vec{a}_3$$

where n_1, n_2 and n_3 are three integers.



Red (shorter) vectors: \vec{a}_1 and \vec{a}_2

Blue (longer) vectors: \vec{b}_1 and \vec{b}_2

\vec{a}_1 and \vec{a}_2 are primitive lattice vectors

\vec{b}_1 and \vec{b}_2 are NOT primitive lattice vectors

$$\vec{b}_1 = 2\vec{a}_1 + 0\vec{a}_2$$

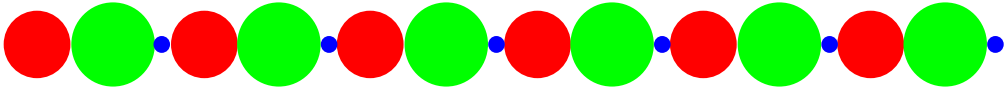
Integer coefficients

$$\vec{a}_1 = \frac{1}{2}\vec{b}_1 + 0\vec{b}_2$$

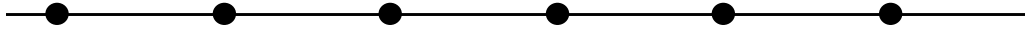
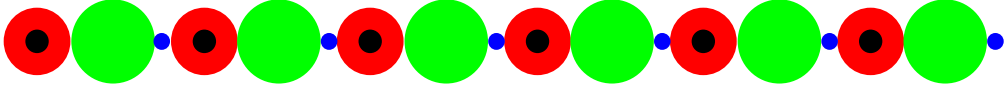
noninteger coefficients

1D crystal
3 atoms/periodicity

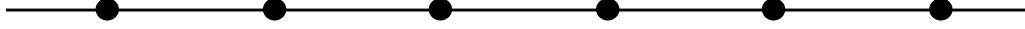
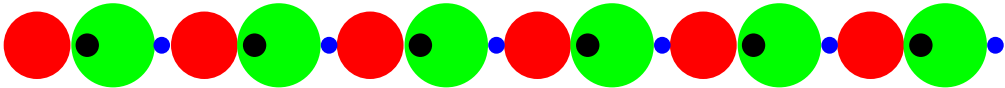
1D crystal
3 atoms/periodicity



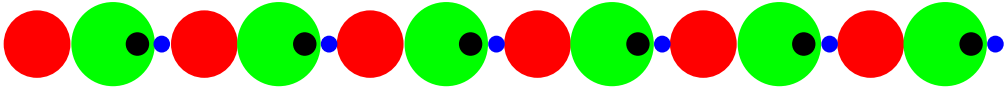
Choice I:



Choice II:



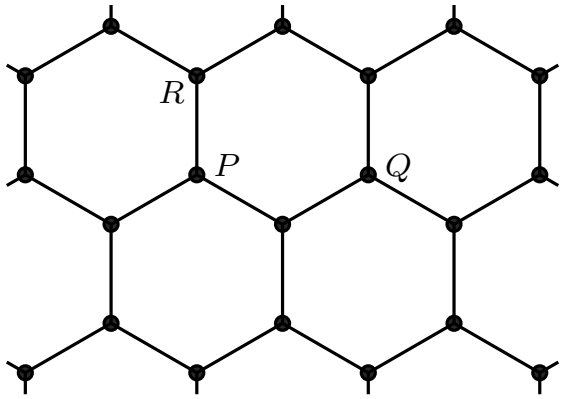
Choice III:



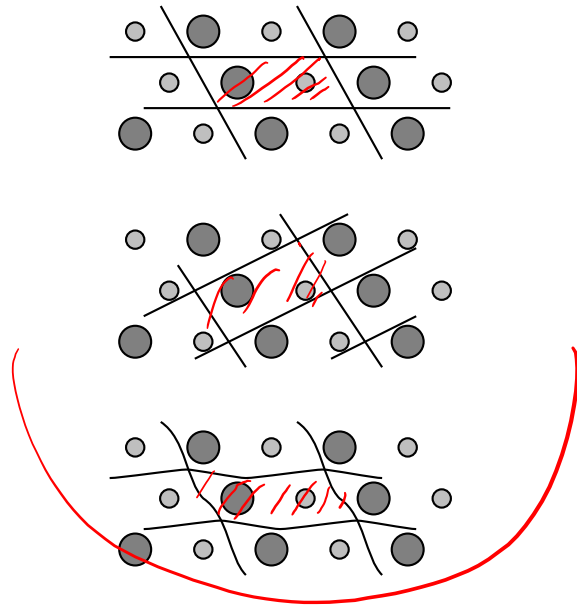
Choice I:

Choice II:

Choice III:



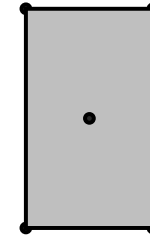
not
a lattice!



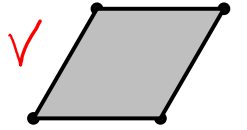
A unit cell is a region of space that has identical units stacked up and it tiles all the space

Primitive u.c. for a periodic crystal
should contain exactly one lattice point

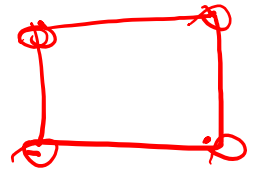
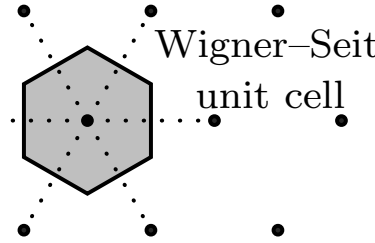
A conventional
unit cell



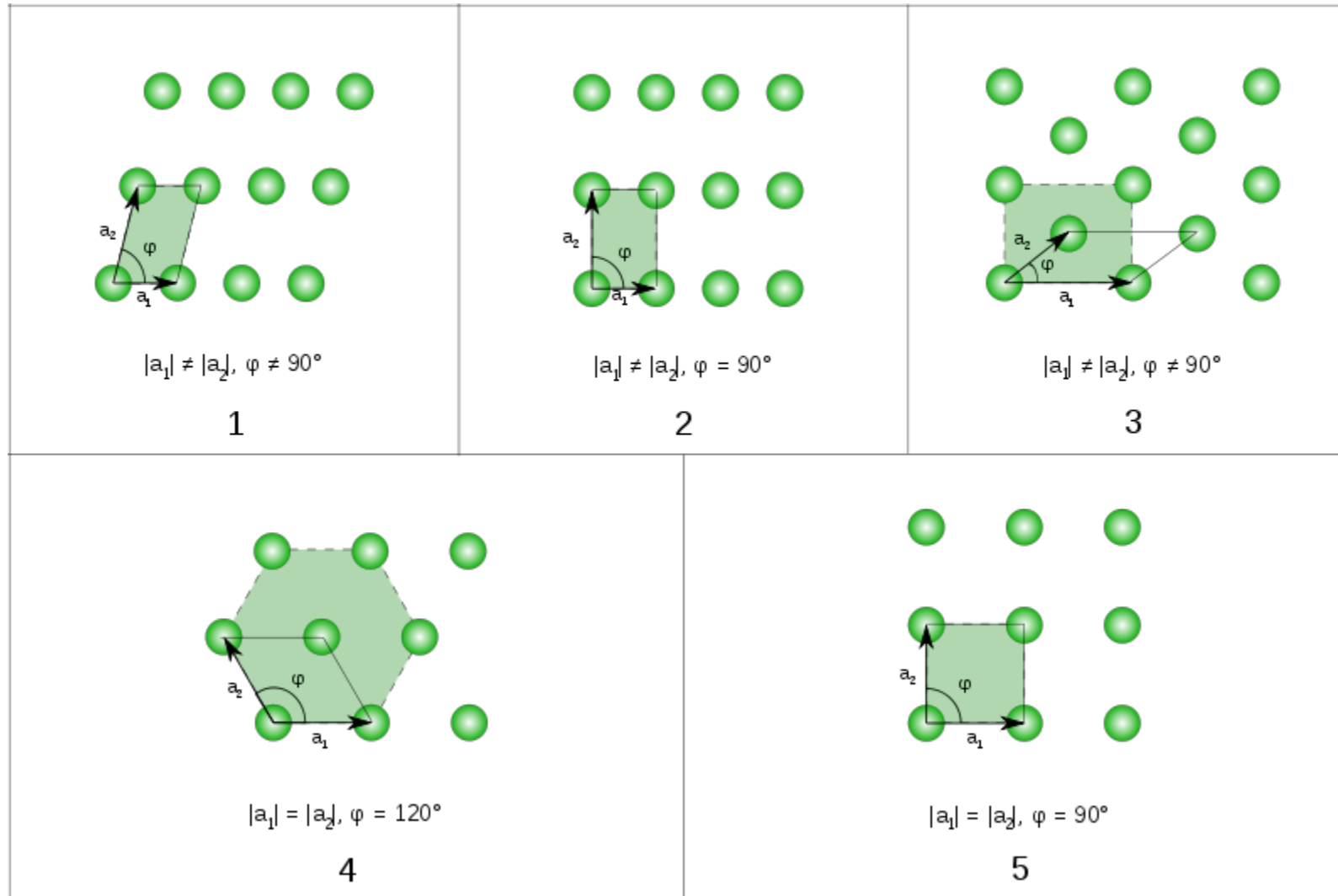
A primitive
unit cell



Wigner-Seitz
unit cell

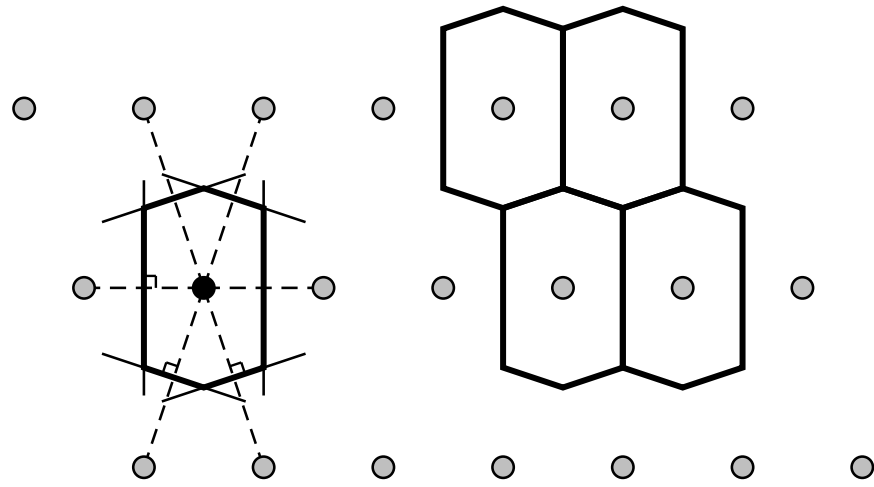


5 Bravais lattices in 2D

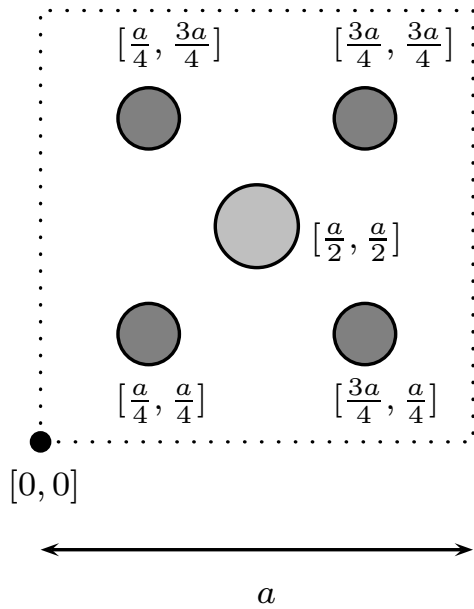
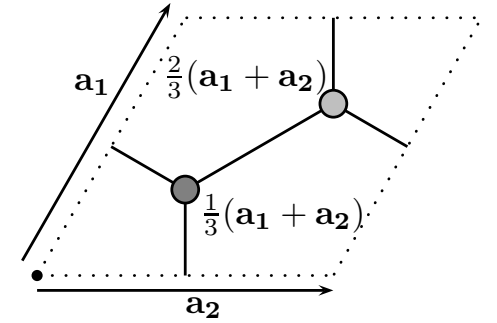
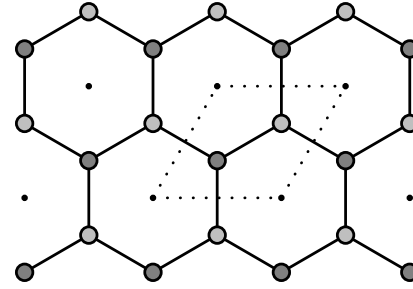
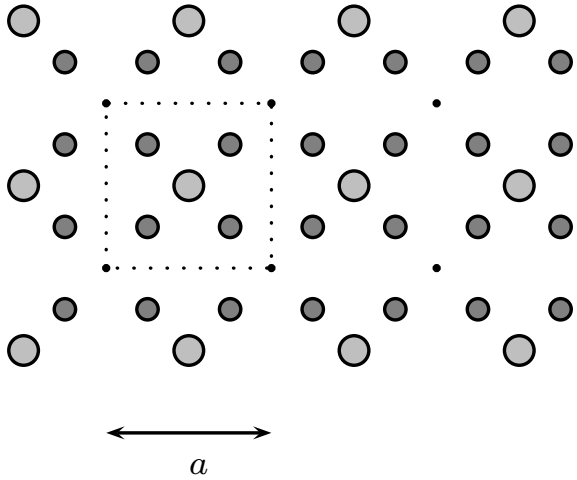


http://en.wikipedia.org/wiki/Bravais_lattice

Wigner Seitz construction



Basis and location of atoms in unit cell

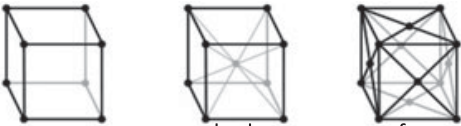


To remember: **CRYSTAL = LATTICE + BASIS**

· Lattices in Three Dimensions

14 Bravais lattices in 3D

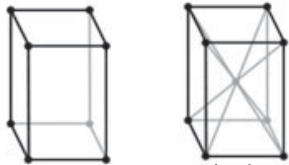
cubic:



simple body centered face centered

cubic:
all sides equal
all angles 90°


tetragonal:



simple body centered

tetragonal:
only two of three side lengths equal
all angles 90°

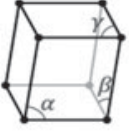
orthorhombic:




simple body centered face centered base centered

orthorhombic:
no two sides equal
all angles 90°

simple monoclinic




base centered monoclinic



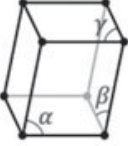
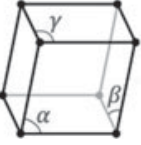
monoclinic:
angles between primitive lattice vectors :
 $\alpha = 90^\circ$ $\beta = 90^\circ$
 $\alpha \neq 90^\circ$
only two right angles

hexagonal

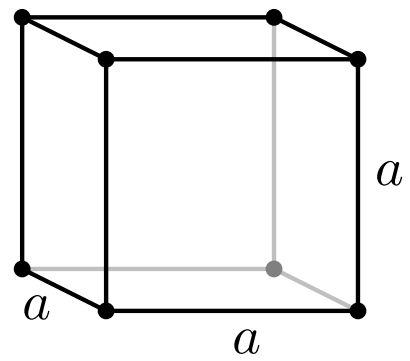
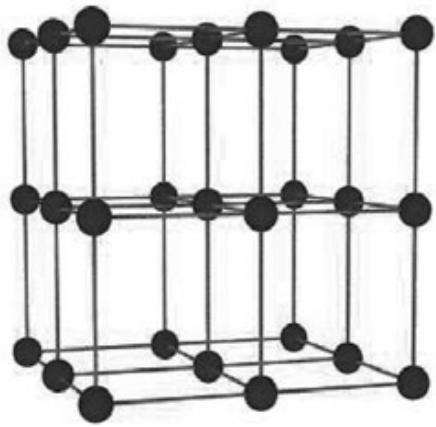


triclinic:

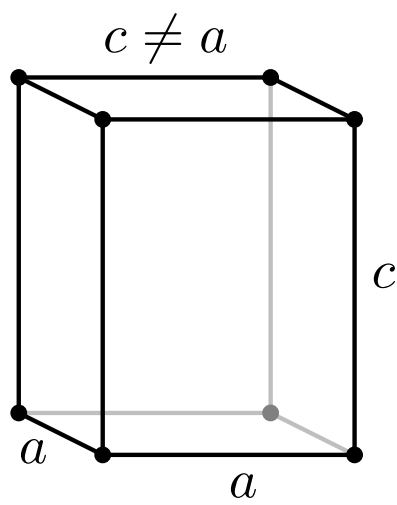
no right angles
no two sides equal

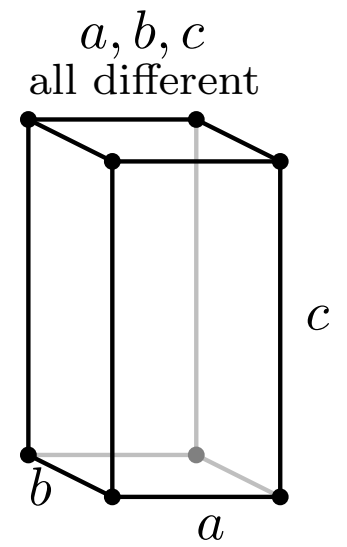
rhombohedral:
all side lengths equal.
all angles equal, but not right angles.



Cubic
unit cell

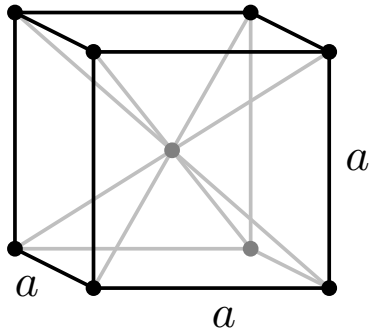


Tetragonal
unit cell

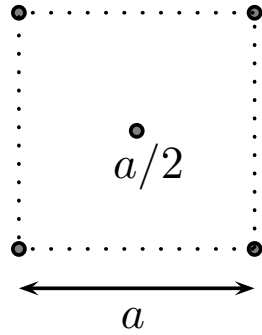


Orthorhombic
unit cell

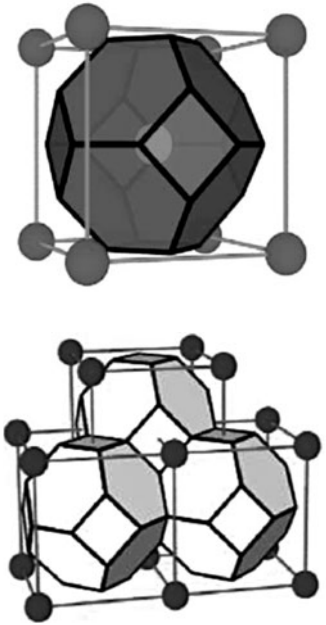
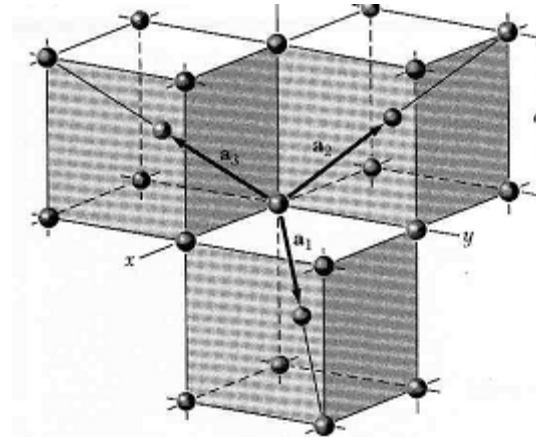
The Body-Centered Cubic (bcc) Lattice



Body-centered cubic
unit cell



Plan view



Lattice sites: $a(l \hat{x} + m \hat{y} + n \hat{z})$

Lattice point per conventional cell: $1 = 8 \times \frac{1}{8}$

Volume (conventional cell): a^3

Volume (primitive cell) : a^3

Number of nearest neighbors: 6

Nearest neighbor distance: a

Number of second neighbors: 12

Second neighbor distance: $\sqrt{2}a$

Coordinates of the sites: (l, n, m)

For the site $(0,0,0)$,

6 nearest neighbors: $(\pm 1, 0, 0)$, $(0, \pm 1, 0)$ and $(0, 0, \pm 1)$

12 next nearest neighbors: $(\pm 1, \pm 1, 0)$, $(0, \pm 1, \pm 1)$ and $(\pm 1, 0, \pm 1)$

Packing fraction

Packing fraction:

We try to pack N spheres (hard, cannot deform).

The total volume of the spheres is $N 4 \pi \frac{R^3}{3}$

The volume these spheres occupy $V > N 4 \pi \frac{R^3}{3}$ (there are spacing)

Packing fraction = total volume of the spheres / total volume these spheres occupy

$$\begin{aligned} \text{Packing fraction} &= \frac{N 4 \pi \frac{R^3}{3}}{V} = \frac{4 \pi \frac{R^3}{3}}{V/N} = \frac{4 \pi \frac{R^3}{3}}{\text{Volume per site}} \\ &= \frac{4 \pi \frac{R^3}{3}}{\text{Volume of a primitive cell}} \end{aligned}$$

High packing fraction means the space is used more efficiently

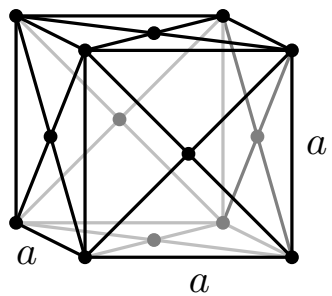
Packing fraction of simple cubic

$$\begin{aligned} \text{Packing fraction} &= \frac{4 \pi \frac{R^3}{3}}{\text{Volume of a primitive cell}} \\ &= \frac{4 \pi \frac{R^3}{3}}{a^3} = \frac{4 \pi}{3} \left(\frac{R}{a}\right)^3 = \frac{4 \pi}{3} \left(\frac{a/2}{a}\right)^3 = \frac{\pi}{6} \approx 0.524 \end{aligned}$$

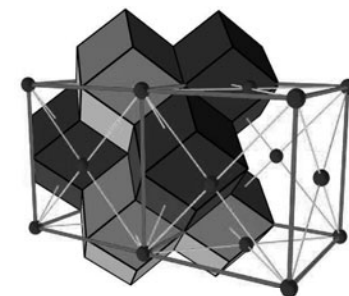
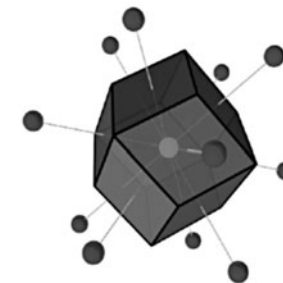
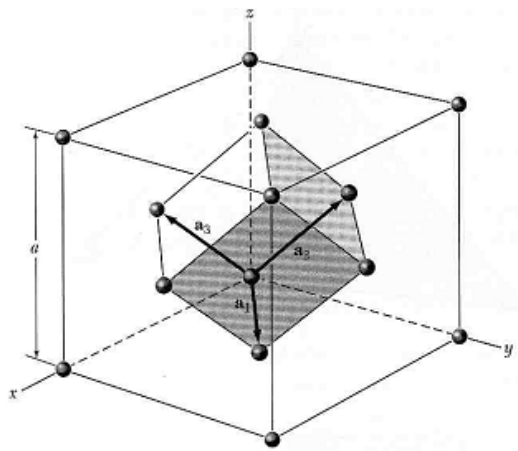
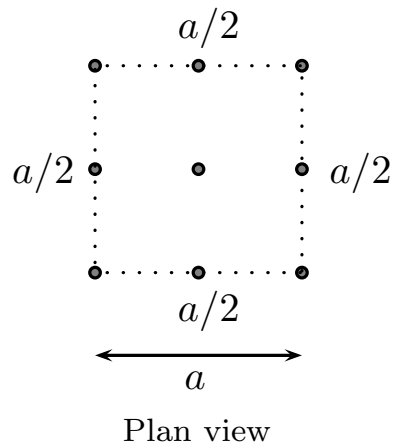
Nearest distance = 2 R
R = Nearest distance/2 = a/2

- About half (0.524=52.4%) of the space is really used by the sphere.
- The other half (0.476=47.6%) is empty.

The Face-Centered Cubic (fcc) Lattice




Face-centered cubic
unit cell

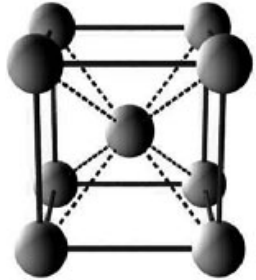


Some Real Crystals

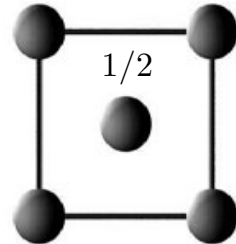
Sodium (Na)

Lattice = Cubic-I (bcc)

Basis = Na at [000] 





Plan view
unlabeled points at $z = 0, 1$

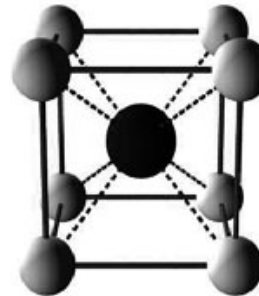


Caesium chloride (CsCl)

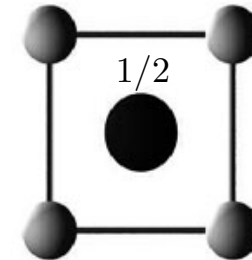
Lattice = Cubic-P

Basis = Cs at [000] 

and Cl at $[\frac{1}{2}\frac{1}{2}\frac{1}{2}]$ 



Plan view
unlabeled points at $z = 0, 1$



We can mark any unit cell by three integers: lmn

$$\vec{t} = l\vec{a}_1 + m\vec{a}_2 + n\vec{a}_3$$

Coordinates of an atom:

We can mark any atom in a unit cell by three real numbers: xyz .

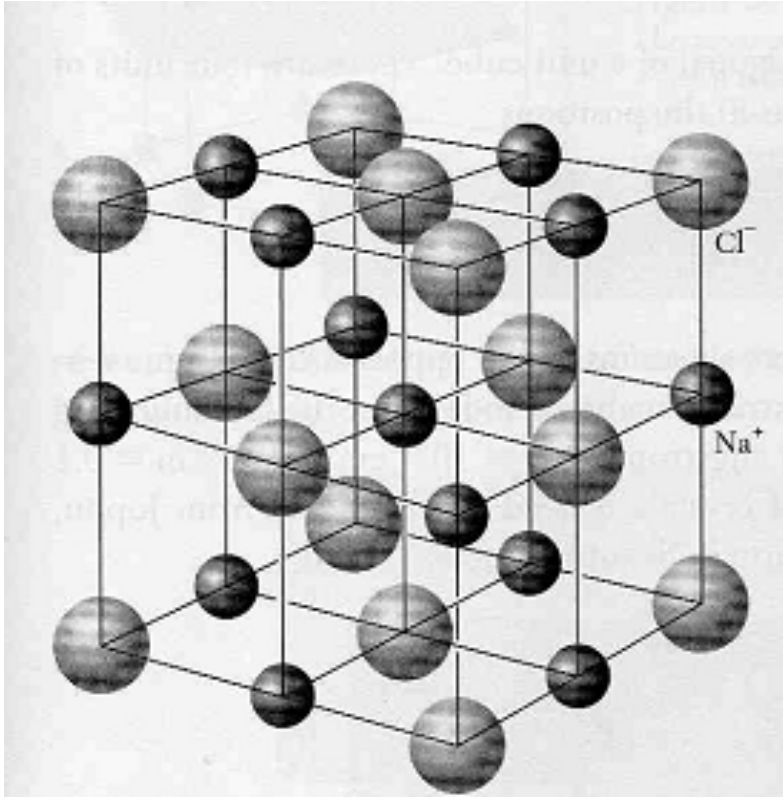
The location of this atom: $x\vec{a}_1 + y\vec{a}_2 + z\vec{a}_3$

Notice that $0 \leq x < 1$ and $0 \leq y < 1$ and $0 \leq z < 1$

Q: Why x cannot be 1?

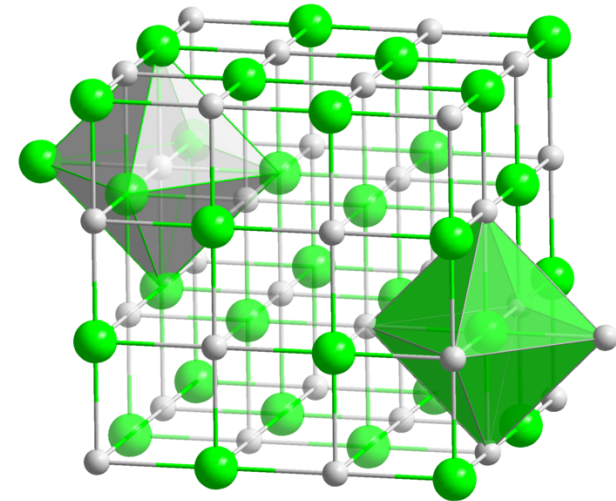
A: Due to the periodic structure. 1 is just 0 in the next unit cell

Sodium Chloride



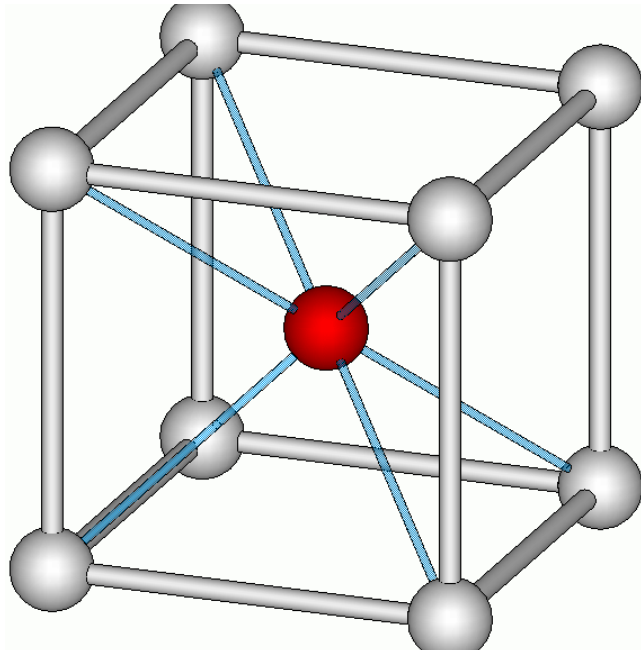
Face-centered cubic lattice
Na⁺ ions form a face-centered cubic lattice
Cl⁻ ions are located between each two neighboring Na⁺ ions

Equivalently, we can say that
Cl⁻ ions form a face-centered cubic lattice
Na⁺ ions are located between each two neighboring Na⁺ ions



Primitive cells

Cesium Chloride



Simple cubic lattice

Cs⁺ ions form a cubic lattice

Cl⁻ ions are located at the center of each cube

Equivalently, we can say that

Cl⁻ ions form a cubic lattice

Cs⁺ ions are located at the center of each cube

Coordinates:

Cs: 000

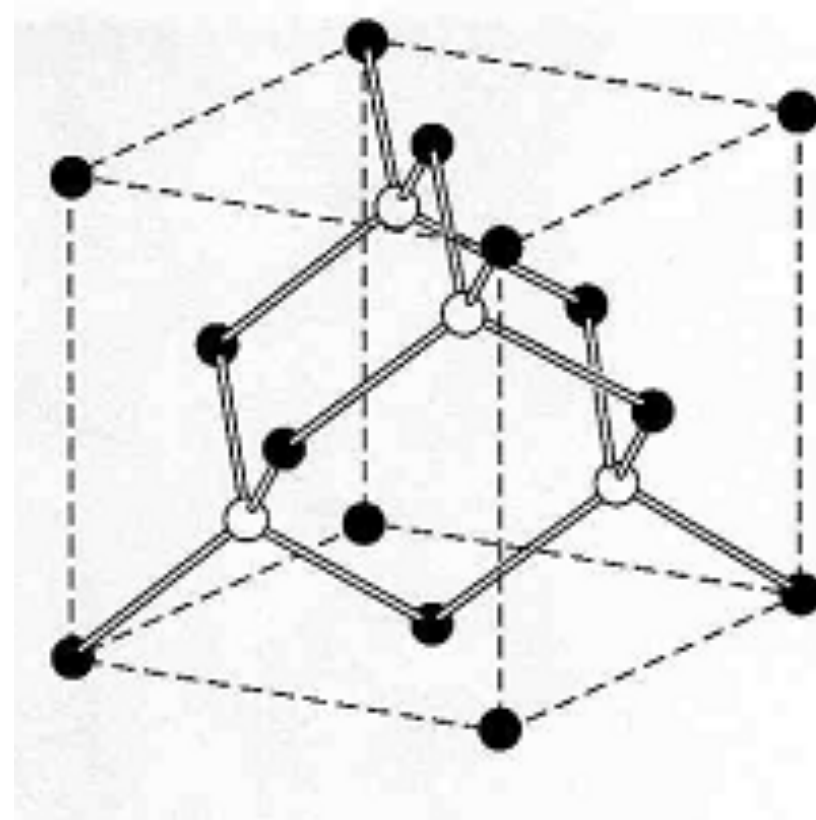
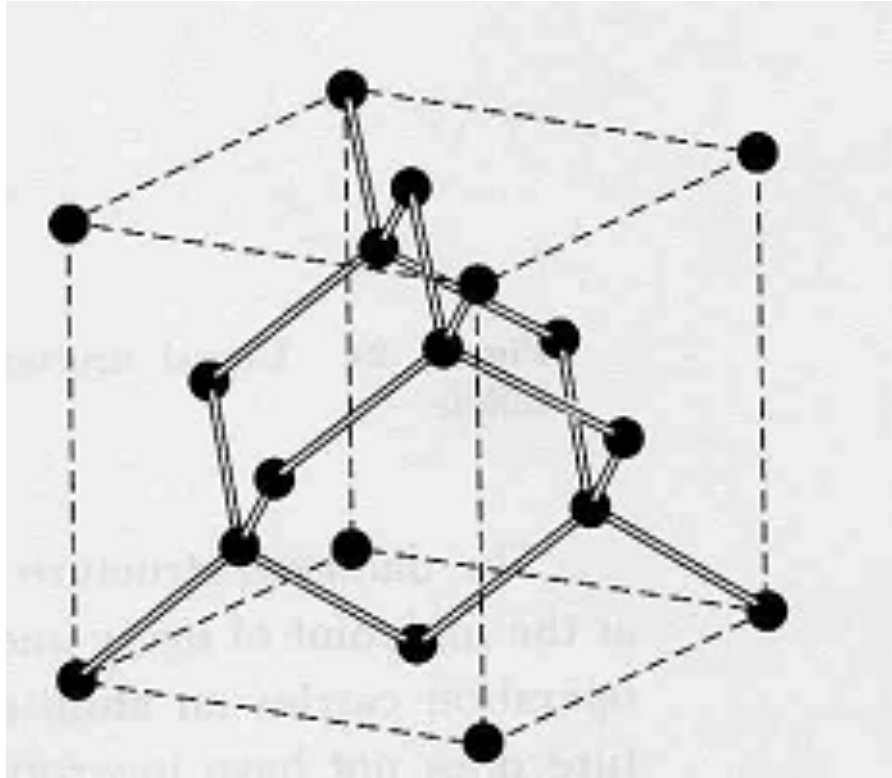
Cl: $\frac{1}{2} \frac{1}{2} \frac{1}{2}$

Notice that this is a simple cubic lattice

NOT a body centered cubic lattice

- For a bcc lattice, the center site is the same as the corner sites
- Here, center sites and corner sites are different

Diamond is not a Bravais lattice



Same story as in graphene:

We can distinguish two different type of carbon sites (marked by different color)

We need to combine two carbon sites (one black and one white) together as a (primitive) unit cell

If we only look at the black (or white) sites, we found the Bravais lattice: fcc