Lecture 1 - Ionic structures

Course Summary

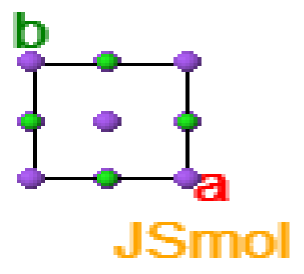
This course aims to introduce you to the importance of ionic materials in many applications.

What do I want you to do?

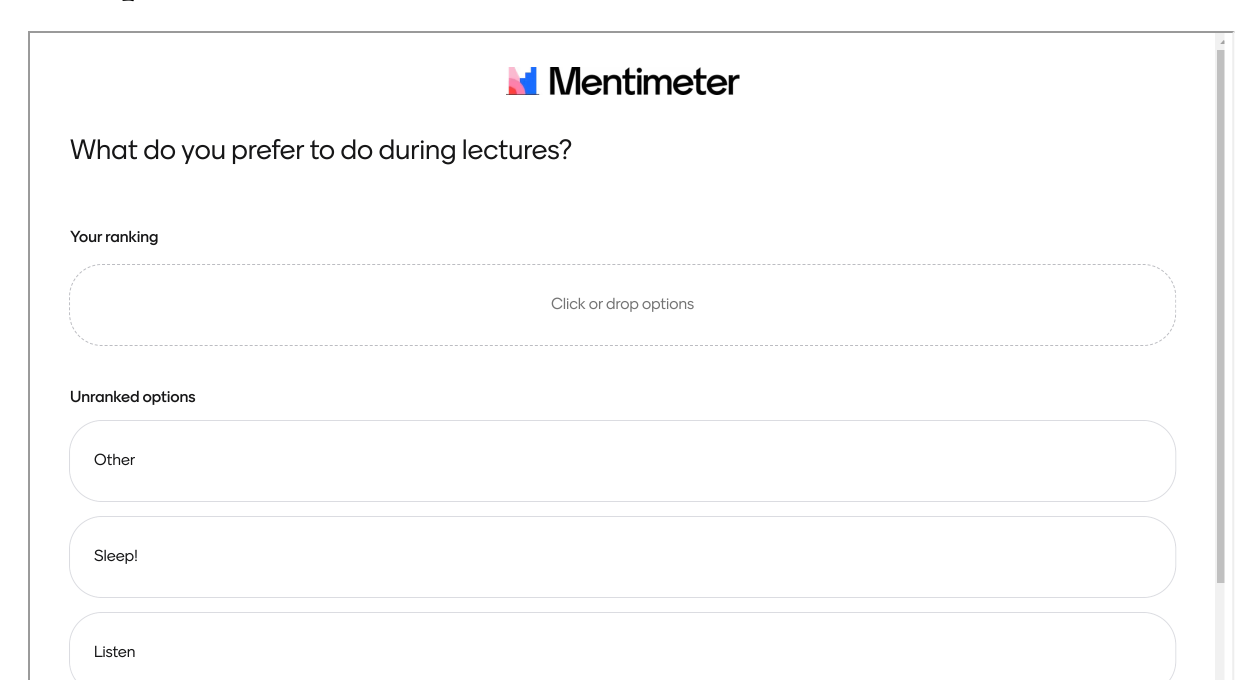
- Revise basic crystallography and ionic solids
- Try to understand examples, don't memorise them
- If in doubt ask questions!

Lecture Notes

- Interactive HTML notes
 - Detailed instructions on overview page
 - Explore the jmol structures!
 - We'll have live quizzes during lectures please engage
- PDF notes also on Learn if needed



Test poll!



Results

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Let's get you thinking in 3D!

Picture a cube. Imagine touching one corner, and then also touch every corner that is two edges away from it.

| Mentin | neter | | |
|---|-------|--|--|
| you join those corners by new edges, what shape do you get? | | | |
| Hexagon | | | |
| Tetrahedron | | | |
| Square | | | |
| Triangle | | | |

Results

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Lecture 1 Summary

- Types and applications of ionic materials
- Crystallography recap
- Lattice energy and ionic bonding
- Close-packing and ionic structure types

- Many inorganic solids
 - \circ e.g. $\mathrm{Na}^{+}\mathrm{Cl}^{-}$ and $\mathrm{Mg}^{2+}\mathrm{SO}_{4}^{2-}$

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 - ∘ ammonium acetate NH₄⁺CH₃COO[−]
 - o chlorphenirammonium maleate (active part of Piriton®)

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- (in)organic salts
 - $\circ\,$ Mono-/Di-/Tri-Sodium citrate $\mathrm{Na_{x}C_{6}H_{8\,-x}O_{7}}$
 - collectively used as E331 in food
 - x can be varied from 1–3

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- Ionic liquids
 - Either organic or inorganic, liquid below 100 °C

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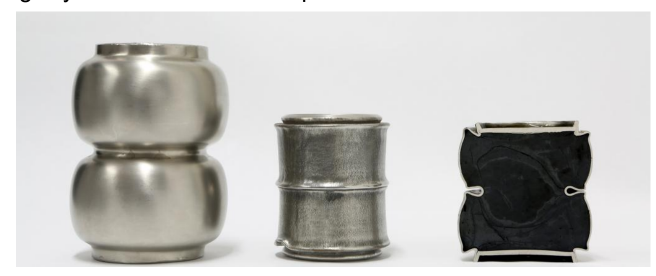
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- High melting points due to Coloumbic energy (see <u>later</u>)
- Electrically insulating
 - Electronegativity differences promote localised electrons
- Usually hard, and often robust to harsh conditions
 - ∘ e.g. Synroc* is used to encapsulate nuclear waste

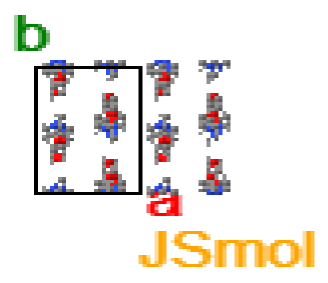




We can divide solids into two categories:

Molecular (e.g. paracetamol)

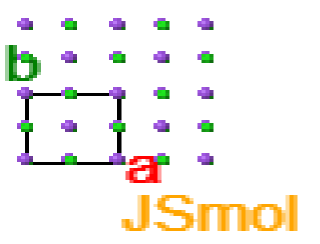
- Strong intramolecular bonds
- Weaker intermolecular interactions



We'll concentrate on infinite materials.

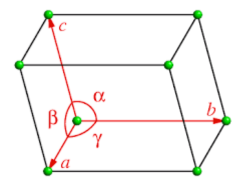
Infinite (e.g. NaCl)

- Strong bonds between all atoms
- No discrete molecules



Recap on crystal structure

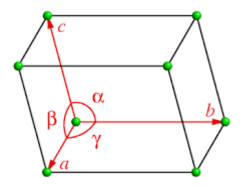
Periodic solids can be described by a unit cell



- Defined by lengths (a, b, c) and angles (α, β, γ)
 - 'Lattice parameters'

Recap on crystal structure

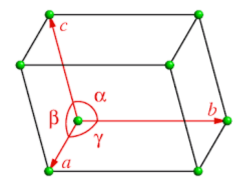
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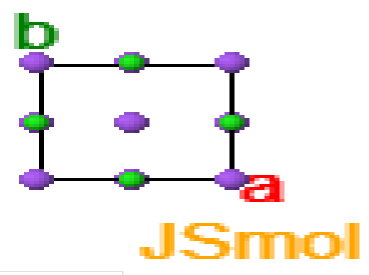
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- Atom positions defined by fractional position along lattice directions

Example: Sodium chloride



| Cubic structure | $a=b=c=5.62$ Å, $lpha=eta=\gamma=90^\circ$ | | | | |
|-----------------|--|-------------|-------------|---------------|------------------------|
| Spacegroup | $\mathrm{Fm}\bar{3}\mathrm{n}$ | n (#225, p | oint group | $o = O_h$) | |
| Na atoms at: | (0 0 0) | (1/2 1/2 0) | (1/2 0 1/2) | (0 ½ ½) | (all symmetry-related) |
| Cl atoms at: | (1/2 0 0) | (0 ½ 0) | (0 0 ½) | (1/2 1/2 1/2) | (all symmetry-related) |

Because of symmetry, we only need to define one Na and one CI position.

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• $\frac{1}{r}$ dependence makes long-range interactions important

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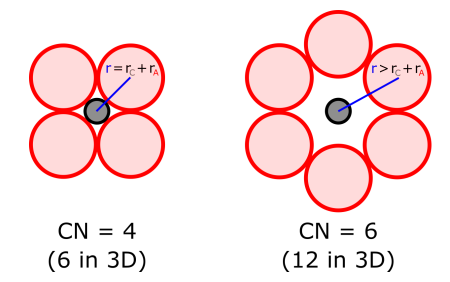
For example NaCl:

$$egin{align} E_{ ext{Madelung}} &= \sum_{i
eq j} rac{q_i q_j}{4\pi\epsilon_0 r} \ &= rac{ ext{N}_{ ext{A}} q_i q_j}{4\pi\epsilon_0 r} igg(6 - rac{12}{\sqrt{2}} + rac{8}{\sqrt{3}} - rac{6}{2} + rac{24}{\sqrt{5}} - \dots igg) \ &\simeq rac{ ext{N}_{ ext{A}} q_i q_j}{4\pi\epsilon_0 r} imes 1.74756 \ \end{aligned}$$

Ionic Structures

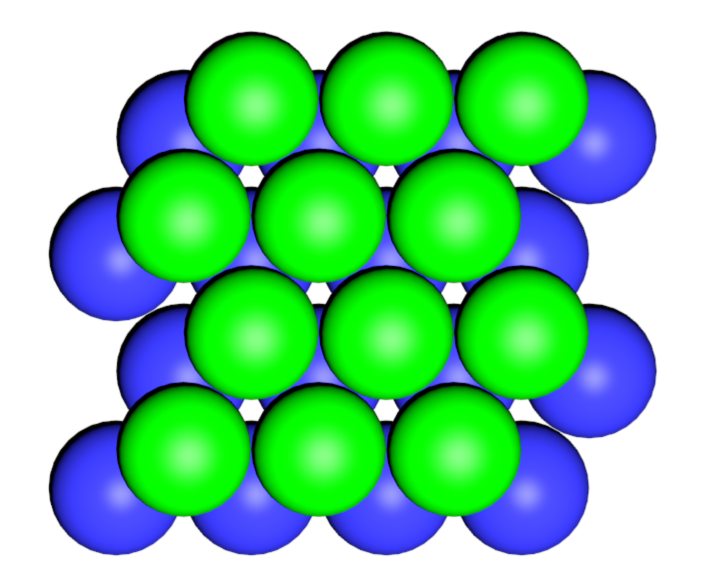
Generally, structures **maximise cation-anion** interactions (-ve energy) while **minimising like-charge** interactions (+ve energy)

- Maximise cation-anion coordination number
 - Ideally, ions should be densely packed



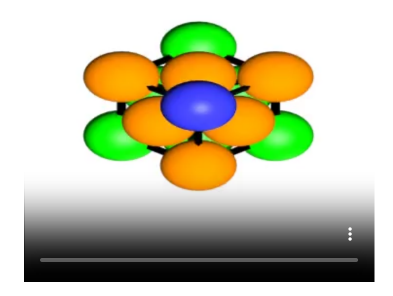
In many materials, the optimum is found when the largest ion (often oxide) is **close- packed**

Close packing

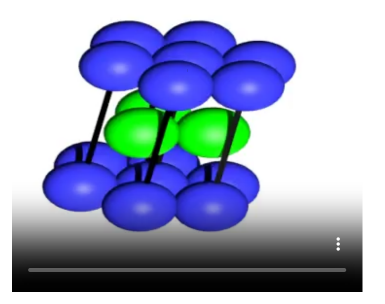


Close packing

Face-centered cubic (FCC) ... ABCABC ...

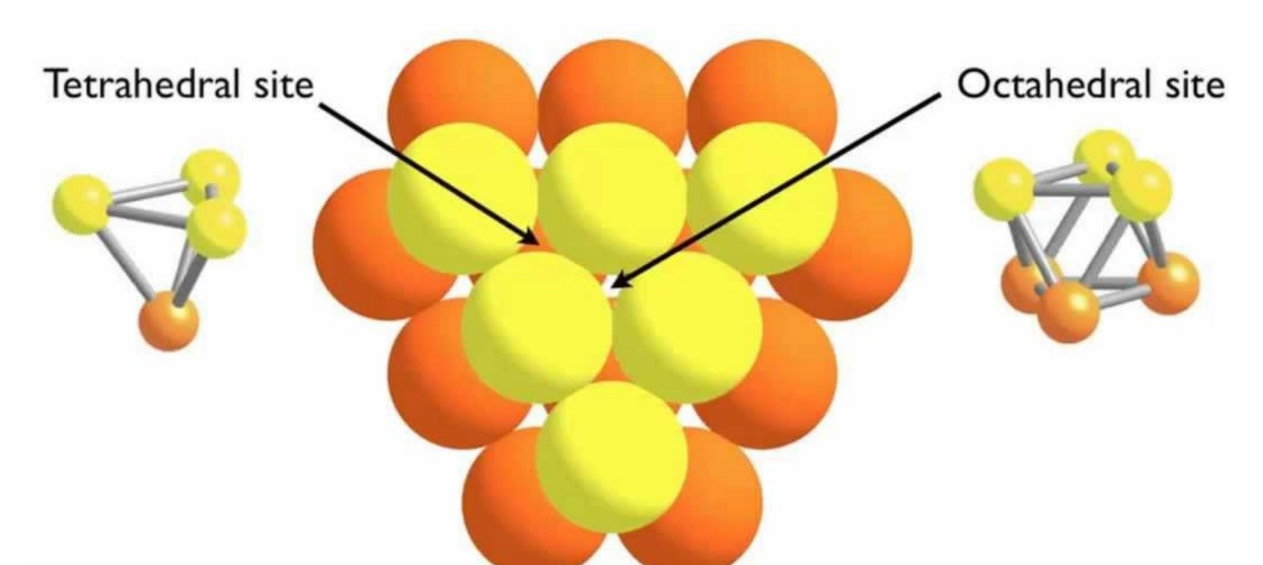


Hexagonal close-packed (HCP) ... ABABAB ...

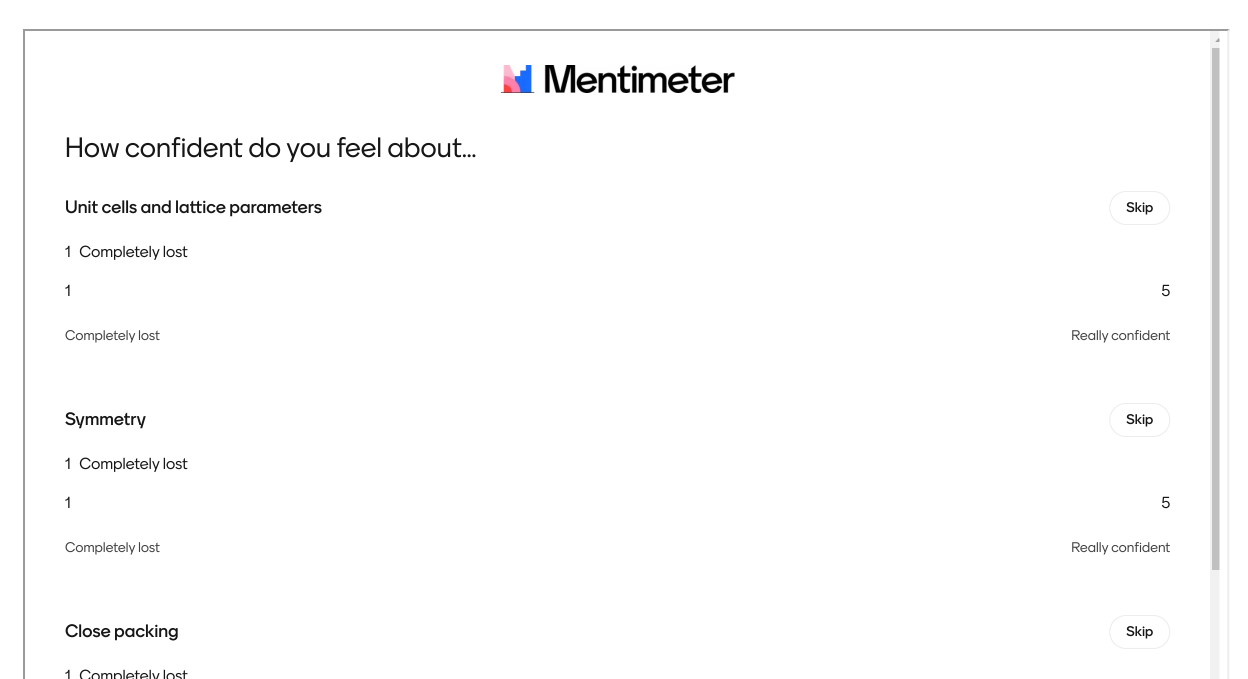


Holes

CP arrangements of large (an)ions [X] leave 'holes' within the structure, which can be occupied by smaller (cat)ions [M]



How are you getting on? Vote



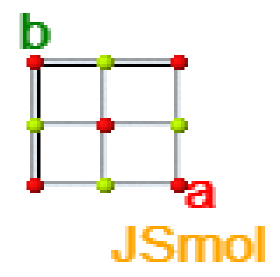
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How are you getting on? Results

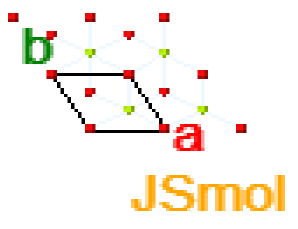
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Octahedral holes

One hole per cp ion - both are 6-coordinate



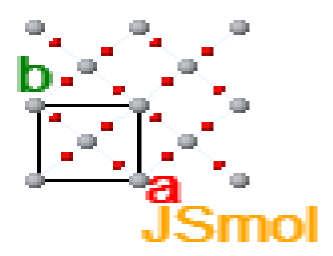
Rock salt (NaCl) structure



Nickel Arsenide structure (e.g. FeS)

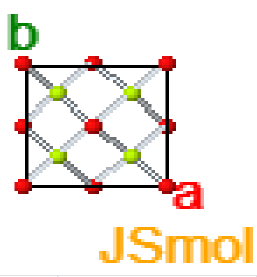
Rutile

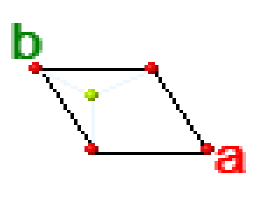
Although not strictly close-packed, rutile (${
m TiO_2}$) is distorted HCP with ${
m Ti}^{4+}$ filling half the octahedral holes CN = 6 / 3



Tetrahedral holes

Two holes per cp ion







| Holes filled | FCC Type | CN(A/X) | HCP Type | CN(A/X) |
|--------------|------------------------------|---------|----------------|---------|
| All | Fluorite (CaF ₂) | 4/8 | (not possible) | - |
| Half | Zinc-blende (ZnS) | 4/4 | Wurtzite (ZnS) | 4/4 |

Which structure type?

Generally, the structure formed depends on the ratio of ionic radii

• Smaller cations will prefer lower coordination numbers

| $rac{r^+}{r^-}$ | Cation C.N. | MX Structure | MX ₂ Structure |
|------------------|-------------|----------------------------|-------------------------------|
| 0.7 - 1.0 | 8 | CsCl | CaF_2 |
| 0.4 - 0.7 | 6 | NaCl | ${ m TiO}_2$ |
| 0.2 - 0.4 | 4 | ZnS (Wurtzite/Zinc-blende) | Anti-fluorite (e.g. Li_2S) |

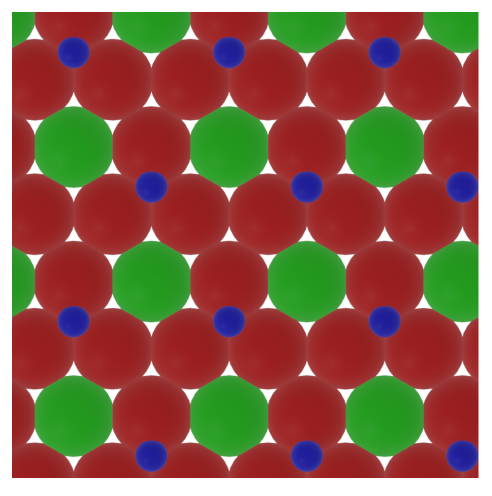
These are only approximate 'rules', and other binary structures exist (e.g. CdI_2 , CdCl_2 , PbO , etc...)

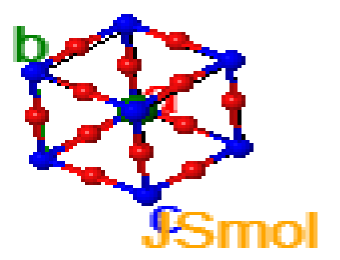
Very difficult to predict!

Beyond binary compounds

With 3 or more elements, structures become much more complicated! An important one is perovskite, ABX_3

• $r(A) \simeq r(X)$, so can be considered as FCC AX_3 layer with B filling 25% of octahedral holes:





Lecture recap

- Variety of ionic materials with a range of applications
- Revision of basic crystallography
 - Unit cells, symmetry
- Electrostatic interaction hold ionic crystals together
 - Long-ranged
 - Aim to maximise cation-anion interactions
- Close-packing of anions often most stable
 - Ratio of ionic radii suggests which structure is adopted
 - Beyond binary compounds, predicting structures is hard!

Feedback



What did you like or dislike about this lecture?

Short answers are recommended. You have 200 characters left.

200

You can submit multiple responses

Submit

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