

Lecture 2 - defects

Lecture summary

- Introduction to defects
- Types of defect
- Intrinsic and extrinsic defects
- Defect equations

Defects

All crystals contain defects of some sort, for example:

- Missing atoms (*vacancies*)
- Atoms in the 'wrong' place
 - *interstitials* (between lattice sites) or *substitutions* (different atom types)
- Extended defects of lines or planes of atoms

Defects

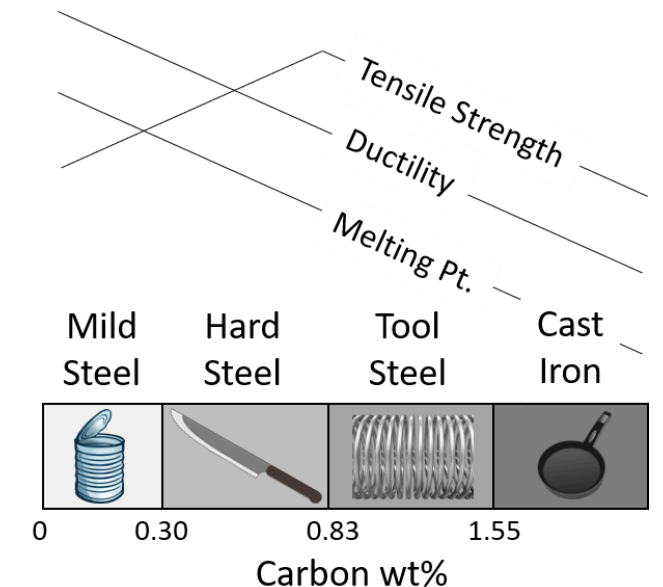
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Defects are often the source of interesting properties



~1% Substitution in Al_2O_3

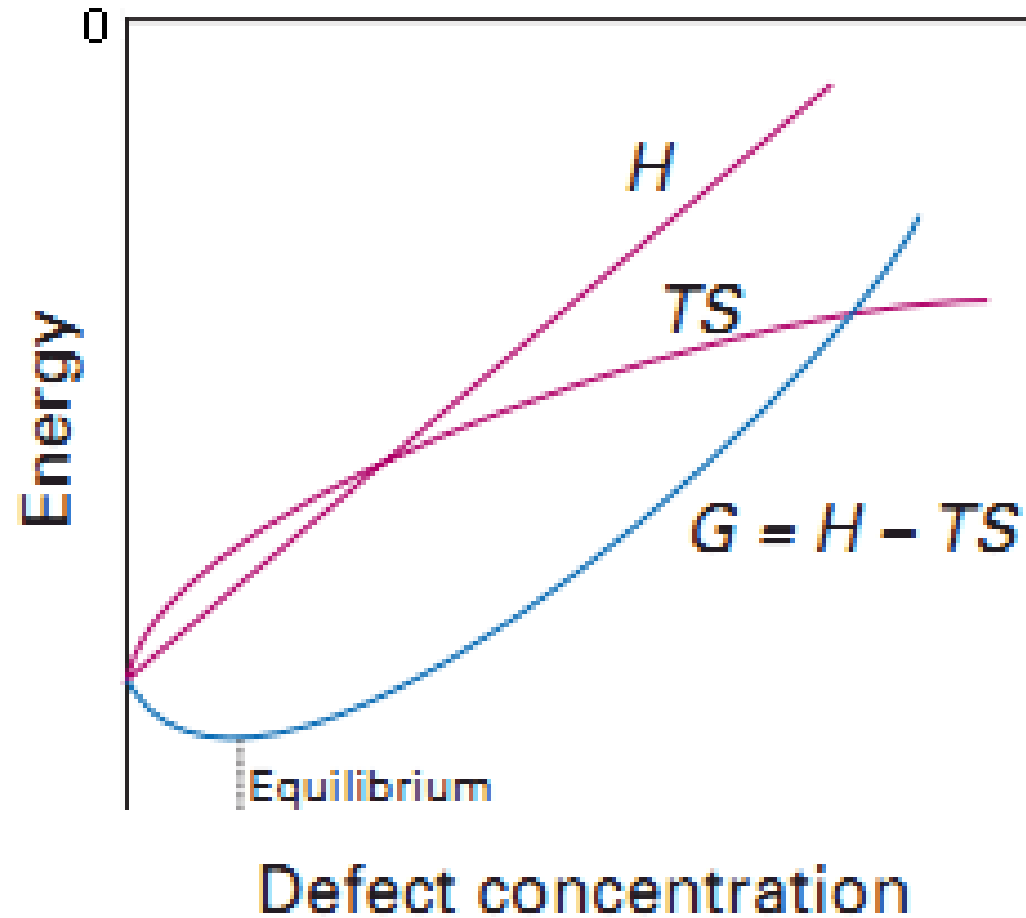


Effect of interstitial carbon on iron properties

Defect amounts

The number of defects is a fine balance of entropy and enthalpy


- Defects gain entropy but have a (often large) formation energy



Minimum in ΔG depends on structure and bonding, but typically $\ll 1\%$.


Types of defect

The three most common defect types in ionic solids are:

Vacancy	Interstitial						Substitution					
M X M X M X	M X M X M X							M X M X M X				
X M X M X M	X M X M X M							X M X M X M				
M X M X M X	M X M X M X							M X M X M X				
X M X  X M	X M X ^X M X M							X M X M X M				
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Additionally, defects can be either

- *intrinsic* (maintaining stoichiometry) or
- *extrinsic* (non-stoichiometric)

Intrinsic defects

Two of the most common stoichiometric defects are:

Schottky

- Charge-balanced combination of anion and cation vacancies

Frenkel

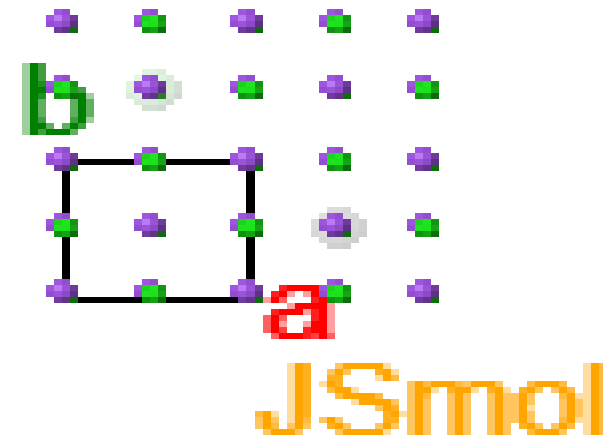
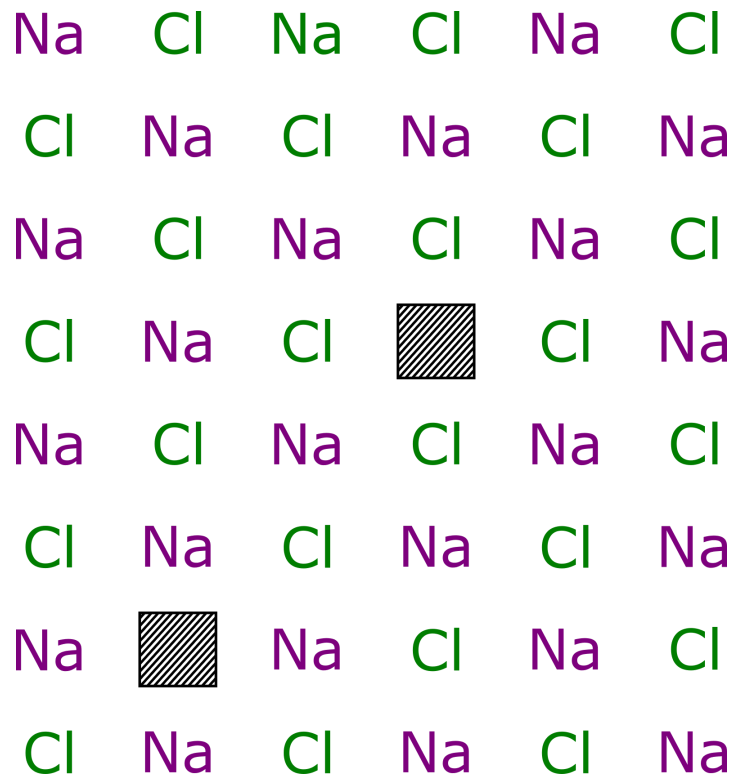
- Ions displaced to interstitial sites

Defects observed depend on both structure type and atoms involved.



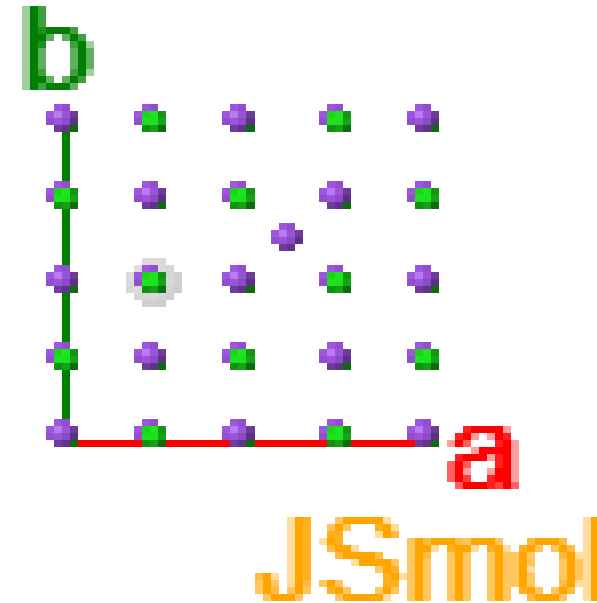
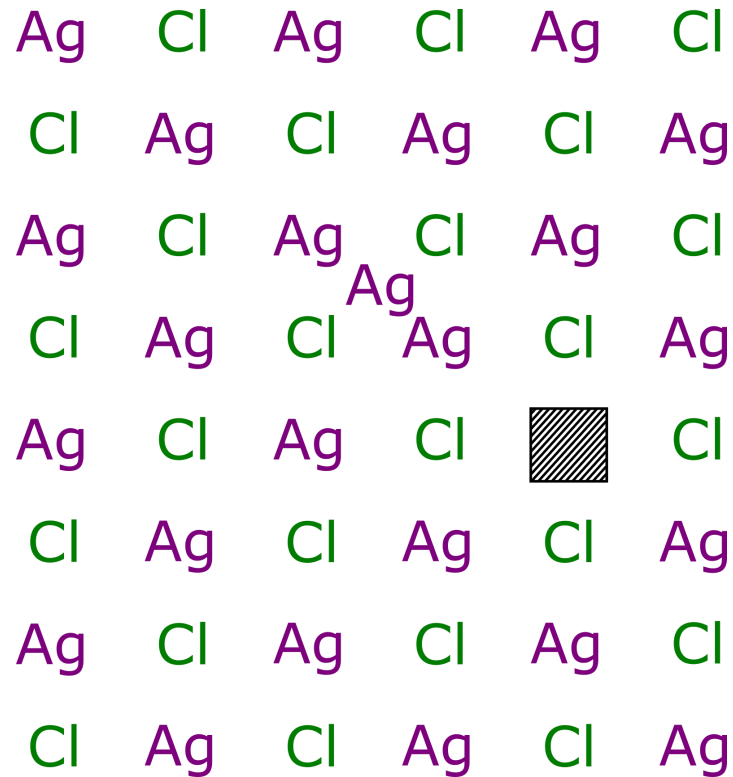
Schottky defects

- Typically occur when anions and cations have similar size (e.g. NaCl structure)
- Reduced density compared with the ideal material
- e.g. NaCl - equal numbers of Na and Cl vacancies



Frenkel defects

- Smaller ion normally displaced
- Only one ion type shows defect
- e.g. AgCl (NaCl-type)
 - Smaller Ag^+ ion displaced to tetrahedral holes in CCP Cl^- structure



Defect equations

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- For example:
 - Na vacancy in NaCl: V_{Na}'
 - Ag interstitial in AgCl: Ag_i^\bullet

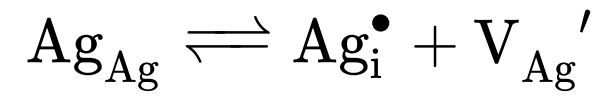
Defect equations (2)

like normal, defect equations must balance in terms of:

- composition
 - vacancies are not treated as an atom
- charges
- **sites**
 - *specified* atomic positions cannot be created or destroyed
 - interstitials are ignored in balancing

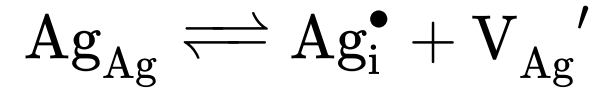
Examples

AgCl interstitial formation again:



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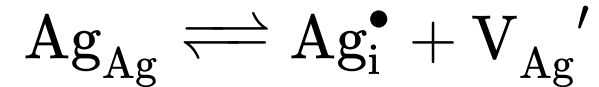


NaCl Schottky formation:



Examples

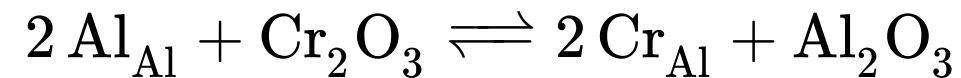
AgCl interstitial formation again:



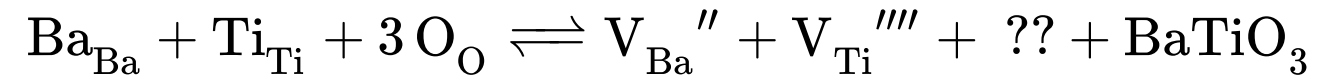
NaCl Schottky formation:



Easily extended to substitutions, e.g. substituting Al^{3+} with Cr^{3+} in Al_2O_3 (ruby):



Quick test - BaTiO₃ Schottky Formation

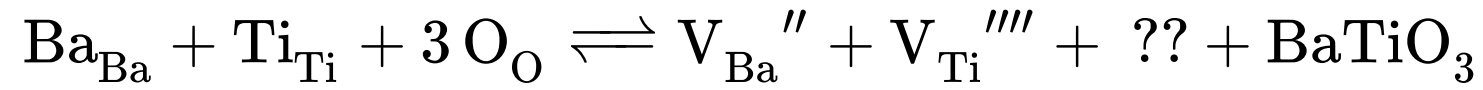


What is missing?

$3 \text{V}_{\text{O}}^{\bullet\bullet}$

$3 \text{O}_{\text{i}}''$

Submit



wooclap

Quiz results will be available here
after the lecture

Ionic Substitution

- Ions of similar size can often replace each other

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- While an integer number are substituted across a crystal, the average can be non-stoichiometric
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 - i.e. Ruby is $\text{Al}_{2-x}\text{Cr}_x\text{O}_3$ ($0 \leq x \leq 2$)
- Substitution can dramatically affect properties:
 - e.g. $\text{La}_{2-x}\text{Sr}_x\text{CuO}_4$:
 - semiconducting for $x = 0$
 - superconducting (below 40 K) for $x = 0.15$

Extrinsic defects

Substitution can also drive formation of defects, e.g. doping NaCl with CaCl₂:

Overall synthesis reaction:



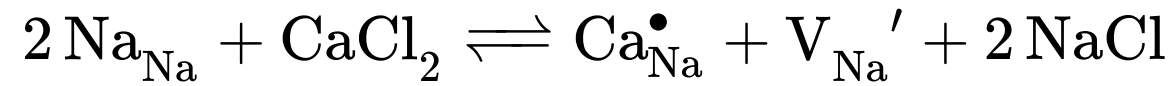
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Kroger-Vink notation:



More complex example

Sometimes, substitution (or 'doping') can give rise to multiple potential defects.

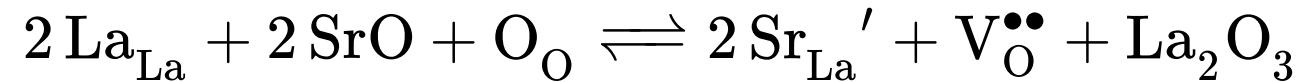
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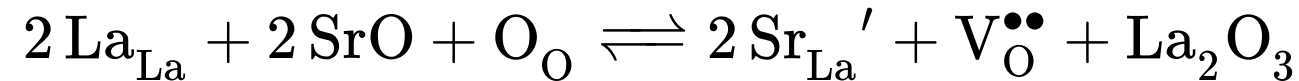


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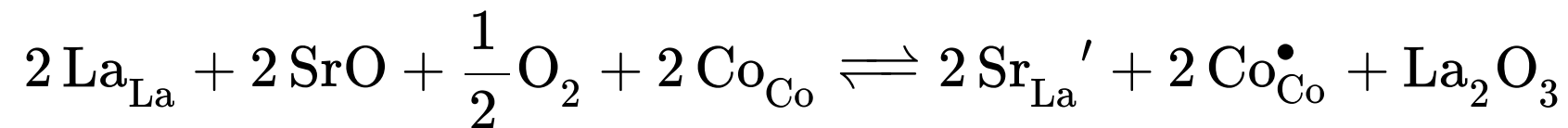
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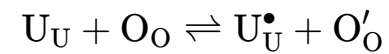
- or by oxidising Co^{3+} to Co^{4+}



Quiz 2 - UO₂ Extrinsic defects

Nuclear fuel (fluorite-type UO₂) can oxidise in air to form UO_{2+δ}.

What is the correct KV mechanism for UO₂ oxidation?



Submit

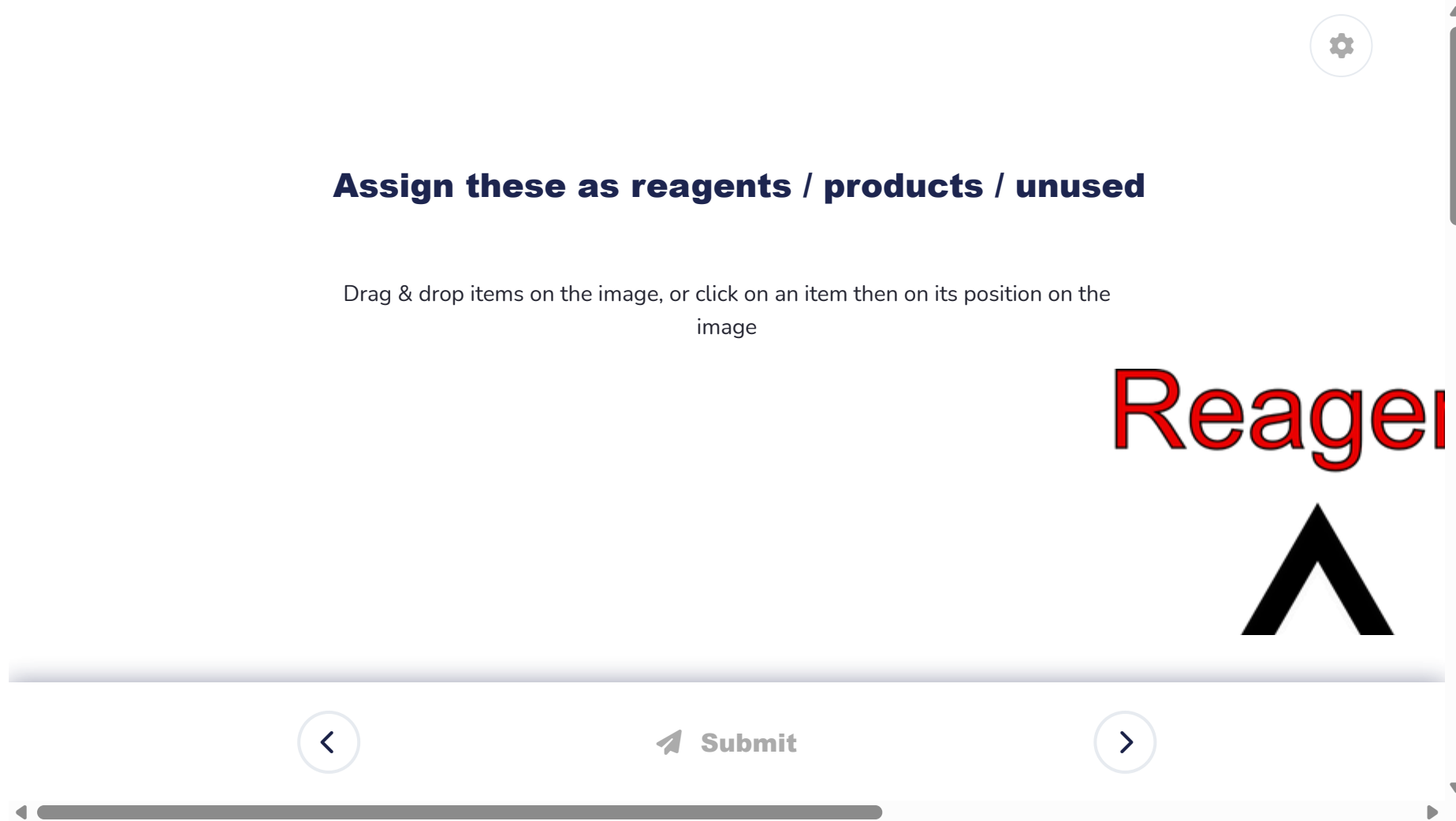
Results - UO₂ defects

wooclap

Quiz results will be available here
after the lecture

Quiz 3 - More Extrinsic defects

At high pressure, oxygen vacancies in Mg_2SiO_4 can react with H_2O to form new defects.



Assign these as reagents / products / unused

Drag & drop items on the image, or click on an item then on its position on the image

Reagent

Submit

Results - Extrinsic defects

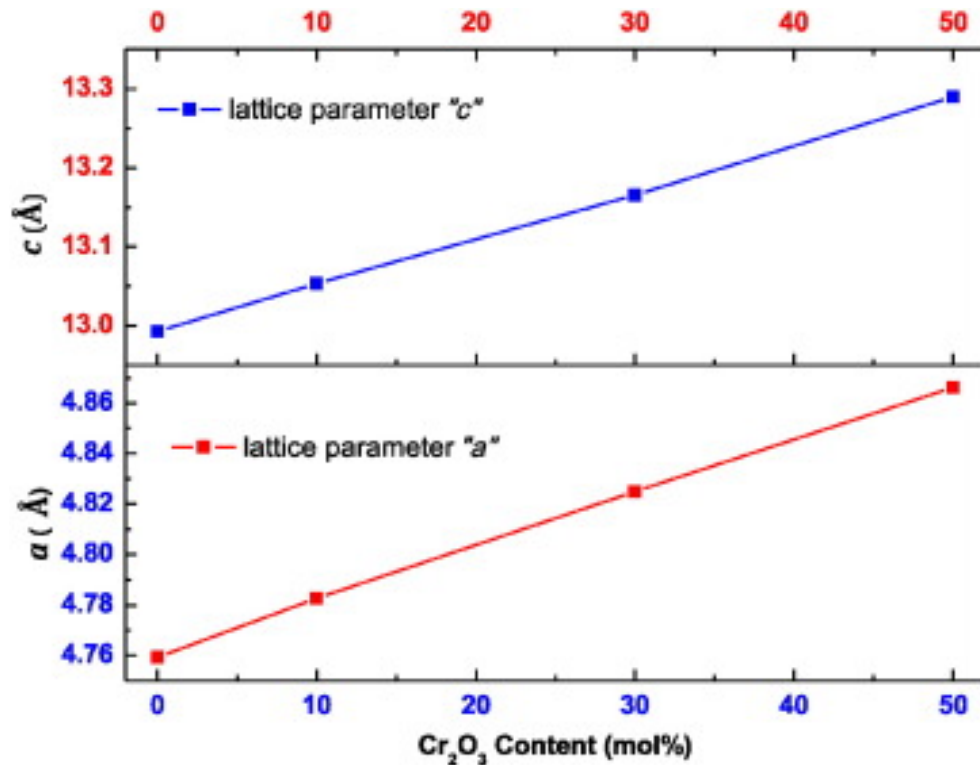
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Solid solutions

Frequently, substitutional defect concentrations can exceed 1%

- known as a 'solid solution'
- Very important for tuning properties *via* synthesis
- Often useful to think of the "average ion" properties at each site
 - e.g. ionic radius, resulting in *Vegard's Law*
 - Lattice parameter is weighted average of the end-members, e.g. $\text{Al}_{2-x}\text{Cr}_x\text{O}_3$:



Non-stoichiometry

Some materials are naturally non-stoichiometric even without extrinsic defects

- Very common in transition metal compounds
 - multiple oxidation states available
- Example: FeO (wustite, NaCl structure) cannot actually form stoichiometrically at ambient pressure
 - Actually Fe_{1-x}O , with $0.05 \leq x \leq 0.15$

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Note: From cation:anion ratio alone you cannot determine the defect type(s)

e.g. Fe:O ratio of 0.9 could equally be $\text{Fe}_{0.9}\text{O}$ or $\text{FeO}_{1.11}$!

Lecture recap

- Crystals are never perfect!
 - defects favoured at higher temperature
- Three main types of defect:
 - vacancy (called Schottky if stoichiometry maintained)
 - interstitial (called Frenkel if stoichiometry maintained)
 - substitution or extrinsic
- Kroger-Vink notation is a way to write defect equations
- Some materials can form solid solutions and/or non-stoichiometric compositions
- If defects order, this can lead to new stoichiometric structure types

Feedback



What did you like or dislike about lecture 2?

Write your answer...



 **Submit**



