



Metal Speciation

ENV-200 Weeks 3 and 4

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We discussed the hard-soft classification scheme and how it allows us to predict the reactivity between metals and ligands.

We used the equilibrium approach to model metal speciation.

We performed an experiment to determine the hardness of water.

Today, we will discuss precipitation and dissolution reactions.

And we will figure out what caused the Flint water crisis!



Metal speciation

Precipitation and dissolution

You should be able to

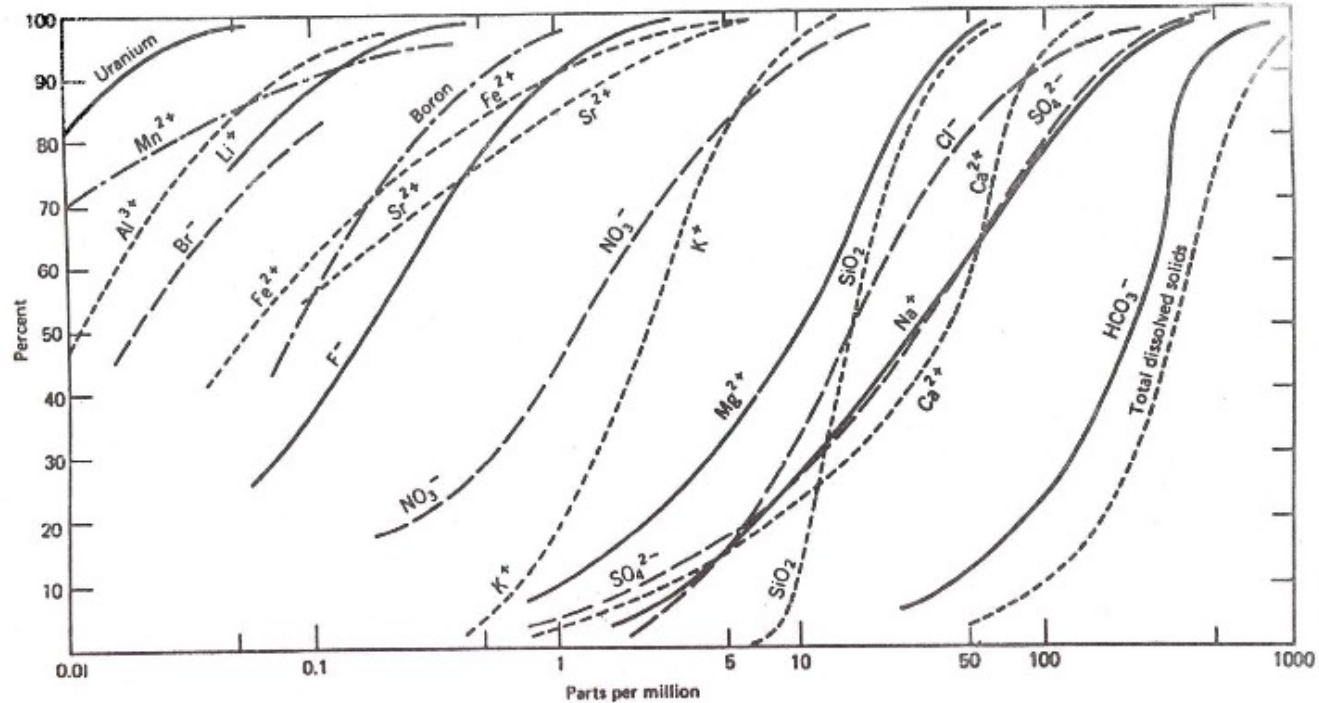
1. formulate solubility equilibria and assess if a solution is undersaturated or oversaturated with respect to a solid phase.
2. determine which solid phases regulate the concentrations of elements in natural waters.
3. assess the solubility of solid phases using log-log diagrams.
4. optimize conditions in engineering applications based on your understanding of metal solubility.

- Water treatment: metal hydroxides are used for coagulation and flocculation processes (e.g., removal of natural organic matter, removal of phosphate, etc.)
- Removal of toxic metals from natural waters: precipitation and sorption to mineral surfaces (especially metal oxides) can remove metals from waters
- Passivation of pipes: formation of mineral passivation layer in water pipes can prevent corrosion

Composition of natural waters

- Waters acquire their chemical characteristics by
 - Dissolution of minerals
 - Chemical reactions with solids, liquids, and gases
- Variation in water composition can be explained by
 - Environmental history of the water
 - Chemical reactions of the rock-water-air system
- First approximation
 - Fresh water: CO_2 of the atmosphere vs. mineral rock
 - Seawater: acid-base titration of acids of volcanoes vs. bases of rocks

Composition of natural waters



- Cumulative curves showing the frequency distribution of constituents in natural waters
- Many constituents show little variation, e.g., dissolved silica: $10^{-3.8}$ to $10^{-3.2}$ M; H^+ : $10^{-6.5}$ to $10^{-8.5}$ M

- Weathering encompasses a variety of processes that decompose rocks
 - Mechanical weathering: fragmentation or loss of materials without chemical change
 - Chemical weathering: chemical reaction of minerals in rocks and soils with acidic and oxidizing substances
- Important for
 - Water composition
 - Soil fertility
 - Biological diversity
 - Agricultural productivity

- Humans have played a big role in increasing weathering
 - Mining & extracting buried fossil fuels increases chemical & mechanical weathering (e.g. acid mine drainage)
 - Agriculture increases chemical weathering due to more plant growth
 - Construction & clearing land for agriculture increases mechanical weathering
- Result: significant quantities of dissolved materials are added to surface water
- Humans have increased the global mechanical weathering rate by a factor of 2
- This results in the accumulation of more sediments in rivers & deltas

Chemical weathering: sequence of mineral stability

<u>Mineral</u>	<u>Reaction equation</u>	<u>Solubility</u>
Rock salt (Halite)	$\text{NaCl (KCl)} \rightleftharpoons \text{Na}^+(\text{K}^+) + \text{Cl}^-$	very good
Gypsum (Anhydrit)	$\text{CaSO}_4 \cdot 2 \text{H}_2\text{O} \rightleftharpoons \text{Ca}^{2+} + \text{SO}_4^{2-} + 2 \text{H}_2\text{O}$	good
Calcite	$\text{CaCO}_3 + \text{H}_2\text{CO}_3 \rightleftharpoons \text{Ca}^{2+} + 2 \text{HCO}_3^-$	intermediate
Dolomite	$\text{CaMg}(\text{CO}_3)_2 + 2 \text{H}_2\text{CO}_3 \rightleftharpoons \text{Ca}^{2+} + \text{Mg}^{2+} + 4 \text{HCO}_3^-$	intermediate
Aluminum silicates		
primary minerals + $\text{H}_2\text{CO}_3 \rightarrow$ Base cations + H_4SiO_4 + HCO_3^- $\text{Al}(\text{OH})_3$ + secondary minerals		small
Quarz	$\text{SiO}_2 + 2 \text{H}_2\text{O} \rightleftharpoons \text{H}_4\text{SiO}_4$	small
Gibbsite	$\text{Al}(\text{OH})_3 + 3 \text{H}^+ \rightleftharpoons \text{Al}^{3+} + 3 \text{H}_2\text{O}$	very small
Goethite	$\text{FeOOH} + 3 \text{H}^+ \rightleftharpoons \text{Fe}^{3+} + 2 \text{H}_2\text{O}$	very small

Weathering rates depend on relative concentrations in rock but in general the order is: $\text{Ca} > \text{Na} > \text{Mg} > \text{K} > \text{Si} > \text{Fe} > \text{Al}$

- Dominant form of chemical weathering: carbonation reaction
 - Plant roots and microbes release CO_2 through respiration
 - $\text{H}_2\text{O} + \text{CO}_2 \rightleftharpoons \text{H}_2\text{CO}_3$
 - Local H_2CO_3 concentrations in soils can be much higher than concentrations for water in equilibrium with the atmosphere
- Plants and microbes further increase weathering by releasing acids that lower pH and can chelate metals (increased dissolution rate, increased solubility, and increased mobility of metals)
- Other factors that influence chemical weathering rate include
 - Rock type (see previous slide)
 - Climate (faster at higher temperature and precipitation)
 - Tropical forests > temperate forests
 - Forests > grasslands > deserts

Incongruent dissolution: only some of the constituents of the primary mineral are dissolved

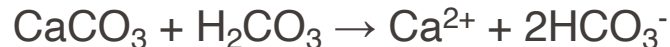
- Example: carbonic acid and silicate rocks



- Primary mineral albite ($\text{NaAlSi}_3\text{O}_8$) is converted to a secondary mineral kaolinite ($\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4$)

Congruent dissolution: no initial solids remain after dissolution

- Example: carbonic acid and limestone



Examples of dissolution reactions

Consider the dissolution of apatite:



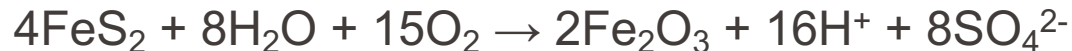
- a. Is the dissolution reaction congruent or incongruent?
- b. Where did we encounter apatite before?

Apatite is the only primary mineral with significant phosphorus content. Phosphorus is often a limiting nutrient for plant growth.



Examples of dissolution reactions

Consider the dissolution of pyrite:



- a. Is the dissolution reaction congruent or incongruent?
- b. Why is the water orange-red?

This oxidative dissolution reaction accounts for the acidity of runoff in many mining operations (acid mine drainage)



- Solubility: maximum concentration of a solute that can dissolve in a solvent at a given temperature.
- Link to hydrolysis: Formation of a precipitate can be considered the final stage in the formation of polynuclear complexes

- Equilibrium equation
- Solubility product K_{s0}
 - index 0: only simple ions are involved
 - solid phases have activity = 1
- Ion activity product IAP
 - product of actual ion activities



$$K_{s0} = \frac{\{M(aq)\}^n \{X(aq)\}^m}{\{M_n X_m(s)\}} = \{M(aq)\}^n \{X(aq)\}^m$$

$$IAP = \{M(aq)\}_{actual}^n \{X(aq)\}_{actual}^m$$

$IAP < K_{s0}$: undersaturation (solid phase cannot precipitate, solid phase that is present will dissolve)
 $IAP = K_{s0}$: in equilibrium
 $IAP > K_{s0}$: oversaturation, solid phase will precipitate

Exercise 1: Solubility of anhydrite and gypsum



Determine the state of saturation of seawater of normal salinity with respect to anhydrite ($\text{CaSO}_4(\text{s})$).

$$\{\text{Ca}^{2+}\} = 2.4 \cdot 10^{-3} \text{ M}$$

$$\{\text{SO}_4^{2-}\} = 1.9 \cdot 10^{-3} \text{ M}$$

$$K_{\text{s}0} = 4.2 \cdot 10^{-5}$$

Until now, we have been using ion concentrations instead of activity. However, concentration is, strictly speaking, only applicable to ideal solutions under standard conditions (no interactions between species). In electrolyte solutions (solutions with $I \neq 0$) this is not the case! Various interactions between species can occur, most importantly electrostatic effects (attraction between ions with opposite charge). These interactions influence the behavior of the ions and don't allow to treat every ion in the solution independently. To accurately do equilibrium calculations in electrolyte solutions, we use ion activity instead of ion concentration to account for this non-ideal behavior under non-standard conditions.

Standard conditions: $T = 25\text{ }^{\circ}\text{C}$, $P = 1\text{ atm}$, $I = 0$
 $I = 0 \rightarrow$ most common source for non-ideal behavior

activity of compound i is indicated by $\{ i \}$
concentration of compound i is indicated by $[i]$

Ionic strength I expresses the concentration of electrolytes (cations and anions) in solution:

$$I = 0.5 \sum C_i z_i^2$$

where C_i = concentration of the ion i and z_i = charge of the ion i

Example 1: 1 mM NaCl:

$$I = 0.5 * (C_{\text{Na}^+} * z_{\text{Na}^+}^2 + C_{\text{Cl}^-} * z_{\text{Cl}^-}^2) = 0.5 * (10^{-3} * 1^2 + 10^{-3} * (-1)^2) = 10^{-3} \text{ M}$$

Example 2: 1 mM CaCl_2 :

$$I = 0.5 * (C_{\text{Ca}^{2+}} * z_{\text{Ca}^{2+}}^2 + C_{\text{Cl}^-} * z_{\text{Cl}^-}^2) = 0.5 * (10^{-3} * 2^2 + 2 * 10^{-3} * (-1)^2) = 3 * 10^{-3} \text{ M}$$

Typical ionic strength of freshwater: 0.004 – 0.02 M

Typical ionic strength of seawater: 0.7 M

Activity coefficient is the relation between activity and concentration:

$$\{i\} = \gamma_i \cdot [i]$$

γ_i = activity coefficient

γ_i is a function of the charge of the ion and the total ionic strength

Several methods exist to calculate γ_i . The most common one is the Davies equation (valid up to $I = 0.5$ M):

$$\log \gamma_i = -A \cdot z_i^2 \left(\frac{\sqrt{I}}{1 + \sqrt{I}} - 0.2 \cdot I \right)$$

$$A = 1.82 \cdot 10^{-6} (\epsilon T)^{-3/2}$$

ϵ = dielectric constant

use $A = 0.5$ for water at 25 °C

$$\text{If } I = 0 \rightarrow \log \gamma_i = 0 \rightarrow \gamma_i = 1 \rightarrow [i] = \{i\}$$

Exercise 2: Ion activity



What are the activity and concentration of Na^+ in a solution of 1 mM NaCl?

Exercise 3: Solubility in natural waters



- Which factors influence metal solubility?
- Does an increase/decrease of these factors have a positive or negative effect on solubility?



Recall general equations for soluble hydroxo species

Soluble hydroxo species:



$$K_n = \frac{\{\text{Me}(\text{OH})_n^{(m-n)+}\}}{\{\text{Me}(\text{OH})_{n-1}^{(m-n+1)+}\}\{\text{OH}^-\}}$$

Or:



$$^*K_n = K_n \cdot K_w = \frac{\{\text{Me}(\text{OH})_n^{(m-n)+}\}\{\text{H}^+\}}{\{\text{Me}(\text{OH})_{n-1}^{(m-n+1)+}\}}$$

We can express reactions both in terms of OH^- (K has no prefix) or H_2O (K has prefix *)

m = valence of the metal; n = number of hydroxides

Equilibrium of metal hydroxide with metal ion:



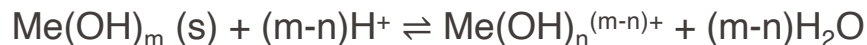
$$K_{s0} = \{\text{M}^{m+}\} \{\text{OH}^-\}^m \text{ or } {}^*K_{s0} = \frac{\{\text{M}^{m+}\}}{\{\text{H}^+\}^m}$$

Equilibrium with soluble hydroxo species:



$$K_{sn} = \{\text{Me}(\text{OH})_n^{(m-n)+}\} \{\text{OH}^-\}^{(m-n)}$$

Or:



$${}^*K_{sn} = \frac{\{\text{Me}(\text{OH})_n^{(m-n)+}\}}{\{\text{H}^+\}^{(m-n)}}$$

Dissolved species: $\{\text{Me}\}_T = \{\text{Me}^{m+}\} + \sum \{\text{Me}(\text{OH})_n^{(m-n)+}\}$

Exercise 4: Multiple solid phases

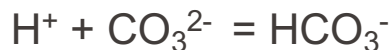


Consider an anoxic water at pH 6.8 and $\{\text{HCO}_3^-\} = 10^{-4} \text{ eq L}^{-1}$ (assume $\gamma_i = 1$).

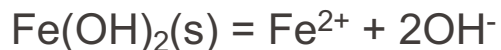
Is the solubility of Fe(II) dominated by siderite ($\text{FeCO}_3(\text{s})$) or $\text{Fe}(\text{OH})_2(\text{s})$?
Note that the solid that gives the lowest concentration of soluble Fe(II) controls the solubility for a given set of conditions.



$$\log K_{s0} = -10.4$$



$$-\log K_2 = +10.1$$

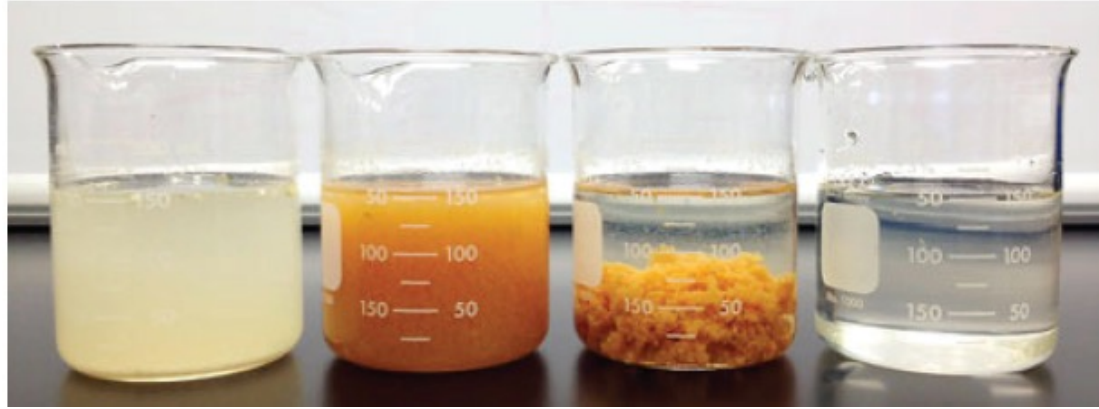


$$\log K_{s0} = -14.5$$



$$-2\log K_w = +27.8$$

Exercise 5: Flocculation



Flocculation is a process by which a chemical coagulant added to the water acts to facilitate bonding between particles, creating larger aggregates which are easier to separate. The method is widely used in water treatment plants.

Example by Urs Von Gunten

Exercise 5: Flocculation



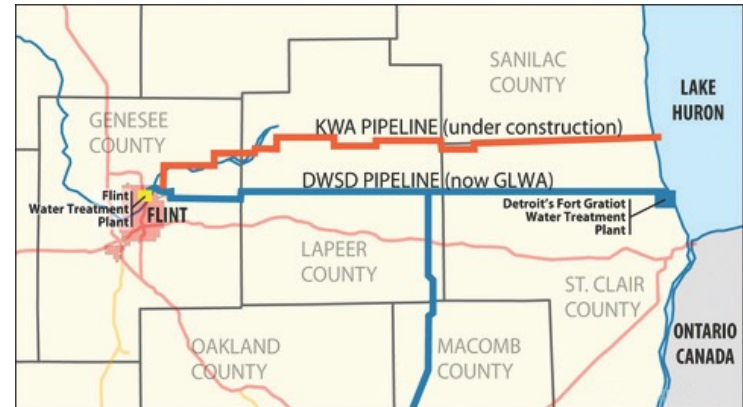
Aluminium is often used as a flocculant in water treatment. To achieve the best results, the pH of minimum solubility should be used. To identify the pH of minimum solubility, sketch the aluminum speciation as a function of pH in a log-log plot.



Environmental engineering challenge: Flint water crisis



Why did the switch to Flint's river water cause this catastrophe?



Images: *The Flint Water Project*; Map: Regina H. Boone/Detroit's Free Press/Zuma Press

Phosphate (PO_4^{3-}) was added to precipitate all metal cations and inhibit corrosion in the Detroit plant.

- a. What reactions can phosphate (use Na_3PO_4) undergo with $\text{FeCl}_2(\text{aq})$ and $\text{Pb}(\text{NO}_3)_2(\text{aq})$ in solution?
- b. What effect does the addition of PO_4^{3-} have on pH?

Flint's water supply was switched to the city's own water treatment plant on the Flint River. Phosphate was NOT added to the Flint River water in the new plant.

- a. What effects does this have on the passivation layer in the pipes and why?
- b. In the absence of a passivation layer, what other chemical reactions can occur between the water and metals in the pipes?

- The composition of natural waters is in part dependent on the history of mineral dissolution reactions.
- Chemical weathering describes the chemical reaction of minerals with acid and oxidizing substances.
- A comparison between the ion activity product and the solubility product of a solid phase allows us to estimate whether or not the solid phase will form under given conditions.
- The solubility of metals in natural waters is affected, amongst other factors, by temperature, salinity, pH, and complexation.
- The Flint water crisis was caused by a shift in water chemistry that led to the dissolution of the mineral passivation layer on the pipes, thereby exposing the underlying Pb and Fe.