## Unit 1

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## 1.1 Lecture 1: Introduction/Key Trends

3/29: • Largely asynchronous, but synchronous discussions, office hours, and tests.

- Refer to the Canvas site for all information; it's the class hub.
- To understand transition metal trends and properties, start with *atomic* properties and trends.
- Electronegativity: The energy that an atom will gain when it gains an electron.
  - Fluorine really wants to gain an electron; thus, it has high electronegativity.
    - Do you mean electron affinity?
  - Increases across a row; decreases down a column.
    - Transition metals, in general, are an exception to this rule.
    - This is because of the lanthanide contraction.
  - Discontinuities in the transition metals (Mn and Ni) correspond to half-filled and filled d shells, respectively.
    - Extra stability means less of a desire to gain an electron.
- **Ionization potential**: The energy required to remove an electron.
  - Varies with the identity of an element and its oxidation state.
  - Increases across a row; decreases down a column.
- Size:
  - Many different definitions (depending on the specific thing you're interested in, one may be more useful than another). For example,
    - Atomic radius: Specific to an element.
    - Ionic radius: Specific to an oxidation state; as in salts and coordination complexes.
    - Covalent radius: Distance that one would expect for a bond; varies with bond order.
  - Decreases across a row; increases down a column (a notable exception to the latter trend follows from the lanthanide contraction).
  - Things that affect size:
    - Oxidation state.
    - Spin state (high spin [larger; this is because the  $e_g$  orbitals are antibonding, and antibonding electrons both push the bounds of the atom and weaken bonds, increasing the covalent radius] vs. low spin [smaller]).

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• Lanthanide contration: In the transition metals, there is a small/no increase in size between the second and third rows.

- This is because the shell added in between contains the f orbitals, which are small, do not extend past the noble gas core, and do not provide good shielding.
- Z goes up a lot with little shielding, so the 5d orbitals are contracted; thus, 4d/5d orbitals are similar in size.
- Oxidation state: The number of electrons a metal center is below its valence.
  - Typically, the maximum oxidation state is defined by the d-count for the 0-valent metal.
- Having discussed four trends, how are they related?
  - As oxidation state increases, "electronegativity" and ionization potential will increase, and the radius will decrease.
    - This is because removing an electron  $\Rightarrow$  reduces shielding  $\Rightarrow$  higher positive charge  $\Rightarrow$  all orbitals decrease in energy  $\Rightarrow$  all orbitals decrease in size (hence radius decreases, too).
    - Watch out for discontinuities such as Mn<sup>2+</sup>.
- $\bullet$  Magnetic properties: Unique to the transition metals and the f block.
  - Consider  $\operatorname{Fe}^{\operatorname{II}} \operatorname{L}_{6}^{2+} (d^{6})$ .
  - Possible states: Low spin (S = 0), intermediate spin (S = 1; rare), and high spin (S = 2).
  - We predict which state dominates by the:
    - Pairing energy.
    - Ligand field stabilization energy.
- Pairing energy: The energy cost of putting two electrons in the same orbital. Also known as PE.
  - Trends with orbital size/radius.
  - Decreases down a column.
- Ligand field stabilization energy: Also known as LFSE.
  - Can be thought of in terms of crystal field theory.
    - Extra thoughts on Figure VI.10 of Labalme (2021): Donating negative charge to a free metal ion in a spherically symmetric fashion uniformly raises the energy of the *d* orbitals by increasing repulsions and size.
    - Low-spin LFSE:  $6 \cdot -4 \operatorname{Dq} + 3 \operatorname{PE} = -24 \operatorname{Dq} + 3 \operatorname{PE}$ .
    - Intermediate-spin LFSE:  $5 \cdot -4 \text{ Dq} + 1 \cdot 6 \text{ Dq} + 2 \text{ PE} = -14 \text{ Dq} + 2 \text{ PE}$ .
    - High-spin LFSE:  $4 \cdot -4 \operatorname{Dq} + 2 \cdot 6 \operatorname{Dq} + 1 \operatorname{PE} = -4 \operatorname{Dq} + 1 \operatorname{PE}$ .
    - Thus, the energy difference between the low-spin and high-spin configurations is  $20 \,\mathrm{Dq} + 2 \,\mathrm{PE}$ . It follows that if  $10 \,\mathrm{Dq} > 1 \,\mathrm{PE}$ , then the complex will be low spin; and if  $10 \,\mathrm{Dq} < 1 \,\mathrm{PE}$ , then the complex will be high spin.
    - This also explains why the intermediate spin state is rare: if  $\Delta_o$  is large enough to make  $10 \,\mathrm{Dq} > 1 \,\mathrm{PE}$ , then it will likely take the complex all the way to a low-spin configuration (and vice versa for high spin).
    - Lastly, this means that Fe<sup>II</sup> is a good **spin-crossover** ion.
- Spin-crossover (ion): An ion that can have both high- and low-spin states.
  - The graph of the magnetic moment  $\chi T$  of Fe<sup>II</sup> vs. temperature T (see Figure 1.1) moves from S=0 at the bottom left to S=2 at the top right.
  - $-d^2$  ions are never spin-crossover ions: -8 Dq + 0 PE for high spin vs. -8 Dq + 1 PE for low spin.

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Figure 1.1: Magnetic moment vs. temperature for the Fe<sup>II</sup> ion.

- Dq and PE values depend on:
  - Ligand field strength.
    - $\Delta O_h$  increases as  $\sigma$  donation increases.
    - $\blacksquare$   $\Delta O_h$  increases as  $\pi$  acceptance increases.
    - $\Delta O_h$  decreases as  $\pi$  donation increases.
    - To what extent do we need to have the spectrochemical series memorized?
  - Metal center.
    - Larger, more diffuse metals (i.e., second- and third-row transition metals) have better overlap with the ligands, giving rise to larger  $\Delta O_h$ .
    - Note that pairing energy decreases in second- and third-row transition metals (due to the larger orbitals).
    - These two factors imply that second- and third-row transition metals are almost always low spin.
  - Oxidation state.
    - As oxidation state increases,  $\Delta O_h$  increases (due to better energy matching, higher "electronegativity," the role of electrostatics, and the electron configuration [ $d^5$  is almost always high spin, and  $d^6$  is often low spin]).
    - Why do  $d^5$  and  $d^6$  exhibit the above behavior? Shouldn't the stabilized orbitals be split more?
    - See the below spectrochemical series for metals.
  - Geometry.
    - In a  $\sigma$ -only sense, lower coordination numbers tend to have smaller LFSEs.
    - $\blacksquare T_d < C_{4v} \approx D_{3h} < O_h < D_{4h}.$
    - $\bullet$   $\Delta_{\text{sq. pl.}} \approx 1.74 \, \Delta O_h$ .
- Spectrochemical series for metals (not as precise as the one for ligands, but a decent approximation):

$$\mathrm{Mn^{2+} < Ni^{2+} < Co^{2+} < Fe^{2+} < V^{2+} < Fe^{3+} < Co^{3+} < Mn^{4+} < Mo^{3+} < Rh^{3+} < Ru^{3+} < Pd^{4+} < Ir^{3+} < Ir^$$

- Hard/soft acid-base theory:
  - Common Lewis acids:
    - $\blacksquare$  Proton:  $H^+$ .
    - Molecules with no octet: AlCl<sub>3</sub>, BR<sub>3</sub> (boranes), BeH<sub>2</sub>.
    - Metal cations: Na<sup>+</sup>, Ti<sup>4+</sup>.
    - $\blacksquare$   $\pi$  acids: CO<sub>2</sub>, CO, PR<sub>3</sub>.
  - Common Lewis bases:
    - $\blacksquare$  Carbanions: CR<sub>3</sub>.

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- $\blacksquare$  Hydrides: KH, NaH, LiAlH<sub>4</sub>.
- Amines, amides, and phosphines: NH<sub>3</sub>, PR<sub>4</sub>, NH<sub>2</sub><sup>-</sup>.
- $\blacksquare$  OH<sub>2</sub>, SR<sub>2</sub>, OH<sup>-</sup>.
- $\blacksquare$  Halides: F<sup>-</sup>, Cl<sup>-</sup>, Br<sup>-</sup>.
- Carbonyl: CO (means CO is amphoteric).
- $\blacksquare$  Olefins:  $C_2H_4$ .
- Distinguishes hard vs.  ${\rm soft}^{[1]}.$

<sup>&</sup>lt;sup>1</sup>Hard vs. soft is the basis for the solubility rules!