

## Topic IX

# Reactions and Mechanisms

### IX.1 Module 46: Ligand Substitution in Octahedral Complexes

3/8:

- Suggested reading: Chapter 12.
- Having learned about structure, isomerism, and electronic structure, let's now talk about reactivity.
- What makes transition metals unique is the reactions of the metal center (i.e., the first coordination sphere).
- **Substitution reaction:** A reaction where a ligand in the first coordination sphere is exchanged.
- Factors that affect the rate of substitution:
  - Role of the entering group.
  - Role of the leaving group.
  - Nature of the other ligands in the complex.
  - Effect of the metal center.
- Stable/unstable are thermodynamic terms, and labile/inert are kinetic terms.
  - $[\text{Co}(\text{NH}_3)_6]^{3+}$  is unstable but inert wrt. aquation (large  $K_{\text{eq}}$ , but slow to react).
  - $[\text{Ni}(\text{CN})_4]^{2-}$  is stable but labile wrt. exchange (small  $K_{\text{eq}}$ , but equilibrium is established quickly).
- **Potential energy landscape:** A thermodynamic representation of a reaction by a (potentially multidimensional) potential energy surface along its reaction coordinate.
- Thermodynamics and kinetics are two different things, but they can be related.
- Potential energy landscapes with small  $\Delta G$  and large  $\Delta E_A$  are slow to react.
  - Increasing  $\Delta G$  (perhaps by destabilizing the reactant) can speed up the reaction.
  - Increasing  $\Delta G$  often occurs at the expense of  $\Delta E_A$  (the activation energy literally decreases as the free energy change increases).
- Consider the reaction  $[\text{Co}(\text{NH}_3)_5\text{X}]^{2+} + \text{H}_2\text{O} \longrightarrow [\text{Co}(\text{NH}_3)_5(\text{H}_2\text{O})]^{3+} + \text{X}^-$ .
  - The more bulky/less electronegative the X-type ligand, the higher the equilibrium constant and reaction rate.
- But how do we adequately compare different reactions when so many parameters are entangled together?

- One good way to compare kinetics of transition metals is with water exchange rate constants (light/heavy water being exchanged in metal coordination spheres), since this eliminates thermodynamic consternation.
  - There is massive variation among the metals.
  - The fastest (the alkali metals) run up against the **diffusion limit**.
  - Metals with larger atomic radii react more quickly (the alkali and alkaline earth metals follow this pattern nicely, but the transition metals are all over the place).
    - Group 1A: As we go down the group, the cations are getting larger and the charge density decreases, so the  $M^+ - OH_2$  bond is getting weaker and more easily broken.
    - Group 2A: The charge density is larger (doubly charged) so the strength of the bond is greater so the rate of exchange is slower.
  - We define such a reaction as having a labile half life if the half life is less than 1 minute; otherwise, it is inert.
- **Diffusion limit:** At most, one molecular collision can result in one successful reaction; thus, reaction rate is bounded above by how many molecular collisions can physically occur in a given period of time at a given temperature.
- You cannot prove mechanisms; you can only disprove other plausible scenarios. *Also known as diffusion limitation.*
- Possible mechanisms for  $ML_5X + Y \longrightarrow ML_5Y + X$  where  $X = H_2O$  and  $Y = \text{anion}$  or vice versa:
- Associative (A): Via a 7-coordinate intermediate.



- Corresponds to organic  $S_N2$ .
- First step (RDS) is slow since the incoming Y causes steric hindrance.
- The transition state is either a monocapped octahedron or pentagonal bipyramidal, depending on how the Y attacks.
- Applying a steady-state approximation for  $[ML_5XY]$ , we have

$$\frac{d[ML_5XY]}{dt} = 0 = k_1[ML_5X][Y] - k_{-1}[ML_5XY] - k_2[ML_5XY]$$

- We can solve the above equation for  $[ML_5XY]$ :

$$[ML_5XY] = \frac{k_1[ML_5X][Y]}{k_{-1} + k_2}$$

- Substituting the above into the rate constant equation for the fast step will give us the rate law in terms of the reactants (where  $k = \frac{k_1 k_2}{k_{-1} + k_2}$ ):

$$\begin{aligned} \text{Rate} &= k_2[ML_5XY] \\ &= \frac{k_1 k_2 [ML_5X][Y]}{k_{-1} + k_2} \\ &= k[ML_5X][Y] \end{aligned}$$

- Thus, the  $S_N2$  mechanism is second-order overall, but first-order in both reactants.

- Dissociative (*D*): Via a 5-coordinate intermediate.



- Corresponds to organic  $\text{S}_{\text{N}}1$ .
- First step (RDS) is slow since its spontaneous elimination of a ligand.
- The transition state is generally square pyramidal, but if it is sufficiently long-lived, it can reorganize itself into the trigonal bipyramidal state.
- The dissociative mechanism predicts that the rate of the overall substitution reaction depends on only the concentration of the original complex  $[\text{ML}_5\text{X}]$ , and is independent of the concentration of the incoming ligand  $[\text{Y}]$ .
- Thus, the overall rate law is

$$\text{Rate} = k_1[\text{ML}_5\text{X}]$$

- We can also derive this with an analogous kinetic analysis to that used for the *A* mechanism, the only difference being that we simplify  $\frac{k_1 k_2 [\text{ML}_5\text{X}][\text{Y}]}{k_{-1}[\text{X}] + k_2[\text{Y}]}$  to the rate law by noting that  $k_2 \gg k_{-1}$  or  $[\text{Y}] \gg [\text{X}]$ .
- Interchange (*I*): As Y begins to bond, X begins to leave, i.e., the bond making to Y and bond breaking to X occur simultaneously.



- Corresponds to organic  $\text{S}_{\text{N}}2$ .
- This is how the majority of ligand substitutions occur.
- It is too simplistic to assume that a first-order rate law implies *D* and a second-order rate law implies *A*. Indeed, most substitution reactions probably involve a mechanism like this, i.e., one that is intermediate between these two extremes.
- In an interchange mechanism, the intermediate involves an association between the original  $\text{ML}_5\text{X}$  complex and the attacking Y ligand. The Y ligand remains outside the coordination sphere of  $\text{ML}_5\text{X}$ , unlike the  $\text{S}_{\text{N}}2$  mechanism, so the intermediate is not seven coordinate. However, it can help weaken the M–X bond.
- Assuming high  $[\text{Y}] \approx [\text{Y}]_0$ , it can be shown that the rate is given by

$$\text{Rate} = \frac{k_2 K_1 [\text{M}]_0 [\text{Y}]_0}{1 + K_1 [\text{Y}]_0}$$

where  $[\text{M}]_0 = [\text{ML}_5\text{X}]_0 + [\text{ML}_5\text{X} \cdot \text{Y}]$  and  $[\text{Y}]_0$  are initial conditions and  $K_1 = \frac{k_1}{k_{-1}}$  is the equilibrium constant for the RDS reaction.

- At high  $[\text{Y}]$  and  $K_1[\text{Y}]_0 \gg 1$ , the rate is first-order in  $[\text{M}]_0 \approx [\text{ML}_5\text{X}]$ .
- At lower  $[\text{Y}]$ , the rate is second order.
- At some point, the kinetic analysis becomes essentially worthless. Indeed, although we speak generally about associative and disassociative reaction mechanisms, the terms *A* and *D* are reserved for situations where 7- and 5-coordinate intermediates have actually been isolated and positively identified. If no intermediates have been isolated or identified, the designations  $I_d$  and  $I_a$  are more appropriate.
  - Two minor variations on the *I* mechanism are  $I_d$  (dissociative interchange) and  $I_a$  (associative interchange).

- If breaking the M–X bond is more important, the mechanism is  $I_d$ .
- If bond formation between  $ML_5X$  and Y is significant, the mechanism is  $I_a$ .
- The difference between  $I_d$  and  $I_a$  is subtle and does not necessarily correspond to whether the observed rate law is first or second order.
- If the rates of a series of comparable substitution reactions are most sensitive to the identity of X, the leaving ligand, then the mechanism is more probably  $I_d$ , and vice versa for  $I_a$ .
- For example, the rate constants for the anation of  $[Cr(NH_3)_5(H_2O)]^{3+}$  by various ligands vary very little. Thus, it is probably  $I_d$ . However, the rate constants for the anation of  $[Cr(H_2O)_6]^{3+}$  by various ligands vary by three orders of magnitude. Thus, it is probably  $I_a$ .
- Solvent (e.g., water) effects in substitution reactions:
  - Many substitution reactions occurring in solvent water may have first-order kinetics regardless of whether their initial steps are primarily  $D$  or  $A$ .
  - For example, this occurs if aquation is a precursor RDS.
  - If  $D$ ,  $\text{Rate} = k[ML_5X]$ .
  - If  $A$ ,  $\text{Rate} = k[ML_5X][H_2O] = k'[ML_5X]$ .
    - The latter equality is valid since water has constant concentration<sup>[1]</sup>.
  - Both mechanisms lead to apparent first-order kinetics.
- Steric factors favoring  $D$  or  $I_d$ :
  - For most octahedral complexes, steric factors inhibit formation of a  $CN_7$  intermediate, which suggests a dissociative mechanism ( $D$  or  $I_d$ ) is more plausible.
    - Even cases showing second-order kinetics may not be  $A$  for this reason.
  - For example, aquation of ammine-halides is second order with a first-order dependence on  $[OH^-]$ .
  - If  $k_2 \gg k_1$ , the rate is approximately  $\text{Rate} = k_2[Co(NH_3)_5X]^{2+}[OH^-]$ .
  - Calls for an alternate mechanism called  $S_N1CB$ , where ligands with lower energy bond as conjugate bases and those with higher  $\Delta_0$  (hence CB).
- Kinetically analyzes  $S_N1CB$ .

## IX.2 Module 47: Substitution and Ligand Field Stabilization Energy

3/10:

- Factors that affect the rate of substitution:
  1. Role of the entering group.
  2. Role of the leaving group.
  3. The nature of the other ligands in the complex.
  4. Effect of the metal center.
- Six factors that support a dissociative mechanism:
  1. The rate of reaction only slightly (within a factor of 10) changes on the incoming ligand.
    - Means that the ligand is not significantly involved in the RDS.
  2. Making the charge on complex more positive decreases the rate of substitution.

<sup>1</sup>Is this what we meant when we said in Labalme (2020) that liquids and solids don't have active mass? In other words, the species should be included in the mass-action expression; they just simply get lumped in with the rate constant generally.

- Increasing the charge increases the bond strength, making it harder for one ligand to spontaneously break away.
- 3. Steric crowding increases reaction rate.
- 4. The volume of activation is positive and reaction rate decreases at high pressure.
  - Especially if the reactant is a gas, for example.
- 5. Reaction rate correlates with Ligand Field Activation Energy (LFAE) predictions.
- 6. Stereochemistry can give additional hints.
  - A messy subject unless you use very well crafted multidentate ligands.
- Simple changes in oxidation state can have massive effects on water exchange rate constants<sup>[2]</sup> (e.g.,  $\text{Cr}^{2+}$  to  $\text{Cr}^{3+}$  results in a shift of 15 orders of magnitude).
- Variations in complex reactivity:
  - Most first row transition metals are labile<sup>[3]</sup>, but  $\text{Cr}^{3+}$  ( $d^3$ ,  $S = \frac{3}{2}$ ) and low-spin  $\text{Co}^{3+}$  ( $d^6$ ,  $S = 0$ ) are usually inert.
  - $d^7$ - $d^{10}$ , with filling of  $e_g^*$  levels, are labile.
    - These configurations tend to have large Jahn-Teller distortions and/or low CFSEs.
    - $d^7$ ,  $d^9$ , and  $d^{10}$  cases are more labile than  $d^8$ .
    - $d^8$  has a  $^3A_{2g}$  ground state, which is immune to Jahn-Teller distortion.
    - With strong-field ligands,  $d^8$  may be square planar, often being inert.
  - Inert octahedral complexes tend to be those with high CFSE; viz.,  $d^3$  low-spin  $d^4$ - $d^6$ .
    - $\text{ML}_6$  complexes of both  $d^3$  ( $^4A_{2g}$ ) and low-spin  $d^6$  ( $^1A_{1g}$ ) are immune from Jahn-Teller distortions and therefore can be perfect  $O_h$ .
    - $\text{ML}_6$   $d^3$  has  $\text{CFSE} = -\frac{6}{5}\Delta_o$  and low-spin  $d^6$  has  $\text{CFSE} = -\frac{12}{5}\Delta_o + 3P$ .
- The most inert complexes among the first row transition metals:
  - $\text{Co}^{3+}$  is primarily dissociative, but with dependence on the incoming ligand  $I_d$ .
  - $\text{Cr}^{3+}$  has a dimorphism where it can be  $I_d$  or  $I_a$ !
    - For example, it is  $I_d$  in the complex  $[\text{Cr}(\text{NH}_3)_5(\text{H}_2\text{O})]^{3+}$  but  $I_a$  in the complex  $[\text{Cr}(\text{H}_2\text{O})_6]^{3+}$ .
- Water exchange rate constants are greatest for Jahn-Teller distorted species.
  - They are smallest for  $d^{3,5,8}$ .
- Variations in complex reality: Explained by the potential energy landscape for a substitution reaction  $\text{ML}_6 \longrightarrow \text{ML}_5\text{Y}$ :
  - $O_h$  to  $C_{4v}$  to “ $O_h$ .”
  - The activated complex is square pyramidal.
  - We can now use the AOM to calculate the LFSE in both the initial  $O_h$  state and the transition  $C_{4v}$  state.
    - The difference between them will be the **ligand field activation energy** or **LFAE**. The LFAE is not particularly useful because it predicts negative activation energies, which are not a thing, but it does allow us to rationalize reactivity trends.
    - If  $\text{LFAE} \leq 0$ , the reaction is labile; otherwise, it is moderate or slow.

<sup>2</sup>We pronounce the units of rate constants, “reciprocal seconds.”

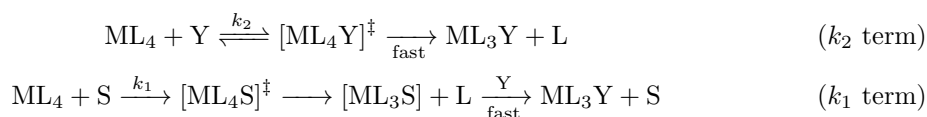
<sup>3</sup>Werner was only able to make his conclusions about coordination chemistry because he happened to be working with the relatively inert  $\text{Co}^{3+}$  complexes.

## IX.3 Module 48: Ligand Substitution in Square Planar Complexes

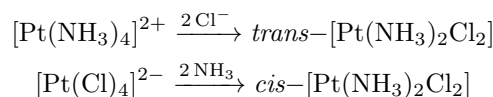
- Although steric factors favor *D*-type mechanisms for octahedral complexes, square planar  $ML_4$  complexes are not so inhibited.
  - For square planar  $ML_4$  complexes, an associative (*A*) mechanism, in which a coordination number 5 (CN5) intermediate is formed, is plausible.
- The  $d^8$  metals are especially likely to form square planar complexes.
- Substitution of square planar complexes, such as  $PtLX_3$ , leads to *trans* and *cis* isomers.
- The rate law is

$$\text{Rate} = k_1[PtLX_3] + k_2[PtLX_3][Y^-]$$

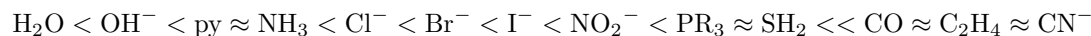
- Suggests two paths, where the first term may be pseudo-first-order due to excess solvent acting as an attacking group.
- If we make L bulkier, we would expect the *D* pathway to dominate ( $k_1$  increases,  $k_2$  decreases).
- However, both  $k_1$  and  $k_2$  decrease with increasingly bulky ligands, suggesting an associative mechanism:



- S means the solvent.
  - Since  $[S] \gg [ML_4]$ , we see pseudo-first order kinetics in the  $k_1$  term in the rate law.
- In square planar complexes, *d*-orbital splitting yields a  $d_{x^2-y^2}$  LUMO.
  - However, it points exactly toward the ligands, so you cannot fill it without displacing a ligand.
  - Indeed, since ligands cannot attack it, they instead attack the higher-lying metal  $p_z$  orbital.
- Note that the stereochemistry stays the same under these reactions.
- An example of the *trans* effect:



- The first ligand takes any of the four spots, but the second substitutes opposite the stronger *trans*-directing ligand.
  - In the above example, chloride is a stronger *trans*-directing ligand than amine groups.
- The *trans* effect:
  - The ratio of *trans* and *cis* isomers is found to vary with the ability of L to act as a *trans*-directing ligand.
  - The increasing order of *trans*-directing ability is



- This ranking comes from a combination of the following two factors.
  - The effect is kinetic rather than thermodynamic.

- Two factors explain the trans effect (both aim at lowering  $\Delta G^\ddagger$ ):
  1. Weakening of the Pt–X bond *trans* to the directing ligand ( $\sigma$ -donor effects).
    - Destabilizes the ground state of the reactants.
    - Pt–A is influenced by *trans*-Pt–T bond because both share Pt  $p_x$  and  $d_{x^2-y^2}$  orbitals. When the Pt–T bond is strong, electron density on those orbitals is shifted away from the Pt–A bond.
    - The more polarizable the L ligand, the better *trans*-director it is; e.g.,  $\text{I}^- > \text{Br}^- > \text{Cl}^-$ .
      - The *trans*-directing ligand polarizes the metal ion, inducing a slight repulsion with the negative electron density on the leaving ligand in the *trans*-position.
    - For example, the Pt–Cl bond is longer in *cis*-[Pt(PMe<sub>3</sub>)<sub>2</sub>(Cl)<sub>2</sub>] than in the *trans* form because the stronger  $\sigma$ -donor ability of PMe<sub>3</sub> weakens the opposite Pt–Cl bond in the *cis* form.
  2. Stabilization of the presumed CN5 intermediate ( $\pi$ -acceptor effects).
    - Stabilizes the activated complex.
    - The strongest *trans*-directors are good  $\pi$ -acceptor ligands.
    - Assuming an *A* mechanism, substitution involves a trigonal bipyramidal transition state.
    - The *trans* intermediate (activated complex) is more favorable for  $\pi$ -acceptor ligands because it permits  $\pi$ -delocalization in the trigonal plane.
      - When the T-ligand engages in a strong  $\pi$ -backbonding, charge is removed from Pt, making the metal center more electrophilic and stabilizing the TBP intermediate.
- The trans effect has applications to synthesis (synthesizing *cis* versus *trans* square planar compounds).
  - For example, *cis*-[Pt(NH<sub>3</sub>)<sub>2</sub>(Cl)<sub>2</sub>] is cisplatin, the first anti-cancer drug.

## IX.4 Module 49: Redox Reactions and Marcus Theory

- 3/12:
- Considers electron transfer (REDOX reaction) of  $[\text{Cr}(\text{H}_2\text{O})_5\text{Cl}]^{3+} + \text{Cr}^{2+} \longrightarrow \text{Cr}^{3+} + [\text{Cr}(\text{H}_2\text{O})_5\text{Cl}]^{2+}$ .
  - Changing the ligands in the coordination sphere affects the rate constant, sometimes dramatically.
  - **Self-exchange reaction:** A REDOX reaction of a compound with itself.
    - These are good model systems, eliminating other thermodynamic factors the same way water exchange reactions do.
  - If we run a self-exchange reaction, radiolabel the coordination centers of one oxidation state, and precipitate the centers of one of the oxidation states after some amount of time, we can perform analyses on how much of what center precipitated (the ratio of labeled  $\text{M}^{n+}$  to  $\text{M}^{m+}$ ).
  - Conclusions (wrt.  $^*\text{Fe}^{\text{III}}(\text{ClO}_4)_3 + \text{Fe}^{\text{II}}(\text{ClO}_4)_2 \longrightarrow ^*\text{Fe}^{\text{II}}(\text{ClO}_4)_2 + \text{Fe}^{\text{III}}(\text{ClO}_4)_3^{[4]}$ ):
    1. Reaction half life is approximately 20 seconds.
    2. Reaction order is 2.
    3. The free energy of activation is approximately  $33 \text{ kJ mol}^{-1}$ .
      - This is derived by recording the rate constant at different temperatures and extracting  $\Delta G^\ddagger$  by running the data through the Arrhenius equation.
  - Rationalizing the results:
    - $\text{Fe}^{2+}$  and  $\text{Fe}^{3+}$  are different in size and bond length.

<sup>4</sup>The starred iron is radiolabeled, i.e., a heavy, radioactive isomer.

- Electron transfer relaxes bond length but costs energy, making the transition state much greater than zero.
  - Essentially, the electron transfer (assumed instantaneous) will destabilize the system, so the solvent will have to reorganize around the coordination complex to lower the energy.
  - Solvent molecules will become random around the reduced complex and oriented around the oxidized one.
- **Reorganization energy:** The key parameter to understanding electron transfer kinetics. *Also known as  $\lambda$ .*

$$\lambda = \lambda_{\text{inner}} + \lambda_{\text{outer}}$$
  - $\lambda_{\text{inner}}$  is the reorganization of bond lengths, angles, the spin state, etc. within the coordination compound.
  - $\lambda_{\text{outer}}$  is the reorganization of the solvent molecules.
  - $\lambda$  is used in many areas of chemistry (it's a foundational concept).
- An example of  $\lambda_{\text{inner}}$ :
  - $\text{Co}^{2+} \longrightarrow \text{Co}^{3+}$  converts from high spin  $d^7$  to low spin  $d^6$ ; since this is significant reorganization of the metal center, self-exchange is slow.
  - $\text{Ru}^{2+} \longrightarrow \text{Ru}^{3+}$  converts from low spin  $d^7$  to low spin  $d^6$ ; since this is minor reorganization of the metal center, self-exchange is fast.

## IX.5 Module 50: Outer- and Inner-Sphere Electron Transfers

- Henry Taube (UChicago) built the foundation for inner-sphere electron transfers.
- Rudy Marcus (CalTech) built the foundation for outer-sphere electron transfers.
- The activation energy for a redox reaction is  $\Delta G^\ddagger = \Delta G_t^\ddagger + \Delta G_i^\ddagger + \Delta G_s^\ddagger$ , where
  - $\Delta G_t^\ddagger$  relates to bringing molecules close;
  - $\Delta G_i^\ddagger$  relates to vibrational energy (reorganization);
  - $\Delta G_s^\ddagger$  relates to solvent reorganization.
- Marcus studies these reactions in terms of weak electronic interactions, where there is no mixing of the reactant and product states.
  - No mixing, molecular orbitals, etc.; instead, electrons transfer from one well-defined state in one molecule to another in another molecule.
- Under these terms, electron transport (to other molecules) can be formulated in terms of Fermi's Golden Rule.
 
$$k_{\text{et}} \propto \rho[\langle \Psi_1 | P_{1 \rightarrow 2} | \Psi_2 \rangle]^2$$
  - Relating back to the Arrhenius equation,  $\rho$  describes the free energy of activation using the **Franck-Condon principle**.
  - In the Arrhenius equation,  $A$  is a pre-exponential factor that combines attempt frequency (approximately  $10^{13} \text{ s}^{-1}$  if no reorganization occurs) and  $x$ -transmission coefficient (the probability that the system crosses the activation barrier as it approaches the top).
- **Franck-Condon principle:** If we have two levels near in energy, they will randomly fluctuate and match up in energy every once in a while; electron transfers occur when they match up.
- We now consider the kinetics of the transfer from the reaction coordinate perspective.



- Potential energy curves of reactants and products by parabolas.

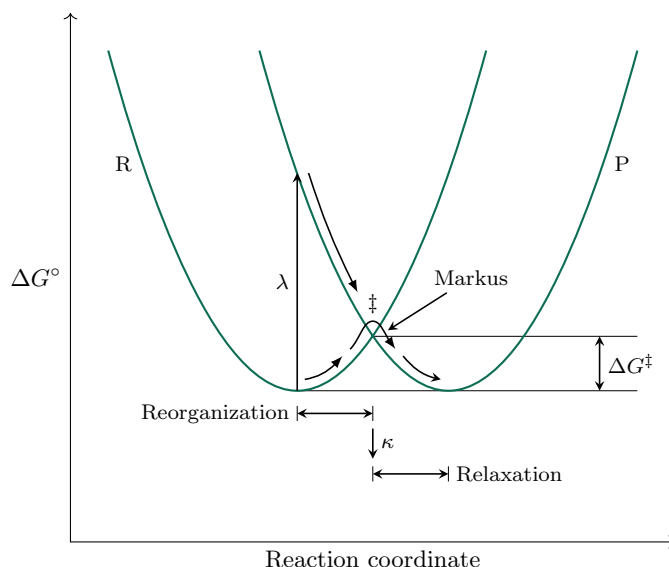
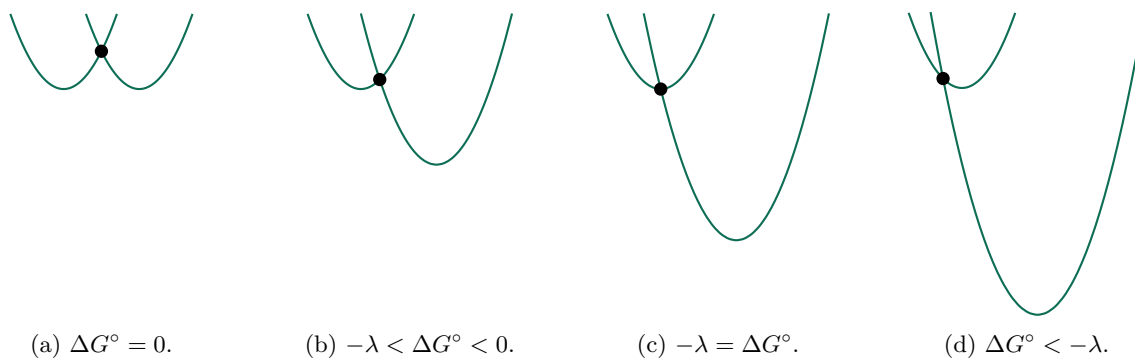


Figure IX.1: Potential energy description of an electron-transfer reaction with  $\Delta G^\circ = 0$ .

- Parabolas come from the representation of potential energy as a function of separation for a harmonic oscillator (if everything's oscillating, classical mechanics predicts a parabolic energy curve).
- The  $x$ -axis (reaction coordinate) represents the change in the geometry of the reactants and products.
- Lieche predicts that the activation energy will be the vertical jump  $\lambda$  from the reactants parabola to the products parabola, and then the energy will fall back down.
  - Conversely, Markus asserts that the reactants will reorganize first to the optimal transfer point (where both parabolas process). This reduces  $\Delta G^\ddagger$  to  $\frac{1}{4}\lambda$ .
- Important parameters needed to describe  $k_{\text{et}}$ .
  1. Reorganization energy  $\lambda$  corresponds to the vertical transition from parabola  $R$  to parabola  $P$ .
  2. Free energy of activation  $\Delta G^\ddagger$ .
  3. Free energy of reaction  $\Delta G^\circ$ .
- Bounds on  $\lambda$ :
  - $\lambda$  is usually on the order of 1.0-1.5 eV for isoenergetic reactions.
  - $\Delta G^\ddagger = \frac{\lambda}{4}$ .
- If we have a thermodynamic driving factor for an electron-transfer reaction, we simply either raise the  $R$  parabola or lower the  $P$  parabola (see Figure IX.1).
- Predictions for  $\Delta G^\ddagger$  based on  $\Delta G^\circ$  and the parabola model (see Figure IX.2):
  - $\Delta G^\circ = 0$  implies  $\Delta G^\ddagger = \frac{\lambda}{4}$ .
  - $-\lambda < \Delta G^\circ < 0$  implies  $\Delta G^\ddagger$  is decreasing (this can be rationalized by normal intuition; it makes sense that as  $\Delta G^\circ$  decreases,  $\Delta G^\ddagger$  would decrease, too).
  - $-\lambda = \Delta G^\circ$  implies  $\Delta G^\ddagger = 0$ .

Figure IX.2: Effect of  $\Delta G^\circ$  and  $\lambda$  on  $\Delta G^\ddagger$ .

–  $\Delta G^\circ < -\lambda$  implies  $\Delta G^\ddagger > 0$ .

- Under the assumption that the shapes of the curves do not change, we have

$$\Delta G^\ddagger = \frac{(\Delta G^\circ + \lambda)^2}{4\lambda}$$

- The above expression can be plugged into the formula for the kinetic rate constant,

$$k_{\text{et}} = Ax \exp\left(\frac{-(\Delta G^\circ + \lambda)^2/(4\lambda)}{RT}\right)$$

where

- $A$  is the attempt frequency (the Arrhenius coefficient).
- $x$  is the transmission coefficient.

- Special cases of  $\Delta G^\circ$ :

- Self-exchange:  $\Delta G^\circ = 0$ .
- Barrierless region:  $\Delta G^\circ = -\lambda$ .
- “Inverted” region:  $\Delta G^\circ < -\lambda$ .

- Experimental verification of Marcus theory:

- This very counterintuitive implication (that in the inverted region, increases in  $\Delta G^\circ$  actually slow the reaction down) needed verification.
- Early attempts faced difficulties because of the diffusion limitation (increasing exothermicity causes  $\log K$  to hit a ceiling).
- Class and Miller (UChicago) eventually found a reaction (intramolecular electron transfer) that could verify Marcus theory (later observed in many other photo and electrochemical systems).

- Back to outer- vs. inner-sphere electron transfer:

- Marcus theory discusses outer-sphere electron transfer, and cannot explain why in some cases Redox reaction rates vary so greatly.
- Taube and Halpern (UChicago) posit that some ligands (i.e., ones with multiple lone pairs or low lying antibonding orbitals) can form a bridge between two metal centers, enabling inner-sphere electron transfer.

- Example:  $[\text{Co}(\text{NH}_3)_5\text{X}]^{n+} / \text{Cr}^{2+}$  in aqueous medium:

- Since chromium is  $d^4$  (hence labile),  $[\text{Cr}(\text{H}_2\text{O})_6]^{2+} \xrightleftharpoons{\text{fast}} [\text{Cr}(\text{H}_2\text{O})_5]^{2+} + \text{H}_2\text{O}$ .
- The undercoordinated chromium ligand can engage in an inner-sphere electron transfer with the chloro ligand:  $[(\text{H}_3\text{N})_5\text{Co}-\text{Cl}-\text{Cr}(\text{H}_2\text{O})_5]^{4+} \xrightarrow{\text{e.t.}} [(\text{H}_3\text{N})_5\text{Co}]^{2+} + [\text{ClCr}(\text{H}_2\text{O})_5]^{3+}$ .
- Lastly, we hydrolyze cobalt:  $[\text{Co}(\text{NH}_3)_5]^{2+} \longrightarrow [\text{Co}(\text{H}_2\text{O})_6]^{2+} + \text{NH}_3$ .
- In the transition state with two metals bridged by a ligand, the reactant and product states mix, forming two **adiabatic** states along the reaction coordinate.
- **Adiabatic** (states): Two state of equal energy.
- State mixing ( $H_{ab} = \langle \Psi_a | \hat{H} | \Psi_b \rangle > 0$ ):
  - With weak electron transfer, we have R and P parabolas.
  - As electron transfer increases in strength, the “inner loop” in Figure IX.1 separates and rises above the bottom loop.
  - The bottom loop is stabilizing and bonding; vice versa for the upper one.
- There exist compounds that can capture the charge-delocalized state.

## IX.6 Chapter 12: Coordination Chemistry IV (Reactions and Mechanisms)

From Miessler et al. (2014).

- 3/11:
- **Transition-state theory:** A theory describing chemical reactions as moving from one energy minimum (the reactants) through higher energy structures (transition states, intermediates) to another energy minimum (the products).
  - **Principle of microscopic reversibility:** The lowest energy pathway going in one direction must also be the lowest energy pathway going in the opposite direction.
    - “Although the complexity of reaction coordinate diagrams can vary widely, the adopted path between the reactants and the products is always the lowest energy pathway available and must be the same regardless of the direction of the reaction” (Miessler et al., 2014, pp. 437–38).
  - **Steady-state approximation:** The concentration of the intermediate is assumed to be extremely small and essentially unchanging during much of the reaction.
    - Allowed by the presence of undetectable intermediates.
  - **Order** (of a reactant): The power of the reactant concentration in the differential equation that describes how its concentration changes with time, which indicates how the reaction rate is tied to a change in that reactant’s concentration.
  - **Rate constant:** A proportionality constant that relates the reaction rate to the concentration of the reactants, which is temperature dependent.
  - The **free energy of activation** can be divided into two components: **enthalpy of activation** and **entropy of activation**.
  - **Volume of activation:** A quantity that offers insight into whether the transition state is larger or smaller than the reactants, derived from examinations of pressure dependence on reaction rates.
  - The rate of reaction depends on the activation energy via the Arrhenius equation

$$k = Ae^{-\frac{E_A}{RT}}$$

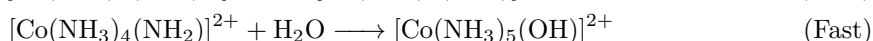
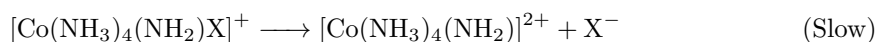
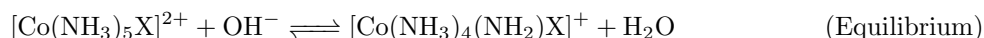
- Miessler et al. (2014) introduces substitution reactions of  $[M(H_2O)_m]^{n+}$  and color-based identification of species.
    - Note that  $[V(H_2O)_6]^{3+}$  has a higher water exchange rate constant than  $[V(H_2O)_6]^{2+}$ , despite the fact that we might expect it to hold onto its ligands more tightly because of its higher oxidation state.
  - **Labile** (compound): Compounds which react rapidly, essentially exchanging one ligand for another within the time of mixing the reactants. *Also known as kinetically labile.*
    - Very low activation energy for ligand substitution.
    - Examples include  $d^1$ ,  $d^2$ , and high-spin  $d^4$  through  $d^6$  compounds, as well as  $d^7$ ,  $d^9$ , and  $d^{10}$  ones.
  - **Inert** (compound): A compound that does not resist ligand substitution but is simply slower to react. *Also known as kinetically inert.*
    - “Inert octahedral complexes are generally those with high ligand field stabilization energies. . . specifically those with  $d^3$  or low-spin  $d^4$  through  $d^6$  electronic structures” (Miessler et al., 2014, pp. 440–41).
    - Strong ligand-field  $d^8$  complexes often form inert square-planar complexes.
  - **Stoichiometric mechanism**: Any one of the substitution reaction categories ( $D$ ,  $A$ , or  $I$ ).
  - **Intimate mechanism**: The distinction between activation processes that are associative and dissociative.
  - A word on notation: In this chapter’s reactions, “X will indicate the ligand that is leaving a complex, Y the ligand that is entering, and L any ligands that are unchanged during the reaction. In cases of solvent exchange, the X, Y, and L may be the same species. Charges will be omitted when using X, Y, and L, but the species may be ions” (Miessler et al., 2014, p. 441).
  - In a rate law, Rate equals the change in the concentration of the product with respect to time,  $d[P]/dt$ .
  - We assume the formation of an **ion pair** or **preassociation complex** in our description of an interchange reaction, instead of just assuming that the interchange occurs in one step.
  - Ion pairs form from ionic reactions while preassociation complexes form from dipole-dipole interactions.
  - Substitution reaction intermediates are detectable when  $k_1, k_{-1} \gg k_2$  since this allows the first (reversible) reaction to reach an equilibrium independent of the second.
    - Determining the concentration of  $[ML_5X \cdot Y]$  facilitates the calculation of  $K_1$ . Otherwise, it must be estimated theoretically.
  - Interchange mechanisms involving preassociation complexes can be difficult to distinguish from  $D$  mechanisms when  $[Y]$  is large since they both tend toward first-order kinetics.
- 3/14:
- **Ligand field activation energy**: The difference between the LFSE of the square-pyramidal transition state and the LFSE of the octahedral reactant. *Also known as LFAE.*
    - LFAEs for trigonal bipyramidal transition states are usually the same or larger than those for square pyramidal ones.
  - Other factors that influence reaction rate:
    - Oxidation state of the central ion: Higher oxidation states lead to slower ligand exchange.
    - Ionic radius: Smaller ions have slower exchange rates.
  - **Aquation**: Substitution by water.
  - **Anation**: Substitution by an anion.

- **Linear free-energy relationship:** A relation between kinetic effects and thermodynamic effects. *Also known as LFER.*

- Observed when the bond strength of a metal-ligand bond (thermodynamic parameter) plays a major role in determining the dissociation rate of a ligand (kinetic parameter).
- Observed when a plot of the logarithm of the rate constants for  $[\text{ML}_5\text{X}]^{n+} + \text{Y}$  substitution reactions, where X is varied but Y is not, versus the logarithm of the equilibrium constants for  $[\text{ML}_5\text{X}]^{n+} + \text{Y} \rightleftharpoons [\text{ML}_5\text{Y}]^{m+} + \text{X}$  is linear.

- **Conjugate base mechanism:** A ligand is deprotonated, the ligand *trans* to it dissociates, and then the deprotonated ligand is reprotonated and a new ligand binds. *Also known as S<sub>N</sub>1CB, substitution, nucleophilic, unimolecular, conjugate base.*

- For example, the mechanism of  $[\text{Co}(\text{NH}_3)_5\text{X}]^{2+} + \text{OH}^- \longrightarrow [\text{Co}(\text{NH}_3)_5(\text{OH})]^{2+} + \text{X}^-$  could be



- This mechanism is common with octahedral cobalt (III) metal centers.
- **Kinetic chelate effect:** Substitution for a chelated ligand is slower than a similar monodentate ligand.
- In dissociative mechanisms, products can have the same or different stereochemistry than the reactants.
  - What happens depends on the stereochemistry of the reactants, the ligands (both to be removed and to be added), and occasionally the concentration of the ligand to be added.
  - A square pyramidal intermediate leads to a retention of configuration. A trigonal bipyramidal one does not (necessarily; different points of attack along the equator lead to different products in varying ratios).
  - “As a general rule, *cis* reactants give a relatively higher percentage of substitution products that retain their *cis* configuration; *trans* reactants often afford a more balanced mixture of *cis* and *trans* substitution products” (Miessler et al., 2014, p. 456).
  - Compounds with multiple chelating rings can interconvert between stereoisomers by a dissociation  $\rightarrow$  rearrangement  $\rightarrow$  reattachment mechanism.

- Square-planar nomenclature: T is the ligand *trans* to the departing ligand X.
- Square-planar axes: Let the plane of the molecule be the *xy*-plane, with the *x*-axis collinear to the T–Pt–X axis.
- Square-planar rate law:

$$\text{Rate} = k_1[\text{Cplx}] + k_x[\text{Cplx}][\text{Y}]$$

- The  $k_2$  term derives from a standard associative mechanism.
- The  $k_1$  term derives from a solvent-assisted *A* mechanism (the solvent substitutes first, slowly; then the incoming ligand replaces the solvent, quickly).
  - This may occasionally lead to a 6-coordinate transition state.
- Determining the effect on reaction rate of ligands other than T:

$$\log k_Y = s \cdot \eta_{\text{Pt}} + \log k_S$$

- $k_Y$  is the rate constant for reaction with Y.
- $k_S$  is the rate constant for reaction with S.

- $s$  is the **nucleophilic discrimination factor** (for the complex).
  - $s := 1.00$  for *trans*-[Pt(py)<sub>2</sub>Cl<sub>2</sub>], and other values are based off of this reference.
- $\eta_{\text{Pt}}$  is the **nucleophilic reactivity constant** (for the entering ligand).

$$\eta_{\text{Pt}} = \log \left( \frac{k_Y}{k_{\text{CH}_3\text{OH}}} \right)$$

- Values are determined via the above equation with kinetic data from the reactions used to determine  $s$ .
- ***trans* influence:** The ground-state, thermodynamic effect where Pt–T  $\sigma$  donation uses a larger contribution of the  $p_x$  and  $d_{x^2-y^2}$  orbitals, leaving less for the Pt–X bond.
- **Inner-sphere reaction:** A redox reaction of transition-metal complexes where the two molecules are connected by a common ligand through which the electron is transferred.
- **Outer-sphere reaction:** A redox reaction of transition-metal complexes where the exchange occurs between two separate coordination spheres.
  - Occur when the ligands of both reactants are tightly held.
  - Primary change upon electron transfer is metal-ligand bond length, affected both by changes in metal oxidation state and use of  $e_g^*$  electrons (adding these increases antibonding character, lengthening bonds; vice versa for removing them).
  - Promoted by the chelate effect.
- Factors that influence electron transfer rates:
  - The rate of ligand substitution within the reactants.
  - The match of the reactant orbital energies.
  - The solvation of reactants.
  - The nature of the ligands.
- Inner sphere mechanisms depend on **quantum tunneling**.
  - “Ligands with  $\pi$  or  $p$  orbitals that can be used in bonding provide good pathways for tunneling. Ligands like NH<sub>3</sub>, with neither extra nonbonding pairs nor low-lying antibonding orbitals, do not provide effective tunneling pathways” (Miessler et al., 2014, p. 463).
- **Quantum tunneling:** A quantum mechanical property whereby electrons can pass through potential barriers that are too high to permit ordinary transfer.
- Three steps in an inner-sphere mechanism:
  1. A substitution reaction that leaves the oxidant and reductant linked by the bridging ligand.
  2. The electron transfer, frequently accompanied by the transfer of the ligand.
  3. A separation of the products.
- Lists specific examples of molecules/ions that are prone to inner- or outer-sphere mechanisms.
- Factors that influence the stability of complexes with different oxidation numbers:
  - LFSE.
  - Metal-ligand bonding.
  - Redox properties of the ligands.
  - Hard/soft character of the ligands.
- Discusses reactions of coordinated ligands.