Lecture 9 – Chapter 15, Section 4

Experiments

• Experimental rate laws
• First order
• Second order
• Method of Isolation
• Method of Initial Rates
Rate Laws can be predicted from

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<tr>
<th>25%</th>
<th>1. Reaction stoichiometry</th>
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<td>25%</td>
<td>2. Phase of the moon</td>
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<td>25%</td>
<td>3. Reaction mechanisms</td>
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<tr>
<td>25%</td>
<td>4. Collision rates</td>
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Understanding Reactions

1. Have some overall reaction
2. Dream up (hypothesize) one or more likely reaction mechanisms
3. Derive rate law corresponding to each mechanism
4. Measure experimental rate law
5. Rule out impossible mechanisms
6. Devise further experiments if necessary
Connecting Rate Law to Experiment

- Reaction rate we had before: \[ \text{Rate} = \frac{1}{d} \frac{\Delta[D]}{\Delta t} \]

- Experimentally we measure [D] at various times

- We would like to convert rate law from \( \Delta \text{concentration} \) just to concentration
Integrated Rate Laws

- Calculus makes this possible (thank goodness for mathematicians). For a first-order rate law...

\[
\text{Rate} = \frac{\Delta [\text{NO}_2]}{\Delta t} = k[\text{NO}_2]
\]

\[
\ln \left( \frac{[\text{NO}_2]}{[\text{NO}_2]_0} \right) = kt
\]

or

\[
[\text{NO}_2] = [\text{NO}_2]_0 e^{-kt}
\]
First-order Rate Laws

• General form of the rate law and integrated rate law

\[
\text{Rate} = \frac{\Delta[A]}{\Delta t} = k[A]
\]

\[
\ln\left(\frac{[A]_0}{[A]}\right) = kt
\]

or

\[
[A] = [A]_0 e^{-kt}
\]
First-order Rate Laws

• Note that the integrated form is linear

\[
\ln \left( \frac{[A]}{[A_0]} \right) = kt
\]

\[
y = \ln \left( \frac{[A]}{[A_0]} \right)
\]

\[
x = t
\]

slope = \(k\)

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>([\text{C}_5\text{HBr}]) (M)</th>
<th>([\text{C}<em>5\text{H}</em>{11}\text{Br}]_0/[\text{C}<em>5\text{H}</em>{11}\text{Br}])</th>
<th>(\ln([A]_0/[A]))</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>0.150</td>
<td>1.00</td>
<td>0</td>
</tr>
<tr>
<td>1.00</td>
<td>0.099</td>
<td>1.52</td>
<td>0.42</td>
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<tr>
<td>2.00</td>
<td>0.073</td>
<td>2.05</td>
<td>0.72</td>
</tr>
<tr>
<td>3.00</td>
<td>0.047</td>
<td>3.19</td>
<td>1.16</td>
</tr>
<tr>
<td>4.00</td>
<td>0.028</td>
<td>5.36</td>
<td>1.68</td>
</tr>
<tr>
<td>5.00</td>
<td>0.022</td>
<td>6.82</td>
<td>1.92</td>
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</table>
Not first order?

- If reaction is not first order, plot will not look linear
Half-life

- Amount of time it takes for $\frac{1}{2}$ of reactant to be used up (or half of product to form)
- For first-order $\frac{1}{2}$-life is independent of concentration
- Always same time to use up $\frac{1}{2}$ of stuff
- Why the number $e$ is so special
- Examples:
  - Radioactivity
  - Photochemistry
  - Some reactions
Half-life

- 50% gone after 5730 years
- 75% gone after 11,460 years
- 90% gone after 19,000 years
Second-order Rate Law

\[ \text{Rate} = \frac{\Delta [A]}{\Delta t} = k[A] \]

or

\[ \frac{1}{[A]} - \frac{1}{[A]_0} = kt \]

or

\[ \frac{1}{[A]} = kt + \frac{1}{[A]_0} \]
A second-order reaction will look linear if we plot $t$ as $x$ and what as $y$?

$$\frac{1}{[A]} - \frac{1}{[A]_0} = kt$$

0% 1. $1/[A]$
0% 2. $1/[A] - 1/[A]_0$
0% 3. $1/[A]_0$
0% 4. $k$
0% 5. $kt$
Second-order Rate Laws

- Integrated form is linear again

\[
\frac{1}{[A]} - \frac{1}{[A]_0} = kt
\]

**Second-Order Kinetic Plot**

- \( y = \frac{1}{[A]} - \frac{1}{[A]_0} \)
- \( x = t \)
- slope = \( k \)

OR

- \( y = \frac{1}{[A]} \)
- \( x = t \)
- intercept = \( \frac{1}{[A]_0} \)
Recall \( 2\text{NO}_2 \rightarrow 2\text{NO} + \text{O}_2 \) reaction

- Two different mechanisms gave two different rate laws
  - Mechanism I predicted: \( \text{Rate}_I = k[\text{NO}_2] \)
  - Mechanism II predicted: \( \text{Rate}_{II} = k[\text{NO}_2]^2 \)
Half-life

- Can derive half-life for second-order reaction
- But, half-life depends on concentration
  - Not nearly as useful as first-order
  - Not usually used, so don’t worry about it
Zero-order

- Zero-order reactions also happen
  \[ \text{Rate} = k \]
  \[ [A]_0 - [A] = kt \]
  or
  \[ [A] = -kt + [A]_0 \]

- Higher-order reactions also common
  \[ y = [A]_0 - [A] \]
  \[ x = t \]
  slope = \( k \)

OR

\[ y = [A] \]
\[ x = t \]
slope = \( -k \)
intercept = \([A]_0 \)
Summary of Integrated Rate Laws

<table>
<thead>
<tr>
<th>Order</th>
<th>Rate Law</th>
<th>Concentration Dependence</th>
<th>Linear Plot</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Rate = $k$</td>
<td>$[A]_o - [A] = kt$</td>
<td>$[A]_o - [A]$ vs. $t$</td>
</tr>
<tr>
<td>1</td>
<td>Rate = $k[A]$</td>
<td>$\ln \left( \frac{[A]_o}{[A]} \right) = kt$</td>
<td>$\ln \left( \frac{[A]_o}{[A]} \right)$ vs. $t$</td>
</tr>
<tr>
<td>2</td>
<td>Rate = $k[A]^2$</td>
<td>$\frac{1}{[A]} - \frac{1}{[A]_o} = kt$</td>
<td>$\frac{1}{[A]} - \frac{1}{[A]_o}$ vs. $t$</td>
</tr>
</tbody>
</table>

Note that we have ignored effects of product concentrations.

This is OK for some reactions, but generally we need to consider concentrations of all components (not right now)
Multiple components

• What if rate law has multiple things in it?
  Rate = \( k[A]^y[B]^z \)

• Different methods used to determine \( y \) and \( z \).
  – Isolation
  – Initial Rates
  – Others
Method of Isolation

- Do experiments with one component in huge excess
  - Concentration is effectively constant

\[ A + B \rightarrow \text{Product} \]
\[ \text{Rate} = k[A]^y[B]^z \]

Initially \( [A]_0 = 5 \times 10^{-3} \text{ M} \) and \( [B]_0 = 1 \times 10^{-5} \text{ M} \)

When reaction is complete…
\( [A]_f = 4.99 \times 10^{-3} \text{ M} \) and \( [B]_f = 0 \text{ M} \)

- Now we can measure concentration of \([B]\) with time and make plots to determine \(z\)
- Repeat with \([B]\) in huge excess to determine \(y\)
Method of Initial Rates

- If possible, measure reaction rate over very short time period
- If time period is short enough, concentrations of all reactants are approximately constant
- Vary initial concentration of one component to determine reaction order

\[ 2\text{H}_2 + 2\text{NO} \rightarrow \text{N}_2 + 2\text{H}_2\text{O} \]

<table>
<thead>
<tr>
<th>Experiment</th>
<th>[H\textsubscript{2}]</th>
<th>[NO]</th>
<th>Initial rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.001 M</td>
<td>0.002 M</td>
<td>1.2 \times 10^{-4} M/s</td>
</tr>
<tr>
<td>2</td>
<td>0.001 M</td>
<td>0.003 M</td>
<td>1.8 \times 10^{-4} M/s</td>
</tr>
<tr>
<td>3</td>
<td>0.002 M</td>
<td>0.002 M</td>
<td>4.8 \times 10^{-4} M/s</td>
</tr>
</tbody>
</table>

Doubling [H\textsubscript{2}] increases rate by 4. So [H\textsubscript{2}] is second-order
Initial Rates

Initial Rate 1: \[1.2 \times 10^{-4} \text{ } M/s = k (0.001M)^y (0.002M)^z\]

Initial Rate 2: \[1.8 \times 10^{-4} \text{ } M/s = k (0.001M)^y (0.003M)^z\]

Divide

\[
\frac{1.2 \times 10^{-4}}{1.8 \times 10^{-4}} \text{ } M/s = \frac{k (0.001M)^y (0.002M)^z}{k (0.001M)^y (0.003M)^z}
\]

\[
2 = \left( \frac{0.002}{0.003} \right)^z
\]

\[
\ln \left( \frac{2}{3} \right) = z \ln \left( \frac{0.002}{0.003} \right)
\]

\[
z = 1
\]
Today

• Finish CAPA #5

Wednesday

• Start on CAPA #6
• Finish reading Chapt 15
• Start thinking about previous chapters…