Today

Review Transition State Theory Arrhenius Theory

Kinetic Mechanisms

Why does a reaciton follow a particular rate law? What is actually happening in the reaction?

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Let's make a new Equation

$$k = A e^{-E_a/RT}$$
 Ink = InA -E_a/RT

let's look at two temperatures

$$Ink_1 = In A - E_a/RT_1$$

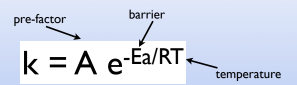
 $Ink_2 = InA - E_a/RT_2$

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Arrhenius Law

The rate constant k is a function of temperature



Arrhenius Law

The higher the temperature the more molecules that have enough energy to make it over the barrier

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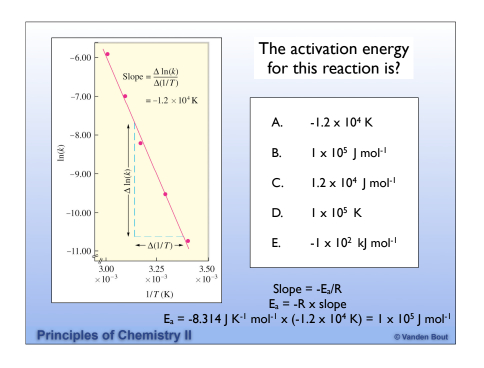
 $Ink_2 = InA - E_a/RT_2$

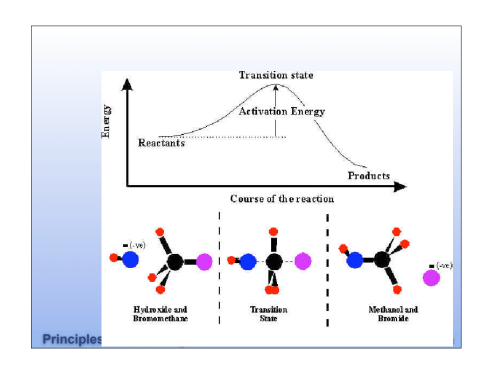
subtract to get a new equation that doesn't have A

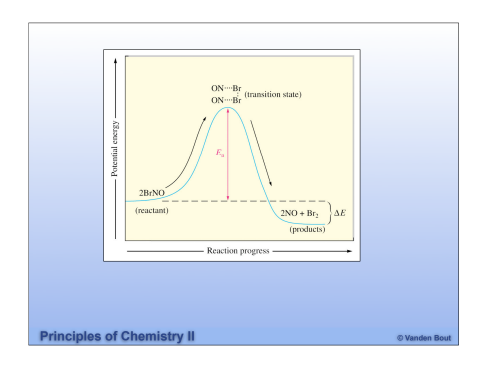
$$ln(k_2/k_1) = \frac{-E_a}{R} \left[\frac{I}{T_2} - \frac{I}{T_1} \right]$$

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Why are reactions faster at higher temperatures?

More molecules have sufficient energy to get over the barrier. BIG EFFECT

More molecules have collisions (but this is a very small effect) that is ignored in Arrhenius view of transition state theory

Why is something first order? second order?

Because of the number of molecules involved in the "key" (slowest) step in the reaction

We need to think about what is actually happening

This is the mechanism of the reaction

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student + quiz + TA

happy student with quiz turned in student + TA + quiz

student with quiz + TA

student with quiz

student with quiz

happy student with completed quiz

student with completed quiz + TA

happy student with quiz turned in

What controls the rate of this reaction?

Nonsense "real world" example

Nonsense "real world" example

student + quiz + TA ———— happy student with quiz turned in

student + TA + quiz → student with quiz + TA

student with quiz → student with completed quiz

student with completed quiz + TA → happy student with quiz turned in

imagine step one was very slow (we passed out preprinted bubble sheets in alphabetical order checking ID before handing you the quiz)

Rate dominated by "handing" out the quiz

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Nonsense "real world" example

student + quiz + TA ——— happy student with quiz turned in

student + TA + quiz → student with quiz + TA

student with quiz → student with completed quiz

student with completed quiz + TA → happy student with quiz turned in

What we want. All steps to be fast except step 2
Then the rate is dominated by step 2
However step 1 still matters

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Nonsense "real world" example

student + TA + quiz \longrightarrow student with quiz + TA

student with quiz \longrightarrow student with completed quiz

student with completed quiz + TA \longrightarrow happy student with quiz turned in

dominated by step 3. You need to have your quiz notarized before turning it in.

All three steps matter

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$$H_2(g) + Br_2(g) \longrightarrow 2HBr(g)$$

What actually happens?

Does a H₂ and a Br₂ molecule collide and react?

Does something else happen?

Step I
$$Br_2 \longleftrightarrow 2Br$$
 intermediates
Step 2 $Br + H_2 \longrightarrow HBr + H$
Step 3 $H + Br_2 \longrightarrow HBr + Br$
Step 4 $2Br \longrightarrow Br_2$
Overall $H_2(g) + Br_2(g) \longrightarrow 2HBr(g)$

Which Step Matters?

The rate is dominated by the slowest step this is called the "rate limiting step"

$$Br_2 \longleftrightarrow 2Br$$
 slow
 $Br + H_2 \longrightarrow HBr + H$ fast
 $H + Br_2 \longrightarrow HBr + Br$ fast
 $2Br \longrightarrow Br_2$ fast

As soon as some Br is formed it reacts very rapidly through the other step.

The rate is dominated by "waiting" for step I

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With this mechanism what is the predicted rate law?

$$Br_2 \longleftrightarrow 2Br$$
 slow

 $Br + H_2 \longrightarrow HBr + H$ fast

 $H + Br_2 \longrightarrow HBr + Br$ fast

 $2Br \longrightarrow Br_2$ fast

What molecules are reactants in the slow step?

only one molecule of Br₂

Therefore rate = $k[Br_2]$

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What if this was the mechanism?

$$Br_2 \longleftrightarrow 2Br$$
 fast
 $Br + H_2 \longrightarrow HBr + H$ slow
 $H + Br_2 \longrightarrow HBr + Br$ fast
 $2Br \longrightarrow Br_2$ fast

$$rate = k[Br][H_2]$$

Can't have Br in the rate law it is an intermediate

where did the Br come from?

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What if this was the mechanism?

$$Br_2 \longleftrightarrow 2Br$$
 fast
 $Br + H_2 \longrightarrow HBr + H$ slow
 $H + Br_2 \longrightarrow HBr + Br$ fast
 $2Br \longrightarrow Br_2$ fast
 $rate = k[Br][H_2]$

You need 1/2 mole of Br₂ to get one mole Br therefore

rate =
$$k[Br_2]^{1/2}[H_2]$$

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What is the rate law expected for the given mechanism?

$$NO_2 + NO_2 \longrightarrow NO_3 + NO$$
 slow
 $NO_3 + CO \longrightarrow NO_2 + CO_2$ fast
 $NO_2 + CO \longrightarrow NO + CO_2$ overall

A. rate =
$$k[NO_2][CO]$$

B.
$$rate = k[NO_3][CO]$$

C.
$$rate = k[NO_2]^2[CO]$$
 two molecules of NO₂ in rate limiting step

D. rate =
$$k[NO_2]^2$$

E. rate =
$$k[NO_2]^2[NO_3][CO]$$

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Mechanisms that involve products

$$Cl_{2} \quad Cl + Cl \qquad fast$$

$$Cl + H_{2}S \quad HCl + HS \qquad fast$$

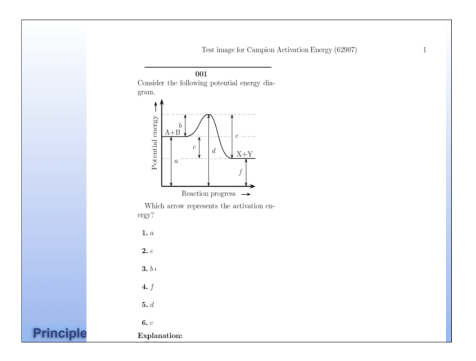
$$Cl + HS \quad HCl + S \qquad slow$$

$$Cl_{2} + H_{2}S \quad 2HCl + S$$

$$rate = k[Cl][HS] \times \frac{[Cl][H_{2}S]}{[HCl][HS]} \times \frac{[Cl_{2}]}{[Cl]^{2}} = \frac{[Cl_{2}][H_{2}S]}{[HCl]}$$

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Test image for Mechanism 13 44 (22135)

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The mechanism proposed for the oxidation of the iodide ion by the hypochlorite ion in aqueous solution is as follows:

1: ClO[−] + H₂O ≈ HClO + OH[−] (fast)

2:
$$I^- + HClO \rightarrow HIO + Cl^-$$
 (slow)

3:
$$\text{HIO} + \text{OH}^- \rightarrow \text{IO}^- + \text{H}_2\text{O}$$
 (fa

3: HIO + OH[−] → IO[−] + H₂O (fast)
What is the rate law for the formation of
HIO implied by this mechanism?

$$\mathrm{rate} = k \; \frac{\left[\mathrm{ClO^{-}}\right] \left[\mathrm{I^{-}}\right]}{\left[\mathrm{H_{2}O}\right]}$$

$$\sum_{\text{rate}} = k \frac{[\text{OH}^-][\text{CIO}^-]}{[\text{I}^-]}$$

3 rate =
$$k \frac{[ClO^-]}{}$$

4 rate =
$$k \frac{[OH^-][I]}{[CIO]}$$

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