	Trial	[A].	[B]	Rate SCAJ	Calculated 'k'
	1	0,150	0-180	0.0016875	0-500
_	2	0.300	0,150	0.0033750	0.500
-	3	0_150	0,300	0.0067500	0,500

Ratezk[A][B]<sup>2</sup>

Now, we'd like to know the value of 'k'. Solve rate law for 'k'.

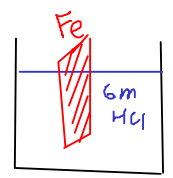
$$k = \frac{R_{a}te}{[A][B]^2}$$
 Plug in each set of data to this equation and calculate 'k'!

The average of these calculated 'k' values equals the rate constant. (For real data, expect some experimental error in these numbers!)

$$Rate = 0.500[A][B]^2$$

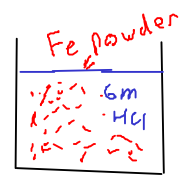


2) SURFACE AREA OF CONTACT BETWEEN REACTANTS



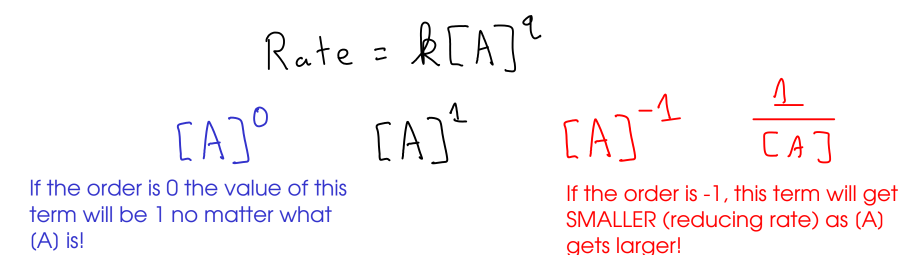
FetzHCI -> Fell, + H2

3 CONCENTRATION OF / PRESENCE OF A CATALYST



(4) TEMPERATURE

- Usually, the reaction rate increases as you increase the concentration of a reactant. This is true if the order for that reactant is greater than zero!

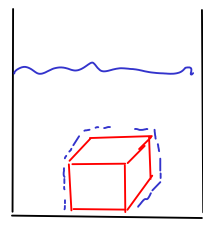


"inhibitor"

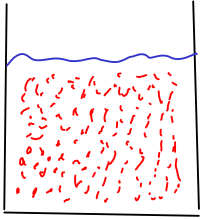
- In most cases, reaction orders ARE positive numbers!

- If a reaction takes place at an INTERFACE, then the reaction rate will depend on the SURFACE AREA of that interface!

Consider a reaction between a liquid and a solid. The reaction can only take place where the liquid and the solid TOUCH each other,



Next, try the same reaction, but break the cube into small fragments first!



The reaction takes place only on the surface of the cube. Here, the reaction takes place on the surface area of each fragment. In total, this is a much LARGER surface than the cube.

- The larger the SURFACE AREA of the INTERFACE between the reactants, the faster the reaction will occur!

- This is not an important factor for reactions that take place IN SOLUTION, since there is no interface.

## CONCENTRATION OF / PRESENCE OF A CATALYST

-a CATALYST is a substance that INCREASES the rate of a reaction without being consumed in the reaction.

- Appear in the rate law equation, but are written as "reaction conditions" in traditional chemical equations. Typically, the higher the concentration of catalyst, the faster the reaction.

$$A + B \xrightarrow{c} D$$
  
 $R_{a} + e = k [A]^{P} [B]^{4} [C]^{T}$   
The catalyst shows up in the rate law  
like the reactants!

- Work via many different mechanisms. Some provide surfaces for reactants to bind to and react. Some react with reactant molecules to leave them in a state more suitable for the main reaction (and are then regenerated during the main reaction). Some catalysts bind to and bend reactants into favorable orientations for reactions, etc.

- Biological catalysts are usually called ENZYMES.

- We observe that chemical reactions proceed FASTER at HIGHER temperatures.

... but sometimes reactions are run at low temperatures for other reasons - like safety, decomposition of desired products at high temperatures, formation of competing undesirable products at high temperatures, etc.

Note: Changing temperature affects the RATE CONSTANT, k !

## **REACTION RATE THEORY**

- We'd like to have an explanation for how the four factors we discussed affect the reaction rate!

- COLLISION THEORY and TRANSITION STATE THEORY attempt to explain reaction rate.

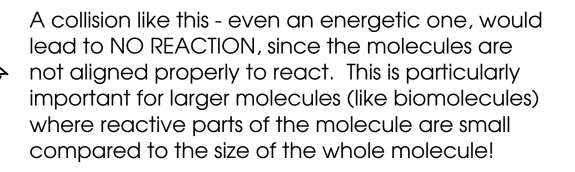
## COLLISION THEORY

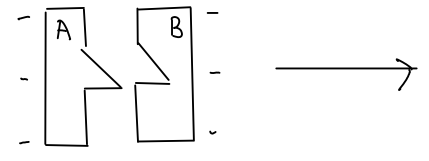
- states that for a reaction to occur between atoms or molecules, three things must occur:

 $(\mathbf{I})$  Reactants must COLLIDE with each other.

(2) Colliding reactants must hit each other with the correct ORIENTATION to react.

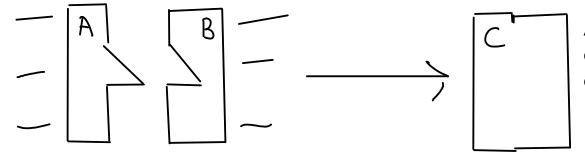
3 Colliding reactants must also hit each other with ENOUGH ENERGY to react (called the ACTIVATION ENERGY)





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A collision where molecules don't hit each other very hard will not lead to a reaction. There isn't enough ENERGY available for the molecules to react with one another.



A collision where molecules hit each other with the correct orientation AND enough energy may lead to a reaction!



- Increasing the concentration of reactants increases THE NUMBER OF COLLISIONS that occur in a reaction mixture - increasing rate.

- Increasing SURFACE AREA provides more opportunities for reactant molecules to COLLIDE - increasing rate.

- Increasing temperature INCREASES THE ENERGY (and number) of collisions, since temperature is proportional to the average kinetic energy of molecules. More collisions will have the ACTIVATION ENERGY needed to react, so rate increases.

- Some catalysts work by bending reactant molecules into ORIENTATIONS favorable for reaction - making it easier for large molecules to react with one another.

... but what about this ACTIVATION ENERGY? Let's look at TRANSITION STATE THEORY

- States that when reactant molecules collide, they first form a TRANSITION STATE which then decomposes and/or loses energy to form product molecules.

- A TRANSITION STATE is a high-energy state: It may be an unstable combination of several reactant molecules or an excited (high energy) state of a single product molecule. The transition state breaks down and loses energy to form the products of the reaction.

- The ACTIVATION ENERGY is the energy required to form the transition state, and it acts as a barrier to reaction.

- We model the rate constant "k" using collision and transition state theory with the ARRHENIUS EQUATION:

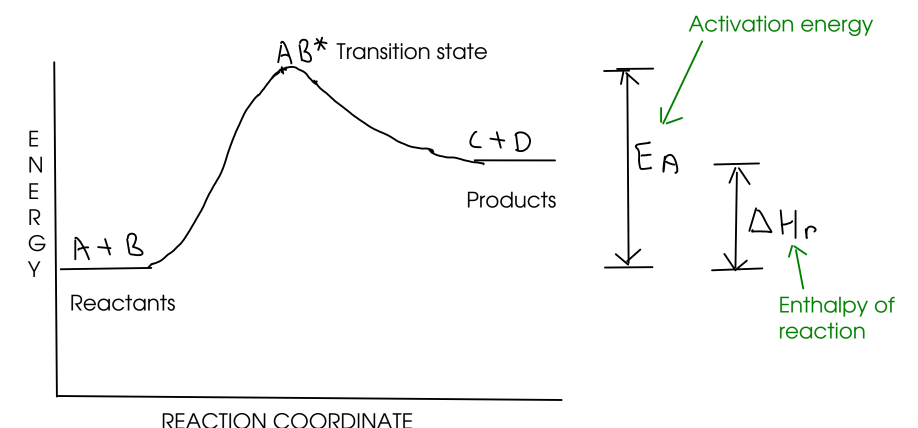
$$\begin{array}{l}
\frac{-E_{\alpha}}{R \times T} & \text{Ea} = \text{ACTIVATION ENERGY} \\
R = \text{ideal gas constant} \\
T = \text{absolute temperature} \\
A = \text{frequency factor ... the fraction of collisions with the right orientation to react}
\end{array}$$

... this equation allows us to calculate the rate constant for different temperatures if we have values for "A" and "Ea". We can get these values by examining a reaction at two (or more) different temperatures.

<sup>%</sup> ENERGY DIAGRAMS

- graphically, we can look at transition state theory via an ENERGY DIAGRAM

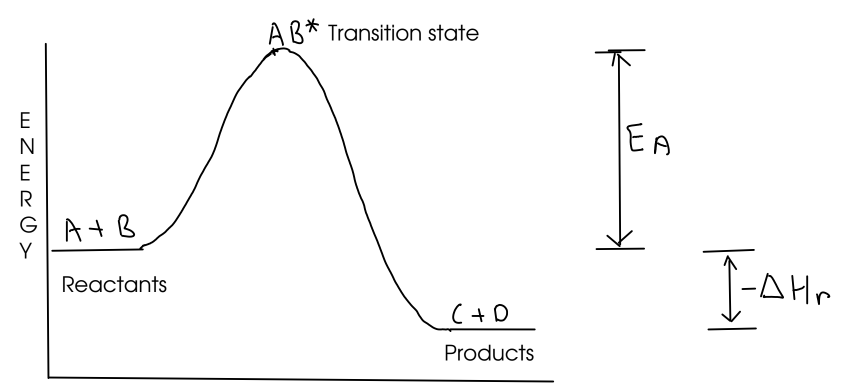
For an ENDOTHERMIC REACTION, the products have a higher energy than the reactants  $A + B \rightarrow C + O$ 



- a measure of how far the reaction has proceeded

For an EXOTHERMIC REACTION, the products have a lower energy than the reactants

 $A + B \rightarrow C + O$ 

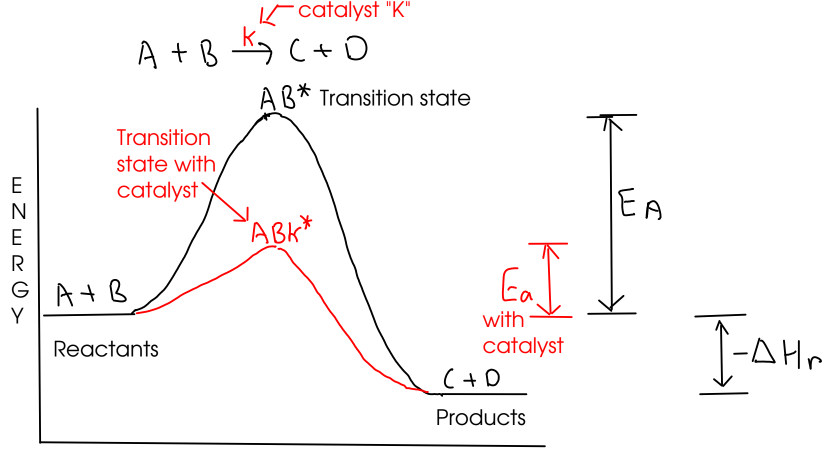


REACTION COORDINATE

- Whether a reaction is endothermic OR exothermic, there is still an activation energy barrier that must be crossed in order to react.

- This explains why a pile of wood that's exposed to air doesn't just burst into flames. Even though the cdombustion of wood is EXOTHERMIC, there's still an activation energy barrier preventing the reaction from occurring without an initial input of energy - a "spark"! <sup>®</sup> CATALYSTS?

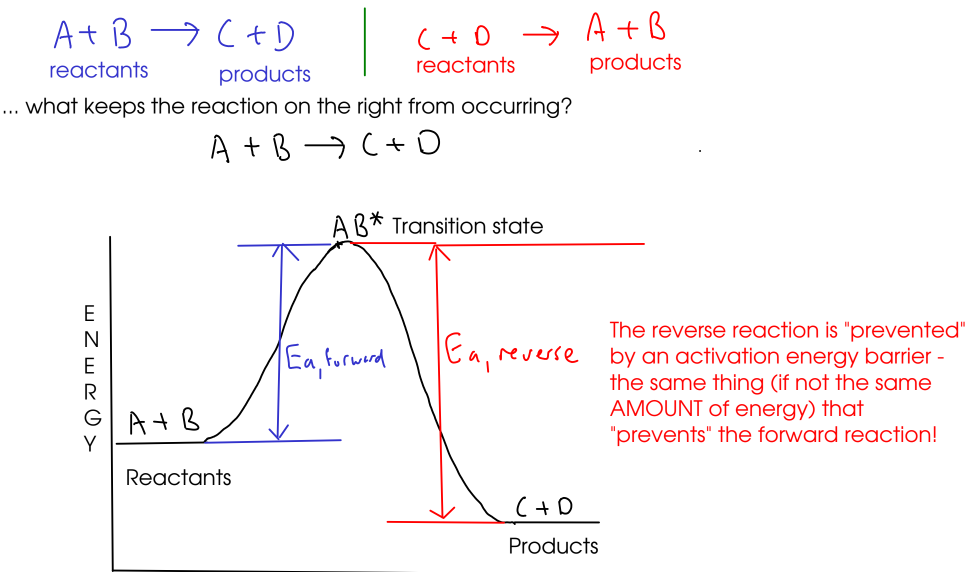
- So how does a catalyst fit into this picture? A catalyst LOWERS the activation energy for a reaction.



**REACTION COORDINATE** 

- The exact mechanism by which a catalyst lowers the energy of the transition state may be simple ... or complex. As we mentioned before, some catalysts hold molecules so that it's easier for reactants to come together, some react with reactant molecules to produce an intermediate that reacts more easily with other reactants to make the final product, etc.

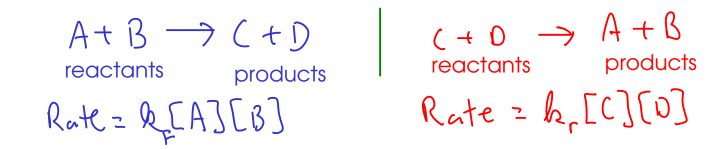
WHAT KEEPS A REACTION FROM GOING BACKWARDS?



REACTION COORDINATE

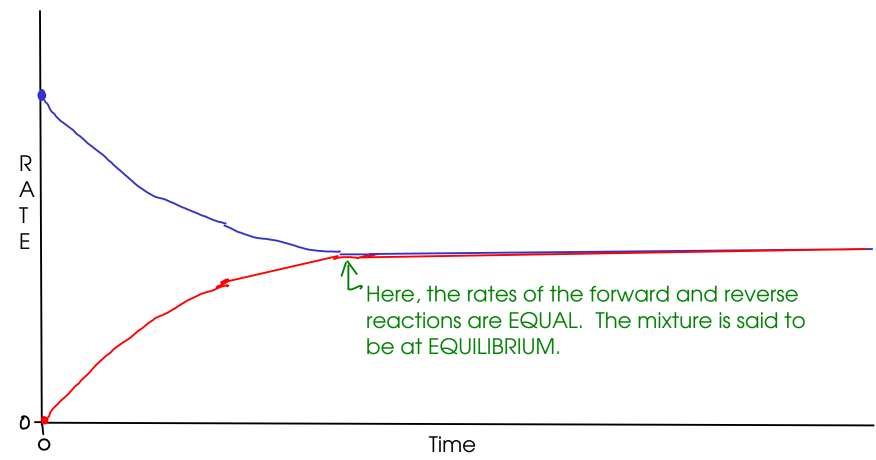
So what really happens during a reaction? Both forward and reverse reactions occur!

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- Let's look at the RATES of both the forward and reverse reactions over time.



- Initially, the mixture is all A and B. As C and D are formed, the rate of the reverse reaction increases while the rate of the forward reaction decreases. Eventually, these rates become equal.

- At EQUILIBRIUM, the concentrations of A, B, C, and D stop CHANGING. The reaction doesn't stop, but it appears stopped to an outside observer.

101 DESCRIBING EQUILIBRIUM

$$\frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \swarrow \frac{A + bB}{T} \swarrow \frac{A + bB}{T} \swarrow \frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \rightleftharpoons \frac{A + bB}{T}$$

$$\frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \end{gathered}$$

$$\frac{A + bB}{T} \Huge \frac{A + bB}{T}$$

$$\frac{A + bB}{T} \Huge \frac{A + bB}{T}$$

$$\frac{A + bB}{T} \lor \frac{A + bB}{T}$$

$$\frac{A + bB$$

based)

- At equilibrium, the ratio above equals a constant number - the EQUILIBRIUM CONSTANT. The equilibrium constant depends on TEMPERATURE, but not on other factors.

- Not all reactants and products are included in the equilibrium constant expression!