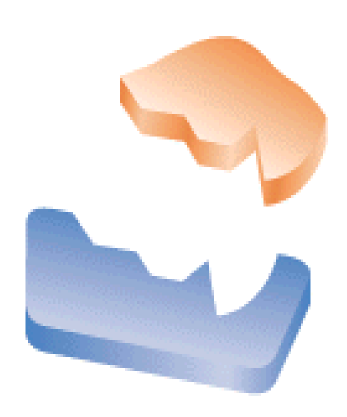
Enzymes and Coenzymes I

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What are Enzymes?

- Enzymes are <u>biological catalysts</u> that speed up the rate of a reaction without being changed in the reaction
- All enzymes are protein in nature
- But all proteins are not enzymes
- Substance upon which the enzymes act are called <u>substrates</u>
- Enzyme converts substrates into product(s)

Substrate

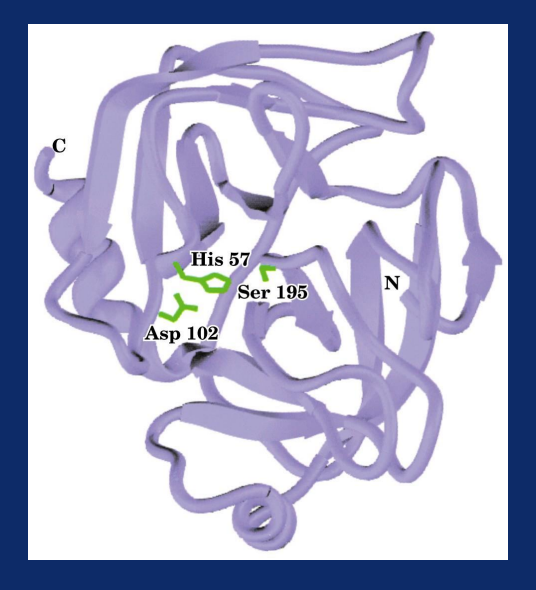


Enzyme



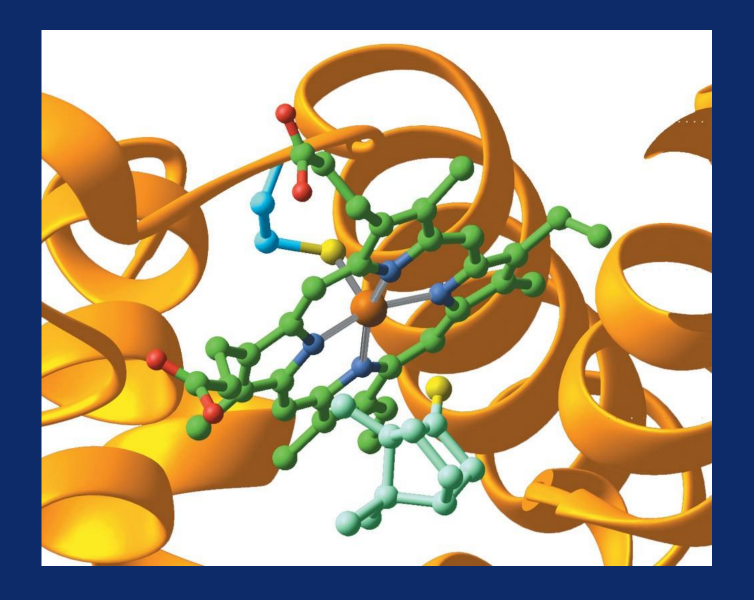
Properties of Enzymes

- Active site-
 - The region of enzyme that binds with the substrate and where catalysis occurs
 - All enzymes have one or more active sites
- Specificity-
 - Enzymes bind to their specific substrates in the active site to convert them to product(s)
- Regulation-
 - Enzymes can be activated or inhibited so that the rate of product formation responds to the need of the cell



Structure of trypsin enzyme

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An enzyme with its active site

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Enzyme specificity

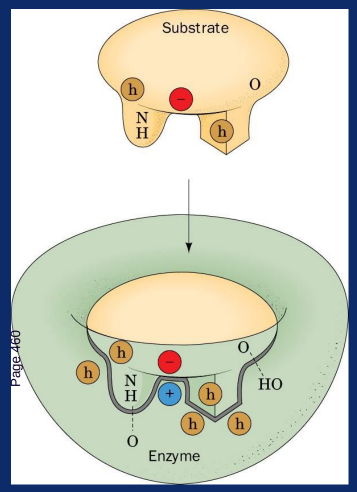
- Enzymes are highly specific
- Interact with only one or a few of the substrates
- Catalyze only one type of reaction

Enzyme-substrate binding

- Two models have been proposed
 - Lock and key binding
 - Induced fit binding

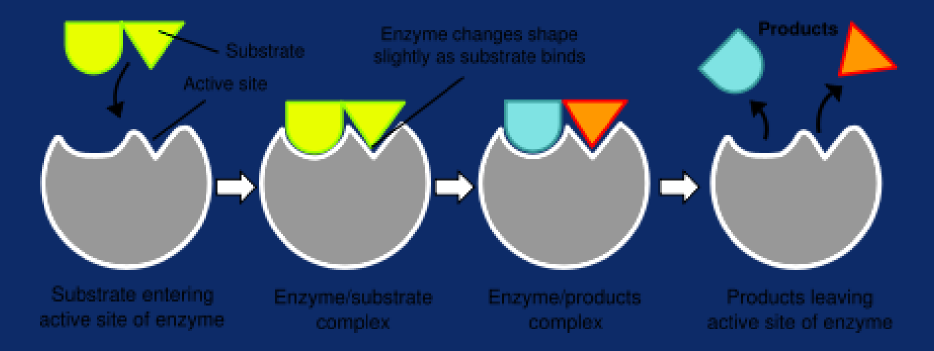
Lock and key binding

The enzyme has an active site that fits the exact dimensions of the substrate



Induced fit binding

 After the binding of substrate, the enzyme changes its shape to fit more perfectly with substrate



Classification of Enzymes

Classified into six types according to the reaction catalyzed

| Classification | Type of Reaction Catalyzed |
|--------------------|--|
| 1. Oxidoreductases | Oxidation-reduction reactions |
| 2. Transferases | Transfer of functional groups |
| 3. Hydrolases | Hydrolysis reactions |
| 4. Lyases | Group elimination to form double bonds |
| 5. Isomerases | Isomerization |
| 6. Ligases | Bond formation coupled with ATP hydrolysis |

Enzyme nomenclature (Naming)

- Enzyme nomenclature is based on the rules given by IUBMB (International Union of Biochemistry and Molecular Biology)
- EC 3.4.17.1 (carboxypeptidase A)
 EC = Enzyme Commission
 Class.Subclass.Subsubclass.Enzyme number

Holoenzymes

 Some enzymes require non-protein groups to become active

- The inactive form of enzyme without its non-protein part is called an apoenzyme
- Apoenzyme (inactive) + nonprotein part = Holoenzyme (active)

Cofactors and Coenzymes

- If the non-protein part is a metal ion such as Cu²⁺, Fe³⁺, Zn²⁺, etc., it is called cofactor
- If small organic molecules, known as coenzymes such as NAD+
 - Prosthetic groups
 - Cosubstrates

Coenzymes

 Coenzymes that are permanently associated with an enzyme known as prosthetic groups e.g. FAD

 Coenzymes that only temporarily associate with an enzyme known as cosubstrates e.g. NAD

Apoenzyme (inactive) + Cofactor/coenzyme = Holoenzyme (active)

Ribozymes, Isoenzymes and zymogens

- Ribozymes are RNAs with enzyme activity
- Isoenzymes are enzymes that catalyze the same chemical reaction but they have slightly different structures
- Zymogens are inactive enzyme precursors that require a biochemical change to become active e.g. cleavage of a peptide blocking the active site

Enzymes decrease activation energy of a reaction

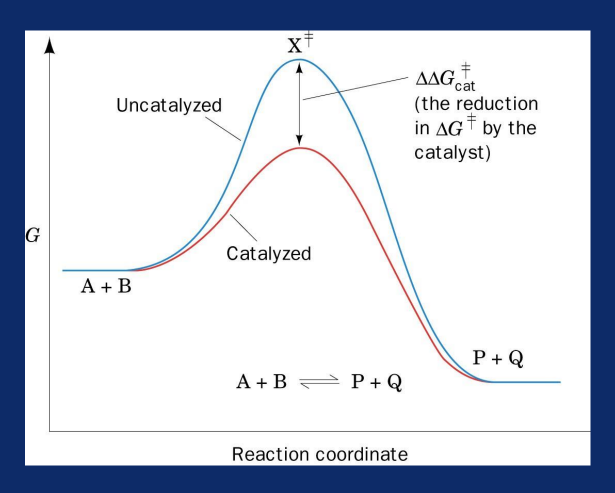
- In every chemical reaction, the reactants pass through a <u>transition state</u> that has greater energy than that of the reactants or products alone
- The difference in energy between the reactants and the transition state is called the <u>activation energy</u>
- If the activation energy is available then the reaction can proceed forming products

 An enzyme reduces the <u>activation</u> <u>energy</u> required for a reaction

 It provides an alternative transition state of lower energy called the enzymesubstrate complex and thus speeds up the reaction

 Enzymes decrease the activation energy but they do not alter the change in the <u>free energy</u> (△G)

The effect of a catalyst on the transition state diagram of a reaction



Enzyme Activity or Velocity

 Velocity is the rate of a reaction catalyzed by an enzyme

Enzyme activity is expressed as:
 mmoles of product formed/min/mg enzyme

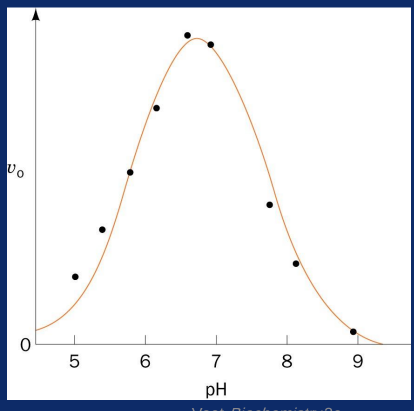
Factors that affect enzyme activity

- Effect of temperature
 - Every enzyme has an optimal temp. for catalyzing a reaction
 - The rate of an enzyme reaction initially increases with rise in temperature
 - At high temp. enzymes are denatured and become inactive
 - In humans most enzyme have an optimal temp. of 37°C

Factors that affect enzyme activity

- Effect of pH
 - Effect of pH on the ionizable groups in the active site of enzyme or in the substrate affect catalysis
 - Every enzyme has an optimal pH for catalyzing a reaction
 - Most enzymes have highest activity between pH 6 and pH 8
 - -Pepsin has highest activity at pH 2

Effect of pH on the initial rate of the reaction catalyzed by most enzymes (the bell-shaped curve)



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Factors that affect enzyme activity

- Effect of [E] and [S]
 - The reaction velocity increases initially with increasing [S]
 - At low [S], the reaction rate is proportional to [S]
 - Further addition of substrate has no effect on enzyme velocity (v)
 - The rate of an enzyme reaction is directly proportional to the conc. of enzyme if the substrate concentration [S] is higher than enzyme

Enzyme kinetics

 The model of enzyme kinetics was first proposed by Michaelis and Menten in 1913 and later modified by Briggs and Haldane

 The Michaelis Menten equation describes the relationship of initial rate of an enzyme reaction to the [S]

Product E+S $\stackrel{k_2}{\rightleftharpoons}$ ES $\stackrel{k_2}{\rightleftharpoons}$

Initial rate of enzyme reaction

Pre-steady state kinetics

- When an enzyme is mixed with high concentration of substrate, there is an initial short period of time (a few hundred microseconds) during which intermediates leading to the formation of product gradually build up

Steady state kinetics

- After initial state, the reaction rate and the concentration of intermediates change slowly with time called steady state reaction
- An intermediate is said to be steady state when its rate of synthesis is equal to its rate of degradation

Michaelis Menten Equation

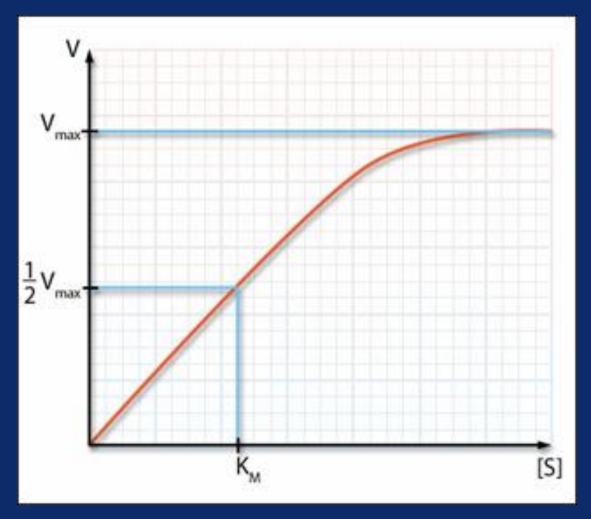
•It measures the initial velocity (ν_{o}) of an enzyme reaction

[S] = substrate concentration

 V_{max} = maximum velocity

 K_m = Michaelis constant

Initial velocity v_0 of a simple Michaelis–Menten reaction versus the substrate concentration[S]



K_m (Michaelis Constant)

• $K_{\rm m}$ is the substrate concentration at which the initial rate is one-half of the maximum rate (½ $V_{\rm max}$)

 It is the [S] required to saturate half of all of the active sites of an enzyme The K_m value of a substrate depends on its affinity with the enzyme

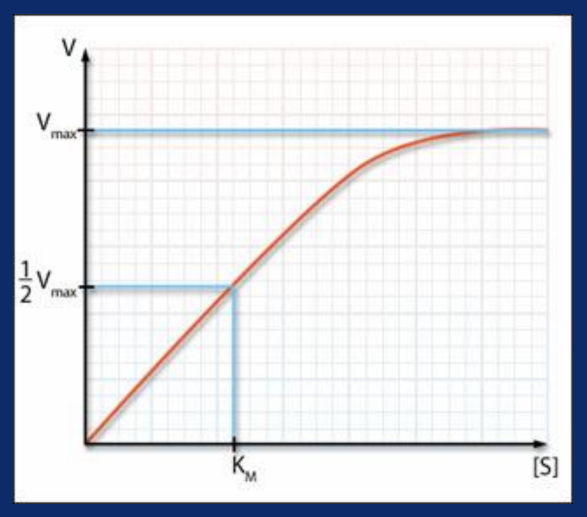
- High K_m means low affinity with enzyme (more substrate needed to saturate the enzyme)
- -Low K_m means high affinity with enzyme (less substrate needed to saturate the enzyme)

Lineweaver-Burk plot

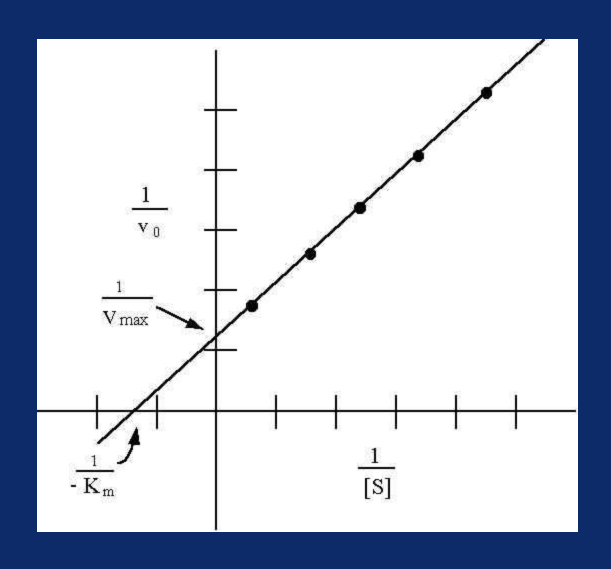
 Also called the double-reciprocal plot, obtained by taking reciprocals of the Michaelis Menten equation

 It is plotted to calculate the Km and Vmax values and to determine the mechanism of action of enzyme inhibitors

Initial velocity v_0 of a simple Michaelis–Menten reaction versus the substrate concentration[S]



Lineweaver-Burk plot



References

 Illustrated Reviews in Biochemistry by Lippincott