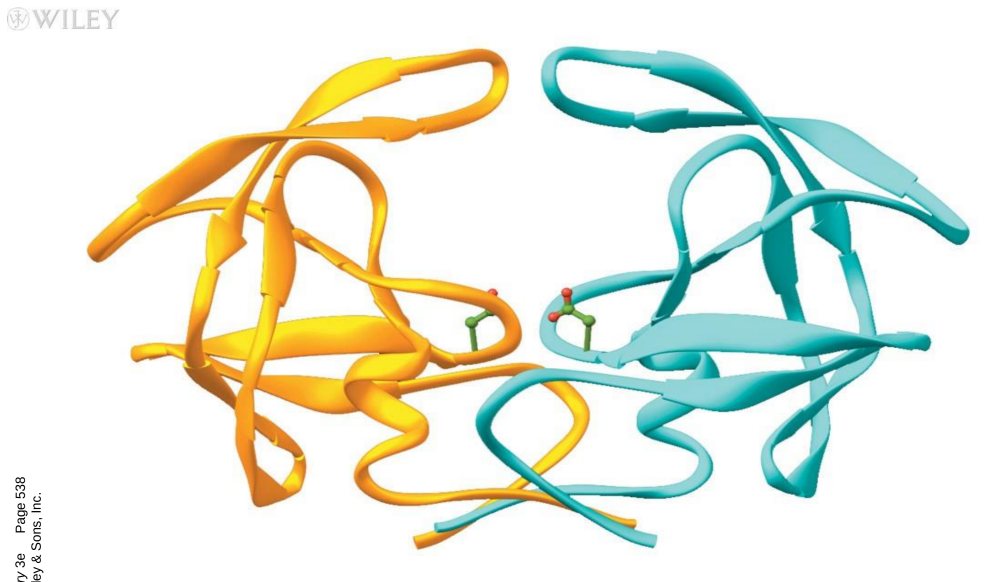
Enzyme inhibition

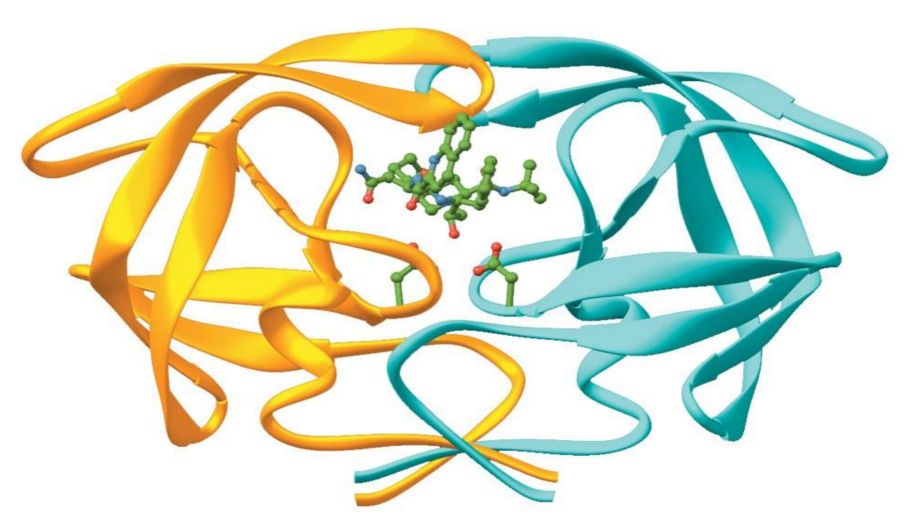
Inhibition is a process by which the enzyme activity is regulated or controlled or stopped

To inhibit means to stop enzyme activity



An enzyme without inhibitor





An enzyme with inhibitor

*K*_i (Inhibitor constant)

*K*_i is a measure of the affinity of inhibitor for enzyme

Also called dissociation constant

Enzyme inhibition

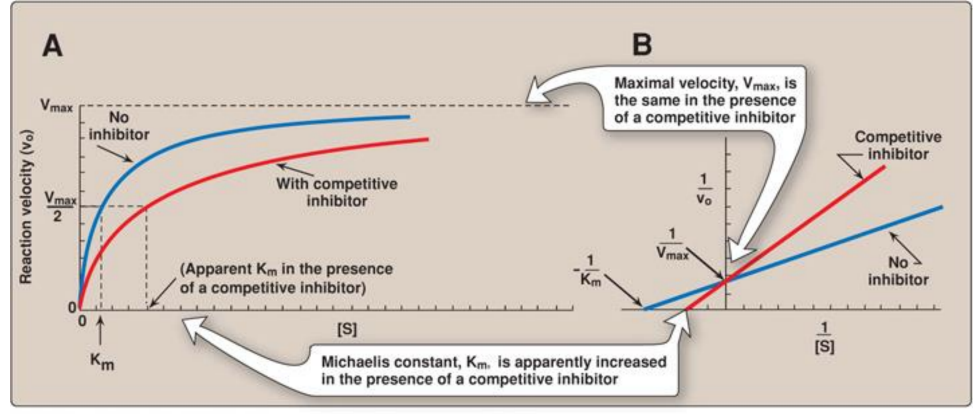
There are three types of enzyme inhibition:
Competitive
Noncompetitive
Uncompetitive

Competitive inhibition

The inhibitor is a structural analogue (similar) that competes with the substrate for binding to the active site of enzyme
 Two reactions are possible:

 $\overline{E} + S \leftrightarrow \overline{ES} \rightarrow E + P$ and $E + I \leftrightarrow \overline{EI}$

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Competitive inhibition

In competitive inhibition, V_{max} is unchanged in the presence and the absence of inhibitor

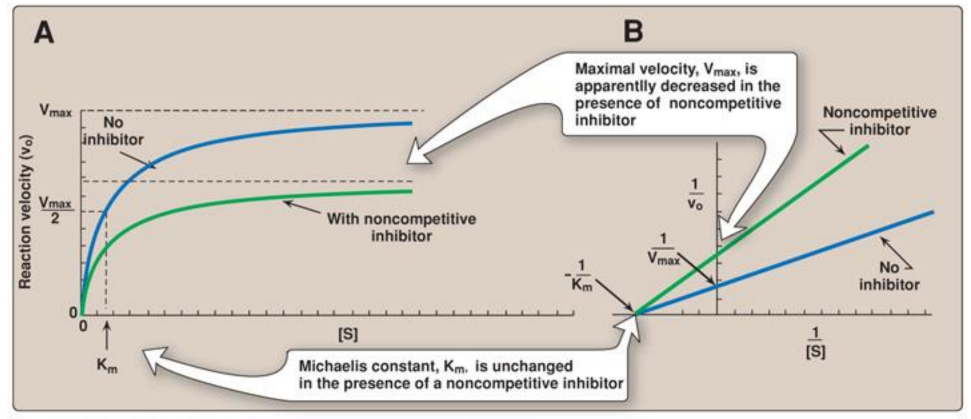
 The value of K_m is increased because S and I compete for binding at the same site

A higher [S] is required to achieve halfmaximal velocity

Noncompetitive inhibition

- The inhibitor does not have structural similarity to the substrate
- The inhibitor binds to the enzyme at a site away from the substrate binding site
- No competition exists between the inhibitor and the substrate
- The inhibitor can bind to a free enzyme or to an enzyme-substrate complex
- In both cases the complex is catalytically inactive

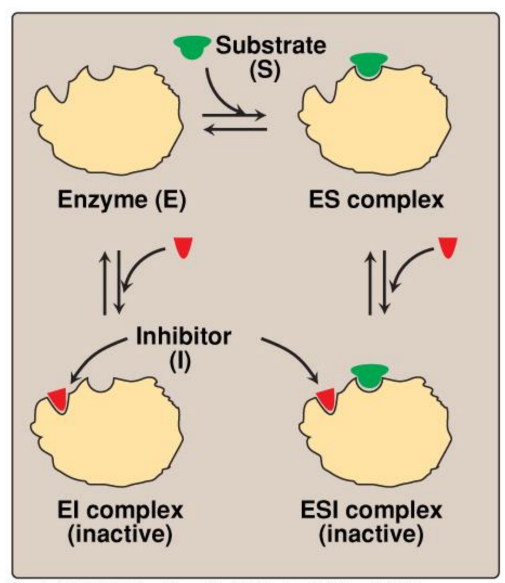
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Noncompetitive inhibition

ES + I ↔ ESI (inactive) E + I ↔ EI (inactive)
The value of V_{max} is decreased by the inhibitor
K_m is unchanged because the affinity of S for E is unchanged



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Noncompetitive inhibition

Regulation of enzyme activity

- Regulatory enzymes usually catalyze the first or an early reaction in a metabolic pathway
- They catalyze a rate limiting reaction that controls the overall pathway
- They may also catalyze a reaction unique to that pathway known as committed step

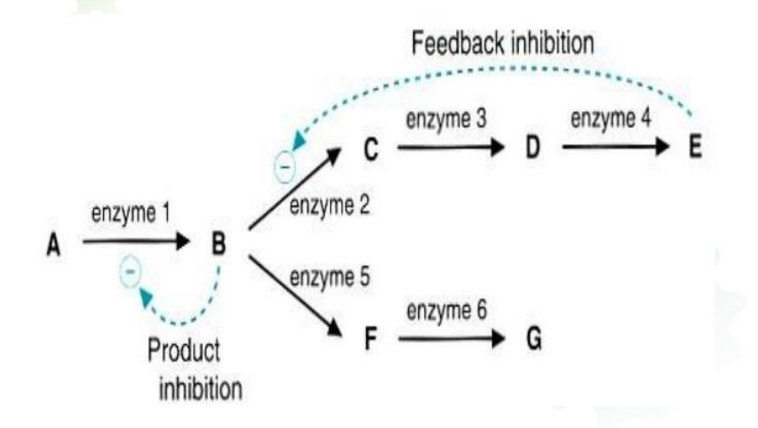
Feedback inhibition:

 When the end-product of a metabolic pathway exceeds its conc. limit, it inhibits the regulatory enzyme to normalize the pathway (feedback inhibition)

- Feed positive activation:
 - When the end-product of a metabolic pathway is below its conc. limit, it activates the regulatory enzyme to normalize the pathway

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Feedback Inhibition



Types of regulation

Allosteric enzyme regulation

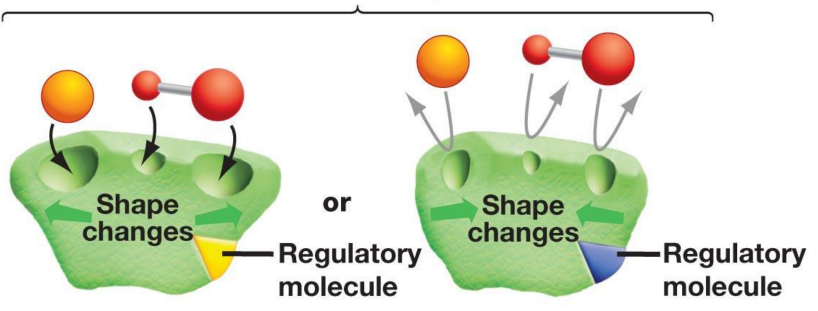
Enzymes in metabolic pathways are regulated by certain compounds (ligands)
These ligands do not bind to active site
They bind to other site (regulatory site) on the enzyme (allosteric enzymes)
The term "allosteric" came from Greek word "allos" meaning "other"

Cooperative binding
 Binding of a ligand to a regulatory site affects binding of the same or of another ligand to the enzyme
 This is called cooperative binding

- Binding of a ligand causes a change in the active site of enzyme
- This causes a change in the binding affinity of enzyme for the substrate



(b) Allosteric regulation



Allosteric activation

The active site becomes available to the substrates when a regulatory molecule binds to a different site on the enzyme.

Allosteric deactivation

The active site becomes unavailable to the substrates when a regulatory molecule binds to a different site on the enzyme.

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The effect of a ligand may be positive (activation) or negative (inhibition) Positive: increased E, S affinity ◆ Negative decreased E, S affinity Most allosteric enzymes are oligomers (two or more polypeptide chains or subunits)

The subunits are known as protomers

Two types of interactions occur in allosteric enzymes:

- ♦ Homotropic
- ♦ Heterotropic
- Homotropic: Effect of one ligand on the binding of the same ligand
- A regulatory enzyme controlled by its own substrate
- Heterotropic: Effect of one ligand on the binding of a different ligand

Enzymatic diagnosis and prognosis of diseases

Enzymes are used clinically in three ways: As indicators of enzyme activity or conc. in body fluids (serum, urine) in the diagnosis/prognosis of diseases As analytical reagents in measuring activity of other enzymes or compounds in body fluids

As therapeutic agents

The most commonly used body fluids for measuring enzyme activity are serum and plasma

There are:

Plasma-specific enzymes

Nonplasma-specific enzymes

Serum markers in the diagnosis of diseases

Heart disease
Pancreatic diseases
Liver diseases

Take home message

 Enzymes are essential for all biochemical reactions in the body

- A number of diseases are treated by inhibiting specific enzymes
- Many enzymes are used as biomarkers for diagnosis of diseases

References

Lippincott's Biochemistry
5th Edition, pp 53-68, Lippincott Williams & Wilkins, New York, USA