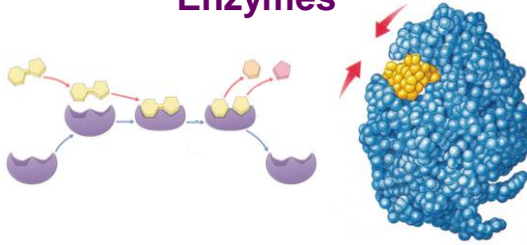


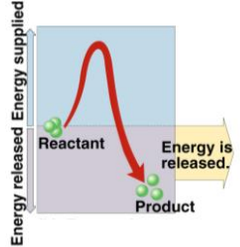
Chapter 8.3 – 8.5

Enzymes



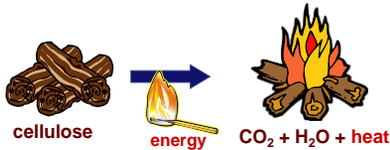
Spontaneous reactions?

- If reactions are “downhill”, why don’t they just happen spontaneously?
 - ♦ because covalent bonds are stable



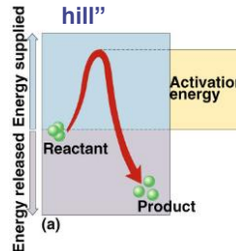
Activation Energy

- Breaking down large molecules requires an initial input of energy
 - ♦ activation energy
 - ♦ large biomolecules are stable
 - ♦ must absorb energy to break bonds



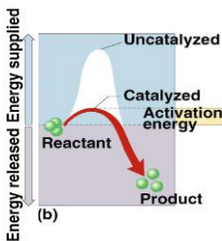
Activation Energy

- the amount of energy needed to destabilize the bonds of a molecule
 - ♦ moves the reaction over an “energy hill”



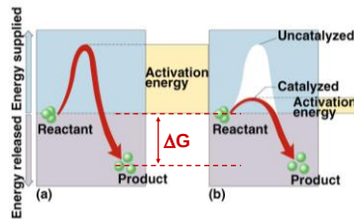
Reducing Activation Energy

- Catalysts
 - ♦ reducing the amount of energy to start a reaction



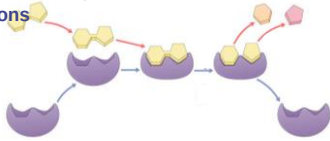
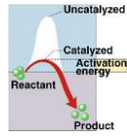
Catalysts

- So what’s a cell to do to reduce activation energy?
 - ♦ **get help!** ... chemical help... **ENZYMES**



Enzymes

- **Biological catalysts**
 - ◆ proteins (& RNA—ribozymes!)
 - ◆ **facilitate chemical reactions**
 - increase rate of reaction without being consumed
 - reduce activation energy
 - don't change free energy (ΔG) released or required
- ◆ required for most biological reactions
- ◆ **highly specific**
 - thousands of different enzymes in cells
- ◆ 'control' reactions



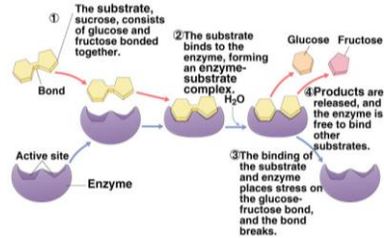
Enzymes & Substrates

substrate

- reactant which binds to enzyme
- enzyme-substrate complex: temporary association

product

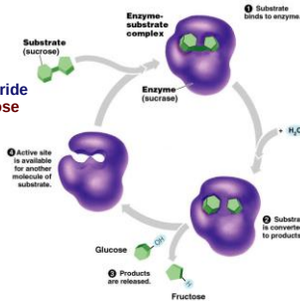
- end result of reaction



Enzymes & Substrates

Enzyme + substrates → products

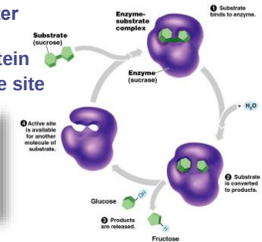
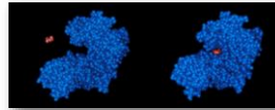
- ◆ **sucrase**
 - enzyme breaks down sucrose
 - binds to sucrose and breaks disaccharide into fructose & glucose
- ◆ **DNA polymerase**
 - enzyme builds DNA
 - adds nucleotides to a growing DNA strand



Lock and Key Model

Simplistic model of enzyme action

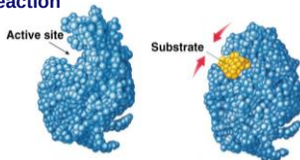
- ◆ 3-D structure of enzyme fits substrate
- ◆ **Active site**
 - ◆ enzyme's catalytic center
 - ◆ pocket or groove on surface of globular protein
 - ◆ substrate fits into active site



Induced Fit Model

More accurate model of enzyme action

- ◆ 3-D structure of enzyme fits substrate
- ◆ as substrate binds, enzyme changes shape leading to a tighter fit
 - "conformational change"
 - bring chemical groups in position to catalyze reaction



How does it work?

Variety of mechanisms to lower activation energy & speed up reaction

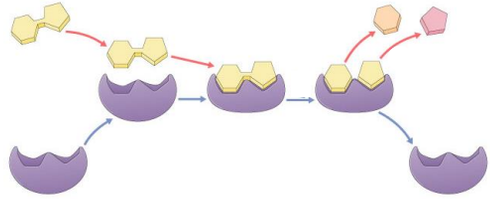
- ◆ active site orients substrates in correct position for reaction
 - enzyme brings substrate closer together
- ◆ active site binds substrate & puts stress on bonds that must be broken, making it easier to separate molecules
- ◆ groups near the active site can add a chemical charge for re-dox reactions

Specificity of Enzymes

- Reaction **specific**
 - ◆ each enzyme is substrate-specific
 - due to fit between active site & substrate
 - ◆ substrates held in active site by weak interactions
 - H bonds
 - ionic bonds
 - ◆ enzymes named for reaction they catalyze
 - **sucrase** breaks down sucrose
 - **proteases** break down proteins
 - **lipases** break down lipids
 - **DNA polymerase** builds DNA
 - **pepsin** breaks down proteins (**polypeptides**)

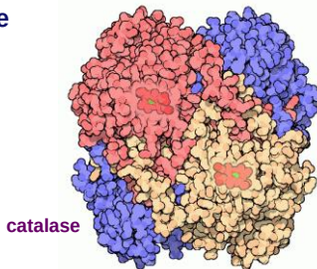
Reusable

- Not consumed in reaction!
 - ◆ single enzyme molecule can catalyze thousands or more reactions per second
 - ◆ enzymes **unaffected** by the reaction

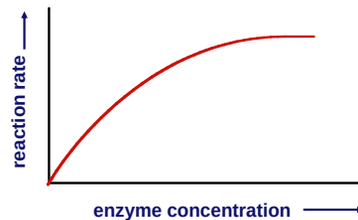


Factors Affecting Enzymes

- Enzyme concentration
- Substrate concentration
- Temperature
- pH
- Salinity
- Activators
- Inhibitors

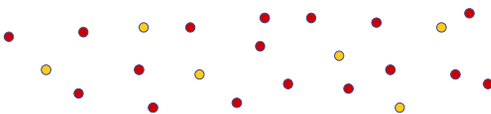


[Enzyme]

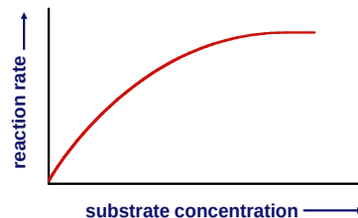


[Enzyme]

- Effect on rates of enzyme activity
 - ◆ as \uparrow enzyme = \uparrow reaction rate
 - more enzymes = more frequently collide with substrate
 - ◆ reaction rate levels off
 - substrate becomes limiting factor
 - not all enzyme molecules can 'find' substrate

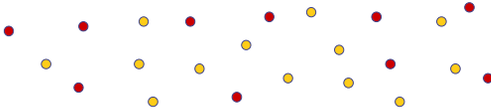


[Substrate]

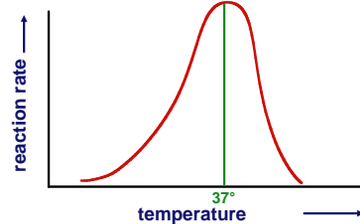


[Substrate]

- **Effect on rates of enzyme activity**
 - ◆ as ↑ substrate = ↑ reaction rate
 - more substrate = more frequently collide with enzymes
 - ◆ reaction rate levels off
 - all enzymes have active site engaged
 - enzyme is **saturated**; is the limiting factor
 - maximum rate of reaction



Temperature

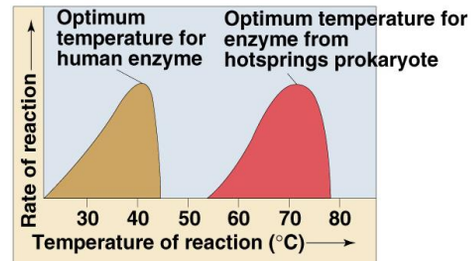


Temperature

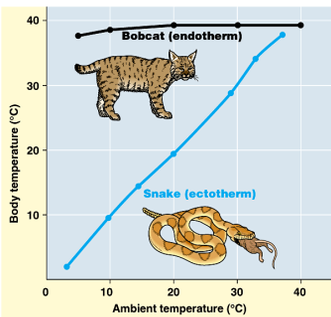
- **Effect on rates of enzyme activity**
 - ◆ **Optimum T°**
 - greatest number of molecular collisions
 - human enzymes = 35°- 40°C (body temp = 37°C)
 - ◆ **Increase beyond optimum T°**
 - increased agitation of molecules disrupts bonds
 - ◆ H, ionic = weak bonds
 - **denaturation** = lose 3D shape (3° structure)
 - ◆ **Decrease T° below optimum T°**
 - molecules move slower
 - decrease collisions

Temperature

- **Different enzymes functional in different organisms**

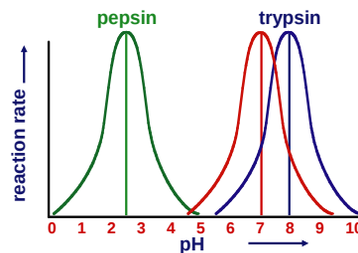


How do ectotherms do it?



ISOZYMES!
different enzymes with different chemical compositions and physical properties, but control the same reaction.

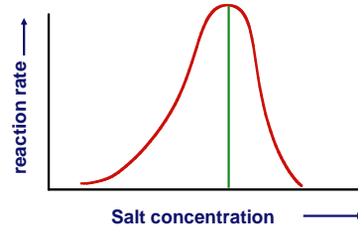
pH



pH

- **Effect on rates of enzyme activity**
 - ♦ **pH changes**
 - changes charges (add or remove H⁺)
 - disrupt bonds, disrupt 3D shape
 - ♦ affect 3^o structure
 - ♦ **most human enzymes = pH 6-8**
 - depends on localized conditions
 - pepsin (stomach) = pH 3
 - trypsin (small intestines) = pH 8

Salinity (salt concentration)

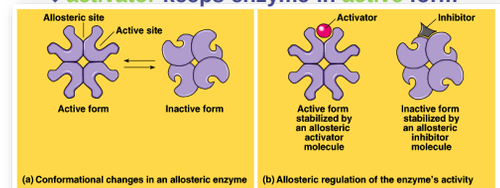


Salinity (salt concentration)

- **Effect on rates of enzyme activity**
 - ♦ protein shape (conformation)
 - depends on attraction of charged amino acids
 - ♦ salinity changes
 - change [inorganic ions]
 - changes charges (add + or -)
 - disrupt bonds, disrupt 3D shape
 - ♦ affect 3^o structure
 - ♦ **enzymes intolerant of extreme salinity**
 - **Dead Sea is called dead for a reason!**

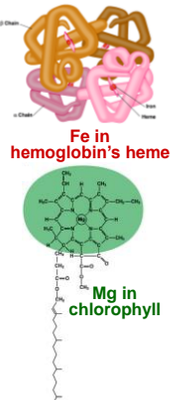
Action of Allosteric Control

- **Inhibitors & Activators**
 - ♦ regulatory molecules attach to allosteric site (**not active site**) causing conformational (shape) change
 - ♦ **inhibitor** keeps enzyme in **inactive form**
 - ♦ **activator** keeps enzyme in **active form**



Activators

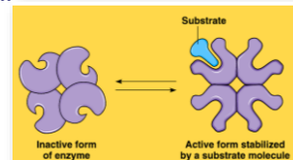
- **Compounds which help enzymes**
- **Prosthetic groups**
 - ♦ non-amino acid groups bound to enzymes
 - heme group in hemoglobin
- **Cofactors**
 - ♦ non-protein, small inorganic compounds & ions
 - Mg⁺⁺, K⁺, Ca⁺⁺, Zn, Fe, Cu
 - bound in enzyme molecule
- **Coenzymes**
 - ♦ non-protein, organic molecules
 - bind temporarily or permanently to enzyme near active site
 - ♦ many vitamins
 - NAD (niacin; B3)
 - FAD (riboflavin; B2)
 - Coenzyme A



Substrate Cooperativity

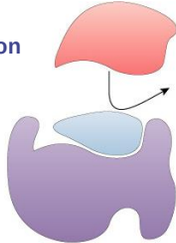
- **Substrate acts as an activator**
 - ♦ substrate causes conformational change in enzyme
 - induced fit
 - ♦ favors binding of substrate at 2nd site
 - ♦ makes enzyme more active & effective
 - ex: hemoglobin

- 4 polypeptide chains:
 - bind 4 O₂;
 - 1st O₂ binds
 - makes it easier for other 3 O₂ to bind



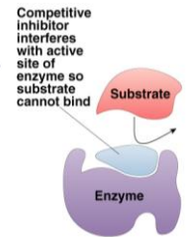
Inhibitors

- Regulation of enzyme activity
 - ◆ other molecules that affect enzyme activity
- Selective inhibition & activation
 - ◆ competitive inhibition
 - ◆ noncompetitive inhibition
 - ◆ irreversible inhibition
 - ◆ feedback inhibition



Competitive Inhibitor

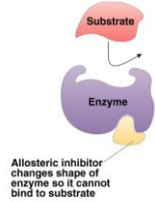
- Effect
 - ◆ inhibitor & substrate “compete” for active site
 - ex: **disulfiram** (Antabuse) to overcome alcoholism
 - ◆ overcome by increasing substrate concentration
 - saturate solution with substrate so it out-competes inhibitor for active site on enzyme
 - ex: **methanol poisoning**



(a) Competitive inhibition

Non-Competitive Inhibitor

- Effect
 - ◆ inhibitor binds to site other than active site
 - **allosteric site**
 - called **allosteric inhibitor**
 - ◆ ex: some anti-cancer drugs (methotrexate & FdUMP) inhibit enzymes involved in synthesis of nucleotides & therefore in building of DNA = stop DNA production, stopping abnormal division
 - ◆ ex: heavy metal poisoning
 - ◆ ex: cyanide poisoning
 - causes enzyme to have a conformational shape change
 - renders active site unreceptive

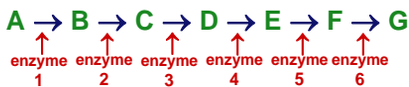


(b) Noncompetitive inhibition

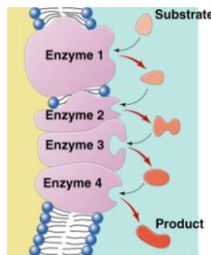
Irreversible Inhibition

- Inhibitor permanently binds to enzyme
 - ◆ competitor
 - permanently binds to active site
 - ◆ allosteric
 - permanently changes shape of enzyme
 - ex: nerve gas, sarin, many insecticides (malathion, parathion...)
 - ◆ DIFP (diisopropylphosphorofluoridate) is an...
 - ◆ acetylcholinesterase inhibitor—doesn't breakdown the neurotransmitter, acetylcholine, which is vital for muscle contraction

Metabolic Pathways

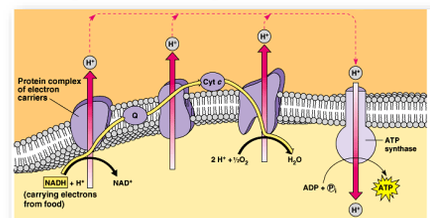


- Chemical reactions of life are organized in pathways
 - ◆ divide chemical reaction into many small steps
 - efficiency
 - control = regulation



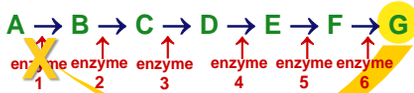
Efficiency

- Groups of enzymes organized
 - ◆ if enzymes are embedded in membrane they are arranged sequentially
- Link endergonic & exergonic reactions



Feedback Inhibition

- Regulation & coordination of production
 - ◆ product is used by next step in pathway
 - ◆ final product is inhibitor of earlier step
 - allosteric inhibitor of earlier enzyme
 - feedback inhibition
 - ◆ no unnecessary accumulation of product



G is an allosteric inhibitor of enzyme 1

Feedback Inhibition

- Example
 - ◆ synthesis of amino acid, isoleucine from amino acid, threonine

