Metabolism

Bioenergetics:

- It is the study of the energy changes accompanying biochemical reactions.
- Biologic systems are essentially isothermic and use chemical energy to power living processes.

Free energy:

- G is energy that can do work when temperature and pressure are uniform, as in a living cell.

The free energy change ($\Delta G0$)

- It is the difference between the free-energy content of the products and the free-energy content of the reactants under standard conditions. It depends on the nature and concentration of initial reactants and the final products.

Stages of chemical reactions

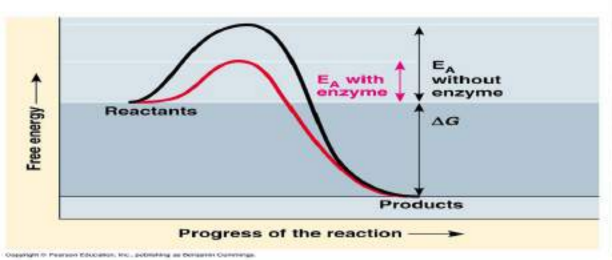
1- Activation Energy:

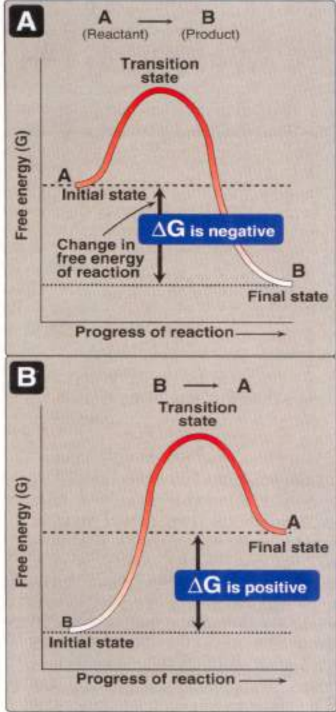
- Amount of energy that reactant molecules must absorb to start a reaction.
- This energy is usually provided in the form of heat absorbed by the reactant molecules form the surroundings.

2- Transition State:

- Unstable condition of reactant molecules that have absorbed sufficient free energy to react.

3- Products





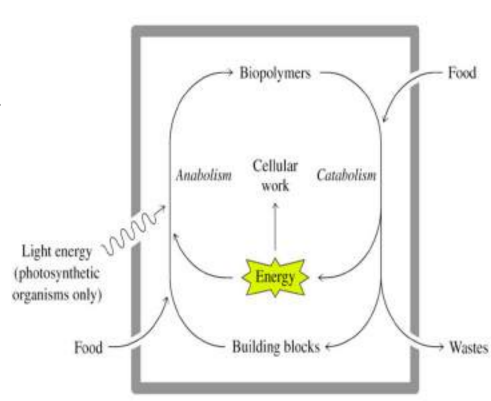
Metabolic pathways can be grouped into two pathways:

1- Catabolic reactions:

- Degrade molecules to create smaller molecules and energy (oxidation reactions releasing free energy to be transformed to ATP)

2- Anabolic reactions:

- Synthesize molecules for cell maintenance, growth and reproduction (reduction reactions utilizing energy in ATP molecules)
- Catabolism and anabolism are tightly coupled together by energy.



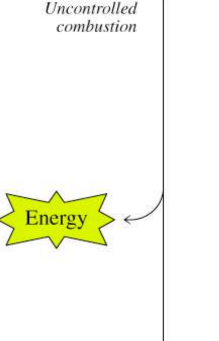
Metabolism proceeds by discrete steps

- Multiplestep pathways permit control of energy input and output
- Catabolic multi-step pathways provide energy in smaller stepwise amounts
- Each enzyme in a multi-step pathway usually catalyzes only one single step in the pathway
- Control points occur in multistep pathways
- Metabolic pathways are regulated to permit organisms to respond to changing conditions
- Most pathways are irreversible
- Flux: flow of material through a metabolic pathway which depends upon: 1-Supply of substrates
 - 2- Removal of products
 - 3- Pathway enzyme activities

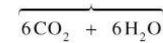
Single-step versus multi-step pathways

Multistep

Glucose + 6O₂



A multistep enzyme pathway releases energy in smaller amounts that can be used by the cell



Levels of metabolism regulation

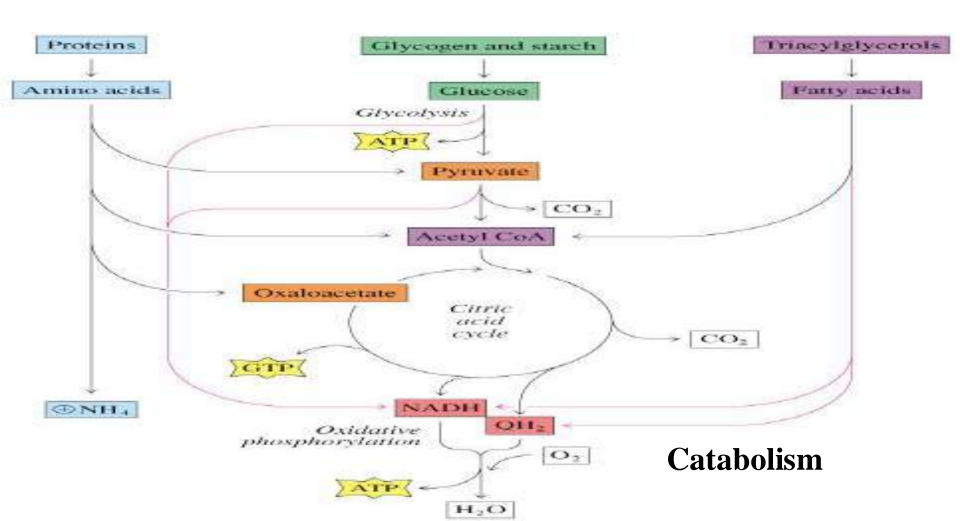
- 1- Nervous system.
- 2- Endocrine system.
- 3- Interaction between organs.
- 4- Cell (membrane) level.
- 5- Molecular level

Stages of metabolism

Catabolism

- **Stage I**. Breakdown of macromolecules (proteins, carbohydrates and lipids to respective building blocks.
- **Stage II**. Amino acids, fatty acids and glucose are oxidized to common metabolite (acetyl CoA)
- **Stage III**. Acetyl CoA is oxidized in citric acid cycle to CO₂ and water. As result reduced cofactor, NADH₂ and FADH₂, are formed which give up their electrons. Electrons are transported via the tissue respiration chain and released energy is coupled directly to ATP synthesis.

- Catabolism is characterized by convergence of three major routs toward a final common pathway.
- Different proteins, fats and carbohydrates enter the same pathway-TCA cycle.

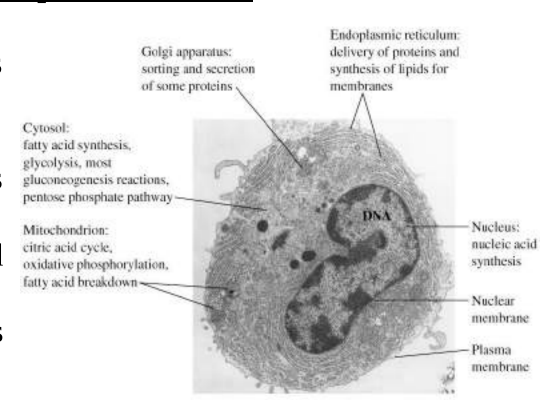


Anabolism can be divided into stages

- Anabolic pathways are characterized by divergence.
 - Monosaccharide synthesis begin with CO₂, oxaloacetate, pyruvate or lactate, then, polysaccharides.
 - Amino acids are synthesized from acetyl CoA, pyruvate or keto acids of Krebs cycle, then, proteins
 - Fatty acids are constructed from acetyl CoA, then, fats

Compartmentation of metabolic processes in cell

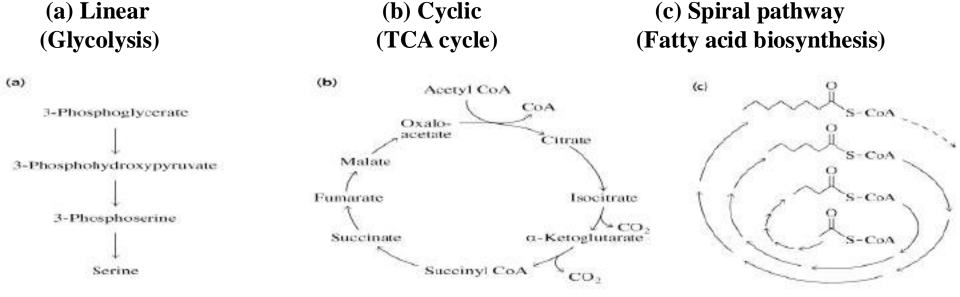
- permits:
- 1- Separate pools of metabolites within a cell
- 2- Simultaneous operation of opposing metabolic pathways
- 3- High local concentrations of metabolites such as fatty acid synthesis enzymes (cytosol), fatty acid breakdown enzymes (mitochondria)



The chemistry of metabolism

- There are about 3000 reactions in human cell.
- All these reactions are divided into six categories:
- 1- Oxidation-reduction reactions
- 2- Group transfer reactions
- 3- Hydrolysis reactions
- 4- Nonhydrolytic cleavage reactions
- 5- Isomerization and rearrangement reactions
- 6- Bond formation reactions using energy from ATP

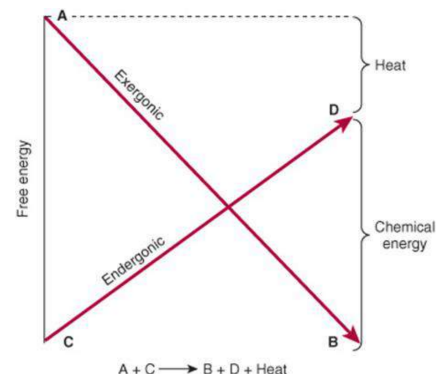
Metabolic pathway may be:



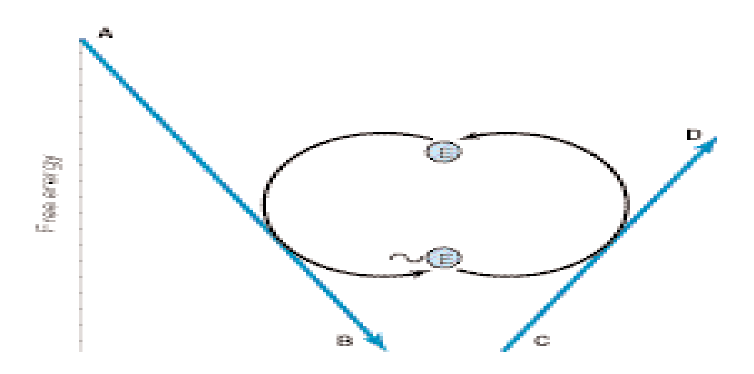
- In practice, an endergonic process cannot exist independently but must be a component of a coupled exergonic- endergonic system. One possible mechanism of coupling could be observed if **common obligatory intermediate** (I) took part in both reactions.

$$A + C \rightarrow I \rightarrow B + D$$

- The coupling concept is provided by forming an intermediate carrier through dehydrogenation/hydrogenations reactions



- An alternative method of coupling an exergonic to an endergonic process is to synthesize a compound of high-energy potential in the exergonic reaction and to incorporate this new compound into the endergonic reaction, thus affecting a transference of free energy from the exergonic to the endergonic pathway.
- -The biologic advantage of this mechanism is that the compound of high potential energy, (~E), unlike I in the previous system.



The Intermediate value for the free energy of hydrolysis of ATP

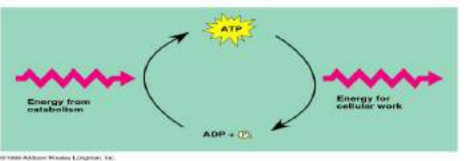
- The tendency of each of the phosphate groups to transfer to a suitable acceptor may be obtained from the $\Delta G0$ of hydrolysis at 37 °C.
- -The value for the hydrolysis of the terminal phosphate of ATP divides the list of energy compounds into two groups:
- Low-energy phosphates, exemplified by the ester phosphates found in the intermediates of glycolysis, have $\Delta G0$ values smaller than that of ATP, while in high-energy phosphates the value is higher than that of ATP.
- High-energy phosphates compounds, including ATP, are usually anhydrides (1-phosphate of 1,3-biphosphoglycerate), enolphosphates (phosphoenolpyruvate), and phosphoguanidines (creatine phosphate, arginine phosphate).

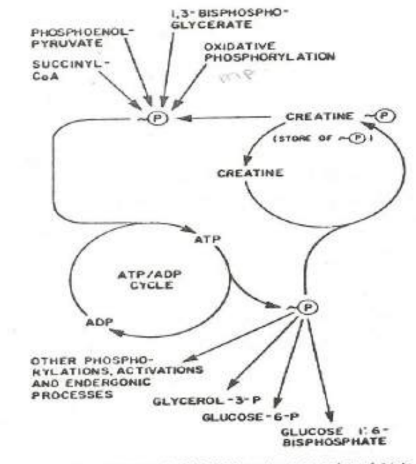
-The high free energy change on hydrolysis of ATP is due to relief of charge repulsion of adjacent negatively charged oxygen atoms.

- Other "high-energy compounds" are thiol esters as coenzyme A (acetyl-CoA), acyl carrier protein, amino acid esters involved in protein synthesis, S- adenosylmethionine (SAM), UDPGlc (uridine diphosphate glucose), and PRPP (5-phosphoribosyl-1-pyrophosphate).

Compound	kJ/mol	kcal/mol
Phosphoenolpyruvate	-61.9	-14.8
Carbamoyl phosphate	-51.4	-12.3
1,3-Bisphosphoglycerate	-49.3	-11.8
Creatine phosphate	-43.1	-10.3
$\mathbf{ATP} \rightarrow \mathbf{ADP} + \mathbf{Pi}$	-30.5	-7.3
$ADP \rightarrow AMP + Pi$	-27.6	-6.6
Pyrophosphate	-27.6	-6.6
Glucose 1-phosphate	-20.9	-5.0

- ATP powers cellular work by coupling exergonic reactions to endergonic reactions
- A cell does three main kinds of work: Mechanical, Transport and Chemical
- -To do work, cells manage energy resources by energy coupling, the use of an exergonic process to drive an endergonic one
- ATP is the cell's energy shuttle providing energy for cellular functions





Role of ATPIADP cycle in transfer of highenergy phosphate

Sources of ATP.

ATP levels are maintained through several processes:

1. Adenylate kinase.

- ATP has two "high-energy" phosphate groups β and γ .
- Splitting off the γ phosphate group of ATP yields ADP and inorganic phosphate. ATP \longrightarrow ADP + pi
- Splitting off both high-energy groups in one step yields AMP and inorganic pyrophosphate (ppi). **ATP AMP** + **ppi**
- Adenylate kinase, an enzyme found in all tissues, catalyzes a transfer of the energy-rich phosphate bond from one ADP molecule to another, giving ATP and AMP.
- The conversion is very rapid in muscle and liver.

2ADP \leftarrow ATP + AMP

- -AMP levels are crucial in adjusting the balance between carbohydrate and fatty acid metabolism in varying physiological situations.
- AMP is an active intracellular signal substance.
- -AMP is also an activator of glycogen mobilization and, therefore, sugar metabolism.

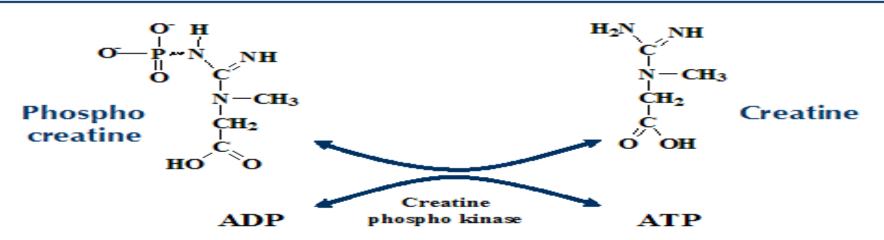
2. Creatine Phosphokinase / Phosphocreatine.

- Most of our body tissues contain phosphocreatine at concentrations approximately three times that of ATP.
- Phosphocreatine is a reserve source of high-energy phosphate.
- -This reserve can be transferred to ADP, thus forming ATP to replace that used by working muscle.
- -While the creatine phosphokinase reaction is the most rapid ATPyielding reaction we possess, the amount of ATP which is produced is quite small.
- Muscle tissues have about 5 mmol/l ATP and approximately 17-20 mmol/l of creatine phosphate.
- Under extreme work (sprinting, for example) the phosphocreatine reserves are used up in about 30-40 seconds.
- However, "seconds do count" in sport. During those few seconds muscles can and do work with "explosive force".

TABLE 1.1 THE ENERGY SYSTEMS AND THEIR APPROXIMATE CONTRIBUTIONS TO VARIOUS DURATIONS OF EXERCISE AT MAXIMAL INTENSITY (1)

ENERGY SYSTEM	0–10 seconds	
Phosphocreatine system		
Phosphocreatine system and glycolytic system (slow)	10-30 seconds	
Glycolytic system (fast)	30 seconds-2 minutes	
Glycolytic system (fast) and oxidative system	2-3 minutes	
Oxidative system	< 3 minutes and rest	

Creatine Phosphate-ATP interaction



1.3 CHARACTERISTICS OF THE PHOSPHOCREATINE SYSTEM

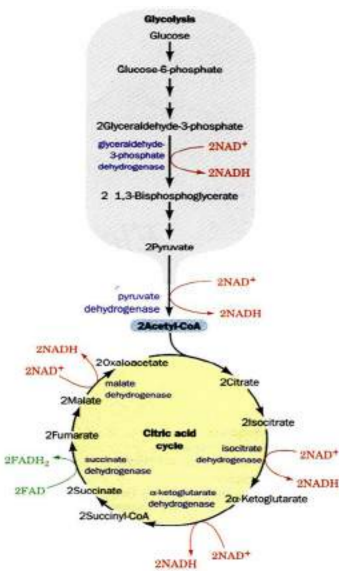
- It involves only one chemical step.
- 2. It is catalyzed by the enzyme creatine kinase (CK).
- 3. Its chemical reaction is very fast.
- One ATP is generated per phosphocreatine molecule.
- The reaction lasts for 5 to 10 seconds at maximal intensity.
- 6. It is anaerobic.
- Fatigue is associated with the depletion of phosphocreatine.
- It is the dominant energy system in speed and explosive power events.

3. Anaerobic Metabolism.

- This is a rapid cytosolic formation of ATP driven by oxidation of glucose (or glucosyl groups from glycogen) to pyruvate and lactate.
- ATP formation through cytosolic glycolysis proceeds with a speed equal to about 50% of that we see using creatine phosphate and creatine phosphokinase.
- Only two ATP molecules result for each glucose molecule that is processed, three ATPs are formed for each glucosyl group that derived from glycogen.
- The disadvantage is that a lot of lactic acid is produced and accumulates in the working muscle and lipids cannot be used as substrates for anaerobic metabolism.
- Muscles exhaust their stored glycogen and take up so much glucose from the blood resulting in hypoglycemia and CNS malfunction.

4. Aerobic Metabolism.

- All of our cells, with the important exception of blood cells, contain mitochondria which use oxygen and form water while oxidizing our "food".
- Mitochondrial substrate is acetyl-CoA coming from all types of nutrients
- Approximately 40% of the energy released in this process is trapped in the terminal phosphate group in ATP.
- The rest of the energy in acetyl-CoA escapes as heat to maintain our temperature
- It is a slow process, produces 10 moles of ATP for each mole of acetyl-CoA.

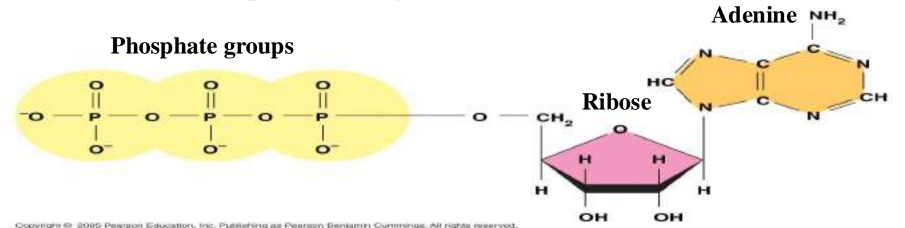


ATP (adenosine triphosphate)

- Compounds with high energy bonds release large quantities of free energy on hydrolysis.
- -The most important parts of the ATP molecule are its two phosphodiester bonds (high energy bonds).
- Breaking down either of these bonds is accompanied by the release of energy (7.3 Kcal/mol for each bond).

Importance of ATP as a source of energy:

- 1- Synthesis of macromolecules: such as DNA and RNA, protein,etc
- 2- Support the endergonic reaction in metabolic pathways.
- 3- Important for active transport across membranes.
- 4- Important for muscle contraction...etc.
- 5- transmission of impulses along neurons.



Electron transport chain (ETC)

Oxidation reduction reactions (Redox reactions)

- Commonly the oxidation reactions are accompanied by reduction reactions and they are called redox reaction.
- Redox reactions are accompanied by energy liberation, necessary for the cells.
- In the redox reaction. H_2 is oxidized while, O_2 is reduced, and if occurs it will be accompanied by a massive energy explosion.

$$2H_2 + O_2 \longrightarrow 2H_2O + energy$$

- Instead of massive energy is liberated, hydrogen must be transferred to oxygen in gradual steps. Thus, small fractions of energy are liberated and stored for further use

Redox Potential (electron affinity)

- Oxygen has the highest electron affinity i.e. highest redox potential.
- Hydrogen has the lowest electron affinity i.e. lowest redox potential.

Redox chain:

- It is a chain of different compounds of increasing redox potential between hydrogen and oxygen.

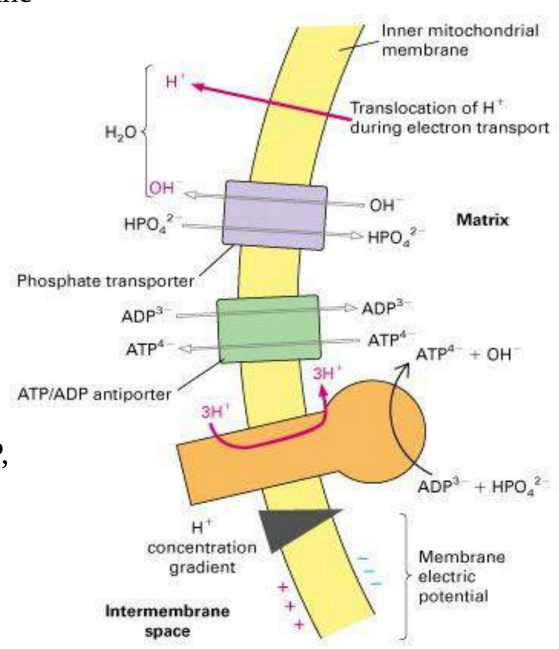
$$A \rightarrow B \rightarrow C \rightarrow D....$$

- Each component of redox chain has a redox potential higher than hydrogen and lower than oxygen.
- During hydrogen (H⁺ and electron) transfer through different components of the redox chain, energy is liberated in steps and in small amounts to be utilized.

Electron transport chain (ETC)

- It is a system of electron carriers located in the innermitochondrial membrane, oxidizes the reduced cofactors by transferring electrons in a series of steps to O₂ (the terminal electron acceptor).
- Free energy released by these oxidation reactions is used to drive the synthesis of ATP.
- Each component of the chain can accept electrons from the preceding carrier and transfer them to the following one.
- A variety of substances (carbohydrates, fatty acids and amino acids) can use respiratory chain as a final pathway as they give electrons to the oxidized NAD⁺ and FAD⁺ to form the energy rich reduced coenzymes NADH+H⁺, FADH₂.
- NADH+H⁺ and FADH₂ give hydrogen and a pair of electrons to electron carriers collectively, called the respiratory chain components.

- Outer mitochondrial membrane is permeable to most ions as
 O₂, CO₂, NH₃ and monocarboxylic acids.
- Di- and tricarboxylic acids need special transporters.
- ATP and ADP need special transporter to allow ADP in and ATP out of mitochondria.
- Inner membrane is impermeable to most ions & molecules: H⁺, Na⁺, K⁺, ATP, ADP, pyruvate.
- Matrix contains enzymes for oxidation of pyruvate., A.A.s , F.A.s and TCA.



Organization of Electron transport chain

- -The inner mitochondrial membrane contains four enzymatic complexes (I, II, III, IV) and complex V catalyses ATP synthesis, arranged in order of increasing electronegativity (weakest to strongest)
- Each complex accepts or donates electrons to relatively mobile electron carriers as coenzyme Q and cytochrome C.
- Oxidative phosphorylation starts by entry of electrons into the respiratory chain.
- Most of these electrons arise by the action of dehydrogenases that collect electrons from catabolic pathways and pass them to the electron acceptors NAD and FAD.
- As electrons are passed down the respiratory chain, they lose much of their free energy.

- Part of this energy can be captured and stored by the production of ATP from ADP and inorganic phosphate (Pi).
- The process is called oxidative phosphorylation.
- The remainder of the free energy not trapped as ATP is released as heat.

Components of the respiratory chain

- -With the exception of coenzyme Q, all members of this chain are proteins.
- All are embedded in the inner mitochondrial membrane.

Complex I

- Contains an enzyme called NADH dehydrogenase.
- Its coenzyme is FMN (can accept two hydrogen atoms to become FMNH₂)
- It contains several iron and sulfur atoms (iron sulfur protein).
- NAD+ is reduced to NADH+H⁺ by dehydrogenases that remove hydrogen atoms from their substrates.

Complex II

- The entry point of FADH₂ (its coenzyme is FAD).
- Contains an enzyme called: flavo protein dehydrogenase e.g. succinate dehydrogenase of TCA and acyl CoA dehydrogenase of β oxidation of fatty acids.
- It contains iron and sulfur atoms (iron sulfur protein).

Complex III

- It is cytochrome reductase complex, or cytochrome bc1 complex"
- Transfers electron from QH₂ to cytochrome C.
- Contains an enzyme cytochrome b.

Complex IV

- This complex contains cytochrome a, a3 and 2 copper atoms.
- Complex IV catalyzes the transfer of electrons from reduced cytochrome C to molecular oxygen.
- The copper atoms are crucial for such a transfer.

Ubiquinone "Coenzyme Q"

- It is a lipid soluble vitamin K derivative
- Coenzyme Q can accept hydrogen ions both from FMNH₂, produced by NADH dehydrogenase (complex I) and from FADH₂ which is produced by (complex II).
- It is freely diffusible between the lipid bilayer of inner mitochondrial membrane.

Cytochromes

- Cytochromes are proteins that contain an iron-containing heme group. This iron oscillates between ferric form (Fe⁺⁺⁺) when it losses an electron, and ferrous form (Fe⁺⁺) when it accepts an electron.
- All are integral membrane proteins with the exception of cytochrome C, a soluble free protein.

TABLE 19–3 The Protein Components of the Mitochondrial Electron-Transfer Chain

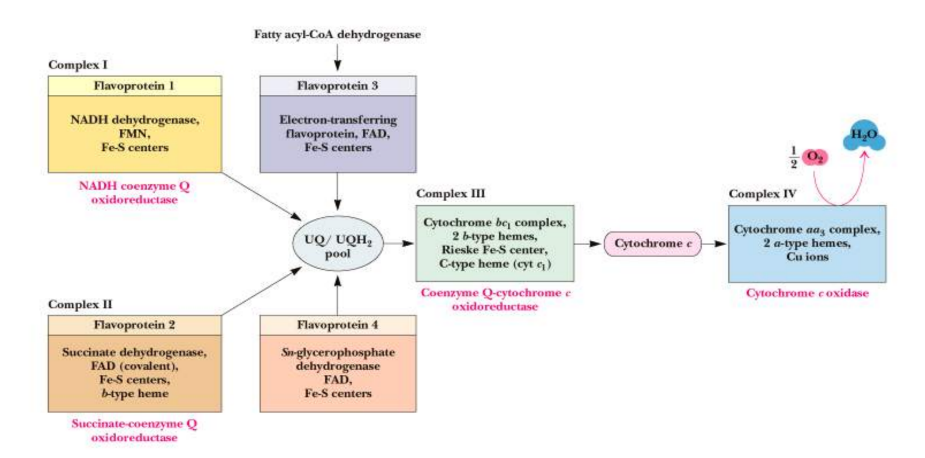
Enzyme complex/protein	Mass (kDa)	Number of subunits*	Prosthetic group(s)
I NADH dehydrogenase	850	43 (14)	FMN, Fe-S
II Succinate dehydrogenase	140	4	FAD, Fe-S
III Ubiquinone cytochrome c oxidoreductase	250	11	Hemes, Fe-S
Cytochrome c [†]	13	1	Heme
IV Cytochrome oxidase	160	13 (3-4)	Hemes; Cu_A , Cu_B

[&]quot;Numbers of subunits in the bacterial equivalents in parentheses.

- Cytochrome a3 contains copper in addition to iron and called cytochrome oxidase, it is the terminal component of the ETC.

[†]Cytochrome c is not part of an enzyme complex; it moves between Complexes III and IV as a freely soluble protein.

Garrett & Grisham: Biochemistry, 2/e Figure 21.4



- As electrons pass down the respiratory chain, they lose much of their free energy.
- Part of this energy can be captured and stored as ATP from {ADP and inorganic phosphate (Pi)}.
- -The process is called oxidative phosphorylation.
- -The non trapped free energy as ATP so, released as heat.

Oxidative phosphorylation

- Oxidative phosphorylation is a coupling process of oxidation and phosphorylation.
- -The flow of electrons from NADH to oxygen (oxidation) results in ATP synthesis by phosphorylation of ADP with inorganic phosphate (phosphorylation), therefore, there is coupling between oxidation and phosphorylation.
- Chemiosmotic hypothesis and membrane transport system can explain synthesis of ATP.

Chemiosmotic hypothesis:

-This hypothesis postulates that the transfer of electrons along the respiratory chain is accompanied by outward pumping of protons across the inner mitochondrial membrane.

Proton pump

- -The transport of electrons down the respiratory chain creates an energy which is used to transport H+ from mitochondrial matrix across inner mitochondrial membrane → inner mitochondrial space.
- -This process is carried out by complexes I, III, IV to create across the inner mitochondrial membrane:
 - An electrical gradient with more positive changes on the outside of the membrane than on the inside.
 - A pH gradient as the outside of the membrane is at lower pH than the inside.
- The energy generated is sufficient for ATP production.

INTERMEMBRANE SPACE

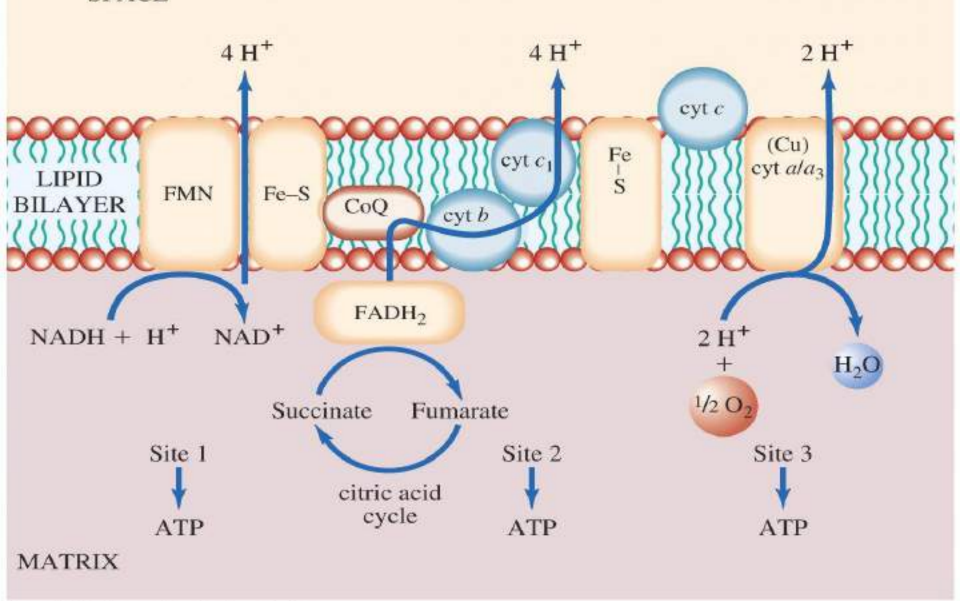


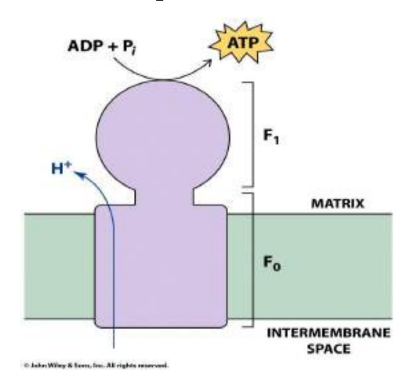
Figure 17-12 Concepts in Biochemistry, 3/e © 2006 John Wiley & Sons

ATP synthase (complex V)

- ATP synthase enzyme presents in the inner mitochondrial membrane, it is a phosphorylating enzyme complex and it is formed of 2 subunits:
 - $\mathbf{F_1}$ subunit which protrudes into matrix.
 - $\mathbf{F_0}$ subunit which presents in the membrane.
- -The energy stored in the electrochemical gradient is used to drive the synthesis of ATP by the movement of protons down the electrochemical gradient using **ATP synthase**.
- -The protons outside the inner mitochondrial membrane can re-enter the mitochondrial matrix by passing through channel (**F0-F1** complex) to pass by ATP synthase enzyme which is present in F1 subunit.
- This results in the synthesis of ATP from ADP + Pi.
- At the same time decreases the pH and electrical gradients.

ATP Synthase

- $\mathbf{F_0F_1}$ ATP Synthase uses the proton gradient energy for the synthesis of ATP
- Large transmembrane protein complex
- Faces into the mitochondrial matrix spans the IMM
- Composed of a "knob-and-stalk" structure
- F_0 (stalk) has a <u>proton channel</u> which spans the membrane.
 - Forms a proton pore
 - Membrane-spanning portion integral membrane protein
 - Made up of 4 different subunits
 - F₀ subunit composition: a₁b₂c₉₋₁₂ (c subunits form cylindrical, membrane-bound base)

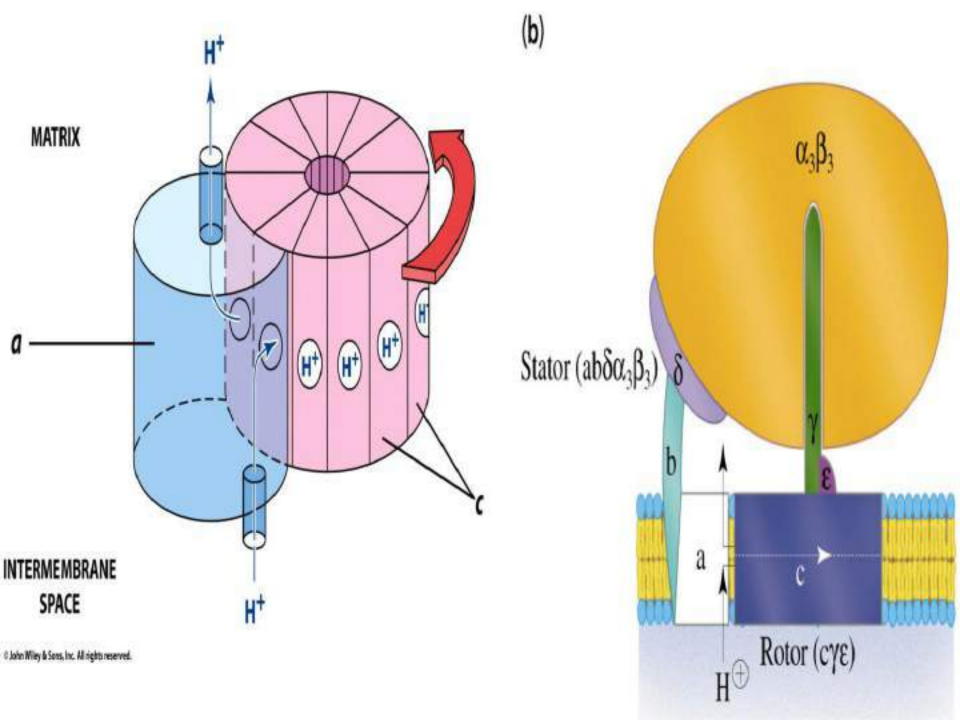


- F₁ (knob) contains the <u>catalytic (ATP-synthesizing) subunits</u>

- Where ATP synthesis takes place
- F_1 knobs: inner face of the inner mitochondrial membrane
 - (subunit composition: $\alpha 3\beta_3 \gamma \delta \epsilon$)
 - $\alpha_3 \beta_3$ oligomer of F_1 is connected to catalytic (C) subunits by a multisubunit stalk of γ and ϵ chains
- Protons passage through F₀ into the matrix is coupled to ATP formation
- Estimated passage of 3 H+/ATP synthesized
- F_0 is sensitive to **oligomycin**, it binds in the channel and blocks H^+ passage, thereby inhibiting ATP synthesis

Mechanism of ATP Synthase

- F₁-F₀ complex serves as the molecular apparatus for coupling H+ movement to ATP synthase.
- There are 3 active sites, one in each β subunit
- Passage of protons through the F_0 channel causes the rotor to spin in one direction and the stator to spin in the opposite direction
- Proton flow \rightarrow C unit rotates \rightarrow γ rotates \rightarrow conformation changes \rightarrow ATP synthesized



Regulation:

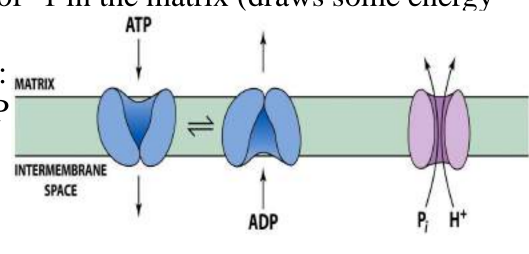
- Electrons do not flow unless ADP is present for phosphorylation
- Increased ADP levels cause an increase in catabolic reactions of various enzymes including:
 - Glycogen phosphorylase

- PFK-1

- Citrate synthase

ATP, ADP and Pi active transport across the inner mitochondrial membrane

- ATP is synthesized in the mitochondrial matrix
- ATP must be transported to the cytosol in exchange with ADP and Pi
- ADP/ATP carrier exchanges mitochondrial ATP⁴⁻ for cytosolic ADP³⁻
- -The exchange causes a net loss of -1 in the matrix (draws some energy from the H+ gradient)
- Adenine nucleotide translocase: MATRIX unidirectional exchange of ATP for ADP (antiport)
- Symport of Pi and H⁺ is electroneutral



The P:O Ratio

P:O ratio = molecules of ADP phosphorylated atoms of oxygen reduced

- Translocation of 3H⁺ required by ATP synthase for each ATP produced
- 1 H⁺ needed for transport of P_i, ADP and ATP
- Net: 4 H⁺ transported for each ATP synthesized

Calculation of the P:O ratio

Complex	I	III	IV
#H+ translocated/2e-	4	4	2

- Since 4 H⁺ are required for each ATP synthesized:

So,
$$P/O = (10 \text{ H}^+/ 4 \text{ H}^+) = 2.5 \text{ ATP/O}$$

For succinate substrate =
$$6 \text{ H}^+/\text{ O} (2e^-)$$

So,
$$P/O = (6 H^{+}/4 H^{+}) = 1.5 ATP/O$$

- It equals zero in presence of uncouplers.

Inhibitors of respiratory chain:

- Are compounds prevent the passage of electrons to bind a component of the chain (the three sites responsible for electrochemical potential difference), blocking the oxidation reduction reaction.
- There are specific sites for binding inhibitors:
- **Site I:** binding with complex I as barbiturates, rotenone (an insecticide) and piercidin A (an antibiotic).
- **Site II:** binding with complex III as antimycin A and dimercaprol.
- **Site III:** binding with complex IV as H₂S, cyanide (CN), carbon monoxide (CO) and sodium azide.
- Because electron transport and oxidative phosphorylation are tightly coupled, inhibition of the respiratory chain also inhibits ATP synthesis.

- **4-** ADP/ATP transporter inhibitors as atractyloside.
- **N.B.** Malonate which acts as competitive inhibitor of succinate dehydrogenase inhibits ETC through complex II.

Cyanide poisoning

- Cyanide is one of the most potent and rapidly acting poisons. Cyanide binds to cytochrome aa3 so, inhibits the oxidative phosphorylation at level of cytochrome oxidase complex (complex IV).
- -The energy production of cells will be blocked resulting in tissue asphyxia especially of central nervous system leading to death.

Uncouplers of oxidative phosphorylation

- -Uncouplers are a group of substances that interrupt (uncouple) oxidation and phosphorylation i.e. oxidation will proceed building proton gradients but will not result in ATP synthesis, so, energy released by electron transport will be lost in the form of heat.
- -This explains the hotness sensation after these substances intake.
- **1- Oligomycin:** This drug binds to the stalk of ATP synthase, closes the H channel and prevents re-entry of protons to the mitochondrial matrix.
- **2-2,4 dinitrophenol:** it increases the permeability of the inner mitochondrial membrane to proton causing decrease in the proton gradient.
- **3- Calcium and high doses of aspirin**: this explains the fever that accompanies toxic overdoses of these drugs.

4- Ionophores : e.g. Valinomycin and Nigericin.

- -They are lipophilic substances and they have the ability to make a complex with cations as potassium "K" and facilitate their transport into mitochondria and other biological membranes.
- -They inhibit phosphorylation because of pH gradient.
- **5- High level of Thyroxine**: as in thyrotoxicosis and bilirubin.
- 6- Snake venoms.
- **N.B.** Uncoupling proteins (UCP) = separate oxidation from ATP synthesis (the synthesis is interrupted) \rightarrow energy from H+ gradient is released as a **heat**

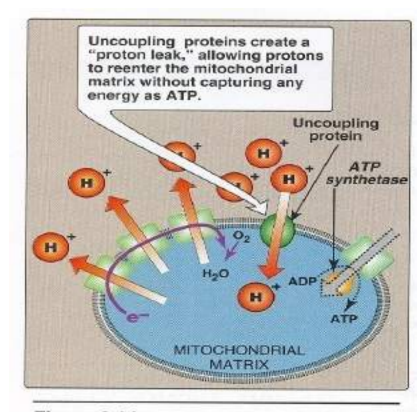


Figure 6.14

Transport of H⁺ across mitochondrial membrane by 2,4-dinitrophenol.

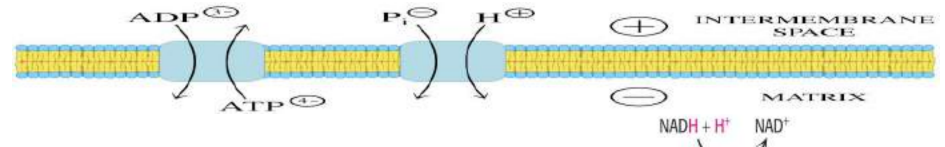
7- Thermogenin: (brown adipose tissues)

- -Thermogenin also called uncoupling protein 1, or UCP1 is an uncoupling protein found in the mitochondria of brown adipose tissue.
- It is used to generate heat by non-shivering thermogenesis.
- Non-shivering thermogenesis is the primary means of heat generation in hibernating mammals and in human infants.
- -The molecular mechanism of UCP1-mediated uncoupling is reasonably well understood; UCP1 allows protons to reenter the mitochondrial matrix without passing through F0-F1 complex (ATP synthase), allowing respiration (and hence heat production) to proceed in the absence of ATP synthesis.
- UCP1 is restricted to brown fat, where it provides a mechanism for the enormous heat-generating capacity of the tissue.

Membrane transport chain

-The inner mitochondrial membrane contains numerous transport proteins (carriers) that permit passage of specific molecules from the

cytosol to the mitochondrial matrix e.g. ADP-ATP carrier (adenine nucleotide translocase) which carries ADP from cytosol into mitochondria, while, carrying ATP from the matrix back to cytosol.



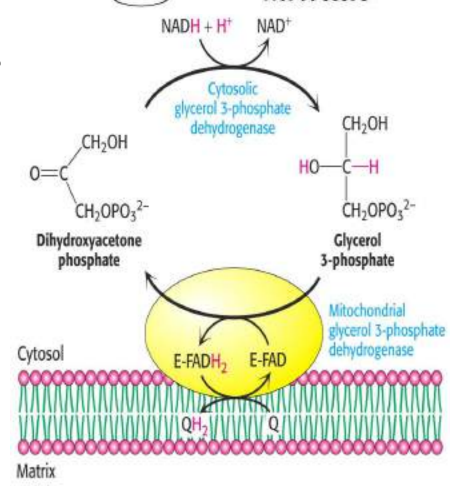
What about NADH from glycolysis?

- NADH made in cytosol
- Can't get into mitochondrial matrix

By 2 mechanisms:

A- In muscle and brain (Glycerol phosphate shuttle)

- Each NADH converted to FADH₂ inside mitochondrion
- FADH₂ enters later in the electron transport chain
- Produces 2 ATP



B- In liver and heart (Malate / aspartate shuttle)

- -NADH oxidized while reducing oxaloacetate to malate by malate dehydrogenase
- Malate crosses membrane
- Malate reoxidized to oxaloacetate
- Malate dehydrogenase
- NAD+ reduced to NADH
- NADH via electron transport yields 3ATP

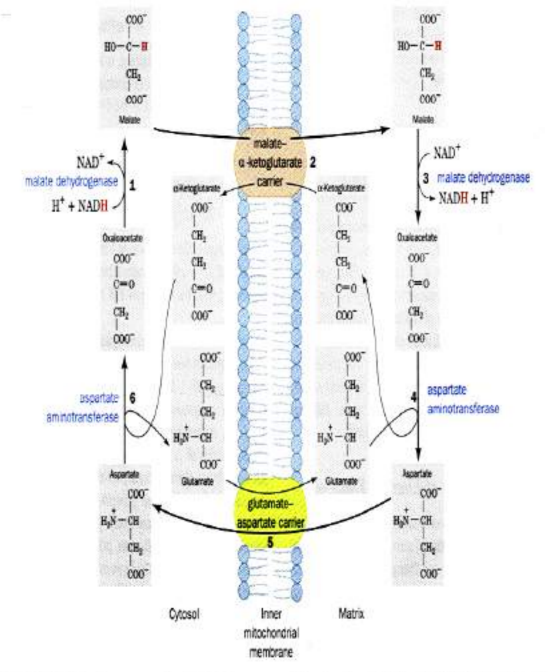


FIGURE 20-7. The malate-aspartate shuttle. The electrons of cytosolic NADH are transported to mitochondrial NADH

(shown in red as hydride transfers) in Steps 1 to 3. Steps 4 to 6 then serve to regenerate cytosolic oxaloacetate.

Inherited defects in oxidative phosphorylation

- Mitochondrial DNA (mtDNA) (37 genes) is maternally inherited as mitochondria of sperm cell do not enter the fertilized ova.
- Mitochondrial DNA (mtDNA) codes for **13 polypeptide** (of total 120) required for oxidative phosphorylation, 22 tRNA and 2 rRNA. (while the remaining are synthesized in the cytosol & are transported into the mitochondria).
- Defects of oxidative phosphorylation usually results from **alteration in mtDNA** (mutation rate 10 times more than that of nuclear DNA).
- -Tissues with greater ATP requirement (as CNS, skeletal muscles. & cardiac muscle, kidney & liver) are most affected by defects in oxidative phosphorylation.

Examples for diseases caused by mutations in mtDNA:

- 1- Mitochondrial myopathies (defective energy production → muscle cramping, weakness and severe fatigue).
- 2- Leber hereditary optic neuropathy (bilateral loss of vision due to optic nerve damage).

Substrate Level Phosphorylation:

- Very small amount of ATP molecules are produced
- Few reactions can form ATP at substrate level: e.g.
- 1. Glycolysis (phosphoglycerate kinase and pyruvate kinase)
- 2.TCA cycle (succinate thiokinase)

Respiratory control:

- -There is no mechanism for storage of ATP and ATP present at any moment is only enough to meet the need of our cell for only few seconds.
- For this reason, there must be an efficient and controlled way for the production of ATP.
- 1- Availability of ADP (ATP/ADP transporter may rate limiting at certain times.
- 2- Availability of electrons (↑ NADH/NAD and/or ↑ FADH2/FAD).
- 3- Availability of O2.
- 4- Insulin