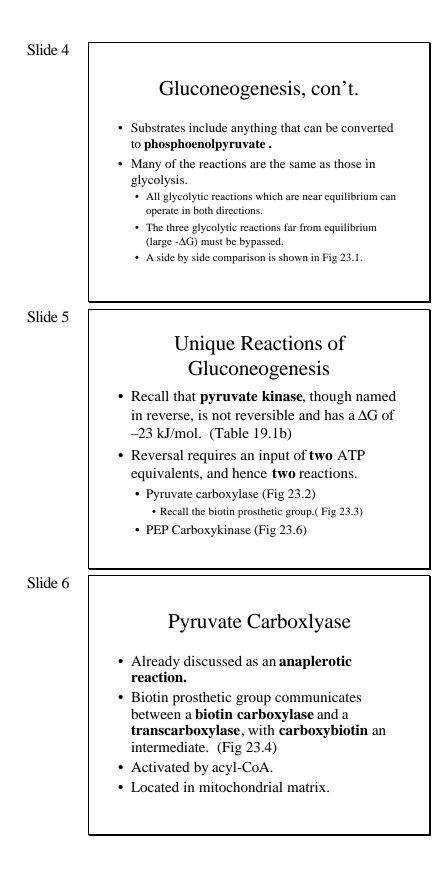
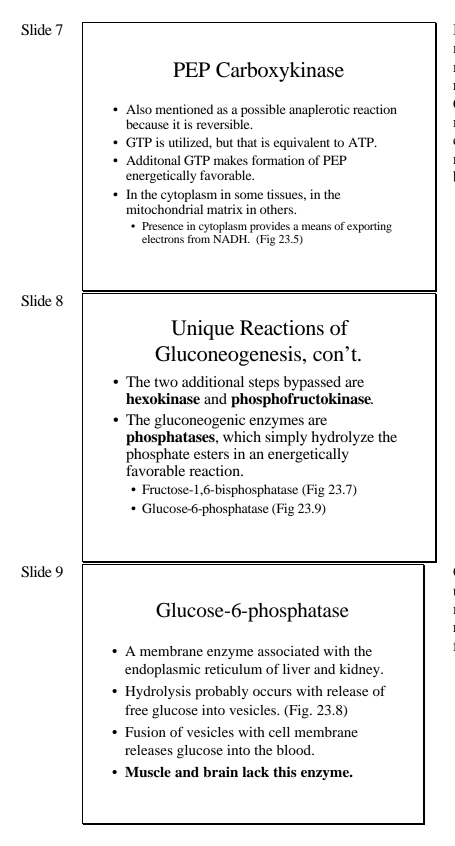


BCH 4054 Fall 2000 Chapter 23 Lecture Notes



Remember it is necessary for the pathways to differ in some respects, so that the overall G can be negative in each direction. Usually the steps with large negative G of one pathway are replaced in the reverse pathway with reactions that have a large negative G in the opposite direction.

Acyl-CoA activation, especially acetyl-CoA, regulates the fate of pyruvate. When acetyl-CoA is low, pyruvate is broken down by pyruvate dehydrogenase. High levels of acetyl-CoA signal the need for more OAA to run the citric acid cycle, or to be converted to glucose.



Because the oxaloacetate made in mitochondria cannot cross the mitochondrial membrane, it is reduced to malate, which can cross. Conversion of the cytoplasmic malate back to oxaloacetate in the cytoplasm produces the NADH needed for reduction of 1,3bisphosphoglycerate.

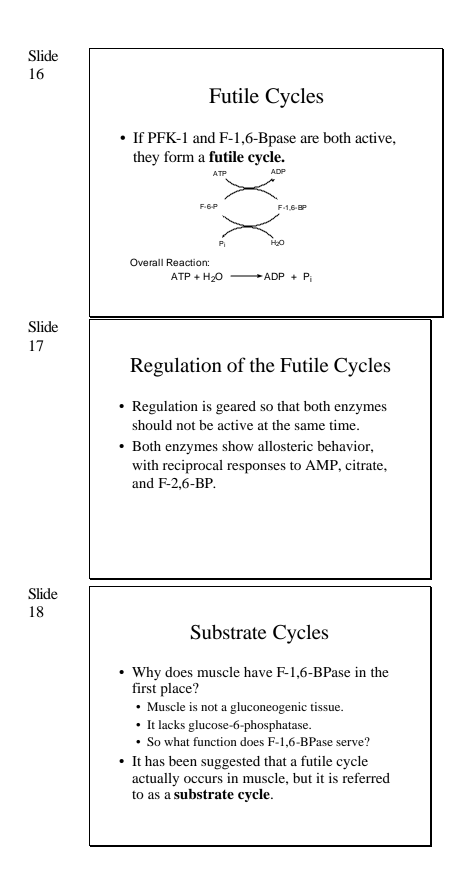
Glucose-6-phosphatase is actually used as a "marker" enzyme to measure the amount of endoplasmic reticulum present in sub cellular fractionation experiments.

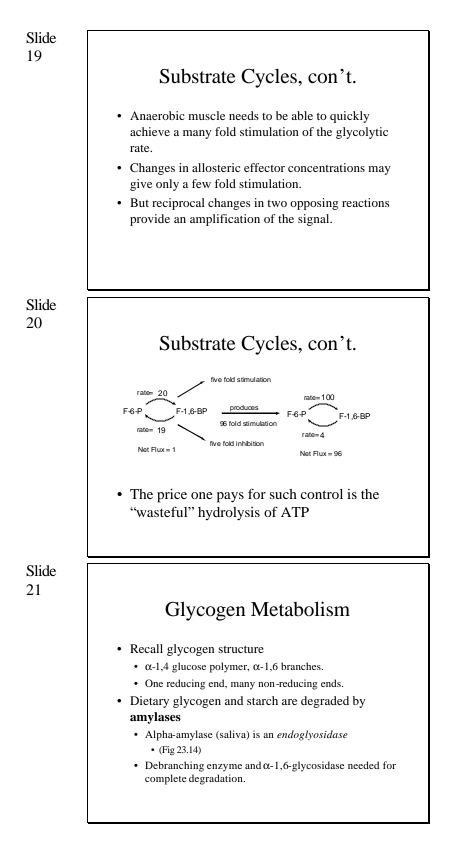
| Slide | |
|-------------|--|
| 10 | Regulation of Hexokinase and Glucose-6-phosphatase |
| | Hexokinase is inhibited by glucose-6- phosphate |
| | Glucose-6-phosphatase has a high Km for glucose-6-phosphate |
| | • Therefore gluconeogenesis is favored by high concentrations of glucose-6-phosphate. |
| | |
| Slide 11 | |
| | Fructose-1,6-Bisphosphatase |
| | Allosterically regulated in a fashion opposite to phosphofructokinase. Citrate stimulates. AMP inhibits, ATP activates. |
| | • Fructose-2,6-bisphosphate (F-2,6-BP) inhibits. (Fig 23.12) |
| | High energy state (high ATP, citrate) stimulates the phosphatase and gluconeogensis. Low energy state (high AMP, low citrate) |
| | stimulates PFK and glycolysis.Hormonal regulation acts through F-2,6-BP. |
| Slide 12 | |
| 12 | Overall Stoichiometry from |
| | Pyruvate |
| | • Recall the stoichiometry of glycolysis was: Glucose + 2 NAD + 2 ADP + 2 $P_i \rightarrow 2$ pyruvate + 2 NADH + 2 ATP |
| | The stoichiometry of gluconeogenesis is: 2 pyruvate + 2 NADH + 6 ATP \rightarrow glucose + 2 NAD + 6 ADP + 6 P _i (where ATP is equivalent to GTP) |
| | • The difference of 4 ATP is sufficient to give the reverse reaction a negative ΔG . |
| | |
| | |

| Slide 13 | The Cori Cycle Vigorous exercise cause muscles to produce lactate. (white muscle, low in mitochondria, is geared for rapid anaerobic glycolysis) The lactate is excreted into the blood and taken up by liver. Liver converts some of the lactate back to glucose. The glucose excreted by the liver can then be used by muscles. See Fig. 23.10 |
|-------------|--|
| Slide | |
| 14 | Regulation of Gluconeogenesis |
| | Fig 23.11 summarizes the reciprocal regulation of glycolysis and gluconeogenesis. High energy status favors gluconeogenesis Low energy status favors glycolysis Acetyl-CoA influences the fate of pyruvate F-2,6-BP favors glycolysis and inhibits gluconeogenesis. |
| Slide | |
| 15 | Hormonal Regulation of F-2.6- Bisphosphatase |
| | F-2,6-BP made by PFK-2 F-2,6-BP degraded by F-2,6-BPase Both activities are on the same protein A bifunctional enzyme (Fig 23.13) Hormonal stimulation results in phosphorylation of this protein. In liver, phosphorylation activates F-2,6-BPase, inhibits PFK-2. (stimulating gluconeogenesis) In muscle, phosphorylation activates PFK-2, inhibits F-2,6-Bpase. (stimulating glycolysis) |

High acetyl-CoA inhibits favors pyruvate to oxaloacetate, either for increased TCA activity or increased gluconeogenesis. Low acetyl-CoA favors breakdown of PEP and pyruvate to form acetyl-CoA.

We will discuss the mechanism of hormone activation a bit later when we discuss regulation of glycogen synthesis and breakdown.

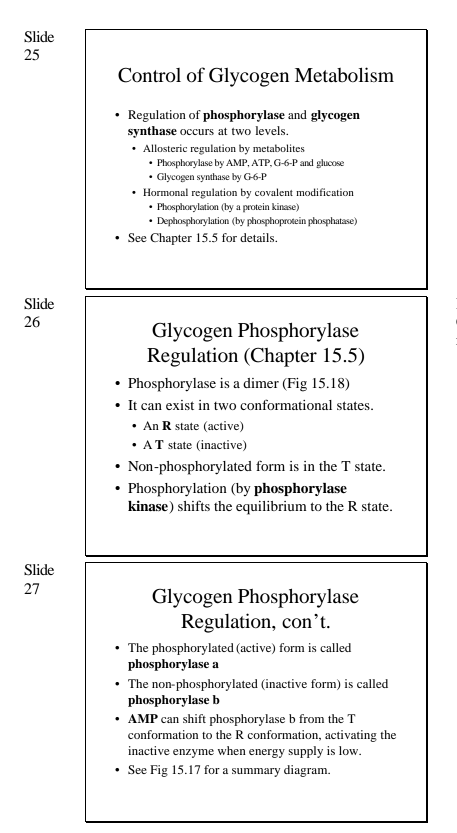




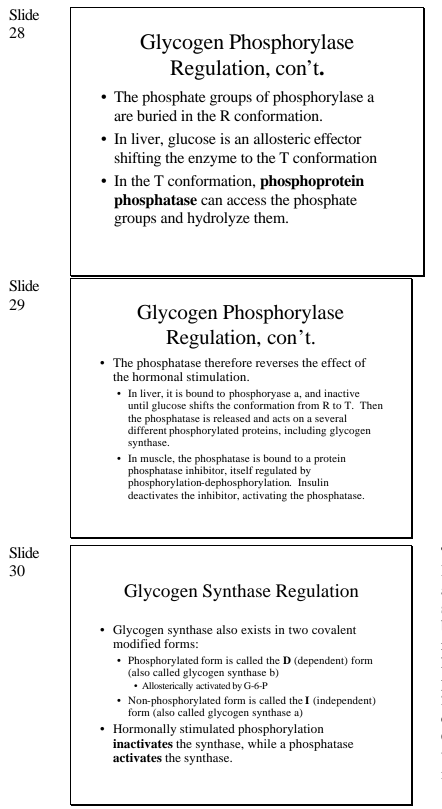
The distinction between alpha and beta amylases is not on the configuration at the anomeric carbon. Beta-amylase is an exopeptidase in plants that cleaves disaccharide (maltose) units from the non-reducing ends of the glycogen chain.

| Slide 22 | |
|-------------|--|
| | Glycogen Metabolism, con't. |
| | Tissue glycogen is degraded by phosphorolysis of the α-1,4 bonds. The enzyme is phosphorylase. A cleavage by phosphate at the non-reducing end. (Fig 23.16) The reaction is reversible. Glucose-1-phosphate is the product. G-1-P is converted to G-6-P by the enzyme phosphoglucomutase Debranching enzyme is needed for complete degradation. |
| Slide 23 | |
| | Glycogen Synthesis |
| | Originally thought to be phosphorylase acting in the reverse direction. Discovery of activation of supers as purcleastide. |
| | Discovery of activation of sugars as nucleotide diphosphate derivates. Uridine diphosphate glucose (Fig 23.17) |
| | • Synthesis of UDP-glucose: UTP+G-1-P ≒ UDP-Glc + P~P _i |
| | Reaction is reversible, and named in reverse: UDP-glucose pyrophosphorylase |
| Slide 24 | |
| | Glycogen Synthesis, con't. |
| | Glycogen Synthase catalyzes the transfer of glucose from UDP-glc to the reducing ends of glycogen, forming an α-1,4 linkage. Fig 23.19 |
| | • A branching enzyme transfers a 6-7 residue chain to the 6 position to form the branch points. (Fig 23.20). |
| | |

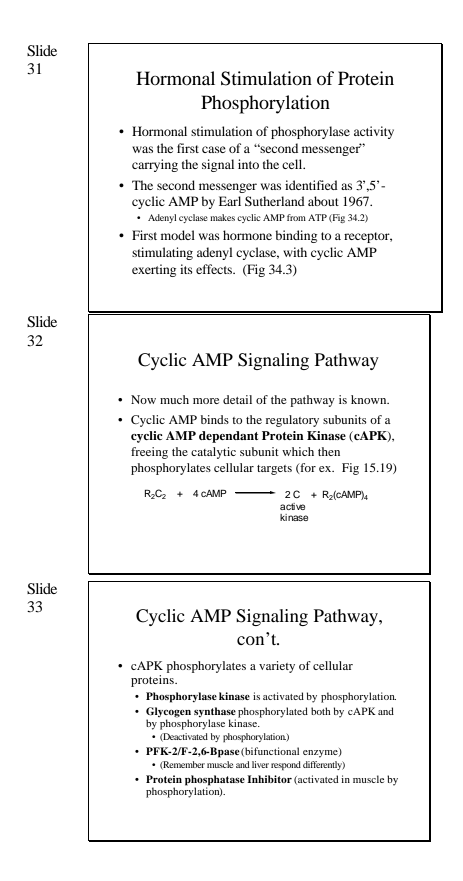
There are many reducing ends on the glycogen on the glycogen polymer, so cleavage can occur at many sites on the same molecule at the same time.

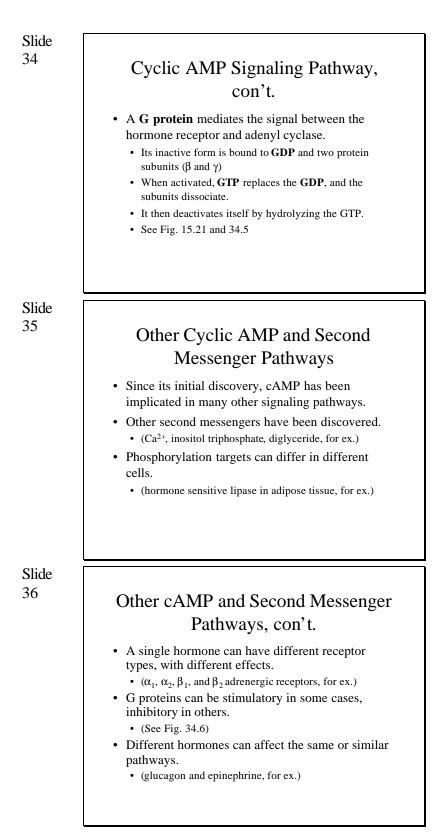


Remember the Monod-Wyman-Changeux model for allosteric regulations. Review it in section 15.4.



Therefore hormonal stimulation leads to activation of phosphorylase and deactivation of glycogen synthase, leading to glycogen breakdown. The enzymes are reciprocally regulated. Recall that hormonal stimulation also leads to phosphorylation of the PFK-2:F-2,6-BPase bifunctional enzyme with different effects in liver and muscle. Glycolysis is promoted in muscle, while gluconeogenesis is promoted in liver.





G in G protein stands for GTPbinding protein.

activating an inhibitory G protein, and a ² receptor activating a stimulatory G protein. Drugs have been developed that can target specific receptor types, some acting as **agonists** (mimicking the hormone), some acting as **antagonists**, blocking the action of the hormone.

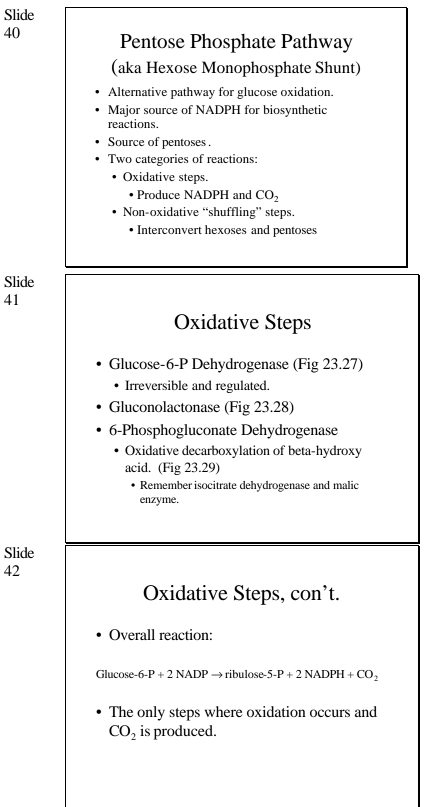
Figure 34.6 shows an ₂ receptor

Chapter 34 attempts to organize the everincreasingly complex signaling mechanisms. We may not get a chance to get into details of all of them, but you should read the chapter for background information.

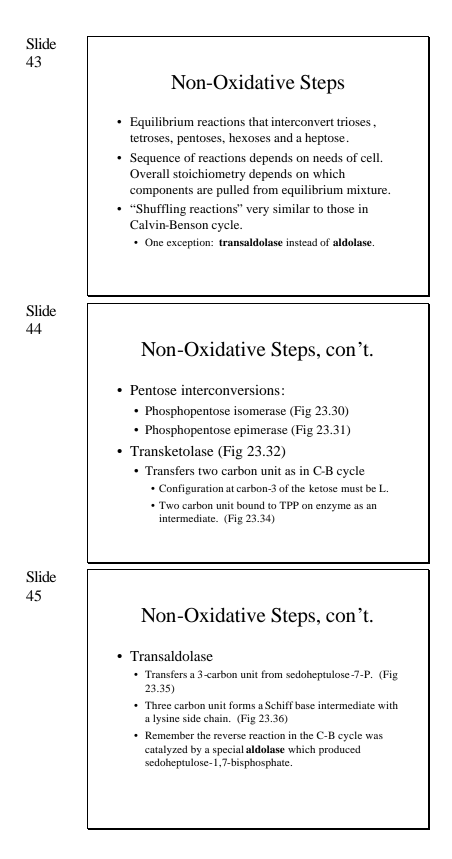
| Slide | |
|-------|---|
| 37 | |
| | Effects of Insulin |
| | |
| | • Made by the alpha cells of the pancreas. |
| | Receptor is a tyrosine kinase. (Chapter 24, page 5, 24) |
| | (Chapter 34, page S-24) Multiple effects one of which is to enable glucose uptake |
| | Multiple effects, one of which is to enable glucose uptake by tissues and to lower blood glucose. Gluconeogenesis is inhibited. |
| | • (See Fig 23.22) |
| | • Part of the effect on glycogen regulation is the activation of phosphoprotein phosphatase , which reverses the effect of cAMP protein kinase. |
| | effect of cAwir proton kinase. |
| | |
| Slide | |
| 38 | |
| 50 | Effects of Glucocorticoids |
| | Elicus Ul Olucocollicolas |
| | Contraction of the formation the |
| | • Steroid hormones act differently by entering the call binding to an intracellular recenter and |
| | cell, binding to an intracellular receptor, and regulating gene activity in the nucleus. |
| | |
| | Cortisol promotes protein degradation in muscle and gluconeogenesis in liver (as well as |
| | and gluconeogenesis in liver (as well as stimulation of urea cycle enzymes to complete |
| | amino acid breakdown). |
| | annio acid oreakdowny. |
| | |
| | |
| | |
| Slide | |
| 39 | |
| 0, | Ca ²⁺ Also Stimulates Glycogen |
| | Degradation |
| | • Phosphorylase kinase is activated both by |
| | phosphorylation and also by Ca^{2+} . |
| | • It has four subunits: α , β , γ , δ |
| | • γ is the active subunit. |
| | • α and β are inhibitors . Inhibition is removed |
| | when phosphorylated by protein kinase . |
| | • δ is a protein called calmodulin , a calcium |
| | binding protein involved in many calcium stimulated reactions. |
| | stillulated reactions. |
| | |
| | |

Insulin also induces synthesis of glycolytic enzymes. It signals the "fed state", stimulating both glycogen and lipid synthesis.

Electrical stimulation of muscle opens calcium channels in the sarcoplasmic which stimulates muscle contraction, so glycogen breakdown can also be coordinated with the muscles need for energy.



G-6-P dehydrogenase is inhibited by NADPH and by fatty acyl-CoA esters.



Notice that transketolase has a broad specificity for both the ketose donor and the aldose acceptor.

