#### BCH 4054 Fall 2000 Chapter 22 Notes



Notice membrane organization of the chloroplast. An internal thylakoid membrane is folded into structures called lamellae that stack to form grana. The interior of the thylakoid vesicles is the "lumen", while the space outside the thylakoid vesicles but inside the chloroplast membrane is called the stroma. Chloroplasts, like mitochondria, contain some DNA that codes for some of its proteins.

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The Light Reaction $H_2$  $H_2$ <

We will see that at least 3 ATP's will be required in the Dark Reaction, so that more than 530 kJ of light energy are needed per mole of oxygen produced.

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Remember, these numbers are **standard free energies**. The actual energy need is greater in order to push the reaction towards completion.





# Harvesting Light Energy, con't.

• Photochemistry occurs generally from the lowest singlet state. (Higher energy states decay rapidly to lowest state).

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# Harvesting Light Energy, con't.

- Need at least 533 kJ of energy per mol of  $O_2$  produced
- Can obtain 171 kJ per mole of quanta
- 533/171 = 3.11; Expected that 4 quanta would be sufficient for each O<sub>2</sub>
- It actually takes 8 quanta per O<sub>2</sub>
  - "Inefficiency" because one needs large overall negative  $\Delta G$  to make reaction irreversible

## Harvesting Light Energy, con't.

- Chlorophyll is transparent to most of visible light.
- Absorption of light in 450-650 nm region accomplished by accessory pigments
  - Beta-Carotene and Phycocyanobilin (Fig. 22.7)
  - Responsible for colorful fall foliage
- Light is trapped by a "pigment system", and energy funneled to a reaction center (Fig 22.9)

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# Trapping Light Energy

- Possible fates of the quantum of absorbed light energy are diagrammed in Figure 22.8
  - Thermal dissipation
  - Fluorescence
  - Exciton transfer (which funnels energy to the reaction center)
  - Transfer of excited electron to another acceptor.
    The excited state has a much lower reduction potential (is a stronger reducing agent) than the ground state.

Increasing light intensity leads to saturation of rate of photosynthesis. If this is done with flashing light, then dark reaction will not be limiting. Still get one CO<sub>2</sub> produced for about every 2500 chlorophyll molecules. This was evidence for a "photosynthetic unit" of about 400 chlorophylls for each electron promoted. Most cholorphyll and the accessory pigments serve as "antennae" which trap the photon. Energy is then transferred to a reaction center, which is a special type of cholorphyll.

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The trick in trapping the energy in this way is to prevent the electron in  $X^-$  from falling back to reduce P<sup>+</sup>. That is accomplished by proper orientation of the components so that Y preferentially reduces P<sup>+</sup>, and the charge is rapidly separated.



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#### Effect of Each Photosystem

- PS I (P<sub>700</sub>) produces:
  - $X^-$ , a strong reductant ( $E_o' \sim -0.6$  volts)
  - $Y^+$ , a weak oxidant ( $E_o' \sim 0.45$  volts)
  - Strong reductant capable of reducing NADP+
- PSII (P<sub>680</sub>) produces:
  - $X^{-}$ , a weak reductant (E<sub>0</sub>' ~ 0.0 volts)
  - $Y^+$ , a strong oxidant ( $E_o$ , ~>+0.8 V volts)
  - Strong oxidant capable of oxidizing water
- The weak reductant of PSII reduces the weak oxidant of PS I (See Fig. 22.11)

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## Organization of Photosystems

- PSI and PSII are macromolecular assemblies found in the thylakoid membrane of the chloroplast.
- The pigment systems are connected by an electron transport chain so that the weak reductant of PSII will oxidize the weak oxidant of PSI.
- The connecting electron transport chain resembles complex III of mitochondria.
- There is also a chlorophyll containing complex that is a **light harvesting complex** (LHC)

Chloroplasts given light at both 680 and 700 nm simultaneously yield more  $O_2$  than the sum of  $O_2$  produced when each wavelength is used alone.

#### The Z Scheme

- Arranging these systems and the intermediate carriers according to reduction potentials creates a pattern resembling a Z laid on its side.
  - See Fig 22.12
- Two quanta of light are necessary to take an electron from a very high reduction potential (species D) to a very low reduction potential (species A<sub>0</sub>)
- A cytochrome complex connects the reductant from PSII to the oxidant from PSI.

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#### Oxygen Evolution

- Oxygen evolution involves a manganese complex on the lumen face of the thylakoid membrane. (Fig. 22.13)
- The complex cycles through 5 oxidation states
- 1 e<sup>i</sup> is removed in each of 4 steps by D, a tyrosyl radical, which is the strong oxidant produced by  $P_{680}^+$ •  $S_0$  to  $S_1$  to  $S_2$  to  $S_3$  to  $S_4$
- Fifth step involves O<sub>2</sub> release
- $S_4 \text{to } S_0$
- Evidence from effect of light flashes on O<sub>2</sub> production (See Fig 22.14)

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# Electron Transport Through a Cytochrome Complex

- Pheophytin, a special chlorophyll missing the  $Mg^{2\star}$  is the weak reductant produced by  $P_{680}^{**}$
- Plastoquinone accepts electrons from pheophytin
  (See Figure 22.15)
- Electrons passed from plastoquinone to plastocyanin, a Cu protein, by the cytochrome b/f complex, containing an Fe/S protein.
  - (See Figure 22.12)
- The complex pumps protons into the lumen via a Q cycle

Note the similarity of the cytochrome b/f complex with complex III of the mitochondrial electron transport chain. In both cases, electrons are passed from a quinone to a peripheral membrane protein, and protons are pumped across the membrane. In this case, the protons are pumped from the stroma to the lumen.

#### NADP<sup>+</sup> Reduction

- P<sub>700</sub><sup>\*</sup> reduces A<sub>0</sub>, a special chlorophyll
- A<sub>0</sub> reduces A<sub>1</sub>, a quinone called **phylloquinone**(phylloquinone is vitamin K<sub>1</sub>)
- Electrons then passed through several Fe/S membrane proteins to a soluble Ferredoxin (Fd)
- · Fd reduces a flavoprotein which reduces NADP+
- NADPH is made on the stromal face
- $P_{700}^{+}$  is reduced at the lumen face by plastocyanin

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#### **ATP** Synthesis

- The proton gradient drives an ATP synthase located in the thylakoid membrane.
- It consists of  $CF_0$ , an integral membrane component, and  $CF_1$ , a peripheral complex, both similar to the  $F_0F_1$  complex of mitochondria.
- CF<sub>1</sub> lies on the stromal (outside) face of the thylakoid membrane.
- ATP is made in the stroma.

The detailed structure, organization, and mechanism of the chloroplast ATP synthase is probably almost identical to that of the mitochondria enzyme except for orientation.

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#### ATP Synthesis, con't.

- Chloroplasts provided early direct evidence for the chemiosmotic hypothesis.
  - See description of Jagendorf and Uribe exp., p 728
- The chloroplast gradient differs in two ways from the mitochondrial gradient
  - Protons are pumped **into** the lumen instead of **out of** the organelle
  - +  $\Delta\psi$  is discharged by movement of  $Mg^{2\scriptscriptstyle +}$  from the lumen into the stroma
    - The proton motive force comes primarily from  $\Delta \! p H$

Chloroplasts were incubated at low pH so the inside equilibrated at that pH. Upon transfer to a solution of high pH to artificially create a proton gradient, and ATP synthesis was observed as the gradient collapsed.

# Cyclic Photophosphorylation

- The dark reaction needs 3 ATP's per two NADPH's reduced.
- The Z scheme may provide one, or less than one, depending on proton stoichiometry.
- Additional ATP's can be made by PSI recycling electrons to the cytochrome chain rather than reducing NADPH.
- This cyclic photophosphorylation produces ATP in the absence of oxygen evolution. (Fig. 22.12 and 22.22)

Proton stoichiometry is not made clear. If the cytochrome complex pumps 2 protons per electron, that amounts to 4 protons per NADPH. Since reduction of CO<sub>2</sub> occurs in the stroma, there is no need to export the ATP for that purpose, so 3 protons per ATP corresponds to about 1.3 ATP per pair of electrons.

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#### Detailed Architecture of Photosynthetic Reaction Centers

- Deisenhofer, Michel and Huber shared the 1984 Nobel Prize for solving the structure of the reaction center from *Rhodopseudomonas viridis*
- This bacterium contains  $P_{870}$ , a bacterial pheophytin, and quinones analagous to the plant pigment systems. (Fig. 22.16)
- It catalyzes a cyclic photophosphorylation in the bacterial membrane. (Fig. 22.17)

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## Architecture of PSI and PSII

- Core structure of PSII is probably similar to the *R*. *viridis* complex. (Fig 22.19)
  - Note the additional antenna chlorophyll proteins, and a cytochrome complex
- X-ray structure of PSI from the cyanobacterium *Synechococcus elongatus* has been solved.
  - Probably a good model for other PSI complexes.
  - Essential features in Fig 22.20



• Longer incubations produced sugar diphosphates, especially ribulose-1,5-bisphosphate (RuBP)

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# The Dark Reaction Early Experiments, con't. • "Cyclic" nature of the dark reaction illustrated by steady state labeling with <sup>14</sup>CO<sub>2</sub>. • Then the reaction is blocked by turning off the light or depleting the CO<sub>2</sub>.

lightof

tim e CO<sub>2</sub> depleted



PSI and ATP synthase are exposed to the stroma, while PSII is buried in the stacked regions of the grana. The cytochrome chain is spread throughout the thylakoid membrane.



Most abundant protein on earth
15% of chloroplast protein
Found in stroma

8 large (55 kD), 8 small (15 kD) subunits
Diagram of X-ray structure (Fig 22.23)
Enzyme bound carboxylated intermediate is cleaved into two PGA molecules (Fig 22.24)



# Ribulose-1,5-Bisphosphate Regeneration

- · Regeneration of RuBP from PGA requires energy in form of NADPH and ATP.
- Reaction pathway is called the Calvin-Benson cycle.
- Overall net reaction converts 3 CO<sub>2</sub> to a triose phosphate. (Or 6 CO<sub>2</sub> to a hexose phosphate)

 $3 C_5 + 3 CO_2 \rightarrow 6 C_3$  $5 C_3 \rightarrow 3 C_5$ net:  $3 \text{ CO}_2 \rightarrow \text{ C}_3$ 

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Note similarity of these reactions to those in glycolysis. NADPH is the reductant for the dehydrogenase, though. The three steps are all at equilibrium, and one gets an equilibrium mixture of the triose phosphates glyceraldehyde-3phosphate and dihydroxyacetone phosphate.

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Aldolase is reversible, acting near equilibrium. It is essentially like the enzyme in glycolysis, except it has a broader specificity. The phosphatase is irreversible, acting far from equilibrium.



Transketolase has a fairly broad specificity for both acceptor and donor. R is a sugar chain with the last hydroxyl phosphorylated, but it can be 2, 3, or 4 carbons. Note the configuration of the third carbon is always L.



## Overall Stoichiometry of the Calvin Benson Cycle • Fig. 22.5 summarizes the reactions we have just gone through. • Table 22.1 shows an overall net reaction in

- producing hexoses as:
  6 CO<sub>2</sub> + 18 ATP + 12 NADPH→ glucose + 18 ADP + 12 NADP<sup>+</sup>
  You should be able to use the reactions we have
- just detailed to show the following stoichiometry is also possible:

 $3 \text{ CO}_2 + 9 \text{ ATP} + 6 \text{ NADPH} \rightarrow \text{triose} - P + 9 \text{ ADP} + 6 \text{ NADP}^+$ 

Either stoichiometry shows that each  $CO_2$  fixed requires 2 NADPH and 3 ATP. Remember the NADPH was used to reduce two 1,3bisphosphoglycerates, 2 ATP's were needed to form the bisphosphoglycerates, and 1 ATP was needed to phosphorylate ribulose-5-phosphate.

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# Slide 40 Photorespiration The oxidase activity leads to photorespiration which is wasteful of ATP and NADPH The phosphglycolate is converted to glycolate, which is oxidized to glyoxalate in peroxisomes, and can be converted to glycine or used in the glyoxalate cycle. Two glycines condense to form serine and CO<sub>2</sub> (See Fig 22.29) Regulation of Rubisco insures that it is only active when the light reaction is running.

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## Regulation of Rubisco

- Rubisco is activated by Mg<sup>2+</sup> and by carbamylation:
   R-NH<sub>2</sub> + CO<sub>2</sub> → R-NH-COOH
- Carbamylation is promoted by high pH and by CO<sub>2</sub>
- The light reaction raises the pH of the stroma (Fig 22.27) as well as increasing [Mg<sup>2+</sup>].
  - An activase also participates in converting rubisco to a form that can be carbamylated, and the activase is also only active in the light.

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# Regulation of the Calvin Cycle

- In addition to the Rubisco regulation, several enzymes are activated by reduction of disulfide bonds.
  - The bonds are reduced by **thioredoxin**, which in turn is reduced by ferredoxin or NADPH (See Fig 22.28)
- The enzymes so activated include **fructose-1,6bisphosphatase, sedoheptulose-1,7bisphosphatase,** and **ribulose-5-P kinase** 
  - (These are the irreversible steps of the pathway)
  - The pathway is active only when light stimulates PSI.

Remember that the electrical component of the proton gradient is neutralized by  $Mg^{2+}$  transport from the thylakoid lumen to the stroma. The book describes also how ribulose-BP inhibits by binding to the inactive, non-carbamylated form of the enzyme, and the activase promotes its removal so carbamylation can occur. The requirement of CO<sub>2</sub> for activation insures that there will be CO<sub>2</sub> around to compete with O<sub>2</sub> and inhibit photorespiration.

Although the Calvin Cycle is referred to as the "dark reaction", it does not occur in the dark. Light must be present to drive the light reaction for the Calvin Cycle to be activated.

#### Starch versus Sucrose Synthesis

- Triose-P in the stroma is converted to glucose-6-P, which is activated to ADP-glucose, the precursor for starch synthesis.
  - Therefore starch is made and stored in the chloroplast
- Triose-P can exit the chloroplast to the cytoplasm, where it can be converted to hexoses as well, but here the hexoses are converted to sucrose.
  - Therefore sucrose is made in the cytoplasm.

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# C-4 Pathway aka Hatch and Slack Pathway

- At high temperatures, photorespiration becomes more of a problem.
- Tropical plants have developed a further protection by using a CO<sub>2</sub> delivery system that increases the CO<sub>2</sub> concentration.
- CO<sub>2</sub> is "fixed" as oxaloacetate in mesophyll cells. PEP + CO<sub>2</sub>  $\rightarrow$  OAA
- Synthesis of PEP "costs" 2 ATP Pyruvate +  $P_i$  + ATP  $\rightarrow$  PEP + AMP + PP  $\rightarrow$  2 $P_i$

The regulation of diverting triose-P in one direction or another is complex, and involves among other things the antiport exchange of triose-P with P<sub>i</sub> across the chloroplast membrane. The book does not cover this topic, so we won't get involved in its details. Fructose-2,6-BP plays a role in the process, though, just as we will see it playing a role in regulation of glycolysis and gluconeogenesis in liver and muscle. (Its concentration decreases during active photosynthesis, causing the rate of glycolysis to drop and diverting the triose-P in the cytoplasm to sucrose synthesis).

Sugar cane and corn are C-4 plants.  $CO_2$  is fixed by the enzyme PEP carboxylase. PEP is made by the enzyme pyruvate-phosphate dikinase, with an unusual mechanism described on page 739. Hydrolysis of two phosphate anhydride bonds are necessary for the reaction to be spontaneous.

#### C-4 Pathway, con't.

- OAA is reduced to malate, which is carried to the bundle sheath cells where malic enzyme releases CO<sub>2</sub>, and can build up the concentration of CO<sub>2</sub> to more effectively compete with O<sub>2</sub>. (Fig 22.30)
   (In some plants, OAA is converted to aspartate instead)
- Also O<sub>2</sub> does not diffuse as readily into the bundle sheath cells.

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# CAM Plants

- CO<sub>2</sub> and O<sub>2</sub> enter plants through pores called **stomata**. Water leaves the same way.
- Succulent plants such as *Crassulaceae* in arid regions do not open their pores during the day, so CO<sub>2</sub> can only enter at night.
- Malate is synthesized as in C-4 plants, but is stored in vacuoles until daylight, when the CO<sub>2</sub> is released for incorporation by the Calvin-Benson pathway.
- This process is referred to as *crassulacean acid metabolism* (CAM).