

## Experiment CM-5: Carbon Dioxide Fixation in Intact Cells

### Exercise 1: Carbon Dioxide Fixation in the Presence of an Uncoupler

**Aim:** To determine the rate of oxygen production by algal cells as an indicator of carbon dioxide fixation. This experiment will be performed with and without ATP synthesis being coupled to the electron transport chain.

#### *Procedure*

1. Turn off the light source used to illuminate the reaction chamber:
  - If the light has a power switch which is separate from the intensity control, use this switch to turn off the light.
  - If the light does not have a separate power switch, plug the light source into a power strip that has a switch. Use the switch on the power strip to turn off the light.
  - While the light is off, set the intensity control of the illuminator to the maximum level (100%).
2. Turn off the stirrer for the chamber. Remove the electrode from the chamber and place it in a beaker of deionized water. Remove the deionized water from the chamber with a plastic-tipped Pasteur pipet. Rinse the chamber two times with deionized water.
3. Make sure no light is reaching the polarograph chamber.
4. Swirl the tube of algae to suspend the cells homogeneously. Fill the chamber with algal cells. Carefully replace the electrode in the chamber and turn on the stirrer.
5. Check the chamber for the presence of bubbles. If bubbles are present, turn off the stirrer, allow the bubble to rise to the top. Remove the electrode and the bubble should burst. Replace the electrode, turn on the stirrer, and check for bubbles, again.
6. Add a 10 $\mu$ l aliquot of HCO<sub>3</sub><sup>-</sup> solution to the chamber through the reagent port. Be careful not to add an air bubble to the chamber.

**Note:** Put the tip of the micropipette down the reagent port and push its plunger to discharge the 10 $\mu$ l of the HCO<sub>3</sub><sup>-</sup> solution into the chamber. Do this carefully so that no bubbles are introduced into the chamber. Remove the micropipette from the chamber before releasing its plunger. If the plunger is released while the tip of the micropipette is still in the chamber, solutions could be siphoned from the chamber.

7. Type Algal Cells in Dark in the Mark box to the right of the Mark button. Position the trace near the bottom of the screen. Click the Record button. Press the Enter key on the keyboard to mark the recording. Record the rate of oxygen production of the algal cells in darkness for one minute. Continue recording.
8. Type Algal Cells in Light at 100 in the Mark box to the right of the Mark button. Press the Enter key on the keyboard as the light is turned on and the algal cell suspension in the chamber is illuminated at the highest intensity.

- Record oxygen production at this intensity for two minutes. Continue recording.
- Type Methylamine Added in the Mark box. Press the Enter key on the keyboard to mark the recording as a 10µl aliquot of methylamine uncoupler solution is added to the chamber through the reagent port. Record the oxygen production of the algal cells exposed to 100% light in the presence of an uncoupler for five minutes.
- Click Stop to halt the recording.
- Select Save in the File menu.
- Remove the electrode from the chamber, and rinse the electrode with deionized water. Make sure there is no water on the electrode before it is replaced in the chamber.
- Remove the fluid from the chamber using a Pasteur pipet with a plastic tip. Rinse the chamber with deionized water about ten times.

### **Exercise 2: Carbon Dioxide Fixation in the Presence of a Calvin Cycle Enzyme Inhibitor**

**Aim:** To determine the rate of carbon dioxide fixation (oxygen production) with and without the Calvin Cycle taking place in the algal cells.

#### ***Procedure***

- Repeat Exercise 1 using Iodoacetamide (IAA) solution in place of Methylamine (MA) solution. Mark the recording at the appropriate times to indicate the absence or presence of light or the presence of the inhibitor.
- Select Save in the File menu.
- Remove the electrode from the chamber, and rinse the electrode with deionized water. Make sure there is no water on the electrode before it is replaced in the chamber.
- Remove the fluid from the chamber using a Pasteur pipet with a plastic tip. Rinse the chamber with deionized water about ten times.

### **Exercise 3: Carbon Dioxide Fixation in the Presence of an Electron Transport Chain Blocker**

**Aim:** To determine the rate of carbon dioxide fixation (oxygen production) with and without a block across the electron transport chains of the algal cells.

#### ***Procedure***

- Repeat Exercise 1 using DCMU (3-(3,4-dichlorophenyl)-1,1-dimethylurea) solution in place of Methylamine (MA) solution. Mark the recording at the appropriate times to indicate the absence or presence of light or the presence of the inhibitor.
- Select Save in the File menu.
- Remove the electrode from the chamber, and rinse the electrode ten times with 70% ethanol and five times with deionized water. Make sure there is no water on the electrode before it is replaced in the chamber.

4. Remove the fluid from the chamber using a Pasteur pipet with a plastic tip. Rinse the chamber ten times with 70% ethanol and five times with deionized water.

#### **Exercise 4: Oxygen Production by Thylakoids in the Presence of an Uncoupler**

**Aim:** To determine the rate of oxygen production by thylakoids, with and without ATP synthesis being coupled to the electron transport chain.

##### ***Procedure***

1. In this exercise, use thylakoids and thylakoid reaction mixture in place of suspended algal cells.
2. Turn off the stirrer for the chamber. Remove the electrode from the chamber and place it in a beaker of deionized water. Remove the deionized water from the chamber with a plastic-tipped Pasteur pipet. Rinse the chamber two times with deionized water.
3. Fill the chamber with the yellow thylakoid reaction media. Carefully replace the electrode in the chamber and turn on the stirrer.
4. Check the chamber for the presence of bubbles. If bubbles are present, turn off the stirrer, allow the bubble to rise to the top. Remove the electrode and the bubble should burst. Replace the electrode, turn on the stirrer, and check for bubbles, again.
5. Make sure no light is reaching the polarograph chamber.
6. Type Thylakoids in Dark in the Mark box to the right of the Mark button. Position the trace near the bottom of the screen. Click the Record button.
7. While in darkness, mix the tube with the thylakoid preparation on the Vortex mixer, so the suspension is even. Use the micropipette to collect a 10  $\mu$ l aliquot of thylakoids from the tube.
8. Press the Enter key on the keyboard to mark the recording as a 10 $\mu$ l aliquot of thylakoids is added to the chamber through the reagent port. Be careful not to add an air bubble to the chamber. Record the rate of oxygen production of the thylakoids in darkness for one minute. Continue recording.
9. Type Thylakoids in Light at 100% in the Mark box. Press the Enter key on the keyboard as the light is turned on and the thylakoid suspension in the chamber is illuminated at the highest intensity. Record oxygen production at this intensity for one minute. Continue recording.
10. Type Methylamine Added in the Mark box. Press the Enter key on the keyboard to mark the recording as a 10 $\mu$ l aliquot of methylamine uncoupler solution is added to the chamber through the reagent port. Record the oxygen production of the thylakoids exposed to 100% light in the presence of an uncoupler for one minute. Continue recording.
11. Turn off the light to put the thylakoids in the dark. Record for another minute. Click Stop to halt recording.
12. Select Save in the File menu.
13. Remove the electrode from the chamber, and rinse the electrode with deionized water. Make sure there is no water on the electrode before it is replaced in the chamber.
14. Remove the fluid from the chamber using a Pasteur pipet with a plastic tip. Rinse the chamber with deionized water about ten times.

### **Exercise 5: Oxygen Production by Thylakoids in the Presence of a Calvin Cycle Enzyme Inhibitor**

Aim: To determine the rate of oxygen production by thylakoids, with and without the Calvin Cycle taking place in the thylakoid suspension.

1. Repeat Steps 1 through 11 from Exercise 4. Mark the recording at the appropriate times to indicate the presence of light and the uncoupler, methylamine.
2. After the recording of the effect of methylamine in Step 11 for one minute, type IAA Added in the Mark box. Press the Enter key on the keyboard to mark the recording as a 10 $\mu$ l aliquot of Iodoacetamide (IAA) inhibitor solution is added to the chamber through the reagent port.
3. Record oxygen production of the thylakoids exposed to 100% light in the presence of iodoacetamide for another minute.
4. Turn off the illuminator to put the thylakoids in the dark. Record for another minute. Click Stop to halt recording.
5. Select Save in the File menu.
6. Remove the electrode from the chamber, and rinse the electrode with deionized water. Make sure there is no water on the electrode before it is replaced in the chamber.
7. Remove the fluid from the chamber using a Pasteur pipet with a plastic tip. Rinse the chamber with deionized water about ten times.

### **Exercise 6: Oxygen Production by Thylakoids in the Presence of an Electron Transport Chain Blocker**

Aim: To determine the rate of oxygen production by thylakoids, with and without a block of the electron transport chain.

1. Repeat Steps 1 through 11 from Exercise 4. Mark the recording at the appropriate times to indicate the presence of light and the uncoupler, methylamine.
2. After the recording of the effect of methylamine in Step 11 for one minute, type DCMU Added in the Mark box. Press the Enter key on the keyboard to mark the recording as a 10 $\mu$ l aliquot of DCMU blocker solution is added to the chamber through the reagent port.
3. Record oxygen production of the thylakoids exposed to 100% light in the presence of DCMU for another minute.
4. Turn off the illuminator to put the thylakoids in the dark. Record for another minute. Click Stop to halt recording.
5. Select Save in the File menu.
6. Remove the electrode from the chamber, and rinse the electrode ten times with 70% ethanol and five times with deionized water. Make sure there is no water on the electrode before it is replaced in the chamber.
7. Remove the fluid from the chamber using a Pasteur pipet with a plastic tip. Rinse the chamber ten times with 70% ethanol and five times with deionized water.

- At the end of the lab period, rinse the electrode and chamber. Then, fill the chamber with deionized water and place the electrode in the chamber until the next time it is used.

## Data Analysis

### *Rate of Electron Transport*

By international consensus, the rate of electron transport in mitochondria is expressed as the rate of oxygen consumption. The units used to express these rates are: moles  $O_2$ /hr/mg protein. Oxygen consumption rates are standardized for comparison of experiments performed in different laboratories around the world. so, the size of the reaction chamber, the amount of mitochondrial protein in the chamber, and the time period for that change in oxygen concentration need to be included in the calculation of the rate.

Early in this experiment, the oxygen probe was calibrated using air-saturated, deionized water at room temperature. Through this calibration, the Y-axis of the recording channel was converted from voltage to oxygen concentration or [Oxygen], which is expressed as  $\mu\text{Molar}O_2$  and abbreviated  $\mu\text{MO}_2$ .

### *Changes in $O_2$ Concentrations in Exercises 1, 2, & 3.*

The rate of change of the oxygen concentration in the polarograph chamber can be measured directly from the recordings by measuring the average slope of the trace (Mean\_dv/dt).

Follow these steps to measure the rates of change of the oxygen concentration during these exercises.

- Scroll through the recording and find the section of data recorded during Exercise 1.
- Use the Display Time icons to adjust the Display Time of the Main window to show the complete experimental run on the Main window. The complete experimental run can also be selected by:
  - Placing the cursors on either side of the complete run; and
  - Clicking the Zoom between Cursors button on the LabScribe toolbar to expand the complete experimental run to the width of the Main window.
- Click on the Analysis window icon in the toolbar or select Analysis from the Windows menu to transfer the data displayed in the Main window to the Analysis window ([Figure CM-5-L1](#)).
- Look at the Function Table that is above the Oxygen Concentration channel displayed in the Analysis window. The functions, V2-V1, T2-T1, and Mean\_dv/dt should appear in the table.
- Once the cursors are placed in the correct positions for determining the change in the oxygen concentration in the chamber, the values of the parameters in the Function Table can be recorded in the on-line notebook of LabScribe by typing their names and values directly into the Journal.

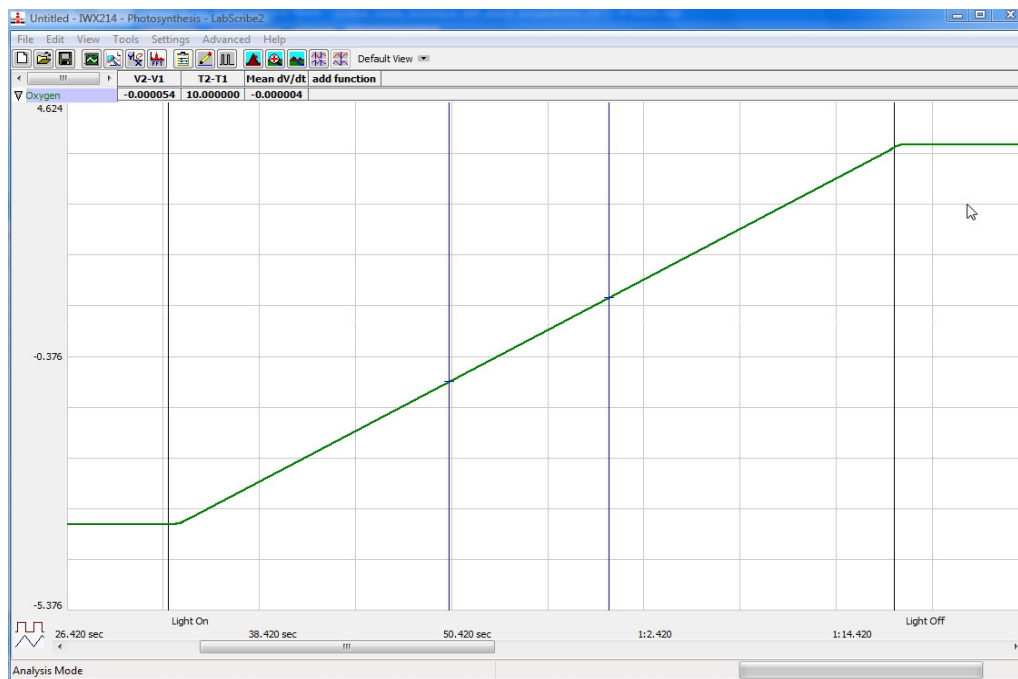


Figure CM-5-L1: Change in oxygen concentration in the polarograph chamber created by thylakoids with whole electron transport chains exposed to 80% light in the presence of an uncoupler. As displayed in the Analysis window, the marks indicate the times when the shutter controlling the light was opened and closed. Cursors are 10 seconds apart. The average rate of change (Mean\_dv/dt) in the oxygen concentration over that period is the 0.331010  $\mu\text{Molar O}_2/\text{sec}$ .

6. The functions in the channel pull-down menus of the Analysis window can also be used to enter the names and values of the parameters from the recording to the Journal. To use these functions:
  - Place the cursors at the locations used to measure the change in oxygen concentration.
  - Transfer the names of the mathematical functions used to determine the change in oxygen concentration to the Journal using the Add Title to Journal function in the Lung Volumes Channel pull-down menu.
  - Transfer the values for the change in oxygen concentration to the Journal using the Add Ch. Data to Journal function in the Oxygen concentration channel pull-down menu.
7. In the Analysis window, use the mouse to click on and drag the cursors to specific points on the recording to measure the following rates:
  - Algal Cells in Dark rate, which is the rate of oxygen production by the algal cells in the absence of light and uncoupler. Set the cursors ten seconds apart on a linear section of the recording of the dark rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
  - Algal Cells in 100% Light rate, which is the rate of oxygen production by the algal cells in the presence of full light. Set the cursors ten seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.

- Algal Cells in 100% Light with Uncoupler rate, which is the rate of oxygen production by the algal cells in the presence of full light and methylamine. Set the cursors ten seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
8. Record the values for these four rates in the Journal using the one of the techniques described in Steps 5 or 6.
  9. Use the same technique described in the previous steps to measure and record these rates from:
    - Exercise 2, where IAA is used in place of Methylamine.
    - Exercise 3, where DCMU is used in place of Methylamine.

### ***Changes in O<sub>2</sub> Concentrations in Exercises 4, 5, & 6***

In Exercises 4, 5, and 6, the rates of change of the oxygen concentration in the polarograph chamber filled with thylakoids were measured in the absence and presence of light, an uncoupler, and inhibitor, and a blocker. Follow these steps to measure the rates of change in the oxygen concentration in the polarograph chamber during these exercises:

1. Scroll through the recording and find the section of data recorded during the first experimental run in Exercise 4, the oxygen production rates with different light intensities in the absence of an uncoupler.
2. Use the same techniques described for the analysis of Exercises 1, 2, and 3 to display the complete experimental run on the Main window, transfer the data to the Analysis window, selecting the correct positions of the cursors, and transferring the measurements to the Journal.
3. In the Analysis window, use the mouse to click on and drag the cursors to specific points on the recording to measure the following rates:
  - Thylakoids in Dark rate, which is the rate of oxygen production by the thylakoids in the absence of light and uncoupler. Set the cursors ten seconds apart on a linear section of the recording of the dark rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
  - Thylakoids in 100% Light rate, which is the rate of oxygen production by the thylakoids in the presence of full light. Set the cursors ten seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
  - Thylakoids in 100% Light with Uncoupler rate, which is the rate of oxygen production by the thylakoids in the presence of full light and methylamine. Set the cursors ten seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
  - Thylakoids in Dark with Uncoupler rate, which is the rate of oxygen production by the thylakoids in the absence of light and the presence of methylamine. Set the cursors ten

seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.

4. For Exercise 5, measure Thylakoids in 100% Light with Uncoupler and inhibitor rate, which is the rate of oxygen production by the thylakoids in the presence of full light, methylamine, and IAA. Set the cursors ten seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
5. For Exercise 6, measure Thylakoids in 100% Light with Uncoupler and Blocker rate, which is the rate of oxygen production by the thylakoids in the presence of full light, methylamine, and DCMU. Set the cursors ten seconds apart on a linear section of the recording of this rate. The value for the Mean\_dv/dt is the average rate of change of oxygen concentration ( $\mu\text{Molar}/\text{sec}$ ) over that ten-second period.
6. Make sure all the changes in oxygen concentration are recorded in the Journal.

### ***Standardizing the Rate of Oxygen Production***

To standardize the rates of oxygen production (or consumption) according to the adopted convention, the volume of the polarograph chamber, the measured rate of change in oxygen concentration, and the concentration of thylakoids in the chamber must be incorporated into the calculations.

1. Multiply the rate of change in the oxygen concentration by the volume of polarograph chamber. The product of this calculation is the number of moles of oxygen produced in one second. For example, if the change in oxygen concentration is 0.333 microMolar  $\text{O}_2$  ( $\mu\text{MO}_2$  or  $10^{-6}$  moles/liter) per second, and the polarograph chamber has a 1.2 milliliter (ml) capacity; then 0.396 nanomoles (nmoles or  $10^{-9}$  moles) of oxygen are produced in one second:

$$(0.333\mu\text{MO}_2/\text{sec})(1.2\text{ml}) = 0.396 \text{ nmoles } \text{O}_2 \text{ produced in one second.}$$

2. Next, the moles of oxygen produced in one second must be converted to an hourly rate. In our example, if the amount of oxygen produced is 0.396 nmoles  $\text{O}_2$  in one second, the hourly production rate is 1.44 micromoles  $\text{O}_2$  ( $\mu\text{moles } \text{O}_2$ ) per hour:

$$(0.396 \times 10^{-9} \text{ moles } \text{O}_2/\text{sec})(60\text{sec}/\text{min})(60\text{min}/\text{hr}) = 1440 \times 10^{-9} \text{ moles } \text{O}_2/\text{hr} = 1.44 \times 10^{-6} \text{ moles } \text{O}_2/\text{hr.}$$

3. Finally, the moles of oxygen produced per hour must be standardized for the amount of thylakoids in the chamber. Clearly, if a chamber contains more thylakoids, the rate of oxygen production will be greater. After the thylakoids are isolated, the laboratory staff determines the concentration of chlorophyll (chl) in the preparation using a spectrophotometer. The concentration of chlorophyll is proportional to the concentration of thylakoids. Then, the lab staff dilutes the preparation with the appropriate volume of buffer to create a stock suspension of thylakoids that has the same approximate concentration of thylakoids in each lab session. In our example, if the concentration of chlorophyll in the stock solution is 2.5 milligrams of chlorophyll per milliliter (mg chl/ml), and 10  $\mu\text{l}$  of thylakoid suspension is added to the polarograph chamber, the amount chlorophyll added to the chamber is 0.025 mg:

$$(2.5 \text{ mg chl}/\text{ml})(0.010 \text{ ml}) = 0.025 \text{ mg chlorophyll in the chamber.}$$



To express the rate of oxygen production properly, the hourly rate of oxygen production must be divided by the amount of chlorophyll in the chamber:

$$(1.44 \times 10^{-6} \text{ moles O}_2/\text{hr}) / (0.025 \text{ mg chl}) = 57.6 \times 10^{-6} \text{ moles O}_2/\text{hr/mg chl} = 57.6 \text{ } \mu\text{moles O}_2/\text{hr/mg chlorophyll.}$$

4. Calculate the rates of oxygen production (or consumption) for each section of each experimental run using the steps presented above.
5. Enter the rates for all the exercises in [Table CM-5-L1](#).

### ***Correction for Non-Zero Dark Rates***

1. Sometimes, before the polarograph chamber is illuminated, thylakoids (or cells) will either produce or consume oxygen at a low rate.
  - If the thylakoids are producing oxygen (+ rate) in the dark, then the recorded rate is greater than the actual light-induced rate of oxygen production. The dark rate is adding itself to the actual rate.
  - If the thylakoids are consuming oxygen (- rate) in the dark, then the recorded rate is less than the actual light-induced rate of oxygen production. The dark rate is subtracting itself from the actual rate.
2. When determining the actual rate of oxygen production (or consumption) attributable to light, the non-zero dark rate, whether positive or negative, must be subtracted from the recorded rates in the same experimental run. For example, if thylakoids consumed oxygen at a rate of  $-9.6 \text{ } \mu\text{moles O}_2/\text{hr/mg chl}$  in the dark and produced oxygen at the rate of  $57.6 \text{ } \mu\text{moles O}_2/\text{hr/mg chl}$  in 100% light, their actual rate of light-induced oxygen production is:
$$(57.6 \text{ } \mu\text{moles O}_2/\text{hr/mg chl}) - (-9.6 \text{ } \mu\text{moles O}_2/\text{hr/mg chl}) = 67.2 \text{ } \mu\text{moles O}_2/\text{hr/mg chl}$$
3. For each experimental run in each exercise, subtract the non-zero dark rate from the recorded light-induced rates to yield the actual light-induced rates. Enter the actual rates in the appropriate columns on the table ([Table CM-5-L1](#)).

### ***Questions***

1. Do algal cells consume or produce oxygen when they are in the dark? What happens to the oxygen production rate when the algal cells and reagents in the chamber are exposed to light?
2. What effect does MA (methylamine) have on the oxygen production rates of algal cells?
3. What effect does IAA (iodoacetamide) have on the oxygen production rates of algal cells?
4. What effect does DCMU (3-(3,4-dichlorophenyl)-1,1-dimethylurea) have on the oxygen production rates of algal cells?
5. What effect does MA have on the oxygen production rates of thylakoids?

6. What effect does IAA have on the oxygen production rates of thylakoids?
7. What effect does DCMU have on the oxygen production rates of thylakoids?
8. How do the oxygen production rates of algal cells in the dark, in the light, and in the light with MA compare to the rates from thylakoids under the same conditions?
9. How do the oxygen production rates of algal cells in the dark, in the light, and in the light with IAA compare to the rates from thylakoids under the same conditions? Remember that MA was added to the chamber before IAA was added.
10. How do the oxygen production rates of algal cells in the dark, in the light, and in the light with DCMU compare to the rates from thylakoids under the same conditions? Remember that MA was added to the chamber before DCMU was added.

**Reference**

Behrens P W, Bingham S E, Hoeksema S D, Cohoon, D L, Cox J C (1989). Studies on the incorporation of CO<sub>2</sub> into starch by *Chlorella vulgaris*. *Journal Applied Phycology* 1: 123-130

**Table CM-5-L1: Rates of Oxygen Production in Cells and Thylakoids Treated with an Uncoupler, an Enzyme Inhibitor, and an Electron Blocker.**

Treatment Exercise	Rate of Oxygen Production (moles O <sub>2</sub> /hr/mg chl)								
	No Reagent			Reagent 1			Reagents 1 & 2		
	Dark	Light	Light-Dark	Reagent 1	Light	Light-Dark	Reagent 2	Light	Light-Dark
1-Algae & Uncoupler				MA					
2-Algae & Calvin Cycle Inhibitor				IAA					
3-Algae & Electron Blocker				DCMU					
4-Thylakoids & Uncoupler				MA					
5-Thylakoids & Calvin Cycle Inhibitor				MA			IAA		
6-Thylakoids & Electron Blocker							DCMU		

## Culturing *Chlorella vulgaris*

1. Obtain a culture of *Chlorella vulgaris* (Beijernick's or similar strain) from a biological supply house or a culture collection. Follow the directions included with the culture or use the following steps.
2. Raise the algae in a 1 liter flask filled with 500ml of mineral salts medium.
3. The flask is kept in a lighted environmental room or chamber with an illumination level of 40-90  $\mu\text{mol photon/square meter/second}$  at 25°C.
4. Agitation of the culture and the carbon source is provided by bubbling CO<sub>2</sub> rich air (2% v/v CO<sub>2</sub>) through the medium.
5. The KNO<sub>3</sub> concentration of the culture medium is set to support a cell density of about 1.7 grams/liter. Measure the rate of oxygen production of the stock culture in an oxygen polarograph illuminated with 100% light intensity. If the slope of the oxygen production line is steeper than 45 degrees, dilute the stock culture with deionized water if the cells will be used within a day.

## Isolation of Thylakoids

1. Wash 16 fresh spinach leaves (*Spinacia oleracea*) in tap water and pat dry with paper towels.
2. Tear the leaves into small pieces (<4 cm per side) and place the pieces in the chilled (4°C) jar of a kitchen blender. Pour 200 ml of cold thylakoid isolation buffer, containing 0.4 grams of sodium ascorbate, on top of the leaves.
3. Homogenize the leaves at low speed for 10 seconds until all the large leaf fragments are in a slurry. Then, homogenize the slurry at high speed for 20 seconds.
4. Filter the homogenate through 8 layers of cheese cloth into a 400 ml beaker. Place the collected filtrate in large centrifuge bottles that can be used in high capacity refrigerated centrifuge (IEC CU5000, for example)
5. Spin the filtrate for 2 minutes at 1000 rpm (~1200xG). Collect the supernatant, and place it in clean centrifuge bottles. Discard the pellet.
6. Spin the supernatant at 2100 rpm (~2500xG) for 10 minutes. Discard the supernatant. Resuspend the pellet in about 5 mls of cold thylakoid isolation buffer (without sodium ascorbate). Add what remains of the 200 ml of thylakoid suspension buffer to the suspension.
7. Spin the suspension at 1500 rpm (~1800xG) for one and a half minutes. Collect the supernatant, and place it in clean centrifuge bottles. Discard the pellet.
8. Spin the supernatant at 2100 rpm (~2500xG) for 10 minutes. Discard the supernatant. Resuspend the pellet in about 2 ml of cold thylakoid isolation buffer (without sodium ascorbate).
9. Dilute a 20  $\mu\text{l}$  sample of the final thylakoid suspension in 2ml of 80% acetone. Filter the solution through Whatman #4 filter paper. Place the filtered solution in a cuvette. Use a cuvette filled with 80% acetone as the blank. Determine the absorbance of the green solution at 663 nm and at 645 nm. The concentration of chlorophyll in the original suspension is calculated from

the following equation, where the dilution factor is 100 (2.0 ml/0.020 ml):

$$\text{mg chl/ml} = ((A_{663})(0.00802) + (A_{645})(0.0202)) \times \text{dilution factor.}$$

## Recipes for Solutions Used in the Experiment

### Chlorella Mineral Salts Medium

Concentration (mMolar)	Chemical	Milligrams/Liter DIH <sub>2</sub> O
	K <sub>2</sub> HPO <sub>4</sub>	100
	MgSO <sub>4</sub> ·7H <sub>2</sub> O	500
	Ca(NO <sub>3</sub> ) <sub>2</sub> ·4H <sub>2</sub> O	625
	KNO <sub>3</sub>	3
	FeSO <sub>4</sub> ·7H <sub>2</sub> O	10
	Disodium EDTA	8
	H <sub>3</sub> BO <sub>3</sub>	2.86
	MnCl <sub>2</sub> ·4H <sub>2</sub> O	1.81
	ZnSO <sub>4</sub> ·7H <sub>2</sub> O	0.22
	Na <sub>2</sub> MoO <sub>4</sub> ·2H <sub>2</sub> O	0.39
	CuSO <sub>4</sub> ·5H <sub>2</sub> O	0.08
	Co(NO <sub>3</sub> ) <sub>2</sub> ·5H <sub>2</sub> O	0.05

Mix all components in 925 ml of deionized water until dissolved. Add deionized water to bring final volume to 1L.

### Sodium Bicarbonate Solution

Concentration (Molar)	Chemical	Grams/100mls DI H <sub>2</sub> O
50	NaHCO <sub>3</sub>	8.4

Need about 0.1 ml per lab group in each lab period.

### Methylamine Uncoupler

Concentration (Molar)	Chemical	Grams/100ml DI H <sub>2</sub> O
3.0	Methylamine	20.26
Dissolve in 100 mls of deionized water. Dispense 0.1 ml per group in capped tubes		

### Iodoacetamide (IAA) Solution

Concentration (Molar)	Chemical	Grams/50ml DI H <sub>2</sub> O
0.50	Iodoacetamide	4.624
Need about 0.05 ml per lab group in each lab period.		

### DCMU Stock Solution

Concentration (mMolar)	Chemical	Grams/50ml 70% EtOH
0.1	DCMU	0.0012
Dissolve in 50 mls of 70% Ethanol. Freeze 25 mls in 1 ml aliquots in capped tubes. Store the remainder at 4°C.		

### Zero-Percent Oxygen Calibration Solution

Concentration (Molar)	Chemical	Grams/50ml DI H <sub>2</sub> O
1.5	Sodium Hydrosulfite	13
Dissolve in 50 mls of deionized water. Store at 4°C.		

### Thylakoid Isolation Buffer

Concentration (mMolar)	Chemical	Grams/Liter DI H <sub>2</sub> O
50	Tricine	
200	Sucrose	
50	NaCl	

Mix all components in 925 ml of deionized water until dissolved. Adjust pH to 7.4-7.5. Add deionized water to bring final volume to 1L. Store in 200ml aliquots in foil wrapped bottles at 4°C. Need about 500 mls per lab period.

### Thylakoid Reaction Buffer

Concentration (mMolar)	Chemical	Grams/Liter DI H <sub>2</sub> O
50	Tricine	
5	MgCl <sub>2</sub> · 6H <sub>2</sub> O	
2	K <sub>3</sub> Fe(CN) <sub>6</sub> KFerricyanide	

Mix all components in 925 ml of deionized water until dissolved. Adjust pH to 7.5. Add deionized water to bring final volume to 1L. Store in 200ml aliquots in foil wrapped bottles at 4°C. Need about 250 mls per lab period.

### Tricine Stock Solution

Concentration (Molar)	Chemical	Grams/100ml DI H <sub>2</sub> O
0.5	Tricine	8.96

Dissolve in 90 mls of deionized water. Adjust pH to 7.5, Add deionized water to bring to final volume of 100ml. Store at 4°C.