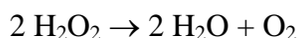


# Enzyme Action: Testing Catalase Activity (Method 1–O<sub>2</sub> Gas Sensor)

Many organisms can decompose hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) enzymatically. Enzymes are globular proteins, responsible for most of the chemical activities of living organisms. They act as *catalysts*, substances that speed up chemical reactions without being destroyed or altered during the process. Enzymes are extremely efficient and may be used over and over again. One enzyme may catalyze thousands of reactions every second. Both the temperature and the pH at which enzymes function are extremely important. Most organisms have a preferred temperature range in which they survive, and their enzymes most likely function best within that temperature range. If the environment of the enzyme is too acidic or too basic, the enzyme may irreversibly *denature*, or unravel, until it no longer has the shape necessary for proper functioning.

H<sub>2</sub>O<sub>2</sub> is toxic to most living organisms. Many organisms are capable of enzymatically destroying the H<sub>2</sub>O<sub>2</sub> before it can do much damage. H<sub>2</sub>O<sub>2</sub> can be converted to oxygen and water, as follows:

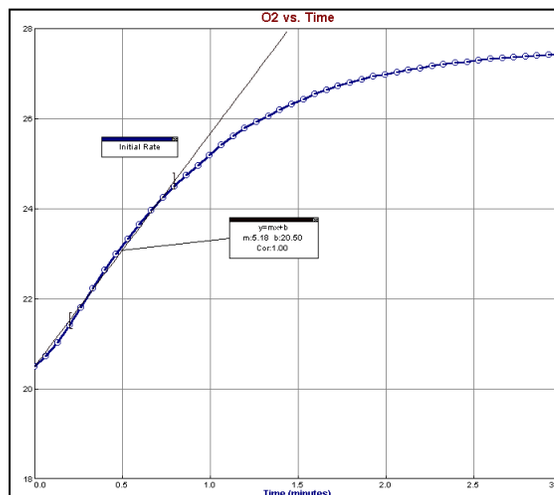


Although this reaction occurs spontaneously, enzymes increase the rate considerably. At least two different enzymes are known to catalyze this reaction: *catalase*, found in animals and protists, and *peroxidase*, found in plants. A great deal can be learned about enzymes by studying the rates of enzyme-catalyzed reactions. The rate of a chemical reaction may be studied in a number of ways including:

- measuring the rate of appearance of a product (in this case, O<sub>2</sub>, which is given off as a gas)
- measuring the rate of disappearance of substrate (in this case, H<sub>2</sub>O<sub>2</sub>)
- measuring the pressure of the product as it appears (in this case, O<sub>2</sub>).

In this experiment, you will measure the rate of enzyme activity under various conditions, such as different enzyme concentrations, pH values, and temperatures. It is possible to measure the concentration of oxygen gas formed as H<sub>2</sub>O<sub>2</sub> is destroyed using an O<sub>2</sub> Gas Sensor. If a plot is made, it may appear similar to the graph shown.

At the start of the reaction, there is no product, and the concentration is the same as the atmosphere. After a short time, oxygen accumulates at a rather constant rate. The slope of the curve at this initial time is constant and is called the *initial rate*. As the peroxide is destroyed, less of it is available to react and the O<sub>2</sub> is produced at lower rates. When no more peroxide is left, O<sub>2</sub> is no longer produced.



## OBJECTIVES

In this experiment, you will

- Use a computer and an Oxygen Gas Sensor to measure the production of oxygen gas as hydrogen peroxide is destroyed by the enzyme catalase or peroxidase at various enzyme concentrations.
- Measure and compare the initial rates of reaction for this enzyme when different concentrations of enzyme react with  $\text{H}_2\text{O}_2$ .
- Measure the production of oxygen gas as hydrogen peroxide is destroyed by the enzyme catalase or peroxidase at various temperatures.
- Measure and compare the initial rates of reaction for the enzyme at each temperature.
- Measure the production of oxygen gas as hydrogen peroxide is destroyed by the enzyme catalase or peroxidase at various pH values.
- Measure and compare the initial rates of reaction for the enzyme at each pH value.

## MATERIALS

Vernier computer interface  
computer  
LoggerPro  
Vernier O<sub>2</sub> Gas Sensor  
10 mL graduated cylinder  
400 mL beaker  
250 mL Nalgene bottle  
3.0%  $\text{H}_2\text{O}_2$

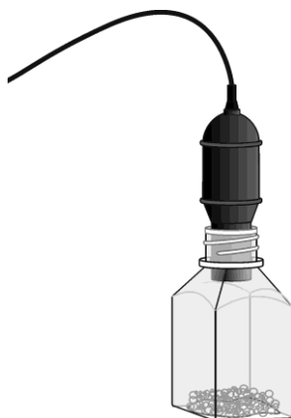
pH buffers  
enzyme suspension  
three 18 × 150 mm test tubes  
test tube rack  
ice  
thermometer  
three dropper pipettes

## PROCEDURE

1. Obtain and wear goggles.
2. Connect the Oxygen Gas Sensor to the computer interface. Prepare the computer for data collection by opening the file “O<sub>2</sub> (O<sub>2</sub>) Enzyme” from the *Advanced Biology with Vernier* folder of LoggerPro.

### Part I Testing the Effect of Enzyme Concentration

3. Place three test tubes in a rack and label them 1, 2, and 3. Fill each test tube with 5 mL of 3.0%  $\text{H}_2\text{O}_2$  and 5 mL of water.
4. Initiate the enzyme catalyzed reaction.
  - a. Using a clean dropper pipette, add 5 drops of enzyme suspension to test tube 1.
  - b. Begin timing with a stopwatch or clock.
  - c. Cover the opening of the test tube with a finger and gently invert the test tube two times.
  - d. Pour the contents of the test tube into a clean 250 mL Nalgene bottle.
  - e. Place the O<sub>2</sub> Gas Sensor into the bottle as shown in Figure 1. Gently push the sensor down into the bottle until it stops. The sensor is designed to seal the bottle without the need for unnecessary force.
  - f. When 30 seconds has passed, click  to begin data collection.



*Figure 1*

5. When data collection has finished, remove the O<sub>2</sub> gas sensor from the Nalgene bottle. Rinse the bottle with water and dry with a paper towel.
6. Move your data to a stored run. To do this, choose Store Latest Run from the Experiment menu.
7. Collect data for test tubes 2 and 3:
  - Add 10 drops of the enzyme solution to test tube 2. Repeat Steps 4–6.
  - Add 20 drops of the enzyme solution to test tube 3. Repeat Steps 4–5.
8. Using the mouse, select the initial linear region of your data on the graph. Click on the Linear Fit button, . Click  and a best-fit linear regression line will be shown for each run selected. In your data table (Table 2), record the value of the slope,  $m$ , for each of the three solutions. (The linear regression statistics are displayed in a floating box for each of the data sets.)
9. To print a graph of concentration vs. volume showing all three data runs:
  - a. Label all three curves by choosing Text Annotation from the Insert menu, and typing **5 Drops** (or **10 Drops**, or **20 Drops**) in the edit box. Then drag each box to a position near its respective curve. Adjust the position of the arrow head.
  - b. Print a copy of the graph, with all three data sets and the regression lines displayed. Enter your name(s) and the number of copies of the graph you want.
10. Determine the rate of reaction for each of the time intervals listed in Table 3 using the procedure outlined in Step 8. Record the rates for all three data runs in Table 3.

### **Part II Testing the Effect of Temperature**

Your teacher will assign a temperature range for your lab group to test. Depending on your assigned temperature range, set up your water bath as described below. Place a thermometer in your water bath to assist in maintaining the proper temperature.

- 0–5°C: 400 mL beaker filled with ice and water.
- 20–25°C: No water bath needed to maintain room temperature.
- 30–35°C: 400 mL beaker filled with very warm water.
- 50–55°C: 400 mL beaker filled with hot water.

## Computer 2

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- Rinse the three numbered test tubes used for Part I. Fill each test tube with 3 mL of 3.0%  $\text{H}_2\text{O}_2$  and 3 mL of water. Place the test tubes in the water bath. The test tubes should be in the water bath for 5 minutes before proceeding to Step 12. Record the temperature of the water bath, as indicated on the thermometer, in the space provided in Table 4.
- Find the rate of enzyme activity for test tubes 1, 2, and 3:
  - Add 10 Drops of the enzyme solution to test tube 1. Repeat Steps 4–6.
  - Add 10 drops of the enzyme solution to test tube 2. Repeat Steps 4–6.
  - Add 10 drops of the enzyme solution to test tube 3. Repeat Steps 4–5.
- Repeat Step 8 and record the reaction rate for each data set in Table 4. Calculate and record the average rate in Table 4.
- Record the average rate and the temperature of your water bath from Table 4 on the class data table. When the entire class has reported their data, record the class data in Table 5.

### Part III Testing the Effect of pH

- Place three clean test tubes in a rack and label them pH 4, pH 7, and pH 10.
- Add 3 mL of 3%  $\text{H}_2\text{O}_2$  and 3 mL of a pH buffer to each test tube, as in Table 1.

pH of buffer	Volume of 3% $\text{H}_2\text{O}_2$ (mL)	Volume of buffer (mL)
pH 4	3	3
pH 7	3	3
pH 10	3	3

- Using the test tube labeled pH 4, add 10 drops of enzyme solution and repeat Steps 4–6.
- Using the test tube labeled pH 7, add 10 drops of enzyme solution and repeat Steps 4–6.
- Using the test tube labeled pH 10, add 10 drops of enzyme solution and repeat Steps 4–5.
- Repeat Steps 8 and 9 to calculate the rate of reaction and print your graph. Record the reaction rate for each pH value in Table 6.

## DATA

### Part I Effect of Enzyme Concentration

Test tube label	Slope or rate (%/min)
5 Drops	
10 Drops	
20 Drops	

**Enzyme Action: Testing Catalase Activity (O<sub>2</sub>)**

Table 3 Time intervals (minutes)					
Rates	0–0.5 min	0.5–1.0 min	1.0–1.5 min	1.5–2.0 min	2.0–3.0 min
5 Drops					
10 Drops					
20 Drops					

**Part II Effect of Temperature**

Table 4	
Test tube label	Slope, or rate (%/min)
Trial 1	
Trial 2	
Trial 3	
Average	
Temperature range: ____ °C	

Table 5 (Class Data)	
Temperature tested	Average rate

**Part III Effect of pH**

Table 6	
Test tube label	Slope or rate (%/min)
pH 4	
pH 7	
pH 10	

**PROCESSING THE DATA**

1. On Page 2 of this experiment file, create a graph of the rate of enzyme activity vs. temperature. Plot the rate values for the class data in Table 5 on the y-axis, and the temperature on the x-axis. Use this graph to answer the questions for Part II.

## **QUESTIONS**

### **Part I Effect of Enzyme Concentration**

1. How does changing the concentration of enzyme affect the rate of decomposition of  $\text{H}_2\text{O}_2$ ?
2. What do you think will happen to the rate of reaction if one increases the concentration of enzyme to twenty-five drops? Predict what the rate would be for 30 drops.

### **Part II Effect of Temperature**

3. At what temperature is the rate of enzyme activity the highest? Lowest? Explain.
4. How does changing the temperature affect the rate of enzyme activity? Does this follow a pattern you anticipated?
5. Why might the enzyme activity decrease at very high temperatures?

### **Part III Effect of pH**

6. At what pH is the rate of enzyme activity the highest? Lowest?
7. How does changing the pH affect the rate of enzyme activity?

## **EXTENSIONS**

1. Design an experiment to determine the effect of boiling the catalase on the rate of reaction.