# 1A. Enzyme Action (Pressure Sensor)

How does the catalyzed decomposition rate of hydrogen peroxide compare with the uncatalyzed spontaneous decomposition rate?

Objectives

* Compare the spontaneous hydrogen peroxide decomposition rate with the catalyzed rate.

Materials and Equipment

|  |  |
| --- | --- |
| * Data collection system | * Pipet, 1-mL |
| * Pressure sensor | * Graduated cylinder, 25-mL |
| * Sampling bottle, 250-mL | * 1.5% Hydrogen peroxide, H2O2, 40.0 mL |
| * Rubber stopper assembly | * Catalase suspension, 2.0 mL |

Safety

Follow these important safety precautions in addition to your regular classroom procedures:

* Wear safety goggles at all times.

Procedure

1. Select Sensor Data in SPARKvue.

2. Connect the pressure sensor to your device.

3. Make sure only the pressure measurement is checked and choose the Graph template.

4. Attach the rubber stopper assembly to the pressure sensor.

5. Use a graduated cylinder to measure 20.0 mL of 1.5% H2O2 and add it to the sampling bottle. Loosely seal the bottle with the rubber stopper assembly and gently swirl to mix the contents.

Note: Always keep the bottle in an upright position when the stopper assembly is attached.

6. Gently grasp the bottle without squeezing and slide it back and forth along the table at a constant speed. Select Start to begin collecting data. Record the initial pressure for spontaneous decomposition in Table 1.

7. Continue to swirl at a medium speed while collecting data. Stop collecting data after 3 minutes. Record the final pressure and time elapsed in Table 1.

8. Pour the bottle contents into a waste container. Thoroughly rinse the sampling bottle.

9. Measure 20.0 mL of 1.5% H2O2 and add it to the bottle.

10. Stir the catalase suspension. Use a pipet to add 2.0 mL of catalase to the bottle. Swirl to mix.

11. Loosely seal the bottle with the stopper and place it on the table. Swirl the bottle without squeezing as before and start collecting data. Record the initial pressure for catalyzed decomposition in Table 1.

12. Stop collecting data after 3 minutes. Record the final pressure and time elapsed in Table 1.

13. Pour the bottle contents into a waste container. Rinse the sampling bottle.

14. Show both runs in SPARKvue and scale the display. Sketch your results in Graph 1. Include numbers, labels, and units on the x- and y-axes. Add a key to identify each run.

15. Calculate the change in pressure for each run. Use the following equation and enter the result in Table 1:

Change in Pressure = Final Pressure - Initial Pressure

16. Calculate the H2O2 decomposition rate for each run. Use the following equation and enter the result in Table 1:

Decomposition Rate = Change in Pressure ÷ Time

Data Collection

Table 1: Comparison of hydrogen peroxide decomposition rate with and without a catalyst

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Decomposition Reaction Type | Initial Pressure (kPa) | Final Pressure (kPa) | Time Elapsed  (s) | Change in Pressure (kPa) | Decomposition Rate  (kPa/s) |
| Spontaneous |  |  |  |  |  |
| Catalyzed |  |  |  |  |  |

Graph 1: Pressure produced from 1.5% hydrogen peroxide with and without a catalyst

A close up of a screen

Description automatically generated

Questions and Analysis

1. Why does the addition of the yeast suspension cause a change in pressure inside the bottle?

2. How much faster is the catalyzed reaction rate compared to the spontaneous decomposition rate?

3. Explain why the reaction is so much faster when an enzyme is present.

4. Is the reaction rate constant for the entire time data is recorded? Support your answer with evidence.

5. If the reaction continued to run, do you predict the reaction rate to be constant? Explain your thinking.

6. What kinds of conditions could you test in your lab classroom that might affect the rate at which catalase breaks down hydrogen peroxide?