**EXPERIMENT 1**

# The Kinetics of a Thiosulfate Solution

INTRODUCTION:

Various approaches are used to study the kinetics of reactions. A usual procedure is to monitor some property, such as intensity of color due to a reactant or product, which changes during the course of a reaction. In the present experiment, the property studied is the precipitation of a product, sulfur, which forms when solutions of hydrochloric acid and thiosulfate are mixed. The acidification of thiosulfate solutions leads to the formation of colloidal sulfur. The rate of this reaction is studied by measuring the time required for the reaction mixture to become so turbid that it ceases to transmit light. The reaction is acid catalyzed and the acid is kept at constant concentration throughout the experiment.

When a solution of thiosulfate is acidified, the following reaction takes place:

## S2O32- (aq) + 2H+ (aq)  H2O + SO2 (g) + S (s) (1)

On mixing a solution of acid, such as HCI, and a thiosulfate solution, there is an initial time-delay, then a precipitate of sulfur appears fairly sharply. The time interval between initial mixing and sulfur precipitation is a measure of the initial rate of reaction (i.e. the amount of time for a given amount of product, sulfur, to form). The color appears after a fixed amount of colloidal sulfur has been produced. Because the amounts of reactant used up in causing this amount of precipitate is small, the reactant concentrations remains essentially constant throughout the time of reaction. Recall that a rate for a chemical reaction is expressed as an amount of product formed (or reactant used up) divided by the time interval of the change:

##  Concentration /  t = [M] / t = [S2O32-] / t (2)

In this experiment you will determine the dependence of the initial rate of reaction on the initial concentration of thiosulfate ion. The concentration of the other reactant, H+, will be kept constant. Recall that, in this case:

## Average Rate = ( [M] /  t) (3)

## Rate law = k [M] m (4)

For Example, suppose the following data was collected for this experiment and we assume (1) the concentration of thiosulfate, at the point of data acquisition, has not significantly changed from its initial concentration and (2) since we are collecting the data at constant product production, the change at that point will be consistent for each trial therefore:

|  |  |  |  |
| --- | --- | --- | --- |
| trial | Minitial = [S2O32-] | Time | ~ Rate |
| 1 | 0.1 M | 160 s | 6.25x10-4 M/s |
| 2 | 0.2 M | 40 s | 5.00x10-3 M/s |
| 3 | 0.4 M | 10 s | 4.00x10-2 M/s |

Using equation (3), we can get an approximate rate of 0.1/16- = 6.25x10-4 M/s. Now using the initial rate method equation 4: (Rate2 = k [M2] m / Rate1 = k [M1] m ) we can calculate the order of m.

(5.00x10-3 M/s/6.25x10-4 M/s) = (.2/.1) m

8 = 2 m

ln(8) / ln(2) = m

m = 3

In a generic graph plotting rate vs concentration, we can see some kinetic rate trends as a function of time. If we plot the above date we would see the graph corresponds to 3rd order curve. Great correlation!

Fig 1 - Kinetic Rate Trends (Order of Rxn) as a Function of Concentration

3rd Order

2nd Order

**1st Order**

Zero Order

Rate (moles/L-sec)

Concentration (M)

In this experiment you will use the initial rate method to determine the order of thiosulfate and then plot the data as either concentration vs time, natural log of concentration vs time or inverse concentration vs time to verify the order obtained by the initial rate method. Then in part 2, you will vary temperature to determine the activation energy of this reaction.

**ONLINE DIRECTIONS**

1. **In canvas is a video of one of the trials for this experiment, watch it first.**
2. **Using the data given in lecture, complete this Laboratory report sheet.**
3. **Turn in the Report sheet filled out AND the excel graphs and data analysis in Canvas.**

List of Chemicals & Equipment per student

70 mL of 1M Na2S2O3 White paper & sharpie, Stopwatch

50 mL of 2M HCl 3 - 250mL Erlenmeyer Flasks

Thermometer Hot plate & Copper pneumatic trough

PROCEDURE:

## Part I: Effect of Thiosulfate Concentration on Reaction Rate

1. Prepare a piece of white paper with a large "X" marked on it (pen or pencil, make it dark).
2. A solution of 1 M sodium thiosulfate, is available (center bench or hood). Add 70 mL of this solution to 210 mL of water to make a solution approximately 0.25 M in thiosulfate. We will refer to this as your "stock thiosulfate solution".
3. Trial 1: Measure 50 mL of this stock thiosulfate solution into a 250 mL Erlenmeyer flask.
4. Carefully noting the time with a stopwatch or clock with a second hand, add about 5 mL of 2 M HCl, (provided for you by the stock room personnel or instructor), make sure it is the same amount each trial.
5. Swirl the contents of the flask vigorously to mix, then place it on the white paper prepared in step (1).
6. Look down vertically through the flask. When the “X" disappears (due to the sulfur precipitate obstructing your vision), note the time.
7. The experiment is now repeated four more times, using progressively more dilute thiosulfate solutions. Note, however, that for each experiment, the disappearance of the "X" corresponds to formation of the same amount of precipitated sulfur in the reaction mixture. The diluted solutions are prepared as follows:

Trial #2: take 40 mL stock thiosulfate solution and dilute to 50 mL by adding water

Trial #3: take 30 mL stock thiosulfate solution and dilute to 50 mL by adding water

Trial #4: take 20 mL stock thiosulfate solution and dilute to 50 mL by adding water

Trial #5: take 10 mL stock thiosulfate solution and dilute to 50 mL by adding water

In each experiment, addition of 5 mL of 2 M HCI solution assures that the concentration of H+ does not vary. Only the initial concentration of thiosulfate varies. Use the data collected above to estimate the order of reaction with respect to thiosulfate solution.

## Part II: Temperature Dependence of Reaction

1. Measure 10 mL of your stock thiosulfate solution (0.25 M) into a 100-mL Erlenmeyer flask.
2. Add 40 mL of water.
3. Measure the temperature of the solution.
4. Noting the time carefully, add 5 mL of 2M HCI, swirl, and place on the marked white paper as in Part 1.
5. Note the time of disappearance of the "X" on the white paper.
6. Repeat the experiment four more times, heating the thiosulfate solution before the addition of HCI to a little over 30o, 40 o, 50 o, and 60 oC so that the reaction can be started as near to these temperatures as possible.
7. Use these data to investigate temperature dependence of the reaction (Report Sheet).

NAME SECTION

REPORT SHEET EXPERIMENT 1

# The Kinetics of a Thiosulfate Solution

General/Pre-Lab Question:

1. Based on Equation (1), write the oxidation and reduction half-reactions for the S2O32-/H+ reaction.

EXPERIMENT DATA:

Part I - Changes due to concentration differences (include units)

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Volume thiosulfate used |  | Time for "X" to disappear, in seconds | Concentration of thiosulfate | Rate of disappearance of "X" |
| 50 mL |  |  |  |  |
| 40 mL |  |  |  |  |
| 30 mL |  |  |  |  |
| 20 mL |  |  |  |  |
| 10 mL |  |  |  |  |

1. Estimate the order of reaction with respect to thiosulfate based on the initial rate method. Show a sample calculation here, the rest of the calculations should be in your lab notebook.

order:

1. Next display your results graphically, in excel or similar program, by plotting (1) concentration vs time, (2) the reciprocal of the concentration against the time, and then (3) plot the natural log of concentration against time and compare your graphs. Attach those graphs to this report sheet. Describe any conclusions/comparisons can you draw from these graphs. How do the answers acquired from the graphs compare to the initial rate method?
2. How would the shape of these graphs change if the order of the reaction were:
   1. higher?
   2. lower?

Part II - Changes due to temperature difference [M] =

Calculate the rate and rate constant per temperature.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Temperature (oC) | Temperature (K) | Time (t) for "X" to disappear | Rate | k |
|  |  |  |  |  |
|  |  |  |  |  |
|  |  |  |  |  |
|  |  |  |  |  |
|  |  |  |  |  |

1. Plot the natural log of k vs. the reciprocal of temperature in Kelvin. Attach that graph to this report sheet. Briefly describe the appearance of the graph. What equation correlates to this graph, (show two variations)?
2. Use this graph to estimate the activation energy (include units). Describe and demonstrate how you analyzed this graph and acquired your Ea value.

Ea =