SL Chemistry Internal Assessment

Candidate code: gbc594

Efficiency per unit mass for five different catalysts of hydrogen peroxide decomposition

Research question

Which catalyst out of Manganese (IV) Oxide, Lead (II, IV) Oxide, Potassium Iodide, Copper (II) Oxide, and Iron (III) Oxide will most rapidly catalyze the decomposition of hydrogen peroxide per unit mass based on the initial rate of reaction measured through the time taken for the catalyzed reaction to produce a specific volume of gas?

Introduction

Background

The "elephant's toothpaste" is one of the first chemical experiments I witnessed as a kid. The reaction happens when high concentrations of hydrogen peroxide rapidly decays into oxygen gas and water in the presence of potassium iodide as a catalyst, where the rapid release of small oxygen bubbles from the solution causes a large volume of foam to emerge. The research question originates from the thought that there could be a more effective

catalyst than potassium iodide for the "elephant's toothpaste" reaction.

Relevant Theory

Hydrogen peroxide, H₂O₂, is an unstable compound that spontaneously decays into water and oxygen through the equation:

$$2H_2O_2 \rightarrow 2H_2O + O_2$$
 [1]

However, the spontaneous decay of hydrogen peroxide is relatively slow, with a half-life of 10-20 hours (FMC Corporation) in open air under standard conditions. This makes spectacles such the "elephant's toothpaste" reaction impossible without a catalyst.

Catalysts are chemicals that increase the rate of certain reactions without undergoing any permanent chemical change itself. Illustrated on *Figure-1* (Burns), the presence of catalysts offer an alternative pathway to the reaction that requires an overall lower activation energy. With reference to the Maxwell-Boltzmann distribution on *Figure-2* (Polypompholyx), this statistically allows a higher proportion of the reactants to have enough energy to react, and hence increases the rate of reaction.

Most catalysts do undergo chemical change during the reaction they catalyze, but the chemical change is reversed after the reaction has finished. Potassium iodide as the catalyst used in the "elephant's toothpaste" reaction is one such example, as can be seen from the following system of equations describing how potassium iodide catalyzes the decomposition of hydrogen peroxide:

$$KI \rightarrow K^+ + I^-$$
 [2]

$$I^{-} + H_{2}O_{2} \rightarrow H_{2}O + OI^{-}$$
 [3]
 $OI^{-} + H_{2}O_{2} \rightarrow H_{2}O + O_{2} + I^{-}$ [4]

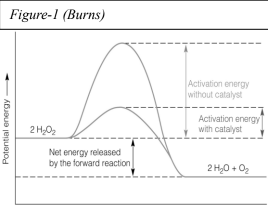
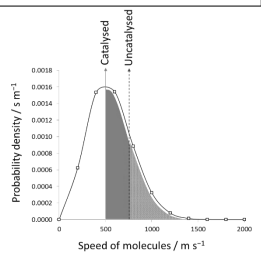


Figure-2 (Polypompholyx)

Progress of reaction -



This shows that hydrogen peroxide initially acts as an oxidizing agent that oxidizes the dissociated iodide ions to

hypoiodite ions, but it later acts as a reducing agent that reduces hypoiodite ions back to iodide ions. Adding the two conversion equations between iodide and hypoiodite together, the iodide and hypoiodite on each side of the equation cancels out to give just:

$$2H_2O_2 \rightarrow 2H_2O + O_2$$
 [5]

, which proves that potassium iodide is a catalyst for the decomposition of hydrogen peroxide, and quantitatively shows that 1 mol of potassium iodide will be able to decompose 2 mol of hydrogen peroxide per catalytic cycle. Besides potassium iodide, a number of transition metal oxides are also catalysts for hydrogen peroxide decomposition. Transition metals' ability to change between different oxidation states also allows catalytic cycles with hydrogen peroxide to occur. Take iron (III) oxide for example:

$$Fe_2O_3 \rightarrow 2Fe^{3+} + 3O^{2+}$$
 [6] (Note that the " \rightarrow " above signifies composition instead of reaction)

$$\begin{array}{c}
2Fe^{3+} + H_2O_2 \rightarrow 2Fe^{2+} + O_2 + 2H^+ \\
+ 2Fe^{2+} + 2H^+ + H_2O_2 \rightarrow 2Fe^{3+} + 2H_2O
\end{array}$$

$$\begin{array}{c}
2H_2O_2 \rightarrow 2H_2O + O_2
\end{array}$$
[7]

As shown, hydrogen peroxide can reduce Fe^{3+} to Fe^{2+} and also oxidize Fe^{2+} back to Fe^{3+} , which creates a catalytic cycle similar to that of potassium iodide, where 1 mol of iron (III) oxide is able to decompose 2 mol of hydrogen peroxide per catalytic cycle.

The catalytic cycle for copper (II) oxide and manganese (IV) oxide largely follow the same logic, which that:

$$CuO \rightarrow Cu^{2+} + O^{2-} [8] \qquad MnO_2 \rightarrow Mn^{4+} + 2O^{2-} [10]$$

$$Cu^{2+} + \frac{1}{2}H_2O_2 \rightarrow Cu^{+} + O_2 + H^{+} \qquad Mn^{4+} + H_2O_2 \rightarrow Mn^{2+} + O_2 + 2H^{+}$$

$$+ Cu^{+} + H^{+} + \frac{1}{2}H_2O_2 \rightarrow Cu^{2+} + H_2O \qquad + Mn^{2+} + 2H^{+} + H_2O_2 \rightarrow Mn^{4+} + 2H_2O \qquad + H_2O_2 \rightarrow H_2O + \frac{1}{2}O_2 \qquad (9)$$

$$= \frac{11}{2}$$

1 mol of copper (II) oxide is only able to decompose 1 mol of hydrogen peroxide per catalytic cycle, whereas 1 mol of manganese (IV) oxide will be able to decompose 2 mol of hydrogen peroxide per catalytic cycle.

The same logic also applies lead (II, IV) oxide, but we have to realize that lead ions exists as both lead (II) and lead (IV) in this compound. The catalytic cycle between Pb^{2+} and Pb^{4+} are as below:

$$\begin{array}{c} Pb^{4+} + H_2O_2 \longrightarrow Pb^{2+} + O_2 + 2H^+ \quad [12] \\ Pb^{2+} + H_2O_2 + 2H^+ \longrightarrow Pb^{4+} + 2H_2O \quad [13] \end{array}$$

Given that:

$$Pb_3O_4 \rightarrow 2Pb^{2+} + Pb^{4+} + 4O^{2-}$$
 [14]

, it can be deduced that the complete catalytic cycle for lead (II, IV) oxide is:

Hence, 1 mol of lead (II, IV) oxide will be able to decompose 6 mol of hydrogen peroxide per catalytic cycle.

Variables

Independent	Definition	Catalysts
Variable		Manganese (IV) Oxide, MnO ₂
	Five different catalysts for hydrogen peroxide decomposition.	Lead (II, IV) Oxide, Pb ₃ O ₄
		Potassium Iodide, KI
		Copper (II) Oxide, CuO
		Iron (III) Oxide, Fe ₂ O ₃

Dependent	Symbol	Definition	Proximate Variables		
Variable	rate of reaction when the catalyst comes into contact with hydrogen peroxide. (cm³ s⁻¹) The instantaneous initial rate will be indicative of how rapidly a catalyst		independent variable Proximate Syn	Symbol	Definition
		contact with hydrogen peroxide. The instantaneous initial rate will be indicative of		V	Volume of gas produced by the reacting system.
				(cm ³)	
				Symbol	Definition
		dependent variable	t	Time taken for the system to produce $V \text{ cm}^3$ of gas.	
			(seconds)		

Controlled Variables	Symbol	Definition	Controlled Value	Reason for control
	m	Mass of the catalyst used	$1.0 \pm 0.1 \; \text{gram}$	The aim of this experiment is to determine the efficiency of each catalyst per unit mass.
	T_0	Initial temperature of the system	$21.0 \pm 0.3 ^{\circ}\text{C}$	Initial temperature influences the initial rate of reaction independent of the catalyst used.
	A	Surface area of the catalyst	All catalysts used are powdered (unable to control quantitatively)	Surface area influences the initial rate of reaction independent of the catalyst used.
	[H ₂ O ₂]	Concentration of hydrogen peroxide solution used in the experiment	$\begin{array}{c} 0.176 \\ \pm \ 0.002 \ mol \ dm^{-3} \\ \text{(See page 4 for calculation)} \end{array}$	Concentration of hydrogen peroxide as a reactant influences the initial rate of reaction independent of the catalyst used.
	<i>V</i> (H ₂ O ₂)	Volume of hydrogen peroxide solution used in the experiment	$200.0 \pm 0.2 \text{ cm}^3$ (Class B 200 cm ³ volumetric flask)	The volume hydrogen peroxide solution determines the length of time that the reaction is going to persist. Since time is measured as the independent variable, this variable should kept constant.

Method

Chemicals and equipment list

List of chemicals		
Chemical	Quantity	Description
Manganese (IV) Oxide	5 grams	Solid, finely powdered
Lead (II, IV) Oxide	5 grams	Solid, finely powdered
Potassium Iodide	5 grams	Solid, finely powdered
Copper (II) Oxide	5 grams	Solid, finely powdered
Iron (III) Oxide	5 grams	Solid, finely powdered
6% Hydrogen Peroxide solution	500 cm ³	Assuming ± 1% uncertainty of concentration
Distilled water	-	-

Safety	precaution	for	hazardous	chemical

The metal oxides involved in this investigation, especially Lead (II, IV) Oxide, are labelled as very toxic for aquatic life with long lasting effects under hazard statement H410 and harmful if inhaled under hazard statement H332 (WSH council). Therefore any liquid contaminated by the compound should be thoroughly filtered before going down the sink, while the powdered solid should be handled with extreme care to prevent it from becoming airborne. Hands are washed thoroughly after handling said chemicals, while rubber gloves and safety goggles are worn at all times during the experiment.

List of agricument		
List of equipment	Ι	T = -
Equipment	Quantity	Uncertainty
500 cm ³ Conical Flask	1	-
	_	
100.0 cm ³ beaker	2	-
Glass vial	5	_
Glass viai		
Stop watch	1	± 0.01 seconds
Class B 20.00 cm ³ pipette	1	$\pm 0.03 \text{ cm}^3$;
and suction bulb		± 0.15%
Class B 200.0 cm ³	1	$\pm 0.2 \text{ cm}^3$;
volumetric flask		$\pm 0.08\%$
100 cm ³ Gas syringe set	1	$\pm 2.5 \text{ cm}^3$
(smallest division of 5 cm ³)		
Goldbell digital	1	± 0.0001 gram
electronic balance		
TFA Pocket-digitemp S	1	± 0.1 °C
Digital Thermometer		
Glass stirring rod	1	-
Molar and pestle	1	-
C1	1	
Clamp stand and clamp	1	-
	l	

Preparing diluted hydrogen peroxide solution

A test trial with manganese (IV) oxide shows that the reaction would be too rapid for any meaningful measurement to be taken using the requested 6% hydrogen peroxide solution, hence the following steps are taken to prepare diluted hydrogen peroxide solutions from which meaningful measurements can be made:

- 1. A 20 cm³ aliquot of the 6% hydrogen peroxide solution is taken with the class B pipette and suction bulb, and transferred into the class B 200.0 cm³ volumetric flask.
- 2. The volumetric flask is filled with distilled water to the 200.0 cm³ mark.

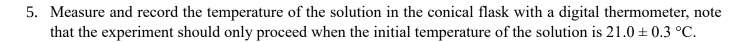
The molar concentration of diluted hydrogen peroxide solution prepared by this method is determined as follows:

6% hydrogen peroxide solution translates to 6 grams of hydrogen peroxide per 100 cm ³ .	
Original solution: $[H_2O_2] = \left(\frac{60 \text{ g} \pm 1\%}{1 \text{ dm}^3}\right) \div 34 \text{ g mol}^{-1}$	Molar mass of H ₂ O ₂ : 34 g mol ⁻¹
$[H_2O_2] = 1.76 \text{ mol dm}^{-3} \pm 1\%$	54 g moi
During step 1: $n(H_2O_2) = \frac{1.76 \text{ mol} \pm 1\%}{1 \text{ dm}^3} \times (0.02 \text{ dm}^3 \pm 0.15\%)$	
$n(H_2O_2) = 0.0352 \text{ mol } \pm 1.15\%$	
After step 2: $[H_2O_2] = \frac{0.0352 \text{ mol } \pm 1.15\%}{0.2 \text{ dm}^3 \pm 0.08\%}$	
$[H_2O_2] = 0.176 \text{ mol dm}^{-3} \pm 1.23\%$	
$[H_2O_2] = 0.176 \pm 0.002 \mathrm{mol}\mathrm{dm}^{-3}$	

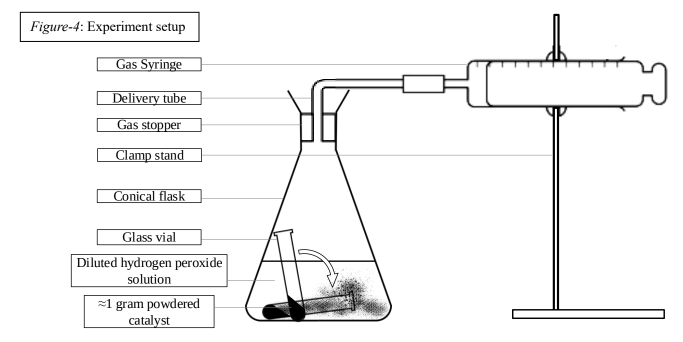
Experiment procedure

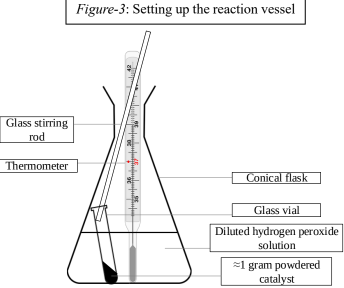
Repeat two trials of the following steps for each of the five catalysts:

- 1. The catalyst should be grinded with a pestle and molar if it contains visible large pieces until finely powdered.
- 2. Measure and transfer 1.0 ± 0.1 gram of the catalyst into a glass vial by using an electronic balance.
- 3. Transfer the prepared 200.0 ± 0.2 cm³ of 0.10 ± 0.01 mol dm⁻³ diluted hydrogen peroxide solution into the 500 cm³ conical flask.
- 4. Carefully lower the glass vial containing the catalyst into the conical flask by using a glass stirring rod as shown in *Figure-3*, make sure the glass vial rests on the wall of the conical flask so that the catalyst does not come into contact with the solution.



- 6. Stopper the conical flask with a gas stopper connected to the gas syringe with a delivering tube as shown on *Figure-4*. Check to make sure that the system is airtight.
- 7. Shake the conical flask to let the glass vial to fall into the hydrogen peroxide solution as demonstrated on *Figure-4*, start timing on the stopwatch the moment that the catalyst is exposed to the solution.
- 8. Measure and record the time taken for the reacting system to produce 10 cm³, 20 cm³, 30 cm³, 40 cm³, 50 cm³, 60 cm³ of gas.





Processing raw results

To evaluate the instantaneous initial rate r_0 of each reacting system, we must first be able to model the rate of reaction r as a function of time t from the raw measurements of t and V. The definition of rate of reaction r measured in terms of the volume of gas produced by a system is:

$$r = \frac{\Delta V}{\Delta t} \quad [16]$$

, hence it can be said that:

$$r = \frac{V_2 - V_1}{t_2 - t_1} \quad [17]$$

, where V_1 and V_2 are two consecutive volumes at which time is measured, while t_1 and t_2 are two consecutive measurements of time at V_1 and V_2 . It should be noted that [17] determines r as the average rate of reaction between the time interval $t_2 - t_1$. Hence if we are to assign a specific value of time to r, this value should be the average time t_{μ} of the interval, where:

$$t_{\mu} = \frac{t_2 + t_1}{2} \quad [18]$$

All raw data are processed using [17] and [18] to evaluate the behavior of the rate of reaction for each reacting system.

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Sample calculation of rate of reaction and average time: $Table-1$, processed data, for $10 \text{ cm}^3 \le V \le 20 \text{ cm}^3$					
Rate of reaction <i>r</i> :	$\Delta V = 20 \text{ cm}^3 - 10 \text{ cm}^3$ $\Delta V = 10 \text{ cm}^3$	$r = \frac{10 \text{ cm}^3}{29.1 \text{ s}}$	Average time t_{μ} :	$t_1 = 32 \text{ s}$	$t_{\mu} = \frac{32 \text{ s} + 61.1 \text{ s}}{2}$
	$\Delta t = 61.1s - 32s$ $\Delta t = 29.1s$	$r \approx 0.3 \text{ cm}^3 \text{ s}^{-1}$		$t_2 = 61.1 \ 0.4 \ s$	$t_{\mu} \approx 47 \text{ s}$
	$\Delta t = 29.18$				

Treatment of errors and uncertainties

To account for both the uncertainty in repeated trials of measurement and the instrument error in individual measurements at the same time, an aggregate uncertainty is calculated for every trial average as follows:

$$Aggregate error = \frac{Trial \ maximum - Trial \ minimum}{2} + \sum Instrument \ error \ [19]$$

, where each measured value's deviation from the average is added to the sum of the instrument error.

Sample calculation of aggregate error: $Table-1$, raw data, Trial average = 32 ± 2 s			
Trial maximum = 34.11 s	Trial average = $\frac{34.11+30.62}{2} \pm \left(\frac{34.11-30.62}{2} + 0.01\right)$ seconds		
Trial minimum = 30.62 s	Trial average = 32.365 ± 1.765 seconds		
Total instrument error = ± 0.01 Trial average $\approx 32 \pm 2$ seconds			

When uncertainties are propagated down into calculated values, the sum of the relative uncertainties associated with each value used in the calculation is taken, and then multiplied by the value of the result to give the propagated absolute error.

Propagated error =
$$\sum \frac{\text{Associated error}}{\text{Value used in calculation}} \times \text{Calculated value}$$
 [20]

Sample calculation of propagated error:
$$Table-1$$
, processed data, $r = 0.3 \pm 0.2$ cm³ s⁻¹

$$t_1 = 32 \pm 2 \text{ s}$$

$$r = \frac{(61.1 \pm 0.4) - (32 \pm 2)}{(20.0 \pm 2.5) - (10.0 \pm 2.5)} \text{ cm}^3 \text{ s}^{-1}$$

$$t_2 = 61.1 \pm 0.4 \text{ s}$$

$$r = \frac{29.1 \pm 8.25\%}{10 \pm 50\%} \text{ cm}^3 \text{ s}^{-1}$$

$$V_1 = 10.0 \pm 2.5 \text{ cm}^3$$

$$r = 0.291 \pm (0.291 \times 58.25\%) \text{ cm}^3 \text{ s}^{-1}$$

$$V_2 = 20.0 \pm 2.5 \text{ cm}^3$$

$$r \approx 0.3 \pm 0.2 \text{ cm}^3 \text{ s}^{-1}$$

Experiment results

Table-1: Data and observations for Manganese (IV) Oxide

Initial conditions and controlled variables, with aggregate error		
Average initial temperature across trials (°C)	21.1 ±0.2	
Average mass of catalyst across trials (grams)	1.019 ±0.003	
Volume of H ₂ O ₂ solution (dm ³)	200.0 ±0.6	
Concentration of H ₂ O ₂ solution (mol dm ⁻³)	0.176 ±0.002	

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Raw data					
Volume of gas V	Time <i>t</i>	Time t (Seconds)			
(cm ³)	Trial 1	Trial 2	Trial average		
$\pm 2.5 \text{ cm}^3$	± 0.01 seconds		with aggregate		
instrument error	instrument error		error		
10.0	34.11	30.62	32 ± 2		
20.0	60.72	61.45	61.1 ± 0.4		
30.0	96.78	93.29	95 ± 2		
40.0	133.92	140.63	137 ± 3		
50.0	193.05	188.23	191 ± 2		
60.0	260.47	256.87	259 ± 2		

Processed data				
Average time t_{μ}	Rate $r (\text{cm}^3 \text{s}^{-1})$			
(Seconds)	with propagated			
with propagated	error			
error				
47 ± 3	0.3 ± 0.2			
78 ± 2	0.29 ± 0.07			
116 ± 5	0.24 ± 0.04			
164 ± 6	0.19 ± 0.03			
225 ± 4	0.15 ± 0.02			

Qualitative Observations

- Catalyst is a finely powdered black solid.
- Instant bubbling when the catalyst is released into the prepared H₂O₂ solution.
- Catalyst has poor solubility, mostly remaining solid in the solution as black suspensions.

Table-2: Data and observations for Lead (II, IV) Oxide

Initial conditions and controlled		
variables, with aggregate error		
Average initial temperature	21.2	
across trials (°C)	±0.1	
Average mass of catalyst	1.016	
across trials (grams)	± 0.002	
Volume of H ₂ O ₂ solution	200.0	
(dm³)	±0.6	
Concentration of H ₂ O ₂	0.176	
solution (mol dm ⁻³)	± 0.002	

Raw data			
Volume of gas V	Time <i>t</i>	(Seconds	s)
(cm ³)	Trial 1	Trial 2	Trial average
± 2.5 cm ³ instrument error	± 0.01	seconds	with aggregate
mstrument error	instrume	ent error	error
10.0	5.82	6.25	6.0 ± 0.2
20.0	10.27	9.78	10.0 ± 0.3
30.0	15.61	13.42	15 ± 1
40.0	21.52	22.63	22.1 ± 0.6
50.0	28.13	30.22	29 ± 1
60.0	35.09	37.56	36 ± 1

Processed data	
Average time t_{μ} (Seconds) with propagated error	Rate r (cm ³ s ⁻¹) with propagated error
8.0 ± 0.5	3 ± 1
12 ± 1	1.9 ± 0.6
18 ± 2	1.7 ± 0.4
26 ± 2	1.5 ± 0.3
33 ± 2	1.4 ± 0.2

Qualitative Observations

- Catalyst is a finely powdered, vividly orange solid.
- Instant rapid bubbling when the catalyst is released into the prepared H₂O₂ solution.
- Catalyst has poor solubility, mostly remaining solid in the solution as orange suspensions.

Table-3: Data and observations for Potassium Iodide, where the system is left undisturbed

Initial conditions and controlled variables, with aggregate error		
Average initial temperature	20.9	
across trials (°C)	±0.2	
Average mass of catalyst	1.043	
across trials (grams)	± 0.003	
Volume of H ₂ O ₂ solution	200.0	
(dm^3)	±0.6	
Concentration of H ₂ O ₂	0.176	
solution (mol dm ⁻³)	± 0.002	

Raw data			
Volume of gas V	Time <i>t</i> (Seconds)		
(cm ³)	Trial 1	Trial 2	Trial average
± 2.5 cm ³ instrument error	± 0.01	seconds	with propagated
ilistrument error	instrume	nt error	error
10.0	30.63	32.67	32 [±] 1
20.0	65.85	70.51	68 ± 2
30.0	103.59	108.92	106 ± 3
40.0	143.13	152.28	148 ± 5
50.0	185.73	191.56	189 ± 3
60.0	230.77	262.81	250 ± 20

Processed data	
Average time t_{μ} (Seconds) with aggregate error	Rate r (cm ³ s ⁻¹) with propagated error
48 ± 3	0.3 ± 0.1
85 ± 5	0.26 \pm 0.07
123 ± 7	0.25 \pm 0.05
164 ± 8	0.23 ± 0.04
210 ± 20	0.22 ± 0.04

Qualitative Observations

- Catalyst is a powdered crystalline white solid.
- The catalyst is highly soluble, the solid catalyst dissolves instantly when released into the prepared H₂O₂ solution, followed by bubbling.
- The originally clear and colourless H₂O₂ solution turns into a clear brown colour after the catalyst is dissolved in it.
- The reactive system is sensitive to motion, where disturbing the system leads to observably more rapid bubbling.

Table-4: Data and Observations for Copper (II) Oxide

Initial conditions and controlled
variables, with aggregate error

Average initial temperature	20.8
across trials (°C)	± 0.1
Average mass of catalyst	0.968
across trials (grams)	± 0.004
Volume of H ₂ O ₂ solution	200.0
(dm^3)	± 0.6
Concentration of H ₂ O ₂	0.176
solution (mol dm ⁻³)	± 0.002

Raw data			
Volume of gas V	Time <i>t</i> (Seconds)		
(cm^3)	Trial 1	Trial 2	Trial average
± 2.5 cm ³ instrument error	± 0.01	seconds	with aggregate
mstrument error	instrumen	t error	error
10.0	179.93	217.67	200 ± 20
20.0	444.61	482.19	460 ± 20
30.0	836.17	1004.02	920 ± 80
40.0	1351.71	1572.23	1500 ± 100
50.0	1942.11	2207.48	2100 ± 100
60.0	2662.65	2905.25	2800 ± 100

Processed data	
Average time t_{μ} (Seconds) with propagated error	Rate r (cm ³ s ⁻¹) with propagated error
330 ± 40	0.04 ± 0.02
690 ± 90	0.022 ± 0.007
1200 ± 200	0.018 ± 0.006
1800 ± 200	0.016 ± 0.004
2400 ± 300	0.014 ± 0.003

Qualitative Observations

- Catalyst is a finely powdered black solid.
- Instant rapid bubbling when the catalyst is released into the prepared H₂O₂ solution.
- Catalyst has poor solubility, mostly remaining solid in the solution as black suspensions.

Table-5: Data and observations for Iron (III) Oxide

Initial conditions and controlled variables, with aggregate error		
Average initial temperature	21.3	
across trials (°C)	± 0.2	
Average mass of catalyst	0.942 + 0.002	
across trials (grams)	+ 0.002	

Volume of H₂O₂ solution 200.0 (dm^3) ± 0.6 Concentration 0.176 of solution (mol dm⁻³) ± 0.002

Attempts to collect data and qualitative observations

- Catalyst is a finely powdered red solid.
- No visual evidence of any reaction occurring after the catalyst is released into the prepared H₂O₂ solution initially, but tiny bubbles begin to emerge after around 600 seconds.
- Catalyst has poor solubility, mostly remaining solid in the solution as red suspensions.
- The amount of gas produced by the system in 1800 seconds is not measureable by the gas syringe used, no data could be collected on the prepared H₂O₂ solution as a result.
- An attempt was made to use the undiluted 1 mol dm⁻³ H₂O₂ solution with the catalyst, but still yielded no measurable result after 1800 seconds, hence measurement for this catalyst is terminated.

Table-6: Data and observations for **Potassium Iodide**, where the system is constantly **swirled**

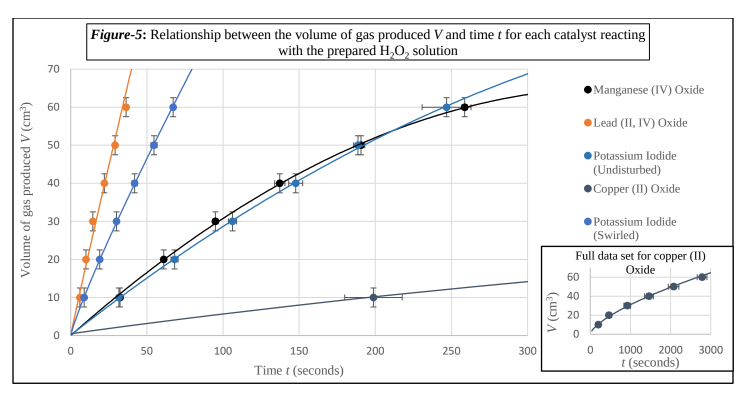
Initial conditions and controlled variables, with aggregate error			
Average initial temperature	21.1		
across trials (°C)	± 0.1		
Average mass of catalyst	1.012		
across trials (grams)	±		
Volume of H ₂ O ₂ solution	200.0		
(dm^3)	± 0.6		
Concentration of H ₂ O ₂	0.176		
solution (mol dm ⁻³)	± 0.002		

Raw data				
Volume of gas V	Time t (Seconds)			
(cm ³)	Trial 1	Trial 2	Trial a	verage
± 2.5 cm ³ instrument error	± 0.01	seconds	with ag	gregate
mstrument error	instrume	ent error	error	
10.0	9.15	8.47	8.8	$^{\pm}$ 0.4
20.0	19.34	18.76	19.1	\pm 0.3
30.0	31.17	29.02	3 <u>0</u>	± 1
40.0	43.68	40.21	42	± 2
50.0	56.72	52.64	55	± 2
60.0	67.33	67.22	67.28	\pm 0.08

Processed data		
Average time t_{μ} (Seconds)	Rate r (cm ³ s ⁻¹) with propagated	
with aggregate	error	
error		
13.9 ± 0.8	1.0 ± 0.4	
25 ± 1	0.91 ± 0.2	
36 ± 3	0.8 ± 0.2	
48 ± 4	0.8 ± 0.2	
61 ± 2	0.8 ± 0.1	

Qualitative Observations

- Catalyst is a powdered crystalline white solid.
- The catalyst is highly soluble, the solid catalyst dissolves instantly when released into the prepared H₂O₂ solution, followed by bubbling.
- The originally clear and colourless H₂O₂ solution turns into a clear brown colour after the catalyst is dissolved in it.
- Bubbling significantly intensifies after swirling begins.



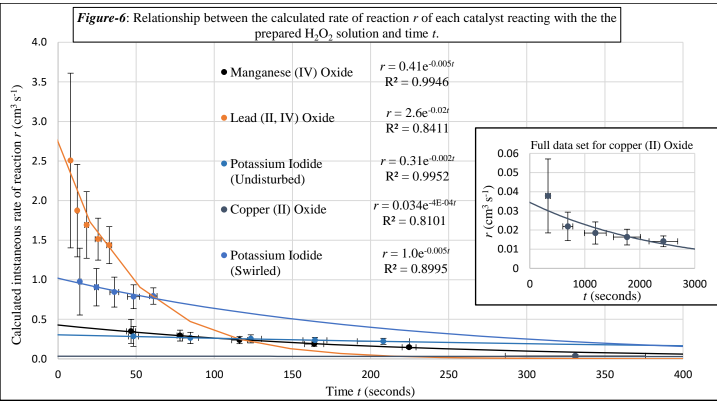


Figure-6 shows that the relationship between the rates of reaction r with time t for each of the reacting system can be accurately modelled as an exponential decay, where exponential trend lines agree with all data points within the acceptable margin of error. The equation for exponential decay takes the format of:

$$y = a e^{-kx}$$
 [21]

, where a and k are real constants. Following this equation, the value of the constant a will be the y-intercept of the trend line, hence the a value of each equation on Figure-6 will represent the initial rate r_0 of each reacting

system. As the y-intercept can be regarded as just another data point along the trend line, Δr_0 , the absolute uncertainty of the initial rate, can be evaluated by taking the average of the relative uncertainty of all calculated r values of data points plotted for a specific reacting system and multiplied by the derived value of r_0 , where:

$$\Delta r_0 = \frac{\sum (\Delta r/r)}{n} \times r_0 \quad [22]$$

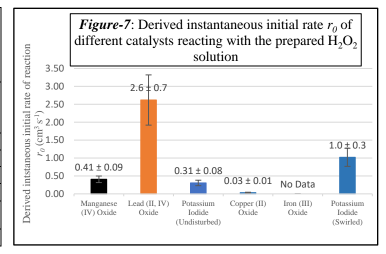
, *n* is the total number of *r* values involved in the calculation. In this way Δr_0 the will be an accurate representation of the overall degree of uncertainty in the set of *r* values that are involved in the calculation of r_0 .

Sample calculation of uncertainty for r_0 : <i>Table-7</i> , Lead (II, IV) Oxide, $r_0 = 2.6 \pm 0.7$ cm ³ s ⁻¹			
All calculated r values for Lead (II, IV) Oxide, data from Table-2: r_0 derived from Figure-6: 2.6 cm ³ s ⁻¹			
Rate r (cm ³ s ⁻¹) with propagated error	1 06 04 02 02		
3 ± 1	$\Delta r_0 = \frac{\frac{1}{3} + \frac{0.6}{2.9} + \frac{0.4}{1.7} + \frac{0.3}{1.5} + \frac{0.2}{1.4}}{\frac{1}{2} \times 2.6}$		
1.9 ± 0.6	$\Delta r_0 = \frac{3 - 2.7 - 1.7 - 1.3 - 1.4}{5} \times 2.6$		
1.7 ± 0.4	A 0.7 3 -1		
1.5 ± 0.3	$\Delta r_0 \approx 0.7 \text{ cm}^3 \text{ s}^{-1}$		
1.4 ± 0.2	$r_0 = 2.6 \pm 0.7 \mathrm{cm}^3 \mathrm{s}^{-1}$		
	• –		

All values of r_0 derived from *Figure-6* are tabulated on *Table-7*.

Table-7: Derived instantaneous initial rate r_0 of different catalysts reacting with the prepared H_2O_2 solution

Catalysts	Derived instantaneous	
	initial rate r_0 (cm ³ s ⁻¹),	
	with average error	
Manganese (IV) Oxide	0.41 ± 0.09	
Lead (II, IV) Oxide	2.6 ± 0.7	
Potassium Iodide (undisturbed)	0.31 ± 0.08	
Copper (II) Oxide	0.03 ± 0.01	
Iron (III) Oxide	No data, Too small to measure	
	given the procedure followed	
Potassium Iodide (swirled)	1.0 ± 0.3	



Conclusion

The experiment suggests that lead (II, IV) oxide will most rapidly catalyze the decomposition of hydrogen peroxide per unit mass out of all five tested catalysts, where it has the highest derived instantaneous initial rate of 2.6 ± 0.7 cm³ s⁻¹. Potassium iodide has the second highest initial instantaneous rate of 1.0 ± 0.3 cm³ s⁻¹ when swirled, but the rate drops to the second lowest at 0.31 ± 0.08 cm³ s⁻¹ when the system is left undisturbed. The margin of error for swirled potassium iodide does not overlap with the margin of error for lead (II, IV) oxide, which means that the results of this experiment definitively shows that lead (II, IV) oxide will most rapidly catalyze the decomposition of hydrogen peroxide out of the five tested catalysts.

An explanation for why lead (II, IV) oxide could most rapidly catalyze the decomposition of hydrogen peroxide is that lead (II, IV) oxide contains lead in both of its most common oxidation states, which allows the both reduction and oxidation of lead to happen simultaneously as seen in the system of equations [15]. Since the forward and backward transition between lead (II) and lead (IV) will have the exact opposite energy profile, the presence of lead in both oxidation states in lead (II, IV) oxide will cause hydrogen peroxide to more readily decompose through the reaction pathway involving the catalyst, as the oscillating simultaneous oxidation and reduction of lead will compensate for some of the activation energy of each phase of reaction in a catalytic cycle.

The high solubility of potassium iodide gives it the highest contact surface area with the reactant out of all five tested catalysts, which explains why it can relatively more rapidly catalyze the decomposition of hydrogen peroxide per unit mass compared to the remaining three tested catalysts. The reason why the rate of hydrogen peroxide decomposition catalyzed by potassium iodide increases when the system is swirled is most likely the

result of a systematic error, which that the hydrogen peroxide and potassium iodide solution might not be well-mixed during experiment. Under standard conditions, potassium iodide with a density of 3.12 g cm⁻³ (NCBI) is more than twice as dense as hydrogen peroxide at 1.44 g cm⁻³ (NCBI), hence it is possible that the aqueous hydrogen peroxide and aqueous potassium iodide in the undisturbed system would be separated by a density gradient, which greatly reduces the contact surface area between the catalyst and the reactants and results in a lower rate of reaction. Measures that can be taken to eliminate this systematic error will be discussed under evaluation.

Out of the remaining three tested catalysts, manganese (IV) oxide will most rapidly catalyze the decomposition of hydrogen peroxide per unit mass, with an instantaneous initial rate of 0.41 ± 0.09 cm³ s⁻¹, followed by copper (II) oxide with 0.03 ± 0.01 cm³ s⁻¹. The decomposition of hydrogen peroxide catalyzed by iron (III) oxide was the least rapid out of all five tested catalysts, as the rate of reaction was too slow for any meaningful data to be collected from the reacting system given the procedure followed. An improvement to the method that enables meaningful data to be collected data for iron (III) oxide is detailed under evaluation.

Although no literature of similar experiments can be found through secondary sources, data from an experiment investigating how rapidly chlorides of manganese, iron, and copper can catalyze the decomposition of hydrogen peroxide per unit mass was available. Since that transition metal ions are responsible for the catalysis of hydrogen peroxide decomposition, while the chloride and oxide ions are just spectators, one can expect similar results in terms of how rapidly a transition metal oxide and chloride can catalyze the decomposition of hydrogen peroxide when compared to the chlorides and oxides of other transition metals. While lead chloride and potassium iodide are not tested in the chloride experiment, the experiment concludes that manganese (II) chloride will most rapidly catalyze the decomposition of hydrogen peroxide per unit mass, followed by copper (II) chloride, while iron (II) chloride is will least rapidly catalyze the decomposition of hydrogen peroxide per unit mass compared to the other two transition metals (*Lee, et al.*). This strongly agrees with the results of this investigation for manganese (IV) oxide, copper (II) oxide, and iron (III) oxide, which suggests that the method of this experiment is largely valid, while the data collected from this experiment is mostly accurate and reliable.

Evaluation<u>Limitations and validity of method</u>

Γ	Ι	Τ	
Limitation	Impact and evidence	Resolution	
The dissolved potassium	The rate of hydrogen peroxide	Stir all reacting systems with an	
iodide solution is not	decomposition increased when the	magnetic stirrer at a constant rate of	
well-mixed.	system is swirled.	rotation in a container with a constant	
The swirling of the	The rate of swirling was not constant and	dimension when the catalyzed reaction	
system catalyzed by	not quantitatively controlled. This aspect	begins.	
potassium iodide was	of the experiment is not reproducible,		
done by hand.	hence leading to a loss of reliability.		
The 5 cm ³ volume	No meaningful data could be collected	Use a more accurate gas syringe.	
graduation of the gas	for iron (III) oxide, while there is a loss		
syringe used was too big.	of precision due to large instrument	Alternatively the experiment could also	
	errors.	be done using gravimetric methods,	
		where the change in mass is measured	
		against time on an electronic balance.	
Only two repeated trials	The error bars for the initial rate of the	Conduct at least three trials of	
of measurement are		measurements for each reacting system,	
conducted for each	relatively large $(2.6 \pm 0.7 \text{ cm}^3 \text{ s}^{-1})$,	so that unconcordant measurements can	
reacting system.	accounting for approximately 27% of the	be rejected and re-measured, which will	
	calculated value.	greatly improve the accuracy and	
		reliability of this experiment.	
Continues on the following page			

Limitation	Impact and evidence	Resolution
Surface area of the	This aspect of the experiment is not	
transition metal oxide	reproducible, hence leading to a loss of	metal oxides, use transition metal chips
catalysts are not	reliability.	that can be cut into regular shapes with
quantitatively controlled.		calculable surface area.
The experiment takes	This means the loss of control over the	To further maintain the constant
place over two weeks,	controlled variable [H ₂ O ₂], leading to	concentration of hydrogen peroxide
which is significantly	inaccurate measurements. However, the	over time, the hydrogen peroxide used
longer than the 10-20	impact of this should be minimal, as the	should not only be stored in an airtight
hour half-life of the	container for hydrogen peroxide is	container, but also placed in a fridge in
spontaneous	always sealed air-tight after use to	the absence of ultra violetlight to
decomposition of	maintain internal pressure. Since the	further slowdown the rate of
hydrogen peroxide in	decomposition of hydrogen peroxide	spontaneous decomposition.
open air under standard	produces oxygen gas, an air-tight	
conditions.	container with a stable internal pressure	
(FMC Corporation)	will sufficiently slow down the rate of	
	decomposition.	

Despite various limitations stated above, the method of this experiment is quite easily reproducible given a similar set of apparatus and material. Since the experiment did provide sufficient data that enables a meaningful conclusion to be drawn, the method of this investigation is overall valid.

Scope for further investigation

The presence of different types of catalyst is not the only variable that can change the rate of hydrogen peroxide decomposition. Further investigations can explore the rate of hydrogen peroxide decomposition under different temperatures and pressures, or the relationship between the rate of hydrogen peroxide decomposition and the intensity of ultraviolet light that the system is exposed to.

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