How does the hardness of water affect the rate of electrolysis?

Chemistry SL Internal Assesment

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August 24, 2025

1 Introduction and background knowledge

Electrochemistry was not invented until the dawn of 1800s, marked by the creation of the first electrolytic cell by Alessandro Volta, and the subsequent discovery of water electrolysis by William Nicholson(Fabbrizzi, 2019). Voltaic pile is considered the first artificial battery that provided continuous current to a circuit, laying a solid foundation for electricity-related studies. The discovery of water electrolysis by Nicholson, on the other hand, testified the composition of water and showed that water can be chemically decomposed. However, electrochemisty is still a relatively new subdiscipline in chemistry, and there are still many unknowns about the mechanism of electrolysis.

That work inspired Faraday's quantitative study on the general pattern of electrolysis, which resulted in Faraday's laws of electrolysis (Stock, 1991). Faraday concluded the amount of deposition is proportional to the amount of electric charge passing through the electrolyte, showing that electrolysis is foundamentally placing back of electrons. The study revealed the electric nature of many substances, formulated terms like ions, and greatly contributed to human's understanding of the submicroscopic world.

Nowadays, electrolysis plays a vital role in chemical industry fields like metal refining, electroplating, and mass production of sodium and chloride. Water electrolysis is also currently the main method for the production of hydrogen and oxygen (Flowers, 2019).

Electrolysis experiments also serve a pivotal role in chemistry education, since it allows students to explore the property of different materials and reveal the atomic nature of substances (e.g. showing that water consists of hydrogen and oxygen atoms).

Research question: How does the hardness of water (measured using MgCl₂ concentration) affect the rate of electrolysis?

2 Hypothesis and reasoning

3 Experiment design

3.1 Variables

- Independent variable: hardness of water (manipulated using the concentration of MgCl₂: 0.2, 0.4, 0.6, 0.8 and 1.0M).
- Dependent variable: Rate of electrolysis.
- Controlled variables: Temperature, atmospheric pressure, voltage, distance between electrodes, material of the electrodes and time duration of the experiment, etc. They are listed in Table 1.

3.2 Materials

- * 1 Iron stand with clamp ($\approx 50 \, \mathrm{cm}$)
- * 1 Side-armed U-shaped tube ($\approx 100\,\mathrm{ml}$), with a pair of cork stoppers penetrated by graphite electrodes.

Variable	Value	Table 1: Controlled variables Reason to control	Method to control	
Temperature	$25^{o}C$	Can significantly affect the rate	Complete the experiment during	
		of chemical reactions.	a short time period, in an air- conditioned room.	
Atmospheric	$1.0\mathrm{atm}$	Might change the volume and	Complete the experiment during	
pressure		pressure of the gas collected	a short time period.	
Voltage	$16\mathrm{V}$	Can affect the current passing	Use the same cell throughout the	
		through the electrolyte.	experiment.	
Distance	$5.0\mathrm{cm}$	Can affect the resistance be-	Use the same setup throughout	
between		tween electrodes.	the experiment.	
electrodes				
Material of	Graphit	e Can affect the conductivity and	Use the same electrodes	
the elec-		chemical property of the elec-	throughout the experiment.	
trodes		trodes.		
Time dura-	$20\mathrm{min}$	Short time leads to inaccuarcy,	Always time 20 min using a stop-	
tion of the		but the concentration chage	watch.	
experiment		and side-effect accumulates over		
_		time.		

^{* 1} Rubber tube.

- * 1 Gas syringe ($\approx 20 \,\mathrm{ml}$).
- * 1 Student power supply ($\approx 16\,\mathrm{V})$ and conducting wire.

3.3 Setup diagram

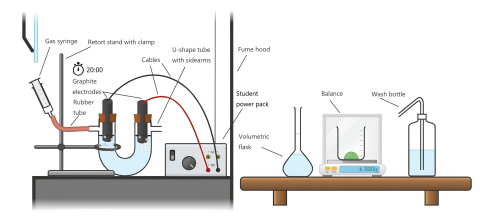


Figure 1: Setup diagram 1

¹This diagram is made using chemix and mspaint.

3.4 Procedure

- 1. Measure $m = 2.06 \,\mathrm{g}$ of MgCl₂·6H₂O in a beaker using analytical balance. Dissolve it with approximately 50 ml distilled water.
- 2. Transfer the solution to a 100 ml volumetric flask. Rinse the beaker with distilled water and put it into the flask as well.
- 3. Slowly add distilled water into the volumetric flask until it is just below the calibration mark. Use teat pipette to slowly add more distilled water until the base of the meniscus reaches the mark.
- 4. Insert the stopper and invert the solution for multiple times to ensure solution homogeneity.
- 5. Transfer the solution to the sidearmed U-shaped tube. Insert the corkstopper fitted with graphite electrodes and attach a syringe to one side arm using flexible tubing.
- 6. Mark the meniscus in the U-shaped tube using thin marker pen.
- 7. Switch the power supply to 16 V and turn it on. Start the timer at the same time.
- 8. Keep adjusting the syringe to make sure the meniscus is on the marked line
- 9. Turn off the power supply when the timer reaches 20 min.
- 10. Repeat steps 1-9 twice to get two additional trials.
- 11. Repeat steps 1-10 with $m = 4.12 \,\mathrm{g}$, $6.18 \,\mathrm{g}$, $8.24 \,\mathrm{g}$ and $10.30 \,\mathrm{g}$.

3.5 Risk Assessment

Conclusion: With PPE, teacher supervision, and basic laboratory discipline, overall risk is low for high school students.

4 Results

4.1 Raw data

4.2 Processed data

Processed data can be found in Table 4. The rate is plotted against the $MgCl_2$ concentration, and the graph is plotted in Figure 2.

4.3 Sample processing

5 Discussion and conclusion

The diagram suggests strong correlation between the rate and the concentration. The best fit line has

Table 2: Materials and Safety Information

Name	Potential Hazards	Measure/Precaution
Distilled	Physical, health, environmental	None
water	hazards: Not Classified	
Magnesium	Nonhazardous per GHS; treat all	Use protective gloves/clothing/eye pro-
chloride	lab chemicals with caution	tection. Avoid inhalation, ingestion, or
		skin/eye contact. First-aid measures
		included.
$\mathrm{Hydrogen}\ ^1$	H220: Extremely flammable gas;	P210: Keep away from ignition sources.
	H280: Gas under pressure	P377: Do not extinguish leaking gas fire
		unless leak can be stopped safely.
Oxygen 1	H270: May intensify fire; oxi-	P220: Keep away from combustibles.
	dizer. H280: Gas under pressure	P244: Valves free from oil/grease. In
		fire: Stop leak if safe.
Chlorine	Acute toxicity (oral, dermal, in-	Avoid breathing vapors; immediate
water 1,2	halation Cat. 5); skin/eye irrita-	medical attention if exposed. First-aid
	tion; inhalation toxicity for chlo-	and dilution instructions provided.
	rine gas	
Student	Below safe voltage (24V DC)	No special precautions. Disconnect
power sup-		power in unexpected events.
ply		

Sources: (Chemical Suppliers Ltd., 2023; Flinn Scientific, Inc., 2014b; Linde plc, 2022; ChemicalBook, 2025; Flinn Scientific, Inc., 2014a)

¹ Product. Other supplier of the same chemical substance is referred to for the MSDS table and hazard assessment. These products are in very low amount (hydrogen and oxygen less than 10ml, chlorine gas less than 1ml) and are not able to cause as much potential hazard, therefore disposal or accidental release is not listed in the table.

² Chlorine may be released as gas, or undergo complicated side reactions in the solution during the experiment. The experiment will therefore be conducted in a fume hood. However, it still share similar property with chlorine water, so chlorine water is used for hazard assessment.

Table 3: Raw data						
Experi-	Trial	Mass (g)	Solution	Gas Volume	Time (s)	
ment No.		(8)	Volume (mL)	(mL)		
1	1	4.0920 ± 0.0001	100.0 ± 0.5	4.2 ± 0.1	1200 ± 1	
1	2	4.0593 ± 0.0001	100.0 ± 0.5	3.9 ± 0.1	1200 ± 1	
1	3	4.0300 ± 0.0001	100.0 ± 0.5	3.2 ± 0.1	1200 ± 1	
2	1	8.1372 ± 0.0001	100.0 ± 0.5	6.9 ± 0.1	1200 ± 1	
2	2	8.1031 ± 0.0001	100.0 ± 0.5	7.0 ± 0.1	1200 ± 1	
2	3	8.1203 ± 0.0001	100.0 ± 0.5	7.1 ± 0.1	1200 ± 1	
3	1	12.1982 ± 0.0001	100.0 ± 0.5	10.0 ± 0.1	1200 ± 1	
3	2	12.2637 ± 0.0001	100.0 ± 0.5	10.0 ± 0.1	1200 ± 1	
3	3	12.1816 ± 0.0001	100.0 ± 0.5	10.0 ± 0.1	1200 ± 1	
4	1	16.2455 ± 0.0001	100.0 ± 0.5	12.9 ± 0.1	1200 ± 1	
4	2	16.2241 ± 0.0001	100.0 ± 0.5	12.8 ± 0.1	1200 ± 1	
4	3	16.2502 ± 0.0001	100.0 ± 0.5	13.0 ± 0.1	1200 ± 1	
5	1	20.1607 ± 0.0001	100.0 ± 0.5	14.1 ± 0.1	1200 ± 1	
5	2	20.3037 ± 0.0001	100.0 ± 0.5	14.9 ± 0.1	1200 ± 1	
5	3	20.3013 ± 0.0001	100.0 ± 0.5	14.3 ± 0.1	1200 ± 1	

Table 4: Processed data				
MgCl ₂ concentration	Rate of electro-			
$(\text{mol } \mathbf{L}^{-1})$	lysis $(10^{-7} \text{ mol s}^{-1})$			
0.200 ± 0.002	1.367 ± 0.051			
0.399 ± 0.001	2.392 ± 0.034			
0.601 ± 0.002	$3.405 {\pm} 0.017$			
0.799 ± 0.001	4.408 ± 0.034			
$0.996 {\pm} 0.004$	$4.931 {\pm} 0.137$			

- \bullet a gradient of $4.591\times10^{-7} \rm Ls^{-1}$
- and a v-intercept of $0.550 \times 10^{-7} \text{mols}^{-1}$.

The error on the independent variable was too insignificant to be shown on the graph. The errorbar on the y-axis is relatively small, showing high precision of the experiment.

The dots and error bars shows strong linear relationship between the rate and the concentration, which can be quantified with a high ${\bf r}^2$ value of 0.989032. The linear correlation suggests that the initial hypothesis is valid.

Not all the error bars are passed through by the best fit line. This might have resulted from the high concentration of the last several groups of trials. As is mentioned in the hypothesis part, high concentration can change the behaviour of the electrolyte and cause side reaction. This will be further discussed in Section 6.

6 Evaluation

The experiment was conducted successfully, providing sufficient data supporting the initial hypothesis. However, the expected deviation from linear relationship

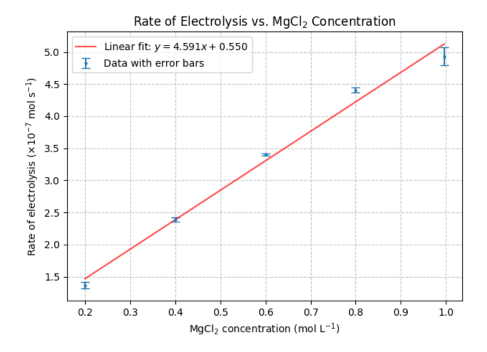


Figure 2: Rate of electrolysis against MgCl₂ concentration

is not significant in the data. There is still room for improvement.

- Imprecise marking: thin marker pen is used to mark the meniscus of the initial solution, which still results in a relatively thick stroke that can lead to unexpected uncertainty. Using a graduated, thin glass ware instead can minimize the uncertainty from adjustment.
- High concentration: a concentration higher than general cases where water electrolysis take place is adopted, due to the limited time and apparatus in the laboratory. That resulted in side reaction, like the production of chlorine taking place. Using more dilute solution, larger electrode and longer experiment time can help stop the side reaction.
- Bubbles' adhesion to electrodes: some gas produced is observed to have delayed or no detachment to the carbon electrodes. This can cause underming of the total gas production,

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