Determining the effect of electrolyte concentration of the voltage produced by galvanic cells

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1 Rationale

1.1 Background

Redox

Redox reactions are reactions that involve the transfer of electrons between atoms. These reactions commonly occur within galvanic cells, which within themselves contain two half cells.

Half Cells

Half cells consists of an electrolyte solution and electrode. When two half cells are connected through a load, and a salt bridge, one of these half cells will become the anode, where oxidation occurs, and the other will become the cathode, where reduction occurs. The oxidised cell will transfer its electrons across a wire, where they will then reach and be accepted by the reduced cell.

Each half cell has its own standard electrode potential (E^0) which dictates the magnitude of electron emission, or attraction it has under standard lab conditions. This is also known as the 'Voltage' of the cell

By measuring the difference in magnitude between half cells, the overall potential between the cells can be quantified.

$$E_{\text{Cell}}^0 = E_{\text{Cathode}}^0 - E_{\text{Anode}}^0$$

For this experiment, the half cells will be made up of zinc and copper, which have E^0 values of -0.7618V, and +0.3419V respectively. Cell's with this composition are also known as "Daniell Cells", and will have a theoretical voltage, under standard conditions, of:

$$(0.3419 - (-0.7618)) = 1.1037V$$

The Nernst Equation

The Nernst equation is a mathematical equation that allows the theoretical voltage potential of a cell to be quantified under non-standard conditions. While factors such as temperature and pressure throughout the experiment were not modified, the reaction quotient (Q) was indirectly altered by the change in $[CuSO_4]$.

The Nernst equation is defined as:

$$E_{\text{cell}} = E^0 - \frac{RT}{nF} \cdot \ln(Q)$$

Where:

- E = Theoretical electrode potential (voltage) of the cell
- $E^0 = \text{Standard electrode potential of the cell}$
- T = Temperature in kelvin
- R =Universal gas constant
- F = Faraday constant
- z = Number of moles of electrons transferred
- Q = Reaction Quotient

By substituting in constants for all values other than $[Cu^{2+}]$, the theoretical voltage can be found as a function of its concentration.

$$\begin{split} E_{\text{cell}} &= (0.3419 + 0.7618) - \frac{8.31446261815324 \cdot 298}{2 \cdot 96485} \cdot \ln \frac{[Zn^{2+}]}{[Cu^{2+}]} \because \text{Constants and environmental conditions} \\ E_{\text{cell}} &= (0.3419 + 0.7618) - \frac{8.31446261815324 \cdot 298}{2 \cdot 96485} \cdot \ln \frac{1}{[Cu^{2+}]} \because [Zn^{2+}] \text{ remains constant} \end{split}$$

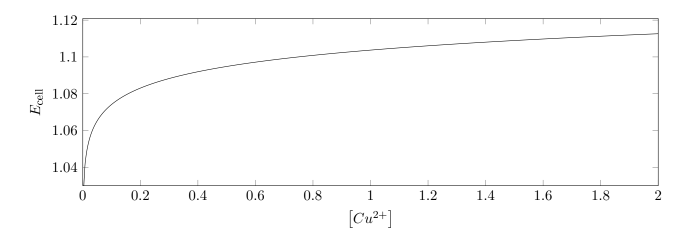


Figure 1: The theoretical relationship between E_{cell} and $[Cu^{2+}]$

Considering Figure 6, the theoretical change in $E_{\rm cell}$ is minuscule in comparison to the change in $[Cu^{2+}]$. When the x-axis doubles from 0.5 to 1, a change of only 0.009V is observed. If the relationship is further considered, with domain [0.5, 2], increasing concentration by a factor of 4 only results in an increase of 0.018V. It was questioned whether this relationship held true, especially as $\lim_{x\to 0}$. As Cu^{2+} increases from 0 to 1, it is theorised that the initial climb may appear more linear. If collision theory is considered, an increase in concentration is usually linearly proportional to the rate of reaction. When the concentration of ions is increased, more reactions should be able to take place per second, therefore increasing the output of the cell. It is hypothesised that at low concentrations, they may be too few ions in solution for the redox reaction to take place as expected. The concentration may rise linearly at low concentrations, but taper off as the reaction becomes 'saturated'. The current model for this process was questioned.

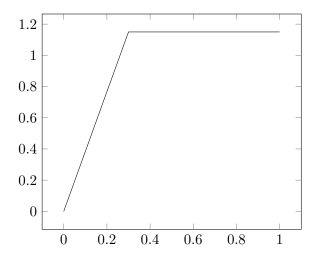


Figure 2: Hypothesized relationship between voltage and $[Cu_SO_4]$

Investigation of $0 < [Cu^{2+}] \le 1$ would theoretically be the most distinguishable from a logarithmic relationship if the observed relationship diverges, due to the distinct curve observed in the area.

Clearly $[Zn^{2+}] \neq 0$ and $[Cu^{2+}] \neq 0$, as this would either result in a division by 0, or $\ln(0)$, which both imply $E_{\text{cell}} = \text{undefined}$. Therefore $[CuSO_4] > 0$ for the duration of the experiment so direct comparison could be made.

1.2 Research Question

How does altering $[CuSO_4]$ at low concentrations (0.1-1 mol) affect its voltage output, and does the observed relationship follow a linear relationship.

2 Methodology

2.1 Modifications

The following modifications were made to the original experiment. The original method can be found in the appendix section.

Half Cell Compositions

The results of original experiment indicated that copper and aluminium produced the largest voltage. It was theorised that after multiple trials, the aluminium would oxidise, skewing results.

It was decided that copper and zinc would be more suitable as although they produced a smaller voltage, zinc would take longer to oxidise and therefore producer more consistent results.

Electrodes

Rather than thin strips, larger plates were bent into a "C" shape, slightly smaller than the diameter of the beaker they would be placed in. This maximised the surface area in contact with the electrolyte solution. Furthermore by reusing the same electrodes, this surface area was constant for all trials.

Measurement devices

A digital multimeter was substituted in place of the conventional analog volt meter. Since the digital multimeter uses digital logic to measure voltage, rather than a magnetic field induced by the flow of current, the digital multimeter should reduce the ions 'used up' during each measurement. Furthermore the digital output measures volts to 3 decimal places, leading to reduced uncertainty and increased precision.

Salt Bridge

The salt bridge was unaltered, and still made from filter paper, however, care was taken to ensure consistency in their size.

Procedure

The electrolyte solutions were pre-mixed, labelled, and stored in sealed containers prior to the start of the experiment. This reduced the potential for errors in procedure, as well as the time it took to conduct the experiment, which reduced the influence of environmental factors, such as changes in temperature, on results.

2.2 Method

Materials

- $30 \text{ml } CuSO_4$ at concentrations 1.00 M, 0.85 M, 0.70 M, 0.55 M, 0.40 M, 0.30 M, 0.20 M, 0.10 M
- 240ml 1M $ZnSO_4$
- 20ml KNO₃
- Petri dish
- 50ml Beaker x2
- Zinc sheet $\approx 4 \text{cm} \times 6 \text{cm}$
- Copper sheet $\approx 4 \text{cm} \times 6 \text{cm}$
- Emery paper
- Filter paper strips $\approx 2 \text{cm} \times 10 \text{cm}$
- Digital Multimeter
- Alligator clips
- 150ml Plastic bottle x16
- Tweezers
- 1L Distilled water

• Emery paper

Procedure

- 1. Pour varying amounts of $CUSO_4$ into 150ml plastic bottles and dilute with distilled water to form solutions with concentrations 1.00M, 0.85M, 0.70M, 0.55M, 0.40M, 0.30M, 0.20M and 0.10M.
- 2. Fill 8 of the 150ml plastic bottles with 30ml of 1M $ZnSO_4$ and set aside with copper solutions.
- 3. Polish electrodes with emery paper until they their entire surface is free of visible oxidation then connect to alligator clips.
- 4. Connect alligator clips to multimeter and set to DC voltage mode.
- 5. Transfer the contents of a $CUSO_4$, and $ZnSO_4$ container into separate beakers.
- 6. Wet salt bridge with KNO_3 and rest over the lip of both beakers, so that it is partially submerged in both.
- 7. Add the electrodes to the solution and record result as soon as value settles.
- 8. Remove electrodes immediately and wash with distilled water. Stir electrolyte solutions with a glass stir rod. Remove salt bridge and dispose of.
- 9. Reconstruct cell with a new salt bridge, add electrodes back, and repeat measurementation so that there are three trials in total for any given concentration.
- 10. Safely dispose of electrolyte solutions, wash equipment, and repeat for next CUSO₄ concentration.

2.3 Risk assessment

While the electrolytes utilised in this experiment are considered to be harmful if ingested or exposed to a persons eyes, there is no risk of injury via inhalation or accidental skin contact as long as appropriate action is taken (ventilation, washing affected area). PPE was worn by all participants to protect their eyes (Lab goggles), body (lab coat) and hands (nitrite gloves). The electrolyte materials are also considered environmental hazards $(CuSO_4, ZnSO_4)$, however since the quantities used were so low, they could safely be disposed of in the sink with after dilution. For other risks or more details, see risk assessment table attached as an appendix.

3 Results

$[CuSO_4]$		Voltage		Mean	sigma
	Trial 1	Trial 2	Trial 3		
1	0.703	0.678	0.661	0.681	0.021
0.85	0.664	0.623	0.488	0.592	0.088
0.7	0.557	0.455	0.472	0.495	0.051
0.55	0.457	0.422	0.413	0.431	0.022
0.4	0.438	0.412	0.416	0.422	0.013
0.3	0.498	0.47	0.476	0.481	0.014
0.2	0.458	0.487	0.399	0.4725	0.0145
0.1	0.405	0.456	0.463	0.441	0.029

Figure 3: Raw data with calculations. Outliers (highlighted in red) not included in calculations

3.1 Analysis of Evidence

Identification of relationships and trends.

Across all trials, the voltage produced by the cell was lower than what the Nernst equation predicted. As $[CuSO_4]$ doubled from 0.1 to 0.2, the average E_{cell} was scaled by a factor of 1.07. As it doubled again from 0.2 to 0.4, it is observed that the average E_{cell} was scaled by a factor of 0.89, which was smaller than the initial value. Clearly further investigation was required.

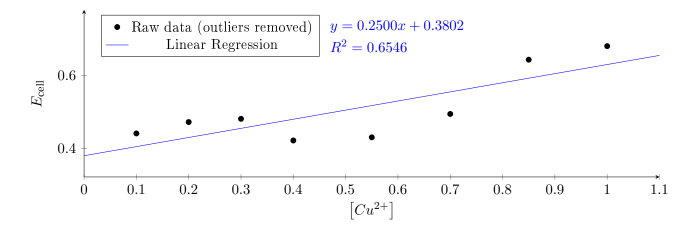


Figure 4: Results (outliers removed) with linear line of best fit.

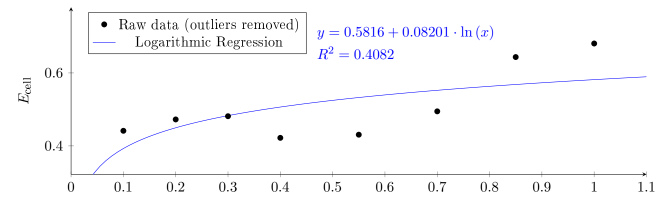


Figure 5: Results (outliers removed) with logarithmic line of best fit.

The scatter plot produced using experimental data suggests a linear relationship. Using desmos, linear and logarithmic lines of best fit were found. Considering Figure 4 and Figure 5, the linear regression showed a greater correlation than the logarithmic.

$$R_{\rm Lin}^2 = 0.6426 > R_{\rm Log}^2 = 0.4082$$

The relationship between $[Cu^{2+}]$ and E_{cell} was found to be linear, with a slope of 0.2500, and vertical shift of +0.3802. The presence of vertical shift indicates divergence in the expected and actual relationships. sised that as $\lim_{[Cu^{2+}]\to 0}$, $E_{\text{cell}}\to 0$, as without another electrolyte to accept ions, the cell would have no potential. This suggests an incorrect hypothesis or error in procedure. Furthermore, it is unknown whether this relationship holds for higher $[Cu^{2+}]$ values.

It should be noted that the data is limited, and does not perfectly match the linear model as the \mathbb{R}^2 value is poor.

$$R_{\rm Lin}^2 = 0.6546 < 1$$

Finding Uncertainty in the gradient

For $[CuSO_4] = 1$ mol

$$\sigma = \pm \frac{\text{max} - \text{min}}{2}$$

$$= \pm \frac{0.703 - 0.661}{2}$$

$$= \pm 0.021$$
Avg = 0.6807

For $[CuSO_4] = 0.1$ mol

$$\sigma = \pm \frac{\text{max} - \text{min}}{2}$$
$$= \pm \frac{0.463 - 0.405}{2}$$
$$= 0.029$$

$$Avg = 0.4413$$

Solve using linear regression between $(Avg + \sigma, Avg - \sigma)$ and $(Avg - \sigma, Avg + \sigma)$ \therefore Gradient_{Max} =

ToDo

- Get max and min slopes for the linear relationship
- Use max and min slopes to find absolute uncertainty in gradient and in offset
- Present this as a final function with \pm uncertainty
- Comment on this

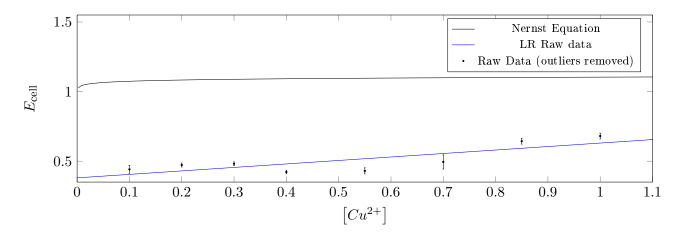


Figure 6: The theoretical relationship between E_{cell} and $[Cu^{2+}]$. (LR stands for Linear Regression)

Figure 7 depicts a trend observed where as trials continued, the voltage of each subsequent trial decreased slightly. Procedure was considered and it was theorised that this was due to reuse of the same electrolyte solution over multiple trials. The number of ions left on solution likely decreased as measurements were taken and and differed due to inconsistent duration of the measurement phase. This was theorised to have lead to a smaller voltage. To accurately consider this trend, and evaluate the reliability of the relationship, the average of all data points against what 'trial' they were collected during was considered. While the change in voltage is only on average -0.02V per trial, considering $R^2 = 0.9922$, the linear regression indicates a very high correlation. However due to the large range of values considered, the uncertainty in this relationship is so high that the minimum trendline has a positive gradient, therefore further investigation is required to confirm this.

4 Discussion

4.1 Improvements

Sources of error/improvements

• Alligator clips in solution (floating voltage)

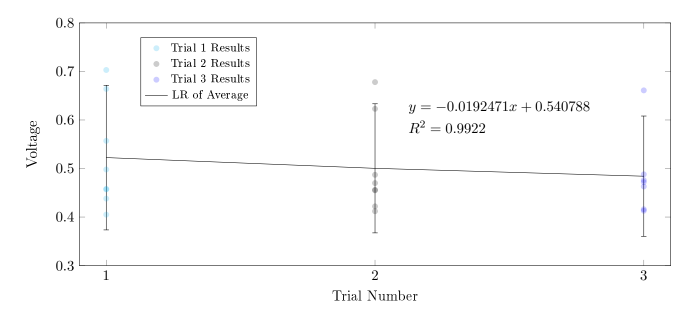


Figure 7: Line of best fit for average voltage over sequential trials (outliers removed)

- Inconsistent measurement time. (outlier is main case of this).
- Alligator clips sitting in electrolyte during experiment.
- Temperature and pressure unknown on day of experimentation. This means that the Nernst equation cannot account for them. Therefore difference in between Nernst equation and actual results n
- increasing number of trials would have allowed voltage drop off to be found and also would have improved reliability after outliers removed.

Observation	Source of error	Solution	
Floating voltage on multimeter in between trials. Voltage fluctuation throughout trials	Multimeter wires in contact with puddle of electrolyte at intermediate connection point between meter and probes. A minor redox reaction may have been occurring on the bench, affecting readings.	Isolate multimeter from experimental setup and ensure probes are only connected to electrodes.	
Inconsistent voltage readings and high electrolyte consumption over relatively short readings	Alligator clips of unknown material were submerged in the half cells electrolyte solution. Likely dissociating and affecting half cell composition unintentionally.	Alter shape of electrodes to a portion of it extends beyond the edge of the beaker so that alligator clips can be attached without being at risk of contacting electrolytes.	
Large voltage drops between trials	Inconsistent measurement times and gaps between trials. During some trials, the multimeter disconnected midway through measurement and when reconnected the voltage reading was significantly lower than other trials.	Perform all step-lens at set time intervals. Take care to ensure setup remains the same throughout all trials.	
Consistently lower voltages than expected	Unknown. Possibly resistance due to small salt bridge or poor wiring, oxide layer on electrodes, incorrect electrolyte concentrations.	Follow procedure with more care. Use more suitable equipment.	

Appendices

Risk Assessment

Item/Substance	Potential Hazards	Standard Handling Procedures	Disposal
Alligator clip with lead	Clip may cause pain and injury if applied to skin.		
Glass beaker	Breakage of beaker. Cuts from	Inspect and discard any chipped or	
	chipped rims.	cracked beakers. Sweep up broken glass	
		with brush and dustpan; do not use fin-	
		gers.	
Metal tweezers	Can be used as a weapon if long and sharply pointed.		
Disposable plastic gloves	May easily be punctured, allowing	Take care not to puncture. Check for	
	entry of liquid. Latex gloves may	punctures before use. Use a type of	
	cause an allergic reaction to some	glove that is suitable for the chemicals	
	people	to be used.	
Lab coat	Flammable. Sleeves may catch on		
	objects and knock them over.		
Safety glasses	Scratched or dirty glasses may hin-	Each person should preferably have	
	der vision, causing headaches dur-	own safety glasses. Check and, if nec-	
	ing prolonged use.	essary, clean glasses before each use	
Filter paper	Flammable. Used filter paper	After use, dispose of residue and filter	Dispose of residue and filter paper
	may contain harmful residues.	paper appropriately.	appropriately.
Copper (sheet)	Not toxic.		<1 kg/day may be placed in the garbage. Larger quantities should be retained for collection by a waste service or metal recycler.
Copper(II) sulfate (>0.94	Toxic. Irritates skin and eyes.	<5 mL/day may be poured down the	service of metal recycler.
M)	Toxic. Ifficates skill and eyes.	drain. Larger quantities should be	
IVI)		placed in a Copper waste container.	
Zinc (pieces)	Not toxic to humans.	praced in a copper waste container.	May be placed in the garbage.
Zinc nitrate (0.79-1 M)	Toxic Irritates skin, eyes and		
Zine nitrate (0.15 1 M)	lungs. Harmful if swallowed.		10 times the volume of water and
	Causes skin irritation. Causes se-		poured down the drain. Larger
	rious eye irritation. Very toxic		quantities should be placed in a
	to aquatic life with long lasting		Zinc waste container.
	effects		Zine waste container
Potassium nitrate (0.1-1	May irritate eyes and skin GHS		<1 L/day may be poured down the
M)	data: Not classified as a hazardous		drain in a stream of water.
•	chemical.		
Wash bottle	May be used to spray others.	Preferably use distilled water. Change	
		water regularly to avoid microbial growth.	