Pre-Lab Assignment

Before coming to lab:

- Read the lab thoroughly.
- Answer the pre-lab questions that appear at the end of this lab exercise. The questions should be answered on a separate (new) page of your lab notebook. Be sure to show all work, round answers, and include units on all answers. Background information can be found in Chapter 20, especially sections 20.3-20.6 and 20.9 in your textbook (Brown and LeMay).
- Follow the guidelines in the "Lab Notebook Policy and Format for Lab Reports" section of the lab manual to complete in your lab notebook the following sections of the report for this lab exercise: Title, Lab Purpose, Procedure and Data Tables.

Purpose

This lab activity has four parts. In Part A and B you will measure the potential of several voltaic cells and compare your measured values to the accepted values that can be found in the tables in your textbook. In Part C, you will construct a simple electrolytic cell, which you can then use to electroplate a nickel (or other coin) with copper. The plated coin will be a souvenir that you can take with you as your reward for successfully completing an entire year of chemistry! In Part D, you will make observations that will allow you to determine the reactions that are occurring in an electrolytic cell when there is more than one possibility.

Background

As you saw in Chem 1A, oxidation–reduction reactions involve a transfer of electrons. In a spontaneous redox reaction, electrons flow from the reactant that is oxidized (reducing agent) to the reactant that is reduced (oxidizing agent). If the two half-reactions are separated by a wire, this flow of electrons, instead of occurring at the surface of the metal, occurs through the external wire and an electric current is generated. This is called a voltaic cell (or galvanic cell), and is exactly how a battery works. Batteries, like the ones found in a flashlight or a calculator, contain oxidizing and reducing substances. As the electrons are transferred, they are run through the device in order to provide the voltage necessary to power the flashlight or calculator.

A good analogy for the flow of electrons is the flow of water. Water flows spontaneously downhill. Dams and waterwheels are examples of ways that the energy of flowing water is tapped to generate power. In order for a redox reaction to serve as a source of power, the reaction must be spontaneous. You will examine these processes in Parts A and B of the lab.

Sometimes we want water to flow uphill. In this case, we need to supply energy in the form of a pump to make this happen. Similarly, if a redox chemical reaction isn’t spontaneous, we can use electricity to make the reaction occur. An electrolytic cell is a device that uses electricity to drive a non–spontaneous redox reaction. For example, water can be separated into hydrogen and
oxygen gas (a non–spontaneous reaction) using electricity. You will examine these processes in Part C and D of the lab.

In short, the field of electrochemistry has two important applications- the use of spontaneous redox reactions to generate electricity, and the use of electricity to force non–spontaneous redox reactions to occur.

Voltaic Cells

In Part A of this lab activity you will measure the potential of several voltaic cells. A typical voltaic cell, such as the one in the figure on the next page consists of two half-cells linked by a wire and a salt bridge. Each half-cell consists of metal electrode in contact with a solution containing a salt of that metal. One half-cell (think beaker in today’s lab) functions as the anode, where the oxidation reaction takes place. Oxidation involves the loss of one or more electron(s) from the metal. For the first voltaic cell you will make and that is diagramed on the next page, the oxidation reaction is:

At the anode: \( \ce{Zn(s) -> Zn^{2+}(aq) + 2e^-} \)  \hspace{1cm} Reaction 1

The other half-cell (think beaker in today’s lab) functions as the cathode, where the reduction reaction takes place. For the first voltaic cell you will make and that is diagramed on the next page, the reduction reaction is:

At the cathode: \( \ce{Cu^{2+}(aq) + 2e^- -> Cu(s)} \)  \hspace{1cm} Reaction 2

Combining these reactions and cancelling electrons, the overall reaction is

\( \ce{Zn(s) + Cu^{2+}(aq) -> Zn^{2+}(aq) + Cu(s)} \)  \hspace{1cm} Reaction 3

In this case, the zinc metal goes into solution as zinc ions, and the copper ions form solid copper. In addition, electrons flow from the anode (where they are released) to the cathode (where they are taken in) via the wire. As the figure on the following page shows, they are also flowing though a voltmeter. Inserting a voltmeter into the circuit between the half-cells permits a measurement of the voltage, or potential difference between the half-cells. In general, this voltage is designated by the symbol, \( E \). When the solutions are 1 Molar (1.0 M) with respect to the ions involved in the oxidation and reduction reactions, the cell is called a standard cell and its voltage is known as a standard potential, \( E^\circ \). Many textbooks and references books contain tables of standard reduction potentials, which show the values of \( E^\circ \) for various reduction half-reactions. The \( E^\circ \) in the tables are all relative to the reduction of hydrogen as shown in the reaction:

\( \ce{2H^+(aq) + 2e^- -> H_2(g)} \)

The potential of this reaction has been assigned a potential of 0.00 Volts (V) and is called the standard hydrogen electrode.

The final piece of a voltaic cell is a salt bridge which allows migration of ions to prevent an imbalance of charge from building up as electrons leave the anode and move to the cathode.
Predicting the Potential of a Voltaic Cell

For today’s lab, you will be predicting the potential that the voltaic cells you construct should generate and then compare your predicted values to your measured values. To order to predict the voltage, you will need to use the table of Reduction Half-Reaction Potentials below. A more complete table is in your book on page 1064.

<table>
<thead>
<tr>
<th>Reduction Half-Reaction</th>
<th>$E^{\text{red}}$ V</th>
</tr>
</thead>
<tbody>
<tr>
<td>Au$^{3+}$ (aq) + 3e$^-$ → Au (s)</td>
<td>+1.50</td>
</tr>
<tr>
<td>Pt$^{2+}$ (aq) + 2e$^-$ → Pt (s)</td>
<td>+1.20</td>
</tr>
<tr>
<td>Ag$^+$ (aq) + e$^-$ → Ag (s)</td>
<td>+0.80</td>
</tr>
<tr>
<td>Cu$^{2+}$ (aq) + 2e$^-$ → Cu (s)</td>
<td>+0.34</td>
</tr>
<tr>
<td>2H$^+$ (aq) + 2e$^-$ → H$_2$(g)</td>
<td>0.000</td>
</tr>
<tr>
<td>Pb$^{2+}$ (aq) + 2e$^-$ → Pb (s)</td>
<td>-0.13</td>
</tr>
<tr>
<td>Sn$^{4+}$ (aq) + 2e$^-$ → Sn (s)</td>
<td>-0.14</td>
</tr>
<tr>
<td>Ni$^{2+}$ (aq) + 2e$^-$ → Ni (s)</td>
<td>-0.28</td>
</tr>
<tr>
<td>Fe$^{2+}$ (aq) + 2e$^-$ → Fe (s)</td>
<td>-0.44</td>
</tr>
<tr>
<td>Zn$^{2+}$ (aq) + 2e$^-$ → Zn (s)</td>
<td>-0.76</td>
</tr>
<tr>
<td>Al$^{3+}$ (aq) + 3e$^-$ → Al (s)</td>
<td>-1.66</td>
</tr>
<tr>
<td>Mg$^{2+}$ (aq) + 2e$^-$ → Mg (s)</td>
<td>-2.37</td>
</tr>
<tr>
<td>Na$^+$ (aq) + e$^-$ → Na (s)</td>
<td>-2.71</td>
</tr>
</tbody>
</table>
Notice that all reactions are listed as reduction reactions in the table. However, in voltaic cells there are always one oxidation half-reaction and one reduction half-reaction taking place. This means that we must “flip” one of the reactions in the table in order to calculate the potential created in the voltaic cell. Recall that voltaic cells are spontaneous processes \( E_{cell} = \text{positive number} \) so nature keep as a reduction reaction the reaction that has the most positive \( E_{\text{red}} \) and will “flip” the other reaction.

### Example: Calculation of the Cell Potential of Standard Voltaic Cells:

Whenever two standard half-cells are joined to create a voltaic cell as in Figure 1, one of the half-cells will function as the anode where oxidation will occur. The other half-cell will serve as the cathode where reduction will occur. Consider the half-reactions for a voltaic cell composed of copper and zinc.

*From the Reduction Half-Reaction Potentials Table:*

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Potential</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Cu}^{2+} \text{(aq)} + 2e^- \rightarrow \text{Cu} \text{(s)} )</td>
<td>+0.34</td>
</tr>
<tr>
<td>( \text{Zn}^{2+} \text{(aq)} + 2e^- \rightarrow \text{Zn} \text{(s)} )</td>
<td>-0.76</td>
</tr>
</tbody>
</table>

When considering which one of these reactions will stay as the reduction reaction and which one will be the oxidation reaction, the species that generates the more positive potential prefers to be reduced. In this example, that is \( \text{Cu}^{2+} \). Therefore, the top reaction will be the reduction half-reaction and the reaction involving zinc will be the oxidation reaction. To determine the voltage, we will use the copper reaction as written in the table, but reverse the zinc reaction.

\[
\begin{align*}
\text{Cu}^{2+} \text{(aq)} + 2e^- &\rightarrow \text{Cu} \text{(s)} + 0.34 \\
\text{Zn} \text{(s)} &\rightarrow \text{Zn}^{2+} \text{(aq)} + 2e^- + 0.76
\end{align*}
\]

Notice that the potential for the reversed zinc reaction is the same value, but opposite in sign. The overall potential for this voltaic cell is the sum of these two values.

\[
E^o_{cell} = 0.34 \text{ V} + 0.76 \text{ V} = 1.10 \text{ V}
\]

Note that the half-cell voltages are never multiplied but are simply added together.

The overall reaction will therefore be:

\[
\text{Zn} \text{(s)} + \text{Cu}^{2+} \text{(aq)} \rightarrow \text{Zn}^{2+} \text{(aq)} + \text{Cu} \text{(s)}
\]

### Calculation of the Cell Potential of a Voltaic Cell NOT at Standard Conditions (1.0 M, 298 K, 1atm):

If the potential that is measured is not a standard cell, then the **Nernst equation** can be used to calculate the standard cell potential.

**Nernst Equation**

\[
E_{cell} = E^o_{cell} - \frac{RT}{nF} \ln Q
\]

Where...

- \( R = 8.14 \text{ J/mol*K} \)
- \( F = \text{Faraday’s constant (96485 Coulombs/mole e\(^{-}\))} \)
- \( T = \text{Temperature (K)} \)
- \( Q = \text{Reaction quotient} \)
- \( n = \text{number of electrons transferred} \)
- \( E^o = \text{standard cell potential (1 M solutions at 298K and 1 atm)} \)
- \( E_{cell} = \text{Cell potential at nonstandard conditions} \)
Electrochemical Reduction of Cu<sup>2+</sup> to Copper on the Surface of a Coin

In Part C of the lab, copper plating will be accomplished by submerging both a coin and a strip of copper in a solution of copper (II) sulfate. At the anode, copper is oxidized (Cu→Cu<sup>2+</sup> + 2e⁻) and at the cathode is reduced (Cu<sup>2+</sup> + 2e⁻→ Cu). The overall process consists of simply moving copper atoms from one electrode to the other. Since this is not a spontaneous process, an electrical current is needed to drive the process. The total amount of charge used in the reaction can be found by multiplying the current (I) times the time (t). Since the current will vary, we will find the area under the line using the area tool in Pasco’s DataStudio.

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**Example: Calculation of the Cell Potential for a Voltaic Cell at Non-standard Conditions**

Consider a voltaic cell made of a Sn electrode in 0.10 M Sn<sup>2+</sup> solution in one half-cell and Al in 0.10 M Al<sup>3+</sup> solution in the other at a temperature of 273 K.

First find the potential at standard conditions (E<sub>0</sub>)

<table>
<thead>
<tr>
<th>Reaction</th>
<th>E&lt;sub&gt;0&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sn&lt;sup&gt;2+&lt;/sup&gt; (aq) + 2e⁻→ Sn (s)</td>
<td>-0.14</td>
</tr>
<tr>
<td>Al&lt;sup&gt;3+&lt;/sup&gt; (aq) + 3e⁻→ Al (s)</td>
<td>-1.66</td>
</tr>
</tbody>
</table>

Between these two reactions, Sn<sup>2+</sup> prefers to be reduced since it is “more positive” (closer to zero in this case). Therefore, the Al<sup>3+</sup> reaction will have to be reversed.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>E&lt;sub&gt;0&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sn&lt;sup&gt;2+&lt;/sup&gt; (aq) + 2e⁻→ Sn (s)</td>
<td>-0.14</td>
</tr>
<tr>
<td>Al (s)→Al&lt;sup&gt;3+&lt;/sup&gt; (aq) + 3e⁻</td>
<td>+1.66</td>
</tr>
</tbody>
</table>

E<sub>cell</sub> = -0.14 V + 1.66 V = 1.52 V

The overall cell reaction after balancing and cancelling electrons will be

Multiply to get e- the same in each reaction

Sn<sup>2+</sup> (aq) + 2e⁻→ Sn (s)    X 3
Al (s)→Al<sup>3+</sup> (aq) + 3e⁻    X 2

### Overall Reaction

3 Sn<sup>2+</sup> (aq) + 6e⁻→ 3 Sn (s)
2 Al (s)→2 Al<sup>3+</sup> (aq) + 6e⁻

Substituting into the Nernst Equation

\[
E_{cell} = E^0_{cell} - \frac{RT \ln Q}{nF} = E^0_{cell} - \frac{RT \ln [Al^{3+}]^2}{nF [Sn^{2+}]^3}
\]

\[
E_{cell} = 1.52 \text{ v} - \frac{8.31 \text{ J/mol*K}(273 \text{ K})}{6 \times 96485 \text{ Coulombs/mole}} \times \ln \frac{[0.1]^2}{[0.1]} = 1.52 \text{ V} - 0.0099V = 1.51 \text{ V (nearly the same as the standard cell potential!)}
\]

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**Electrolytic Reduction of Cu<sup>2+</sup> to Copper on the Surface of a Coin**

In Part C of the lab, copper plating will be accomplished by submerging both a coin and a strip of copper in a solution of copper (II) sulfate. At the anode, copper is oxidized (Cu→Cu<sup>2+</sup> + 2e⁻) and at the cathode is reduced (Cu<sup>2+</sup> + 2e⁻→ Cu). The overall process consists of simply moving copper atoms from one electrode to the other. Since this is not a spontaneous process, an electrical current is needed to drive the process. The total amount of charge used in the reaction can be found by multiplying the current (I) times the time (t). Since the current will vary, we will find the area under the line using the area tool in Pasco’s DataStudio.
An example of the calculations needed for Part C of the lab can be found on pg. 861 of your textbook.

In Part D of the lab, you will run an electrical current through a solution and make observations in order to determine the identity of the products of the electrolysis reaction. Since all the solutions contain water, one of the possibilities is that water will undergo electrolysis. The following half-reactions occur in the electrolysis of water:

**Oxidation half-reaction for water (anode)**

\[2H_2O(l) \rightarrow O_2(g) + 4H^+(aq) + 4e^-\]

**Reduction half-reaction for water (cathode)**

\[2H_2O(l) + 2e^- \rightarrow H_2(g) + 2OH^-(aq)\]

Electrolysis of an aqueous solution may generate products other than oxygen or hydrogen if the electrolyte contains ions that are more easily oxidized or more easily reduced than water molecules. The electrolysis of aqueous silver nitrate (AgNO₃), for example, produces oxygen at the anode and silver metal at the cathode. If you observe this reaction, you will observe the formation of Ag at the cathode since the reduction of silver ions (Ag⁺) to silver (Ag) occurs more readily than reduction of water. At the anion, bubbles are seen which indicates that O₂ gas is being produced. The overall reaction is the sum of the oxidation and reduction half-reactions:

**Oxidation half-reaction (anode):**

\[2H_2O(l) \rightarrow O_2(g) + 4H^+(aq) + 4e^-\]

**Reduction half-reaction (cathode):**

\[4Ag^+(aq) + 4e^- \rightarrow 4Ag(s)\]

**Overall reaction:**

\[2H_2O(l) + 4Ag^+(aq) \rightarrow O_2(g) + 4Ag(s) + 4H^+(aq)\]

**Procedure**

**Part A—Construction of Voltaic Cells and Measurement of Cell Potentials**

1. Obtain about 25 mL of 1 M KNO₃. Cut up pieces of filter paper into six small narrow strips. Soak the strips of filter paper in the KNO₃. These will serve as your salt bridges.

2. Obtain strips of Cu, Zn, Pb, and Ni. Clean the strips of metal by sanding and/or dipping in 3 M HCl.

3. Place approximately 25 mL of 0.1 M CuSO₄ in a 100 mL beaker. Place the strip of Cu in the beaker as well.
4. Place approximately 25 mL of 0.1 M ZnSO₄ in another 100 mL beaker. Place the strip of Zn in this beaker as well.

5. Insert the salt bridge so that it is submerged in the solutions in both beakers.

6. Obtain a multimeter. Connect the black wire to the “-COM” terminal of the multimeter and the red wire to the “+V.Ω.mA” terminal. Turn the multimeter to the 2 Volt DC setting.

7. Before making a measurement, **predict which metal will serve as the anode (oxidation) and which will serve as the cathode (reduction)**—keep in mind the reduction table on page 3 of this lab.

8. Connect the black (negative) electrode to the anode metal strip, and the red (positive) electrode to the cathode metal strip. If you get a negative reading, you have connected the wires backwards and will need to reverse the order.

9. Allow the cell to stand until the voltage reading on the voltmeter has stabilized, then record the highest voltage.

10. Measure the potential of all six possible cell combinations using the method described above—see the table below for all possible combinations. **Use a fresh salt bridge for each cell.**

<table>
<thead>
<tr>
<th>Trial</th>
<th>Trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Cu/ Zn</td>
</tr>
<tr>
<td>2</td>
<td>Cu/ Pb</td>
</tr>
<tr>
<td>3</td>
<td>Cu/ Ni</td>
</tr>
<tr>
<td>4</td>
<td>Zn/ Pb</td>
</tr>
<tr>
<td>5</td>
<td>Zn/ Ni</td>
</tr>
<tr>
<td>6</td>
<td>Pb/ Ni</td>
</tr>
</tbody>
</table>

11. Complete all the questions for each cell as described in the Data Tables and Calculations section of this lab guide.

12. When you have finished, dispose of the metal ion solutions and the filter paper strips in the waste containers provided. Clean the metal strips and return them to the lab cart.

**Part B. Effect of Concentration on Cell Potential**

1. Prepare a copper/zinc voltaic cell as in Part A, using 0.10 M ZnSO₄ solution as before, but replace the 0.10 M CuSO₄ solution with 0.005 M CuSO₄ solution. Record the voltage of the cell. How does the increase in concentration of CuSO₄ affect the voltage of the cell?

2. Prepare a copper/zinc voltaic cell as in Part A, using 0.10 M CuSO₄ solution as before, but replace the 0.10 M ZnSO₄ solution with 0.005 M ZnSO₄ solution. Record the voltage of the cell. Does the increase in concentration of Zn²⁺ ion affect the voltage measured? Why?

**Part C Electrolytic Reduction of Cu²⁺ to Copper on the Surface of a Coin**

You will be using a power amplifier controlled by Pasco DataStudio in order to electroplate copper on to a coin.
1. If it is not already on and connected, connect a ScienceWorkshop 750 interface to the computer, turn on the interface, and turn on the computer.

2. Connect the DIN plug of the Power Amplifier to Analog Channel A on the interface. DO NOT turn on the Power Amplifier yet.

3. In the Chem 1b folder on the desktop, open the file titled “Electroplating”.
   • The DataStudio file has a Graph of current versus time, a Digits display of current, and a Signal Generator window.
   • Data recording is set at one measurement per second (1 Hz) and will stop automatically at 600 seconds (10 minutes).
   • The Signal Generator is set to output 5.00 V DC (direct current). The ‘ON/OFF button is set to ‘Auto’ so the Signal Generator will start automatically when you start to collect data and stop automatically when you stop collecting data.

4. Place 100 mL of 1.0 M copper (II) sulfate solution in a 250 mL beaker such that you will be able to completely submerge a nickel or other coin in the solution when the coin is placed on end.

5. Determine the mass of the coin and record in your notebook.

6. Set up the electrolytic cell as in the figure to the right. Use a strip of copper metal as one electrode, and a coin as the other. Use clamps to suspend the copper strip and the coin on the support rods. Don’t submerge the coin in the solution until you are ready to begin. Keep the alligator clips out of the solution or they will be plated too!

7. Use an alligator clip adapter to attach one end of a red patch cord to one end of the copper strip. This will be the anode. Connect the other end of red patch cord to the positive (red) output terminal of the Power Amplifier.

8. Attach one end of black patch cord to the coin (the cathode). Connect the other end of the black patch cord to the negative (black) output terminal of the Power Amplifier.

9. When you are ready to begin, turn on the switch on the back of the Power Amplifier.

10. Begin recording data by clicking ‘Start’ in DataStudio.
   • The Power Amplifier output will automatically start when data recording begins.
• If the voltage is too high you will see bubbles at each electrode indicating the electrolysis of water.

11. Data recording will continue for ten minutes and then stop automatically.

12. When data recording is complete, turn off the switch on the back of the Power Amplifier.

13. Remove the coin from the solution, pat it dry and record the mass in your notebook.

14. Determine the mass of copper that was electroplated on to your coin.

Analyzing the Data
1. Use the built-in analysis tools in the Graph to find the area under the curve of current versus time by clicking the ‘Statistics menu’ button (Σ) and then select ‘Area’ from the menu.
2. Record the value for the area under the current versus time curve. This is the number of coulombs (ampere • seconds) of electricity.

4. Using the coulombs found, calculate the expected mass of copper produced—see pages 894-895 for an explanation and an example. The electrolytic set-up we used is fairly crude, and there will be some inefficiency in the cell, so don’t expect perfect agreement. Recall that the reaction is

\[ \text{Cu}^{2+} + 2e^- \rightarrow \text{Cu} \]

5. Determine the percent error between the calculated mass of Cu expected to be electroplated and the actual mass of Cu electroplated.

Part D Electrolysis of Solutions

The purpose of this experiment is to identify the products obtained in the electrolysis of aqueous potassium iodide, copper(II) bromide, and sodium chloride solutions. The electrolysis reactions will be carried out in an electrolytic cell consisting of a Petri dish, a 9-V battery, and carbon (pencil lead) electrodes (Figure 1).

1. Place the partitioned Petri dish on a sheet of white paper. Observe that the compartments or segments of the Petri dish are labeled 1, 2, and 3.

2. Carefully pour about 8 mL of 0.5 M potassium iodide solution into the first compartment of the Petri dish until the compartment is one-third to one-half full.

3. Add 3 drops of phenolphthalein solution and stir to mix.

4. Connect the battery cap to the 9-V battery. Carefully attach a “pencil lead” electrode to each alligator clip lead. Caution: Do not allow the electrodes to touch each other.
5. Hold the red (+) lead from the 9-V battery in one hand and the black (–) lead in the other hand. Keeping the electrodes as far apart as possible, dip the pencil lead electrodes into the potassium iodide solution.

6. Let the electric current run for 1–2 minutes while observing any changes in the potassium iodide solution. Record all observations. See the Data Tables and Calculations for guidance. 
*Note: If you do not see any changes, you may need to substitute the 9V battery with the PASCO power supply set the 5.00 V DC. Consult your instructor.*

7. Remove the pencil lead electrodes from the electrolysis solution. Carefully rinse the electrodes with distilled water from a wash bottle and gently pat dry on a paper towel.

8. Add two drops of starch solution to the potassium iodide solution after electrolysis and record observations in the data table.

9. Carefully pour about 8 mL of 0.5 M sodium chloride solution into the second compartment of the Petri dish. Add three drops of phenolphthalein indicator solution and stir to mix.

10. Repeat steps 5–7 for the electrolysis of sodium chloride solution. Record observations as outlines in the Data Table and Calculations section.

11. Carefully pour about 8 mL of 0.2 M copper(II) bromide solution into the third compartment of the Petri dish.

12. Repeat steps 5–7 for the electrolysis of copper(II) bromide solution. Record observations as outlines in the Data Table and Calculations section.

13. Remove the pencil lead electrodes from the alligator clips and disconnect the battery cap from the battery.

14. The electrolysis products may include dilute halogen solutions (chlorine, bromine, and iodine). Working in the hood, carefully pour the contents of the Petri dish into a waste beaker containing sodium thiosulfate solution. Sodium thiosulfate will reduce the halogen waste products. Allow the beaker to stand in the hood overnight.
Data Table and Calculations

Part A — Construction of Voltaic Cells and Measurement of Cell Potentials

<table>
<thead>
<tr>
<th>Trial</th>
<th>Oxidation half-reaction</th>
<th>Reduction half-reaction</th>
<th>Balanced Net reaction</th>
<th>Metal at the anode</th>
<th>Metal at the cathode</th>
<th>Experimental cell potential</th>
<th>Calculated standard cell potential</th>
<th>% difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Cu/ Zn</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Cu/ Pb</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Cu/Ni</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Zn/Pb</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Zn/Ni</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Pb/Ni</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Prepare and complete the following table for each of your six voltaic cells

Example for Trial 1:

Copper/Zinc cell

Oxidation half-reaction

Reduction half-reaction

Balanced Net reaction

Metal at the anode

Metal at the cathode

Experimental cell potential

Calculated standard cell potential

% difference

Part B Effect of Concentration on Cell Potential

For the 0.10 M ZnSO\textsubscript{4}/0.005 M CuSO\textsubscript{4} cell

Anode

Cathode

Experimental cell potential

Calculated cell potential using the Nerst equation
For the 0.20 M ZnSO₄ / 0.005 M CuSO₄ cell

Anode

Cathode

Experimental cell potential

Calculated cell potential using the Nerst equation

Part C Electrolytic Reduction of Cu²⁺ to Copper on the Surface of a Coin

See the Lab Procedure section to determine the data and calculations needed

Part D Electrolysis of Solutions

For the KI solution,

The following oxidation and reduction half-reactions are possible for the electrolysis of potassium iodide solution. The solution contains water molecules, potassium ions (K⁺), and iodide ions (I⁻).

\[ 2\text{H}_2\text{O}(l) \rightarrow \text{O}_2(g) + 4\text{H}^+(aq) + 4e^- \quad 2\text{H}_2\text{O}(l) + 2e^- \rightarrow \text{H}_2(g) + 2\text{OH}^-(aq) \]

\[ 2\text{I}^-(aq) \rightarrow \text{I}_2(s) + 2e^- \quad \text{K}^+(aq) + e^- \rightarrow \text{K}(s) \]

1. What product was formed at the anode in the electrolysis of potassium iodide solution? Explain citing specific evidence from your observations.

2. What product was formed at the cathode in the electrolysis of potassium iodide solution? Explain based on your observations.

3. Write the balance half-reaction that occurred at the anode.

4. Write the balanced half-reaction that occurred at the cathode.

5. Write the balanced chemical equation for the overall redox reaction in the electrolysis of aqueous potassium iodide. Hint: Remember to balance the electrons!

In addition, answer the above five questions for the electrolysis of sodium chloride and electrolysis of copper (II) bromide trials.
Pre-Lab Assignment

1. Given the reaction: $\text{Al}(s) + \text{Cu}^{2+}(aq) \rightarrow \text{Al}^{3+}(aq) + \text{Cu}(s)$
   a. What is being oxidized?
   b. What is being reduced?
   c. Write the half-cell reaction for the oxidation (include electrons).
   d. What is the standard voltage for this half-cell reaction?
   e. Write the half-cell reaction for the reduction (include electrons).
   f. What is the standard voltage for this half-cell reaction?
   g. Write the balanced equation for the cell reaction.
   h. What is the standard cell voltage for the reaction?
   i. Is this reaction spontaneous under standard conditions?

2. The following data were measured using a nickel electrode as the standard:

   $\text{Pt}^{2+}(aq) + 2e^- \rightarrow \text{Pt}(s)$  +1.20 V
   $\text{Ni}^{2+}(aq) + 2e^- \rightarrow \text{Ni}(s)$  -0.28 V
   $\text{Fe}^{2+}(aq) + 2e^- \rightarrow \text{Fe}(s)$  -0.44 V
   $\text{Al}^{3+}(aq) + 3e^- \rightarrow \text{Al}(s)$  -1.66 V

   A. Which ion is most easily reduced?
   B. Which metal is most easily oxidized?
   C. The platinum and iron electrodes are connected in a battery.
      1) Which is the anode? cathode?
         Anode:_________________________  Cathode:_________________________
      2) Which is oxidized? reduced?
         Oxidized:_________________________  Reduced:_________________________
      3) What will the battery voltage be?
      4) Write a balanced net ionic equation for the reaction that takes place.
      5) What happens to the mass of the anode electrode? Does it increase or decrease?
      6) What happens to the mass of the cathode electrode? Does it increase or decrease?
3. Write the equation for the oxidation of water

4. Write the equation for the reduction of water.

5. Calculate the mass of zinc that will be deposited if a current of 0.40 Amps is applied for 25 minutes to a ZnSO₄ solution.
Post-Lab Questions

1. Standard voltaic cells contain 1.0 M $\text{Metal}^{2+}$ solutions, but we used 0.10 M $\text{Metal}^{2+}$ solutions in Part A of the lab. Use the Nernst equation to prove that the cell potentials we measured in this lab, $E_{\text{cell}}$, are equal to the standard cell potential, $E^°_{\text{cell}}$. Show your work.

2. Using Figure 1 on page 2 as your guide, complete the diagram below for cells 2 and 6 (see table on pg 11). You instructor will guide you through an example of this diagram so that you understand how to complete each “box”.

3. Rank the four metals used in Part A (Cu, Zn, Pb, and Ni) in terms of ease of oxidation from easiest to hardest.

4. Given three reasons why your measured voltages may differ from the theoretical voltage in Part A.
5. From the data below, determine what reaction will happen at the anode and what reaction will happen at the cathode for a 1.0 M CdBr₂ solution. In addition, determine the minimum voltage required for the onset of the electrolysis reaction.

\[
\begin{align*}
O_2(g) + 4H^+(aq) \,(10^{-7} \text{ M}) + 4e^- & \rightarrow 2H_2O(l) \quad E^0 = 0.816 \text{ v} \\
2H_2O(l) + 2e^- & \rightarrow H_2(g) + 2OH^- (aq) \,(10^{-7} \text{ M}) \quad E^0 = -0.414 \text{ v} \\
Br_2(s) + 2e^- & \rightarrow 2Br^- (aq) \quad E^0 = 1.09 \text{ v} \\
Cd^{2+}(aq) + 2e^- & \rightarrow Cd(s) \quad E^0 = -0.403 \text{ v}
\end{align*}
\]

6. You only measured the mass gained by the coin in Part C. What do you think was the change in the mass of the copper electrode? Did it increase or decrease?

7. A voltaic cell is constructed with a Cu/Cu^{2+} half-cell and a Ag/Ag^{+} half-cell. If the observed potential of the cell, \(E_{\text{cell}}\), is 0.40 V, calculate the ratio of Cu^{2+} to Ag^{+} present using the Nernst Equation.