Determination of an Unknown Chloride

The determination of a soluble chloride salt concentration is a classic titrametric analysis. A titration involves delivering a measured amount of a solution whose concentration is known accurately (the titrant) into a solution whose concentration is not known (the titrate). The purpose of the titration is to determine the number of moles of titrate present. When the reaction is complete, some physical change is observed, indicating the endpoint of the titration. The endpoint of a titration occurs when stoichiometric ratios of reactants are present and must be determined accurately.

In the titration in this lab, a dilute solution of silver nitrate with a known concentration acts as the titrant. It is added to a salt solution with an unknown amount of chloride, i.e. the titrate. Silver chloride, a white insoluble solid will precipitate from the solution. In order to detect when all the AgCl has been precipitated, another reagent is used as an indicator. The indicator in this lab, potassium chromate, is yellow and reacts with silver ions to form a bright orange silver chromate precipitate. This solid is slightly more soluble than the silver chloride so it does not form until essentially all the chloride has precipitated from the solution.

\[
\text{Ag}^+(aq) + \text{Cl}^-(aq) \rightleftharpoons \text{AgCl(s) at low silver ion concentration}
\]

\[
2 \text{Ag}^+(aq) + \text{CrO}_4^{2-}(aq) \rightleftharpoons \text{Ag}_2\text{CrO}_4(s) \text{ at higher silver ion concentration}
\]

All standard solutions must first be standardized using a primary standard because of potential evaporation. A primary standard is a solid that is stable and does not pick up water. The primary standard in this experiment is purified sodium chloride.

In this lab you will perform six titrations. In the first three titrations you will use a known amount of a pure NaCl sample to determine the exact concentration of an approximate 0.05 M silver nitrate solution. The endpoint is the first permanent orange-red color of Ag₂CrO₄. From this information one can determine the concentration of the AgNO₃. The last three titrations will allow you to find the percentage of chloride in your salt when used in conjunction with the average silver nitrate concentration.

Note: Silver is a heavy metal toxin and should never be flushed down the drain. Dispose of all silver waste (silver nitrate and silver chloride) in the waste bottles provided.

PROCEDURE:
Part A: Standardizing the Silver Nitrate Solution

1. Clean a 50 mL buret with soap and water, then rinse well with water.

2. Fill your buret with silver nitrate from an amber bottle. To prevent contamination, never add anything to the amber bottle. Fill the buret to the 0.00 mL mark with the AgNO₃ solution. Drain 5 mL from the buret into a beaker (to remove air bubbles) and fill to 0.00 mL again. Note: AgNO₃ is the only solution that will be placed in your buret! Also, do not dispose of AgNO₃ in the sink – place this heavy metal in a waste container.

3. Use an analytical balance to weigh three 0.1000 - 0.1200 gram samples of purified NaCl. Record exact mass.

4. Add about 50 mL of distilled water to each sample in a 125 mL Erlenmeyer flask (or larger) to dissolve the NaCl sample. Add about three drops of indicator (K₂CrO₄).
5. Titrate with 0.05 M AgNO₃ solution as you continually swirl the flask to a lovely peach end point. As you add the silver nitrate solution initially in short bursts you will see the orange-red color form and disappear as the solution is swirled. As you approach the end point (which should be between 20-40 mL) the color should begin to persist. At this point you should be adding the solution dropwise. Read the buret to the nearest 0.01 mL. Stop when the sample has a permanent faint peach color.

6. Repeat the titration with the second and third samples.

**Part B: Determination of Percent Chloride**

1. Obtain an unknown chloride salt and record the ID number in your lab notebook. Use an analytical balance to weigh **three** 0.1000 - 0.1200 gram samples.

2. To each sample add 50 mL of distilled water and 3 drops of K₂CrO₄ indicator solution in an Erlenmeyer flask. Titrate each sample with the standardized silver nitrate solution as in part A.

3. When done, place excess AgNO₃ in the waste container and rinse the buret with water before leaving the lab.

**CALCULATIONS:**

For Part A, calculate the molarity of the silver nitrate solution ([AgNO₃]) for each titration. Calculate the Parts Per Thousand (PPT) for [AgNO₃] using the "Parts Per Thousand" handout in the “Lab Notes” of the Companion. If your PPT is greater than 30 for the three trials, consider omitting a deviant molarity value to improve your PPT.

For Part B, calculate the percent chloride. *(Note: Use the average molarity of AgNO₃ as determined in part A.)* Average your three percent chloride values and find the PPT for the %Cl values. As in Part A, if one trial is quite different from the other two, report data from all three trials, but only average two trials.

**POSTLAB QUESTIONS:**

1. a) What is the largest source of error in this experiment? Why?  b) Describe a scenario whereby a percent chloride determination could be of use in an industrial setting. Explain.

2. How would the following hypothetical errors affect the calculated % chloride (increase, decrease or no change)? Explain.
   a. The pure sodium chloride was left open in the scale room and absorbed moisture.
   b. The calculated molarity of the silver nitrate solution was 5% too high.
   c. Two mL of AgNO₃ are added beyond the chromate end in titrating the unknown chloride.

3. 35.46 mL of a silver nitrate solution was used to reach the chromate end point with a 50 mL solution containing 0.1165 g of pure NaCl. What is the molarity of the AgNO₃ solution?

4. How many mL of the silver nitrate solution used in question 3 will react with 0.2595 g of BaCl₂ dissolved in 50 mL of water?

5. A solid chloride sample weighing 0.09969 g required 18.25 mL of 0.05205 M AgNO₃ to reach the chromate end point. What is the % chloride in this sample?