Experiment 2: The Qual Scheme Cation Group I

Note About Notation:

When describing chemical reactions by writing chemical formulas and chemical equations, it is important to balance the equations

Therefore in Proc 1 we try to keep the excess chloride ion concentration low. In Proc 2 we take advantage of the relative large solubility of PbCl₂ and its favorable temperature coefficient to separate PbCl₂ from AgCl and Hg₂Cl₂ which we have no chance of dissolving in mere hot water. In Proc 3 final testing for lead involves two separate reactions which each form a precipitate.

$$Pb^{2+} + 0.2M K_2CrO_4 \rightarrow \underline{PbCrO}_4$$
 yellow

$$Pb^{2+} + 3M H_2SO_4 \rightarrow \underline{PbSO}_4$$
 white

Since K_2CrO_4 solution is vivid yellow the color of the test solution will be yellow even without the presence of lead: there must be a precipitate for a positive lead test. Note here, as with most of the qual

 $Ag(NH_3)_2^+$

Procedure 2

Put 20 drops of water in with the precipitate from Proc 1 and place the tube in your boiling water bath, stirring for 1 min. The next operations should be carried out quickly. Get a balance tube ready and spot an available centrifuge before taking the sample out of hot water. Then quickly spin about 30 sec. and decant quickly to minimize cooling. The decantate will contain most of the PbC1₂ if present. Test it according to Proc 3. Wash the remaining precipitate with 10 drops of hot water (in the same manner) but discard the wash. Save the precipitate for analysis according to Proc 4.

Procedure 3

Divide the decantate from Proc 2 into two equal parts.

To one part add 2 drops of $0.2M \text{ K}_2\text{CrO}_4$. If Pb^{2+} is present a yellow precipitate is produced. To the other part add 2 drops of $3M \text{ H}_2\text{SO}_4$, a white precipitate will form if Pb^{2+} is present in solution. The chromate test is more sensitive, there must be precipitate, significant cloudiness at the least.

Procedure 4

Add 5 drops of 15M NH_3 (hood) to the washed white precipitate from Proc 2. Immediate conversion of the precipitate to dark grey or black indicates the presence of mercurous ion, Hg_2^{2+} . If at this stage some white cloudiness persists, it usually is $PbC1_2$, which was not washed out in Proc 2 and will require a little extra centrifugation. Centrifuge and separate the clear decantate.

Take the decantate to the hood and add 16M HNO₃ dropwise (a very vigorous reaction) until a white precipitate appears or until the solution is acid to litmus. The white precipitate is AgC1, appearing as the ammonia complex is decomposed by acid, and proves the presence of Ag⁺.

Endnote:

When an aqueous solution is created which contains ammonia $(NH_{3\,(aq)})$ molecules, ammonium cations $(NH_{4\,(aq)}^+)$, and hydroxide anions $(OH_{(aq)}^-)$, the following equilibrium is established.

$$NH_{3 (aq)} + H_{2}O_{(1)} \hat{i} NH_{4 (aq)}^{+} + OH_{(aq)}^{-}$$

This equilibrium is strongly shifted to the left. Thus, regardless of how the solution was prepared, ammonia is by far the major component (other than water). Hence the solution should be labeled as aqueous ammonia, rather than as aqueous ammonium hydroxide. You will find that under the hood, it is erroneously labeled as 15M ammonium hydroxide. Don't be confused if the lab manual calls for 15M ammonia.