E-Learning Material

on

Engineering Chemistry Practical

of

1st/2nd semester of all Engineering Branches





State Council for Technical Education and Vocational Training, Odisha Bhubaneswar-751012 Odisha

E-Learning Material

Engineering Chemistry Practical

of 1st/2nd semester of all Engineering Branches of Diploma courses of SCTE&VT, Odisha

Contents Written by

Sri. Papun Mirddha, Lecturer in Chemistry, GP, Boudh.

Reviewed and validated by

Sri. Prabodh Kumar Satapathy, Lecturer in Chemistry, G.P. Bhubaneswar

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| 2 | Preparation and Study of Properties of Amonia Gas | | | |
| 3 | Crystallization of Copper Sulphate from Carbonate Powder | | | |
| 4 | SIMPLE ACID BASE TITRATION (i) ACIDIMETRY (ii) ALKALIMETRY | | | |
| 5 | TEST FOR ACID RADICALS (KNOWN) (i) CARBONATE (ii) SULPHIDE (iii) CHLORIDE (iv) NITRATE (v) SULPHATE | | | |
| 6 | TEST FOR BASIC RADICALS (KNOWN) (i) AMMONIUM (ii) ZINC (iii) MAGNESIUM (iv) ALUMINIUM (v) CALCIUM (vi) SODIUM (vi) POTASSIUM | | | |
| 7 | TEST FOR UNKNOWN ACID RADICALS. | | | |
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EXPERIMENT NO-01

AIM OF THE EXPERIMENT:

Preparation and study of physical and chemical properties of carbon dioxide gas.

APPARATUS REQUIRED:

1.Woulf'sbottle.

- 4. Rubbercork.
- 2.Thistlefunnel. 5. Gas jar withlid.
- 3. Deliverytube.

6. Few testtubes.

CHEMICALS REQUIRED:

- 1.Marblechips (CaCO₃). 4.Magnesiumribbon.
- 2.Dil.Hydrochloricacid (HCl). 5.Limewater.

3.Litmuspaper. 6.Phenolphthaleinsolution.

THEORY:

In laboratory carbon dioxide gas is prepared by the action of dilute hydrochloric acid (HCl) upon marble chips $(CaCO_3)$ in a woulf's bottle. It is collected by upward displacement of air. Carbon dioxide is heaviour in nature.

CHEMICAL EQUATIONS:

 $CaCO_3 + 2HCI \rightarrow CaCl_2 + H_2O + CO_2$

LABORATORY DIAGRAM:

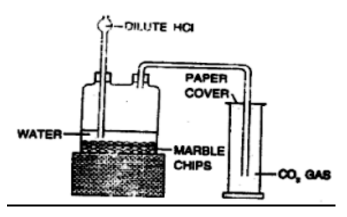


Fig 1.1 Laboratory Preparation of carbon dioxide gas.

PROCEDURE:

1. Take a woulf's bottle fitted with rubber cork, thistle funnel and delivery tube. Examine thatitis perfectly airtight. In case of air leakage, use melted paraffin wax or grease.

2. Introduce few small marble chips into the woulf's bottle by opening one of itsmouths.

3. Nowpoursomewaterintothewoulf'sbottlethroughthethistlefunnelsoastocover the marble chips.

4. Insert the thistle funnel more into the woulf's bottle such that its extremeendremainsinside the water.

5. Now add little quantity of the dil. Hydrochloric acid through the thistle funnel. Do not add excess amount of acid at a time to exhaust the marble chips before the experiment iscompleted.

6. Then collect the carbon dioxide gas in the gas jar by upward displacement of air. Test the collected gas in the jar by showing a burning splinter at the mouth of gasjar.

7. Study the properties of carbon dioxide gas by collecting the gas in different test tubes.

OBSERVATION:

PHYSICAL PROPERTIES

| SL. | EXPERIMENT | OBSERVATION | INFERENCE |
|-----|---|-------------|-----------|
| 1. | Observe the colour ofthe gas | | |
| 2. | Observe the odour of the gas | | |
| | Enter a glowingsplinter into a test tube full of CO_2 gas. | | |
| | Invert the test tube full of CO ₂ gas over another empty test tube containing air. Then add little lime to the testtube containing air initially. | | |

| Collect the gas in a test tube half- filled with water. Shake the test tube vigorously by putting the thumb at its mouth and remove the thumb and observe the level/volume of waterin the test tube. | |
|---|--|
| | |

CHEMICAL PROPERTIES

| SL NO | EXPERIMENT | OBSERVATION | INFERENCE |
|----------|--|-------------|-----------|
| 1 | A piece of moist blue litmus paper is shown to the gas. | | |
| 2. | Pass the CO ₂ gas through 2-3 ml of dilute solutions of sodium hydroxide (NaOH) containing one dropof phenolphthaleinsolution. | | |
| 3. | a) Pass the gas throughlime water.b) Pass the gas in excess.c) Boil thesolution. | | |
| 4. | Introduce a burning magnesium ribbon into a test tube /gas jar containingcarbondioxide gas. | | |

SAFTY AND PRECAUTION:

- 1. The fittings should beairtight.
- 2. The end of thistle funnel must be remaining deep inside thesolution.
- 3. The shorter end of the delivery tube should remain above the surface of the solutionin

the woulf's bottle.

- 4. The longer end of the delivery tube must reach the bottom of the gasjar.
- 5. Addition of excess of dil. hydrochloric acid should beavoided.
- 6. The gas should be collected after removing air from theapparatus.

ASSIGNMENT QUESTIONS:

- (1) What are the apparatus required for this experiment?
- (2) Write the chemical formula of marble chip.
- (3) Can we use CaCO₃ powder for preparation of CO₂ gas?
- (4) Why marble chips are used instead of CaCO₃ powder?
- (5) Write the chemicals used for preparation of CO_2 gas.
- (6) How can you prepare dilute HCI?
- (7) Hydrochloric acid is a strong or weak acid? Give reason.
- (8) Explain the acidic nature of CO₂ gas?
- (9) What happens when CO₂ gas is passed through alkaline phenolphthalein solution?
- 10. What happens when moist blue and red litmus papers are shown to CO₂ gas? How CO₂ gas is collected?
- 11.What are the apparatus required for preparation of CO₂ gas?

12. How can you prepare CO₂ gas in laboratory?

13. Why moist blue litmus paper turns red on exposure to CO₂ gas?

14.Write two methods of preparation of CO₂ gas.

- 15.What happens when a burning match stick is introduced into a jar containing CO₂ gas?
- 16 .What happens when methyl orange indicator is added to aqueous solution of CO₂ gas?

17. What happens when CO_2 gas is passed through lime water first in less amount and then in excess?

18.Write the reactions involved between CO₂ gas and lime water.

19.What is the formula of lime water?

20. Why lime water turns milky when less amount of CO_2 gas is passed through it? Give Equation.

21. Why milky colour disappears on passage of excess CO_2 gas through lime water? Give Equation.

22. What happens when colourless $Ca(HCO_3)_2$ solution will be warmed strongly? Give Equation.

23. What happens when a burning magnesium is introduced into gas jar containing CO_2 gas? Give Equation.

24.Write two uses of CO₂ gas.

25.What is dry ice?

26. How can you test that CO₂ gas is heavier than air?

27.Can sulphuric acid (H_2SO_4) be used in place of HCl for preparation of CO_2 gas?

28.What type of bonding is present in CO₂?

29.Why smaller pieces of marble chips are required?

30. Why the thistle funnel is inserted deep to the bottom of the Woulfe's bottle?

31. How can you test the solubility of CO₂ gas?

EXPERIMENT NO-02

AIM OF THE EXPERIMENT:

Preparation and study of properties of NH₃ gas.

APPARATUS REQUIRED:

Hard glass test tube
 Delivery tube
 Gas jar
 Card cover
 Glass jar containing CaO (quick lime)
 Bunsen burner
 Rubber cork
 Clamp stand

CHEMICALS REQUIRED:

1. Solid Ammonium Chloride, NH₄Cl

2. Anhydrous Calcium Hydroxide, Ca(OH)₂ or Calcium Oxide (CaO)

THEORY:

Ammonia gas is prepared in laboratory by heating the mixture of ammonium Chloride (NH₄Cl) & Calcium Hydroxide, Ca(OH)₂ paste in 1:3 ratio by weight. The reaction proceeds as:

$$2NH_4Cl + Ca(OH)_2 \rightarrow 2NH_3 \uparrow + CaCl_2 + 2H_2O$$

The gas so formed is collected in the gas jar by downward displacement of air because ammonia gas is lighter than air. The gas cannot be collected under water because it is highly soluble in water. Ammonia gas is dried by passing it through the glass bottle containing CaO.

DIAGRAM:

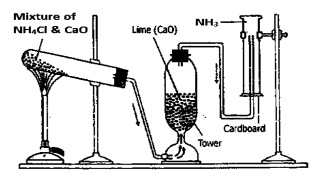


Fig 2.1 Laboratory preparation of Ammonia Gas.

PROCEDURE:

1. Take a hard glass test tube with rubber cork and deliverytube.

2.Mix 1:3 ratio of ammonium chloride and calcium hydroxide and place the mixture into the testtube.

3. Tilt the test tube at 30 degree angle and clamp it to thestand.

4.Attach the rubber cork to the test tube along with delivery tube connected at one end of the test tube and the other end connected to the glass jar containing CaO. The quicklime is present so as to absorb moisture present in the ammonia gas.

5. Make sure that all the connections are airtight to prevent leakage of ammonia gas.

6.Now carefully and gently heat the mixture in the testtube.

7. Then collect the ammonia gas in the gas jar by downward displacement ofair.

OBSERVATION:

PHYSICAL PROPERTIES

| SL. | EXPERIMENT | OBSERVATION | INFERENCE |
|-----|--|-------------|-----------|
| 1 | Color of the gas | | |
| 2 | Odour of the gas | | |
| | Collect the gas in a test tube half- filled with water. Shake the test tube vigorously by putting the thumb at its mouth and remove the thumb and observe the volume of water in the test tube. | | |

CHEMICAL PROPERTIES

| | SL. | EXPERIMENT | OBSERVATION | INFERENCE |
|---|-----|--|-------------|-----------|
| - | | Show a piece of moist red litmus paper to the gas. | | |

| 2 | Pass the gas into the test tube containing copper sulphate solution for short timeat first and then in excess. | |
|---|---|--|
| 3 | Pass the gas into the test tube containing ferric chloride solution. | |
| 4 | Pass the ammonia gas into the test tube containing Nessler's reagent. | |

CONCLUSION:

Ammonia gas is prepared at laboratory by using ammonium chloride (NH_4CI)&Calcium hydroxide ($Ca(OH)_2$). Ammonia gas is basic in nature. It is highly soluble in water.

SAFETY MEASURES:

1. The apparatus must be airtight.

2. The hard glass test tube should be fixed in inclined position towards its mouth in order to prevent crack in it.

3. Heat should be provided uniformly.

4. The gas jar should bedried.

ASSIGNMENT QUESTIONS:

- (1) Write the principle of preparation of ammonia gas in laboratory?
- (2) What is the principle of collection of ammonia gas?
- (3) Write the apparatus required for preparation of ammonia gas?
- (4) What are the chemicals required for ammonia gas?
- (5) While clamping the hard glass test tube, its mouth is present slightly downward. Why?
- (6) Write the physical properties of ammonia gas?
- (7) What is the odour of ammonia gas?
- (8) What happens when a glass rod dipped in conc HCl is shown to the ammonia gas?
- (9) What happens when a test tube filled with ammonia gas is inverted into a trough of water?
- (10) Explain a test to show that ammonia is lighter than air?.
- (11) Which compound is required to dry ammonia gas?
- (12) Why conc. Sulphuric acid is not used to dry ammonia gas?
- (13) Write two tests to show that ammonia is alkaline in nature?
- (14) What happens when ammonia gas is passed through Nessler's reagent? Give equation
- (15) What happens when ammonia gas is passed through ferric chloride solution? Give equation?
- (16) Ammonia gas has rotten egg / pungent/ irritating/ sweet odour.
- (17) Ammonia is sparingly / insoluble / highly soluble in water?
- (18) What happens when ammonia gas is passed through copper sulphate solution in small quantities?
- (19) What happens when ammonia gas is passed through copper sulphate solution in excess?
- (20) Ammonia gas turns blue litmus to red / red litmus to blue?
- (21) Write the uses of ammonia gas?
- (22) What happens when ammonia gas is passed through phenolphthalein solution?
- (23) What is the combustibility property of ammonia?
- (24) Can NaOH or KOH be used in place of CaO or Ca(OH)₂ for preparation of ammonia?
- (25) What is that compound (brown precipitate) which forms when ammonia gas is passed through nessler's reagent?

EXPERIMENT NO-03

AIM OF THE EXPERIMENT:

Crystallization of copper sulphate from copper carbonate.

APPARATUS REQUIRED:

1.Beaker

2.Funnel

3.Glass rod

4.Porcelain basin

5.Tripod stand

6.Wire gauze

7.Bunsen burner

8.Filter paper

9.Filter stand

CHEMICALS REQUIRED:

1.Copper carbonate($CuCO_3$)

2.Dilute sulphuric acid(H₂SO₄)

THEORY:

Copper carbonate reacts with dilute sulphuric acid to form soluble copper sulphate with evolution of carbon dioxide gas. The resulting solution is concentrated by evaporation till the point of crystallization is reached and then cooled to get crystals of copper sulphate pentahydrate (CuSO_{4.5}H₂O) called blue vitriol.

$$CuCO_3 + H_2SO_4 \rightarrow CuSO_4 + CO_2 \uparrow H_2O$$

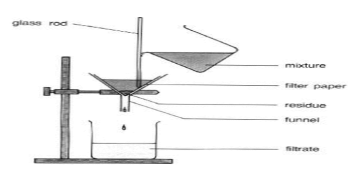


Fig 3.1 Filtration

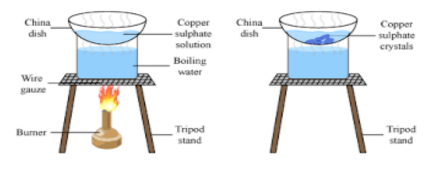


Fig 3.2 Crystallization of copper sulphate .

<u>Procedure</u> :- The preparation of Copper Sulphate crystals from Copper Carbonate involves following steps.

(A) Preparation of Saturated solution :-

- > Take about 20 ml of dilute Sulphuric acid in a beaker.
- Add supplied Copper Carbonate (CuCO₃) power gradually to this acid in small quantities with constant stirring.
- Continue addition of the of the powder till a small quantity of Copper Carbonate (CuCO₃) is left behind.
- > Heat the resulting solution slightly to expel the dissolved CO_2 gas.
- > Take the filter paper and four fold it.
- Prepare a cone of filter paper by taking three folds in one side and one fold in the other side.
- > Take a funnel and insert the cone made by the filter paper in to it.
- > Filter the solution from beaker to the porcelain basin .
- The solution must be transferred from the beaker to the filter paper cone slowly with the help of a glass rod.
- Wash the insoluble component present in the cone with the distilled water so as to make it free from soluble component.

(B) Concentrating the Filtrate :-

- > Evaporate the filtrate in the porcelain basin with constant stirring.
- Continue the process of evaporation till a drop of the liquid solution forms crystals on the tip of glass rod when blown on it. This state is termed as Crystallisation point.

(C) Crystallisation :

- Cool the hot solution (after reaching crystallisation point) slowly in air to start the process of crystallisation.
- If required, keep the hot porcelain basin containing the solution over a beaker full of water for quicker cooling.

- If required, keep the hot porcelain basin containing the solution over a beaker full of water for quicker cooling.
- (D) Drying and Crystals :-
- Decant off the saturated mother liquor present over the crystals after the crystallisation is over.
- Transfer the deep blue crystals present in the porcelain basin to a filter paper and spread to dry.

Precautions :-

- > Minimum amount of dilute sulphuric acid (H_2SO_4) should be used to prepare the solution
- > The solution should be slightly acidic , other wise the salt may get hrdrolysed
- > The solution should not be heated beyond crystallisation point.
- > The concentration of solution must be carried with constant stirring
- Crystals should never be dried by heating.

RESULT:

Color :

Shape:....

Yield: gm

ASSIGNMENT QUESTIONS:

- 1. Define crystallisation.
- 2. Define solubility.
- 3. Define filtration.
- 4. Why the solution is not concentrated or heated to dryness during crystallisation?
- 5. What is decantation?
- 6. Decantation and filtration which is a better process and why?
- 7. Why the saturated solution be cooled slowly?
- 8. What is blue vitriol?
- 9. What are hydrates?
- 10. What are anhydrous salts?
- 11. What is seeding ?
- 12. Aqueous solution of Copper Sulphate solution acidic or basic or neutral?
- 13. What is efflorescence?
- 14. What is the colour and structure of anhydrous Copper Sulphate ?
- 15. Why excess H_2SO_4 acid is not used for dissolving CuCO₃ powder ?
- 16. What is crystallisation point ?
- 17. Define mother liquor .
- 18 Write the reaction between anhydrous $CuCO_3$ and dilute H_2SO_4 .
- 19 Why the CuSO₄ solution be prepared slightly acidic ?
- 20 Can CuO be used instead of CuCO₃ powder for preparation of blue vitriol? If yes, then write the reaction.
- 21 Write two uses of $CuSO_4.5H_2O$?
- 22 Mention two other salts which can be prepared by this method .
- 23 Why a glass rod is used to transfer the solution from beaker to the filter paper cone in the funnel?
- 24 How can one check the crystallisation point ?
- 25 How a filter paper cone be prepared?

EXPERIMENT NO- 4(i)

AIM OF THE EXPERIMENT:

Acidimetry: To determine the strength of unknown acid using standard alkali..

APPARATUSREQUIRED:

1.Burette (50 ml.)

2.Burette Stand with clamp

3.Pipette (10 ml.)

4.Conical flask (100ml.)

5.Measuring flask (250ml.)

6.Glazed porcelain

CHEMICALS REQUIRED:

1.Acid Solution (Unknown Strength)

2.N/10 Alkali Solution (Known Strength)

3.Indicator: Methyl Orange

THEORY:

A known volume of standard alkali solution is titrated against the supplied acid solution of unknown strength in the presence of methyl orange indicator till the colour just changes from pale yellow/ straw yellow to light pink. The volume of the acid required for neutralization is determined. Knowing the volume of both the solutions and the strength of alkali, the strength of acid solution can be calculated by using the principle of equivalency.

$N_aV_a=N_bV_b$

Where,

N_a=Normality of the acid solution.

V_a=Volume of acid solution.

N_b=Normality of alkali solution.

V_b=Volume of alkali solution.

PROCEDURE:

1. Wash the burette, pipette and conical flask thrice with tap water and then rinse with distilled water.

2. Rinse the burette thrice with a few ml. of the given acid solution and reject the washings.

3.Fill the burette with the given acid solution to a convenient level without air bubbles. 4.Remove air bubble (if any) present in it.

5. Rinse the pipette with the alkali solution thrice and reject the washing.

6. Pipette out 10 ml. of alkali solution is into the conical flask.

7. After transferring the acid solution, touch the tip of the pipette to the inner side of the conical flask thrice. Wash the inner sides of the conical flask with a little distilled water.

8.Add one drop of methyl orange indicator to it. Keep the conical flask over a white glazed porcelain tile under the burette.

9.Note the initial burette reading avoiding parallax error.

10.Carry out titration by running alkali from the burette with constant stirring till the colour of the solution just changes from colorless to light pink.

11. Note the final burette reading. Repeat the titration till three concordant values are obtained.

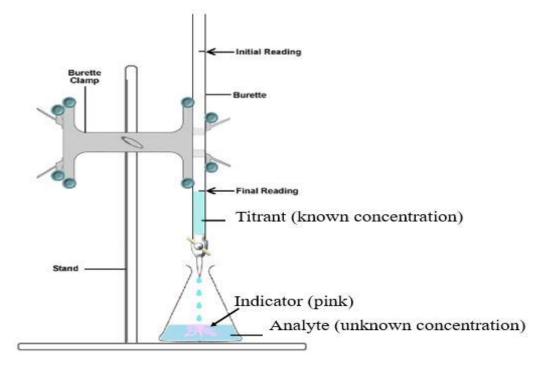


Fig 4.1 Titration

OBSERVATION:

| No of Observations | Volume of alkali (ml.) | Initial burette reading (ml.) | Final burette reading (ml.) | Difference (ml.) | Concordant |
|-----------------------|------------------------|-------------------------------|-----------------------------|---------------------|------------|
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |

CALCULATION:

We know that N_aV_a=N_bV_b

Where,

N_a=Normality of acid solution

V_a=Volume of acid solution consumed

N_b=Normality of alkali solution

V_b=Volume of alkali solution

 $N_a = N_b V_b / V_a = ----- N/10$

CONCLUSION:

From the above titration the strength of unknown acid solution is found to be ------.

PRECAUTIONS:

1.Care should be taken while handling the acid and base.

2. Always rinse the burette and the pipette with the solution which is to be taken in them.

3.Remove the air gap if any, from the burette before titration.

4.Never forget to remove the funnel from the burette before noting the initial reading of the burette and ensure that no drop is hanging from the nozzle.

5. Always read the lower meniscus for all transparent solutions and upper meniscus for the coloured solutions.

6.Never use burette and pipette with a broken nozzle.

7.Never suck a strong acid or an alkali with the pipette, use pipette bulb.

8. Always keep the lower end of the pipette dipped in the liquid while sucking the liquid

ASSIGNMENT QUESTIONS:

- (1) What do you mean by volumetric analysis?
- (2) What is titration?
- (3) Define titrant and titrate?
- (4) What are acidimetry and alkalimetry ?
- (5) Define a standard solution?
- (6) What do you mean by concentration / strength of solutions?
- (7) Define an indicator?
- (8) Name few indicators which are used during acid base titration?
- (9) Define gram equivalent weight and gram equivalent?
- (10) What is molality and normality?
- (11) 10 grams of caustic soda is how much gram equivalents?
- (12) What is the principle of titration?
- (13) Why rinsing is necessary?
- (14) Why conical flask is not rinsed with acid or alkali?
- (15) What is an anti parallax card?
- (16) Why one should not hold the pipette from its bulb?
- (17) Why the last drop of the solution be not blown from its bulb?
- (18) Why one or two drops of indicator should always be used?
- (19) Define the end point or neutralisation point?
- (20) How can you detect the neutralisation point?
- (21) Why mostly decinormal(N/10) solutions are used rather than normal(N) or centinormal(N/100) solutions?
- (22) Which solutions are required to rinse the burette and pipette?
- (23) To prepare 2 litres of solution, how much amount of sodium carbonate is required?
- (24) Why presence of air bubbles is not preferred in the burette while filling the acid solution?
- (25) Why stirring of conical flask and slow addition of acid is required?
- (26) What are the colours of methyl orange and phenolphthalein solutions in acidic, alkali and neutral mediums?
- (27) How the strength of the solution can be determined?

EXPERIMENT NO- 4(ii)

AIM OF THE EXPERIMENT:

Alkalimetry: To determine the strength of unknown alkali using standard acid..

APPARATUSREQUIRED:

1.Burette (50 ml.)

2.Burette Stand with clamp

3.Pipette (10 ml.)

4.Conical flask (100ml.)

5.Measuring flask (250ml.)

6.Glazed porcelain

CHEMICALS REQUIRED:

1. N/10 Acid Solution (known Strength)

2. Alkali Solution (unknown Strength)

3.Indicator: Methyl Orange

THEORY:

A known volume of standard alkali solution is titrated against the supplied acid solution of unknown strength in the presence of methyl orange indicator till the colour just changes from pale yellow/ straw yellow to light pink. The volume of the acid required for neutralization is determined. Knowing the volume of both the solutions and the strength of alkali, the strength of acid solution can be calculated by using the principle of equivalency.

$N_aV_a=N_bV_b$

Where,

N_a=Normality of the acid solution.

V_a=Volume of acid solution.

N_b=Normality of alkali solution.

V_b=Volume of alkali solution.

PROCEDURE:

1. Wash the burette, pipette and conical flask thrice with tap water and then rinse with distilled water.

2. Rinse the burette thrice with a few ml. of the given acid solution and reject the washings.

3. Fill the burette with the given acid solution to a convenient level without air bubbles. 4. Remove air bubble (if any) present in it.

5. Rinse the pipette with the alkali solution thrice and reject the washing.

6. Pipette out 10 ml. of alkali solution is into the conical flask.

7. After transferring the acid solution, touch the tip of the pipette to the inner side of the conical flask thrice. Wash the inner sides of the conical flask with a little distilled water.

8.Add one drop of methyl orange indicator to it. Keep the conical flask over a white glazed porcelain tile under the burette.

9.Note the initial burette reading avoiding parallax error.

10.Carry out titration by running alkali from the burette with constant stirring till the colour of the solution just changes from colorless to light pink.

11. Note the final burette reading. Repeat the titration till three concordant values are obtained.

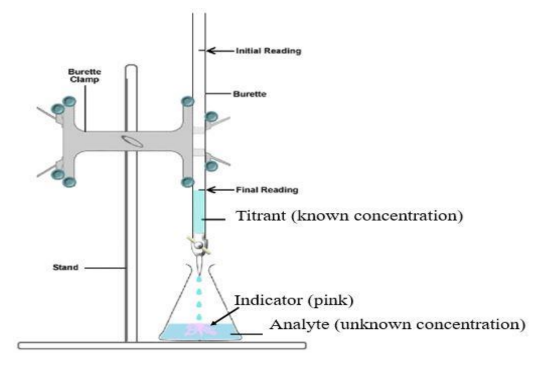


Fig 4.1 Titration

OBSERVATION:

| No of Observations | Volume of alkali (ml.) | Initial burette reading (ml.) | Final burette reading (ml.) | Difference (ml.) | Concordant |
|-----------------------|---------------------------|-------------------------------|-----------------------------|---------------------|------------|
| | | | | | |
| | | | | | |
| | | | | | |
| | | | | | |

CALCULATION:

We know that N_aV_a=N_bV_b

Where,

N_a=Normality of acid solution

V_a=Volume of acid solution consumed

N_b=Normality of alkali solution

V_b=Volume of alkali solution

 $N_a = N_b V_b / V_a = ----- N/10$

CONCLUSION:

From the above titration the strength of unknown acid solution is found to be ------.

PRECAUTIONS:

1.Care should be taken while handling the acid and base.

2. Always rinse the burette and the pipette with the solution which is to be taken in them.

3.Remove the air gap if any, from the burette before titration.

4.Never forget to remove the funnel from the burette before noting the initial reading of the burette and ensure that no drop is hanging from the nozzle.

5. Always read the lower meniscus for all transparent solutions and upper meniscus for the coloured solutions.

6.Never use burette and pipette with a broken nozzle.

7.Never suck a strong acid or an alkali with the pipette, use pipette bulb.

8. Always keep the lower end of the pipette dipped in the liquid while sucking the liquid

ASSIGNMENT QUESTIONS:

- (1) What do you mean by volumetric analysis?
- (2) What is titration?
- (3) Define titrant and titrate?
- (4) What are acidimetry and alkalimetry ?
- (5) Define a standard solution?
- (6) What do you mean by concentration / strength of solutions?
- (7) Define an indicator?
- (8) Name few indicators which are used during acid base titration?
- (9) Define gram equivalent weight and gram equivalent?
- (10) What is molality and normality?
- (11) 10 grams of caustic soda is how much gram equivalents?
- (12) What is the principle of titration?
- (13) Why rinsing is necessary?
- (14) Why conical flask is not rinsed with acid or alkali?
- (15) What is an anti parallax card?
- (16) Why one should not hold the pipette from its bulb?
- (17) Why the last drop of the solution be not blown from its bulb?
- (18) Why one or two drops of indicator should always be used?
- (19) Define the end point or neutralisation point?
- (20) How can you detect the neutralisation point?
- (21) Why mostly decinormal(N/10) solutions are used rather than normal(N) or centinormal(N/100) solutions?
- (22) Which solutions are required to rinse the burette and pipette?
- (23) To prepare 2 litres of solution, how much amount of sodium carbonate is required?
- (24) Why presence of air bubbles is not preferred in the burette while filling the acid solution?
- (25) Why stirring of conical flask and slow addition of acid is required?
- (26) What are the colours of methyl orange and phenolphthalein solutions in acidic, alkali and neutral mediums?
- (27) How the strength of the solution can be determined?

EXPERIMENT NO-5

AIM OF THE EXPERIMENT:

Test for carbonate, sulphide, chloride, nitrate, sulphate radicals (Known)

APPARATUS REQUIRED:

- 1. Test tube holder
- 2. Watch Glass
- 3. Test tubes
- 4. Bunsen Burner

CHEMICALS REQUIRED:

- 1.Given salts
- 2.Various Reagent
- 3. Litmus paper

THEORY AND PROCEDURE:

TEST FOR ACID RADICALS:

<u>Test for Carbonate $(CO_3^{2^-})$:</u>

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|--|-------------------------------------|
| 1.Take 2 ml of dil. HCl or dil. | 1. 1.Effervescences takes | 1.It may be CO_2 from CO_3^{2-} |
| H_2SO_4 , in a clean test tube. | place with the evolution of a | |
| Warm it and add a little of | colourless, odourless gas. | |
| the salt into it. | | |
| 2.Warm the above reaction mixture to get more gas and pass the gas slowly through limewater. | 2. First lime water turns milky and with excess of the gas milkiness disappears. | 2. CO_3^{2} is confirmed. |

Explanation for Carbonate Test:

 $Na_2CO_3 + 2HCI \rightarrow 2NaCI + H_2O + CO_2$

$$Na_2CO_3 + H_2SO_4 \rightarrow Na_2SO_4 + H_2O + CO_2$$

2. Milkyness is due to the formation of $CaCO_3$, and with excess of the gas milkyness disappears due to the formation of water soluble $Ca(HCO_3)_2$

 $\begin{array}{rcl} {\sf Ca}({\sf OH})_2 \,+\, {\sf CO}_2 \,\,\rightarrow & {\sf Ca}{\sf CO}_3 \downarrow \,\,+\,\,{\sf H}_2{\sf O} \\ & {\sf White\ ppt.} \end{array} \\ \\ {\sf Ca}{\sf CO}_3 \,\,+\,{\sf H}_2{\sf O} \,+\,{\sf CO}_2 \,\,\,\rightarrow \,\,{\sf Ca}({\sf HCO}_3)_2 \\ & {\sf Soluble} \end{array}$

Test for Sulphide (S^{2}) :

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|--|--|
| 1.Take 2 ml of dil. HCl or dil. H_2SO_4 , in a clean test tube. Warm it and add a little of the salt into it. | 1. Effervescence takes place with the evolution of a colourless gas with rotten egg smell. | 1. It may be H ₂ S from S ²⁻ |
| 2.Warm the above reaction mixture and show a filter paper soaked with lead acetate to the evolved gas. | 2. The filter paper turns black | 2. S ²⁻ is confirmed. |

Explanation for Sulphide Test:

1.

 $Na_2S + 2HCI \rightarrow 2NaCI + H_2S$

 $Na_2S + H_2SO_4 \rightarrow Na_2SO_4 + H_2S$

2. The black colour is due to the formation of PbS

Test for Chloride (Cl⁻):

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|--|---------------------------------------|
| 1.Take a pinch of salt in a clean and dry test tube and add 2 to 3 drops of conc. H_2SO_4 to it. | 1. Effervescence takes place with the evolution of a colourless gas which fumes in moist air. | 1. It may be HCI from CI ⁻ |
| 2.Warm the above reaction mixture and show a glass rod | Dense white fumes are produced and white solid deposited on the tip of the glass | NH₄CI |

| from dine | | | | |
|--------------------------------|--|--|--|--|
| n of | | | | |
| n dil | | | | |
| due mino | | | | |
| Explanation for Chloride Test: | | | | |
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| | | | | |
| ΝΟ | | | | |
| NO ₂ | | | | |
| NO ₂ | | | | |
| | | | | |

heat it.

| 2. Show a filter paper soaked in | 2. The paper turns black. | 2. May be NO_3^- |
|--|------------------------------------|--------------------------------------|
| freshly prepared FeSO ₄ | | |
| Solution to the above brown | | |
| gas. | | |
| 3.Brown Ring Test: Take 1-2 | | |
| ml. of the salt solution. Add | 3.A brown ring is formed at the | 3.The brown ring is due to the |
| equal volume ofconc. | junction of the two liquid layers. | formation of [Fe(NO)]SO ₄ |
| H_2SO_4 slowly into the test tube. | | |
| Cool the test tube perfectly | | NO ₃ -is confirmed |
| under tap. Then slowly add 2-3 | | |
| ml of freshly prepared FeSO ₄ | | |
| Solution through the sides of | | |
| test tube. | | |

Explanation for Nitrate Test:

| 1. | Cu + | 4HNO ₃ | $\stackrel{\Delta}{\rightarrow}$ | $Cu(NO_3)_2 + 2H_2O + 2NO_2 \uparrow$ |
|----|------|-------------------|----------------------------------|---------------------------------------|
| | | | | Brown gas |

2.
$$2NaNO_3 + H_2SO_4 \rightarrow Na_2SO_4 + 2HNO_3$$

3.

$$6FeSO_4 + 2HNO_3 + 3H_2SO_4 \rightarrow 3Fe_2(SO_4)_3 + 2NO + 4H_2O_4$$

$$FeSO_4 + NO + 5H_2O \rightarrow [Fe(NO)(H_2O)_5]SO_4$$

Brown Complex

<u>Test for Sulphate $(SO_4^{2^-})$:</u>

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|---------------------------------|-----------------------------|
| 1.Take about 1-2 ml of salt | 1. A white precipitate is | 1. SO_4^{2} is confirmed. |
| solution. Acidify with 1-2 ml of | obtained. | |
| dil. HCl add about 1 ml of BaCl ₂ | | |
| solution. | | |
| Add about 1 ml of conc. HCl to | | |
| the above solution and warm it. | | |
| | The precipitate is not soluble. | |

Explanation for Sulphate Test:

The white precipitate is due to the formation of BaSO₄ which is insoluble in conc. HCl.

 $\begin{array}{rl} Na_2SO_4 + \ BaCl_2 \ \rightarrow BaSO_4 \ \downarrow +2NaCl \\ & White \end{array}$

ASSIGNMENT QUESTIONS:

1.What is acid radicals?

2. How do u test for carbonate ?

3. What is the chemical equation involved in the test for sulphate ion ?

4. How do you test for sulphide ion ?

5. How can you test for nitrate ion ?

6. How does a salt become crystalline and amorphous?

7. How do you detect the presence of phosphate ion ?

8. Give the example of chemical reaction which involved for carbonate radicals ?

9. The acid radical carries ------ ion.

10.What is brown ring test ?

EXPERIMENT NO-6:

AIM OF THE EXPERIMENT:

Test for ammonium, zinc, magnesium, aluminium, calcium, sodium & potassium (Known)

APPARATUS REQUIRED:

- 1. Test tube holder
- 2. Watch Glass
- 3. Blow pipe
- 4. Nichrome wire
- 5. Blue glass
- 6.Charcoal cavity
- 7.Test tubes

CHEMICALS REQUIRED:

- 1. Given salts
- 2.Various Reagent
- 3. Litmus paper

THEORY & PROCEDURE:

A.DRY TEST FOR BASIC RADICALS:

I. Heating in a dry Test Tube.

| EXPERIMENT | OBSERVATION | INFERENCE | | |
|------------------------------------|--|---|--|--|
| Heat a small quantity of | a) Water particles condense at | a) Salt contains water of | | |
| supplied salt in a clean, dry test | the cooler part of the test-tube. | crystallisation. | | |
| tube first slowly and then | | | | |
| strongly for about 3 to 4 | b) A sublimate is formed. | b) Volatile Salts | | |
| minutes. | | | | |
| | White sublimate NH ₄ NO ₃ is | is May be NH_4^+ , Hg^{2+} or As^{3+} salts | | |
| | volatile but produces no white | | | |
| | sublimate. | | | |
| | c) The salt decrepitates. | | | |
| | (produces cracking sound) | c) Crystalline Salts. | | |
| | d) Deflagation takes place. | | | |
| | (Catches fire) | | | |
| | e) Infusible mass left. | d) Some nitrate or nitrite salts. | | |

| f) The salt changes colour: | |
|--|---|
| i) Yellow when hot and whi when cold. ii) Red to black when hot an brown when cold. | nd |
| g) The salt is fused on heating and solidified on cooling. | ng f) (i) May be Zinc (Zn ²⁺) salt |
| h) The salt is swelled up of | on ii) May be Fe^{2+} or Fe^{3+} |
| heating. A gas or vapour is evolved. | g) May be alkali or alkaline earth metal salts. |
| (i) A colourless, odourless gain (CO₂) which turns lime wat milky. (ii) A colourless gas (NH₃) with pungent odour which turns realitmus paper blue. | ter h) May be Al ³⁺ ith |
| | (i) May be carbonate salts. |
| | (ii) May be ammonium salts. |
| . Heating in a Charcoal Cavity. | l |

| . Heating in a Charcoar Cavity. | | | |
|--|---|--|--|
| EXPERIMENT | OBSERVATION | INFERENCE | |
| Make a small cavity on a charcoal block. Fill the with the supplied salt. Moisten the salt with a drop of water. Heat the salt strongly with the oxidising | (i) The salt decrepitates or produces cracking sound.(ii) The salt volatilises completely with white fumes. | (i) Crystalline salts. | |
| flame for about three minutes with help of a blow pipe. | (iii) The salt deflagrates (iv) The salt may or may not melt but finally leaves a white infusible incandescent (giving light) residue. (v) The salt melts and sinks into the charcoal cavity on heating and reappears on cooling. | (ii) May be ammonium, arsenic or mercury salts (Perform soda lime and bulb tube test). (iii) May be nitrate or nitrite (iv) Aluminium, zinc, magnesium, | |

| | (vi)The salt is coloured and leaves a coloured residue. | tin or alkaline earth metal salts. (Perform cobalt nitrate test). (v) May be alkali or alkaline on earth metal salts (Perform flame test). (vi) May be chromium, manganese, iron, cobalt, nickel or copper salts. (Perform borax bead test). |
|--|--|---|
|--|--|---|

III. Soda lime Test: (For Volatile Salts)

| Take a pinch of the salt in a a) | | |
|----------------------------------|---|--------------------|
| sodalime (NaOH + CaO) and pr | ammonia is evolved which produces a dense white fume with concentrated HCI. | a) NH₄⁺is present. |

IV. Cobalt Nitrate Test: (For Infusible Salts)

| EXPERIMENT | OBSERVATION | INFERENCE |
|----------------------------------|---------------------------|---------------------------|
| Heat a small quantity of the | a) Blue (infusible) mass. | a) May be aluminium salt. |
| salt in charcoal cavity in the | | |
| oxidizing flame with the help of | b) Green mass. | b) May be Zinc Salt. |
| a blow pipetill an infusible and | | |
| incandescent residue is left. | c) Pink mass | c) May be Magnesium Salt. |
| Moisten in the Residue with a | | d) May be Calaium Salt |
| drop of Cobalt Nitrate solution | d) Grey mass | d) May be Calcium Salt. |
| and heat in the oxidising flame. | | |
| Note the colour of the residue | | |

V. Flame Test: (For fusible Salts)

| OBSERVATION | | INFERENCE |
|--------------|---|--|
| Colour of | the flame | |
| In naked eye | Through Double | |
| | blue glass | |
| / | | |
| , | / | |
| , | , | |
| Green | iv) Bluish Green | (i) May be Sodium salt. |
| | , | |
| | | ii) May be Potassium Salt. |
| | | |
| | | iii) May be Calcium Salt. |
| | | iv) May be Barium Salt |
| | | iv) May be Barium Salt. |
| | | |
| | | |
| | Colour of In naked eye i) Golden Yellow ii) Violet iii) Brick Red iv) Yellowish Green | Colour of the flameIn naked eyeThrough Double blue glassi) Golden Yellowi) Colourlessii) Violeti) Colourlessiii) Brick Redii) Crimson red iii) Light Green |

VI. Charcoal Cavity Reduction Test: (For White salts changing Colour):

| EXPERIMENT | OBSERVATION | INFERENCE |
|------------------------------------|----------------------------------|------------------------|
| Prepare an intimate mixture of | i) White shinning malleable bead | a) May be lead salt. |
| salt charcoal powder and fusion | with lemon yellow incrustation | |
| mixture in 1: 3: 1 proportion. | and the bead marks paper.b) | |
| Take a little of this mixture in a | Red Scale without incrustation. | |
| charcoal cavity and heat it in | | b) May be Copper Salt. |
| reducing flame with the help of | | |
| a blow pipe. | | |

* B.WET TEST FOR BASIC RADICALS:

<u>Test For NH₄+:</u>

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|---|---|
| | 1. Ammonia gas is evolved producing white fumes with a glass rod dipped in conc. HCI. | 1. NH₄ ⁺ is confirmed. |
| 2. To 1-2 ml. of the salt solution, about 1ml dil. NaOH solution is added followed with a little Nessler's reagent. | 2. Brown ppt. is obtained. | 2. NH ₄ ⁺ is confirmed. [This is due to formation of I-Hg-O-Hg-NH ₂] |

Explanation for NH₄⁺Test:

- 1. $NH_4Cl + NaOH \rightarrow NH_3 \uparrow + NaCl + H_2O$
- 2. $NH_4Cl + 2K_2[HgI_4] + 4KOH \rightarrow I Hg O Hg NH_2 \downarrow +KCl + 7KI + 3H_2O$

Brown ppt

Test For Zn²⁺:

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|---------------------------------|---|
| 1. To about 1-2 ml of salt | 1. A white precipitate is | 1. May be Zn ²⁺ . |
| solution solid NH ₄ Cl is added till | obtained. | [It is due to the formation of ZnS] |
| saturation and dil. NH ₄ OH is | | |
| added till ammoniacal and H ₂ S | | |
| is passed through it. | | |
| 2. To about 1-2 ml. of the salt | | 2. Zn ²⁺ is confirmed. |
| solution, dil. NaOH solution is | 2. A white precipitate is first | [The white ppt. is due to the |
| added drop by drop and then in | obtained which is soluble in | formation of Zn(OH) ₂ which gets |
| excess. | excess of the reagent. | dissolved with excess of the |
| | | reagent due to the formation of |
| 3. To about 1-2 ml of salt | | Na_2ZnO_2 . |
| solution few drops of | | 3. Zn ²⁺ is confirmed. |
| $K_4[Fe(CN)_6]$ solution is added. | 3. A white precipitate is | [This is due to the formation of |
| | obtained. | Zinc ferrocyanide] |
| | | |
| | | |

Explanation for Zn²⁺Test:

1. $ZnSO_4 + H_2S \rightarrow ZnS \downarrow (White) + H_2SO_4$

2. $ZnSO_4 + 2NaOH \rightarrow Zn(OH)_2 \downarrow (White) + Na_2SO_4$

- 3. $Zn(OH)_2 + 2NaOH \rightarrow Na_2ZnO_2$ (Sodium Zincate) + $2H_2O$
- 4. $2\text{ZnSO}_4 + \text{K}_4[\text{Fe}(\text{CN})_6] \rightarrow \text{Zn}_2[\text{Fe}(\text{CN})_6] + 2\text{K}_2\text{SO}_4$

Zinc ferrocyanide (White)

Test for Mg²⁺

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|------------------------------------|--|
| 1. To about 1-2 ml of salt | 1. A white precipitate is | 1. May be Mg ²⁺ . |
| solution solid NH ₄ Cl is added till | obtained. | [This is due to the formation of |
| saturation and dil. NH ₄ OH is | | Mg(NH ₄)PO ₄]. |
| added till alkaline and disodium | | |
| hydrogen phosphate solution is | | |
| added to it. | | |
| 2. To about 1-2 ml. of the salt solution about 1 ml of dil. HCl is added followed with 2-3 drops of Magneson reagent and the solution is made strongly alkaline with NaOH solution. | 2. A blue precipitate is obtained. | Mg²⁺ is confirmed. [This is due to the formation of magnesium salt of magneson reagent.] |

Explanation for NH₄⁺Test:

- 3. $NH_4Cl + NaOH \rightarrow NH_3 \uparrow + NaCl + H_2O$
- 4. $NH_4Cl + 2K_2[HgI_4] + 4KOH \rightarrow I Hg O Hg NH_2 \downarrow +KCl + 7KI + 3H_2O$

Brown ppt

Test For Zn²⁺:

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|---------------------------------|---|
| 1. To about 1-2 ml of salt | 1. A white precipitate is | 1. May be Zn ²⁺ . |
| solution solid NH ₄ Cl is added till | obtained. | [It is due to the formation of ZnS] |
| saturation and dil. NH ₄ OH is | | |
| added till ammoniacal and H ₂ S | | |
| is passed through it. | | |
| 2. To about 1-2 ml. of the salt | | 2. Zn ²⁺ is confirmed. |
| solution, dil. NaOH solution is | 2. A white precipitate is first | [The white ppt. is due to the |
| added drop by drop and then in | obtained which is soluble in | formation of Zn(OH) ₂ which gets |
| excess. | excess of the reagent. | dissolved with excess of the |
| | | reagent due to the formation of |
| 3. To about 1-2 ml of salt | | Na_2ZnO_2 . |
| solution few drops of | | 3. Zn ²⁺ is confirmed. |
| $K_4[Fe(CN)_6]$ solution is added. | 3. A white precipitate is | [This is due to the formation of |
| | obtained. | Zinc ferrocyanide] |
| | | |
| | | |

Explanation for Zn²⁺Test:

5. $ZnSO_4 + H_2S \rightarrow ZnS \downarrow (White) + H_2SO_4$

6. $\operatorname{ZnSO}_4 + 2\operatorname{NaOH} \rightarrow \operatorname{Zn}(\operatorname{OH})_2 \downarrow (\operatorname{White}) + \operatorname{Na}_2\operatorname{SO}_4$

- 7. $Zn(OH)_2 + 2NaOH \rightarrow Na_2ZnO_2$ (Sodium Zincate) + $2H_2O$
- 8. $2\text{ZnSO}_4 + \text{K}_4[\text{Fe}(\text{CN})_6] \rightarrow \text{Zn}_2[\text{Fe}(\text{CN})_6] + 2\text{K}_2\text{SO}_4$

Zinc ferrocyanide (White)

Test for Mg²⁺

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|------------------------------------|--|
| 1. To about 1-2 ml of salt | 1. A white precipitate is | 1. May be Mg ²⁺ . |
| solution solid NH ₄ Cl is added till | obtained. | [This is due to the formation of |
| saturation and dil. NH ₄ OH is | | $Mg(NH_4)PO_4].$ |
| added till alkaline and disodium | | |
| hydrogen phosphate solution is | | |
| added to it. | | |
| 2. To about 1-2 ml. of the salt solution about 1 ml of dil. HCl is added followed with 2-3 drops of Magneson reagent and the solution is made strongly alkaline with NaOH solution. | 2. A blue precipitate is obtained. | Mg²⁺ is confirmed. [This is due to the formation of magnesium salt of magneson reagent.] |

Test For Na⁺:

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|-------------|---|
| 1. To about 1-2 ml of salt solution about 1 ml of potassium pyroantimonate solution is added and the inner side of the test tube is scratched with glass rod. | | 1. Na ⁺ is confirmed. [The ppt. is due to the formation of Na ₂ H ₂ Sb ₂ O ₇] |

Explanation for Na⁺Test:

Test For K⁺:

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|----------------------------|---|
| 1. To about 1-2 ml of salt | 1. A Yellow precipitate is | 1. K ⁺ is confirmed. |
| solution solid NaNO ₂ is added | obtained. | [This is due to the formation of |
| till saturation followed with few | | K ₃ [Co(NO ₂) ₆]]. |
| drops Co(NO ₃) ₂ solution. About | | |
| 1 ml of dil. CH ₃ COOH is added | | |
| to it and the solution is kept for | | |
| sometime. | | |

Explanation for K⁺Test:

1. KCl + NaNO₂ \rightarrow KNO₂ + NaCl

2. $Co(NO_3)_2 + 2NaNO_2 \rightarrow Co(NO_2)_2 + 2NaNO_3$

3. $Co(NO_2)_2 + 2KNO_2 + 2CH_3COOH \rightarrow Co(NO_2)_3 + 2CH_3COOK + NO \uparrow + H_2O$

4. $\operatorname{Co}(\operatorname{NO}_2)_3 + 3\operatorname{KNO}_2 \rightarrow \operatorname{K}_3[\operatorname{Co}(\operatorname{NO}_2)_6]$

ASSIGNMENT QUESTIONS:

1.What is dry test?

2. Which type of salt is subjected to sodalime test?

3. Which type of salt is subjected to flame test?

4. What type of wire can be used in the flame test?

5.What is wet test of salt

6.What are the group-I radicals and how they precipitated?

7.For what type of salts charcoal cavity reduction test is performed ?

8.For what type salt cobalt nitrate test is carried out?

9. What is the principle of charcoal cavity reduction test ?

10.Which dry test is applied for infusible salt to identify its basic radicals

EXPERIMENT NO-7

AIM OF THE EXPERIMENT:

Test for unknown acid radicals.

APPARATUS REQUIRED:

- 1. Test tube holder
- 2. Bunsen Burner
- 3. Spatula

CHEMICALS REQUIRED:

1.Given salts

2.Various Reagent

3. Litmus paper

THEORY & PROCEDURE:

PRELIMINARY TEST:

- 1.Salt No-----
- 2.Colour of the salt------
- 3.Structure-----
- 4. Solubility of the salt------

B. TEST FOR ACID RADICALS:

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|----------------------------------|--|
| I. Test with dil. HCI (For CO ₃ ²⁻ , S ²⁻ |) | |
| About 2ml of dil. HCl is taken in a | (a) Effervescence takes place | (a) It may be CO_2 from CO_3^{2-} |
| clean test tube. It is warmed and | with the evolution of a | |
| small amount of the supplied salt | colourless, odourless gas. | 2 |
| is added to it. | (b) Effervescence takes place | (b) May be H_2S from S^{2-} . |
| | with the evolution of a | |
| | colourless gas with the smell of | |
| | rotten egg. | |
| | (c) No effervescence and no | (c) $CO_3^{2^2}$, S ²⁻ are absent. |
| | gas is evolved. | |

| Confirmatory Test for CO ₃ ²⁻ | | | | | | | | |
|--|--|-------------------------------|--|--|--|--|--|--|
| The gas evolved from the reaction of dilute HCI and the salt is passed through lime water | Lime water is turned milky and with excess of gas disappeared. | $CO_3^{2^-}$ is confirmed. | | | | | | |
| Confirmatory Test for S ²⁻ | L | L | | | | | | |
| The gas is passed through lead acetate solution or a filter paper dipped in lead acetate solution is shown to the gas | It is turned into black. | S ²⁻ is confirmed. | | | | | | |

| EXPERIMENT | OBSERVATION | INFERENCE |
|--|---|--------------------------------------|
| II.(a) Test with conc. H ₂ SO ₄ For | | |
| A Pinch of salt is taken in a | (a) Effervescence takes place | (a) May be Cl |
| clean and dry test tube. About 2 to 3 drops at conc. H_2SO_4 | with the evolution of a colourless gas with pungent odour. White | (Confirm by AgNO ₃ test.) |
| was added. Then it was warmed slightly | fumes were produced when a glass rod dipped in conc. NH ₄ OH is shown to the gas. (d) No effervescence and no gas | (d) Cl⁻is absent. |
| | is evolved. | |
| (b)Confirmatory Test for CI (Sil | ver Nitrate Test) | |
| About 1 ml of salt solution taken in a test tube is acidified with dilute HNO_3 and $AgNO_3$ solution is added | (i) Curdy white precipitate soluble in dilute NH₄OH which reappeared on addition of dil.HNO ₃ . | (i) Cl ⁻ is confirmed. |

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|--|-----------|
| III.(a) Test for NO ₃ (Conc. H ₂ SO | ${}_4$ and Copper turning) | |
| | Brown fumes are evolved and solution in test tube is turned green. | |

| (b) Brown Ring Test for NO ₃ ⁻ (Confirmatory Test for NO ₃ ⁻) | | | | | | | |
|---|--|--|--|--|--|--|--|
| To about 1 ml of salt solution taken in a test tube equal volume of concentrated H_2SO_4 is added. It is cooled under tap water. Then freshly prepared FeSO ₄ solution is added slowly. | A brown ring is formed at the junction of the two rings. | NO ₃ ⁻ is confirmed. | | | | | |

| EXPERIMENT | OBSERVATION | INFERENCE |
|---|---|--|
| IV. BaCl ₂ Test for SO ₄ ²⁻ (Confirm | | |
| taken in a test tube is acidified | White precipitate is obtained which is insoluble in concentrated HCI even on boiling | SO ₄ ² is confirmed. |

CONCLUSION:

Acid radical of the salt is detected to be------

EXPERIMENT NO-8

AIM OF THE EXPERIMENT:

Test for unknown basic radicals.

APPARATUS REQUIRED:

- 1. Test tube holder
- 2. Watch Glass
- 3. Blow pipe
- 4. Nichrome wire
- 5. Blue glass
- 6.Charcoal cavity

CHEMICALS REQUIRED:

- 1.Given salts
- 2.Various Reagent
- 3. Litmus paper

THEORY & PROCEDURE:

PRELIMINARY TEST:

- 1.Salt No-----
- 2.Colour of the salt------
- 3.Structure-----
- 4. Solubility of the salt-----

A.DRY TEST FOR BASIC RADICALS:

I. Dry Test tube heating

| EXPERIMENT | OBSERVATION | INFERENCE | | |
|---------------------------------|-----------------------------------|--------------------------------------|--|--|
| In a clean and dry test tube. A | (a) Water vapours condensed at | (a) Salt with water crystallisation. | | |
| pinch of the salt is heated. | the cooler part of the test tube. | | | |
| | (b) Decrepitation took place. | (b) May be crystalline salt | | |

| | (c) Salt is volatilised and white sublimate is formed. | (c) May be Volatile salt of NH_4^+ , As^{3+} and Hg^{2+} . |
|--|--|--|
| | (d) Salt is first melted and finally infusible white mass left. | (d) May be Mg ²⁺ , Al ³⁺ , Zn ²⁺ , Ba ²⁺ , Ca ²⁺ , Sr ²⁺ etc. |
| | (e) Salt is fused on heating and solidified on cooling. | (e) May be alkali or alkaline earth metal. |
| | (f) The colour of the salt is changed | (f) May be salt of Pb ²⁺ , Bi ²⁺ ,Sn ²⁺ etc. The salt is non-volatile. |
| | (i) Yellow when hot and white when cold. | (i) May be Zn ²⁺ salt. |
| | (ii) Yellow when Hot and Cold(iii) Yellowish brown in hot and yellow when cold. | (ii) May be Pb ²⁺ salt. |
| | (iv) Black residue. | (iii) May be Sn ²⁺ or Bi ³⁺ salt. |
| | | (iv) May be Cu^{2+} ,Ni ²⁺ ,Mn ²⁺ or Fe ²⁺ salt. |
| II. Heating in a Charcoal Cavity | L / • | |
| A pinch of the salt is taken in a charcoal cavity and is heated in oxidising flame with a blow | (a) Salt is completely volatilised. | (a) May be salt of NH_4^+ , As^{3+} and Hg^{2+} . (Sodalime test is to be performed). |
| pipe. | (b) An infusible incandescent white mass is obtained. | (b) May be Mg ²⁺ , Al ³⁺ , Zn ²⁺ , Ba ²⁺ , Ca ²⁺ ,Sr ²⁺ ,Sn ²⁺ etc.(Cobalt nitrate test is to be performed). |
| | (c) The salt is fused and sank into the charcoal cavity and reappeared on cooling. | (c) May be alkali or alkaline earth metal salt. (Flame test is to be performed). |
| | (d) Original salt is white and formed a coloured mass. | (d) May be salt of Pb ²⁺ , Bi ²⁺ , Sn ²⁺ Ag ⁺ etc. (Reduction test is to be performed). |
| | (e) Original salt is coloured and formed a coloured mass. | (e) May be Cr³⁺, Ag⁺, Mn²⁺ etc. (Borax bead test is to be performed). |

| watch glass. A little soda lime is that added with a drop of water. C Then it is rubbed. | • | s is evolved and e mixture is not | (a) May be NH4 ⁺ | | |
|--|---------------------------------------|--|---|--|--|
| (1 | | | (1) · · · · · · · · · · · · · · · · · · · | | |
| | . , . | of the residue is wn and there is as. | (b) May be Hg²⁺. (c) May be As³⁺ | | |
| | (c) No gas is e change in colour | evolved and no of residue. | | | |
| IV. Bulb tube test.(for Volatile Sal | lt) | | | | |
| Na_2CO_3 and charcoal powder in for | (i) a white shii formed | nning mirror is | (a) May be Hg ²⁺ . | | |
| prepared. A little of the mixture for | | nning mirror is evolution of a colour. | (b) May be As ³⁺ | | |
| V. Cobalt Nitrate test. (For infusible | ble Salt) | | | | |
| A Pinch of salt is taken in a (i | i)Blue Mass | | (a) May be Al ³⁺ . | | |
| charcoal cavity. It is heated in (i | (ii)Green Mass | | (b) May be Zn ²⁺ | | |
| an oxidizing flame till an infusible mass is obtained. A | (iii)Pink Mass | | (c) May be Mg ²⁺ | | |
| | (iv)Grey Mass | | (d) May be Ca ²⁺ | | |
| added and again heated strongly. | | | (Flame test to be performed.) | | |
| VI. Flame test. (For fusible Salt) | | | | | |
| moisten with concentrated HCI | Colour through naked flame | Colour through double blue | | | |
| , |) Golden Yellow | glass. i) Colourless | (a) May be Na⁺. | | |
| eye and through double blue ii) | i) Violet | ii) Red | (b) May be K ⁺ | | |
| glass. | ii) Brick Red | iii) Light Yellow | (c) May be Ca ²⁺ | | |
| VII. Charcoal Reduction Test. (For | r white salt cha | nging colour) | | | |
| A mixture of salt and fusion (i | (i) White shining without incrusta | malleable bead tion which did | (i) May be Ag⁺. | | |
| is prepared. A little of this n mixture is taken in a charcoal ,. | not mark on pape | er. | | | |

| cavity | and | is | heated | in | а | with lemon yellow incrustation (ii) May be Pb^2 | |
|---------|---------|-----|--------|----|---|---|--|
| reducir | ng flan | ne. | | | | which marked on paper. | |

.WET TEST FOR BASIC RADICALS:

1. WET TESTS FOR BASIC RADICALS (Group Analysis)

| | Experiment | | Observation | | Inference |
|----|--|----|---|----|---|
| 1. | To 1ml. of salt solution in a clean test tube 1 cc. of dil HCI is added. | | A white precipitate is formed. | а. | One of the Gr. I basic radicals (Pb^{2+} , Ag^+ , Hg_2^{2+}) may be present (Analysis of Gr. I basic radicals should be performed) |
| | | b. | No white precipitate is formed | b. | Gr. I basic radicals are absent. |
| 2. | To 1ml. of the supplied salt solution in a clean test tube solid NH"CI is added till saturation followed by addition of dil NH4OH till alkaline. | a. | A precipitate is obtained, (colour should be noted) | | One of the Gr III A basic radicals (Fe³⁺ , AI³⁺, Cr³⁺)may be present (Analysis of Gr III A basic radicals should be performed) |
| | | b. | No precipitate is formed. | b. | Gr III A basic radicals are absent. |
| 3. | Through the contents of the above test tube H ₂ S gas is passed under pressure. | a. | Precipitate is formed (colour should be noted). | a. | One of the Gr III B basic radicals (Zn ²⁺ , Mn ²⁺ , Co ²⁺ , Ni ²⁺) may be present(analysis of Gr III B radicals should be performed) |
| | | b. | No precipitate is formed | b. | Gr III B basic radicals are absent. |
| 4. | To 1 cc of the salt solution is taken in a clean test tube solid NH_4CI is added till saturation followed by addition of dil NH_4OH till alkaline. To this saturated solution of ammonium carbonate is added. | а. | (colour should be noted). | | One of the Gr IV basic radicals (Ba ²⁺ , Sr ²⁺ , Ca ²⁺) may be present(analysis of Gr IV radicals should be performed) |
| | The above basic radicals are absent i | | No precipitate is formed. | | Gr. IV basic radicals are absent. |

The above basic radicals are absent indicating that one of the Gr. V basic radicals may be present. As there is no specific group reagent for Gr. V test for individual radicals should be performed

2. ANALYSIS OF BASIC RADICALS (GROUP WISE)

i) Analysis of Gr. IIIA Basic Radicals (Al³⁺)

| | Experiment | Observation | Inference |
|----|---|---|-----------------------------|
| 1. | 1 - 2 cc of the supplied salt solution is saturated with solid NH ₄ Cl followed by the addition of dil NH ₄ OH solution till alkaline. | A white ppt. is formed. | May be Al ³⁺ |
| 2. | 1 – 2 cc of the supplied salt solution is treated with dil NaOH solution drop wise and then in excess. | A white ppt. of Al(OH) ₃ is formed which dissolved in excess of the reagent. | May be Al ³⁺ |
| 3. | 1 cc of the supplied salt solution, disodium hydrogen phosphate solution is added. | A gelatinous white ppt. of AIPO ₄ is formed which is soluble in dil. HCI solution. | Al ³⁺ confirmed. |

ii) Analysis of Gr. IIIB Basic Radicals (Zn²⁺)

| Experiment | Observation | Inference |
|---|--|-----------------------------|
| 1 – 2 cc of the supplied salt solution is saturated with solid NH₄CI followed by the addition of dil NH₄OH solution till alkaline. Then H₂S gas is passed through it. | A white ppt. is formed. | May be Zn ²⁺ |
| 2. 1 -2 cc of the supplied salt solution is treated with potassium ferrocyanide solution drop by drop and then in excess. | A white ppt is obtained. | May be Zn ²⁺ |
| Dil. NaOH solution is added to 1 cc of the salt solution drop by drop and then in excess. | A gelatinous white ppt. is formed which is soluble in excess of NaOH solution. | Zn ²⁺ confirmed. |

iii) Analysis of Gr. IV Basic Radicals (Ca²⁺)

| Experiment | Observation | Inference |
|--|--|-------------------------|
| 1 – 2 cc of the supplied salt solution is saturated with solid NH₄Cl and then made alkaline with dil | A white ppt. of CaCO ₃ is formed. | May be Ca ²⁺ |

| NH₄OH solution. Then saturated solution of ammonium carbonate [(NH₄)₂CO₃] is added. | | |
|---|--|-------------------------|
| The above ppt. is dissolved in a minimum quantity of dil CH₃COOH. The solution is boiled to remove CO₂ and then ammonium oxalate solution is added to it. | A white ppt. of CaC_2O_4 is formed which is soluble in dil. HCI but insoluble in CH ₃ COOH. | May be Ca ²⁺ |

ii) Analysis of Gr.V Basic Radicals (NH_4^+ , Na^+ , K^+)

Tests for NH₄⁺

| Experiment | Observation | Inference |
|---|---|-----------------------------|
| A small quantity of the salt is treated with soda lime and two drops of water and then the mixture is rubbed in a | A colourless gas having smell of ammonia which produced dense white fumes with a glass rod dipped in conc. NH ₄ OH. There is no change in the | NH₄⁺ confirmed. |
| mortar. | colour of the residue. | |
| Nessler's reagent is added to 1 cc of the salt solution. | A brown ppt. is obtained. | NH₄ ⁺ confirmed. |

Tests for Mg²⁺

| | Experiment | Observation | Inference |
|----|---|--------------------------|-----------------------------|
| 1. | 1 - 2 cc of the supplied salt solution is saturated with solid NH ₄ Cl followed | A white ppt. is formed. | May be Mg²⁺ |
| | by the addition of dil NH ₄ OH solution till | | |
| | alkaline. Then dihydrogen sodium phosphate | | |
| | solution is added to it. | | 2. |
| 2. | 1 cc of the salt solution is acidified with dil. HCl and then treated with a few drops of magneson reagent followed by the addition of excess of dil NaOH solution. | A blue ppt. is obtained. | Mg ²⁺ confirmed. |

Tests for Na⁺

| Experiment | Observation | Inference |
|----------------------------|-------------------------------------|----------------------------|
| 1. Potassium | A white crystalline ppt. is formed. | Na ⁺ confirmed. |
| pyroantimonate solution is | | |
| added to 1 cc of the | | |
| supplied salt solution. | | |

Tests for K⁺

| Experiment | Observation | Inference |
|---|--------------------------|---------------------------|
| 1. 1 cc of the salt solution is treated with two drops of | A yellow ppt. is formed. | K ⁺ confirmed. |
| cobalt nitrate solution followed by the addition | | |
| of solid NaNO ₃ and dil. CH_3COOH solution. | | |

Hence, the basic part of the supplied salt is _____.

EXPERIMENT NO. 09

(Tests for Unknown Salt)

AIM OF THE EXPERIMENT:

Test for unknown basic radicals.

APPARATUS REQUIRED:

- 1. Test tube holder
- 2. Watch Glass
- 3. Blow pipe
- 4. Nichrome wire
- 5. Blue glass
- 6.Charcoal cavity

CHEMICALS REQUIRED:

- 1.Given salts
- 2.Various Reagent
- 3. Litmus paper

THEORY & PROCEDURE:

1. Preliminary Test:

- (a) Salt No:
- (b). Colour of the Salt : Colourless / name of the colour
- (c) . Structure of Salt : Crystalline/ Amorphous
- (d). Solubility:
 - i)Soluble in cold water (if not)
 - ii) Soluble in hot water (if not)
 - iii) Soluble in dilute HCI (if not)
 - iv) Soluble in hot dilute HCl
 - v) If not then salt is insoluble (*Salt soluble in dil HCl implies Gr.I basic radicals absent)

2. DRY TEST FOR BASIC RADICALS

Dry Test Tube heating:

| Experiment | Observation | Inference |
|---|---|--|
| A small quantity of salt is | (a) A sublimate is formed | (a) It is volatile salt, (Soda lime |
| taken in a clean and dry test | (Note the colour of the | test and bulb tube test should |
| tube and heated strongly in | sublimate) | be performed.) |
| the hottest part of the non- luminous flame. | b)Water particles condense at the cooler part of the test | (b) Salt contains water of crystallisation. |
| | (c) Decripitation or cracking | (c) May be crystalline salt. |
| | sound is produced. | |
| | (d) Deflagration takes place. | (d) The salt may be nitrate of alkali or alkaline earth metal. |
| | (e) The salt changes colour. Yellow when hot and white when cold. | (e) It may be Zinc salt. |

| (f) Salt fuses on heating and solidifies on cooling. | (f) May be alkali or alkaline earth metal salt. |
|--|---|
|--|---|

3. SODALIME TEST:

| Experiment | Observation | Inference |
|------------------------------------|--------------------------------------|-------------------------|
| A little of the salt is taken in a | A colourless gas evolved with strong | NH4 ⁺ may be |
| clean watch glass along with | smell of ammonia and colour of the | present. (To be |
| soda-lime and it is rubbed by | mixture is unchanged. | confirmed in the |
| adding two drops of water. | | wet test) |

4. CHARCOAL CAVITY HEATING (OXIDISING FLAME)

| Experiment | Observation | Inference |
|--|---|--|
| A little of the Salt is taken in | a. The salt decrepitates. | a. Maybe crystalline salt. |
| the charcoal cavity and | b. The salt deflagrates. | b. May be NO_3^- salt |
| heated by oxidizing flame with the help of a blow pipe. | c. The salt fuses and sinks into the charcoal cavity. | c. Salt contains alkali or alkaline earth metal. (Flame test should be performed). |
| | d. Infusible incandescent white residue. | d. Cobalt nitrate test should be performed. |

5. COBALT NITRATE TEST

| Experiment | Observation | Inference |
|---|----------------------------|-------------------------------------|
| The salt is taken in the | a. Blue mass is obtained. | a. Al ³⁺ may be present. |
| charcoal cavity and heated in | b. Green mass is obtained. | b. Zn ²⁺ may be present. |
| the oxidizing flame with the | c. Rosy mass is obtained. | c. Mg ²⁺ may be present. |
| help of a blow pipe till an infusible, incandescent white mass is obtained. Then one drop of cobalt nitrate solution is added to it and heated strongly. | d. Grey mass is obtained. | d. Ca ²⁺ may be present. |

6.FLAME TEST

| Experiment | Observation | Inference |
|--|---|-------------------------------------|
| The nichrome wire is cleaned with sand paper and dipped in conc. HCl and shown to non- luminous flame. This process is repeated till no colour is imparted to the flame. Then the wire is moistened with | Persistent golden Yellow coloured flame is seen in naked eye and colourless through double blue glass. b) Violet flame is seen in naked eye and red through a pair of blue glass. | a. Na ⁺ may be present. |
| conc. HCl and a little of the salt is taken by touching to the salt is taken by touching to the | b. Violet flame is seen in naked eye and red through a pair of blue glass. | b. K ⁺ may be present. |
| salt and shown to the oxidizing flame. | c. Brick red flame is observed. | c. Ca ²⁺ may be present. |

7.IDENTIFICATION OF ACID RADICAL

Test for Gr-I acid radicals (Carbonate and Sulphide)

| Experiment | Observation | Inference |
|---|---|---|
| 1 cc dilute HCl taken in d test tube and slightly warmed. To this a pinch of the supplied salt is added. | Effervescence took place with the evolution of a colourless odourless gas is evolved. | Carbonate (CO₃²⁻) may be present (other test should be performed for its confirmation.) |
| | Effervescence took place with the evolution of a colourless odourless gas with rotten egg smell is evolved. | a. Sulphide (S²⁻) may be present (other test should be performed for its confirmation.) |

Test for Gr- II acid radicals (Chloride)

| Experiment | Observation | Inference |
|-----------------------------------|--------------------------------------|---------------------------------|
| A few drops of conc. H_2SO_4 is | A colourless fuming gas with pungent | Cl [−] may be present. |
| taken in a clean and dry test | odour is evolved. | (Other test should |
| tube, a pinch of the supplied | | be performed for |
| salt is added in to it and is | | its confirmation). |
| gently warmed. | | |

Test for Gr- III acid radicals (Nitrate and Sulphate)

TESTS FOR NITRATE (NO₃⁻)

| Experiment | Observation | Inference |
|---|--|---|
| A pinch of the supplied salt is moistened with a few drops of conc. H_2SO_4 is taken in a clean and dry test tube and is gently warmed. | A brown fume with pungent smell is observed. | May be NO₃⁻. (Other test should be performed for its confirmation). |

TESTS FOR SULPHATE (SO₄²⁻)

| Experiment | Observation | Inference |
|--|---|------------------------------|
| 1-2 cc of the salt solution is taken in a clean test tube and is acidified with dil HCl. A few cc of Barium chloride (BaCl ₂) solution is added into it. | A white ppt. is obtained which is insoluble in conc. HCl even on boiling. | SO₄ ^{2−} confirmed. |

CONFIRMATORY TESTS FOR CARBONATE (CO₃²⁻)

| | Experiment | Observation | Inference | |
|----|--|--|---|----|
| 1. | A burning match stick is shown to the evolved gas. | The burning stick extinguished. | CO ₃ ^{2–} may be present. | |
| 2. | A little more salt is added to the above test tube and the evolved gas is passed through lime water with the help of a delivery tube. | At first white turbidity (milk colour) appeared which disappeared with excess passing of the gas. | CO ₃ ²⁻ may present | be |
| 3. | A little more salt is added to the above test tube and the evolved gas is passed through acidified potassium dichromate solution with the help of a delivery tube. | No change of the colour took place. | CO₃ ^{2−} confirmed | |

CONFIRMATORY TESTS FOR SULPHISE (S²⁻)

| Experiment | Observation | Inference |
|--|--------------------------------|----------------------------|
| A filter paper soaked with Lead acetate solution is shown to the mouth of the test tube. | The filter paper turned black. | S ^{2−} Confirmed. |

CONFIRMATORY TESTS FOR CHLORIDE (CI⁻)

| | Experiment | Observation | Inference |
|----|--|--|----------------------------|
| 1. | A glass rod dipped in conc. NH₄OH solution is shown to the gas evolved. | A white dense fume is formed. | Cl⁻may be present. |
| 2. | A pinch of MnO_2 is added to the above test tube and is warmed gently. | A greenish yellow gas is formed which turned starch iodide paper blue. | Cl⁻ may be present. |
| 3. | A pinch of the given salt is taken in a clean and dry test tube and is acidified with dil HNO ₃ solution. And a few drops of silver nitrate (AgNO ₃) solution is added into it. | A curdy white ppt. is formed which is soluble in dil NH ₄ OH and is insoluble in dil HNO ₃ . | Cl [⊤] confirmed. |

CONFIRMATORY TEST FOR NITRATE (NO3-)

| | Experiment | Observation | Inference |
|----|---|--|----------------------|
| 1. | A pinch of the supplied salt and a few copper turnings are taken in a clean test tube. $1 - 2 \text{ cc}$ of 50% conc. H_2SO_4 is added into it and is heated gently. | Deep brown vapours are formed and the solution turned bluish green or green. | May be NO ₃⁻. |
| 2. | A piece of filter paper soaked in FeSO ₄ solution is shown to the | It turned black. | May be NO₃⁻. |

| evolved gas. | | |
|--|---|-----------------------------|
| 1 cc of the supplied salt solution in water is taken in a clean test tube. Equal volume of conc. H₂SO₄ is added in to the test tube. The test tube is cooled under tap water. And equal volume of freshly prepared ferrous sulphate (FeSO₄) solution is added from the side of the test tube. | A brown ring is formed at the junction of the two liquids. The ring disappeared on shaking. | NO₃ ⁻ confirmed. |

8.WET TESTS FOR BASIC RADICALS (Group Analysis)

| | Experiment | | Observation | | Inference |
|----|--|----|---|----|---|
| 1. | To 1ml. of salt solution in a clean test tube 1 cc. of dil HCl is added. | a. | A white precipitate is formed. | a. | One of the Gr. I basic radicals (Pb^{2+} , Ag^{+} , Hg_{2}^{2+}) may be present (Analysis of Gr. I basic radicals should be performed) |
| | | b. | formed | b. | Gr. I basic radicals are absent. |
| 2. | To 1ml. of the supplied salt solution in a clean test tube solid NH"CI is added till saturation followed by addition of dil NH4OH till alkaline. | a. | (colour should be noted) | a. | One of the Gr III A basic radicals (Fe ³⁺ , AI ³⁺ , Cr ³⁺)may be present (Analysis of Gr III A basic radicals should be performed) |
| | | b. | No precipitate is formed. | b. | Gr III A basic radicals are absent. |
| 3. | Through the contents of the above test tube H_2S gas is passed under pressure. | a. | (colour should be noted). | a. | One of the Gr III B basic radicals (Zn ²⁺ , Mn ²⁺ , Co ²⁺ , Ni ²⁺) may be present(analysis of Gr III B radicals should be performed) |
| | | b. | No precipitate is formed | b. | Gr III B basic radicals are absent. |
| 4. | To 1 cc of the salt solution is taken in a clean test tube solid NH_4CI is added till saturation | a. | Precipitate is formed (colour should be noted). | a. | One of the Gr IV basic radicals (Ba ²⁺ , Sr ²⁺ , Ca ²⁺) may be |

| followed by addition of dil NH₄OH till alkaline. To this saturated solution of ammonium carbonate is added. | | | | present(analysis of Gr IV radicals should be performed) |
|--|----|---------------------------|----|--|
| | b. | No precipitate is formed. | b. | Gr. IV basic radicals are absent. |

The above basic radicals are absent indicating that one of the Gr. V basic radicals may be present. As there is no specific group reagent for Gr. V test for individual radicals should be performed.

9.ANALYSIS OF BASIC RADICALS (GROUP WISE)

ii) Analysis of Gr. IIIA Basic Radicals (Al³⁺)

| | Experiment | Observation | Inference |
|----|---|---|-----------------------------|
| 1. | 1 - 2 cc of the supplied salt solution is saturated with solid NH ₄ Cl followed by the addition of dil NH ₄ OH solution till alkaline. | A white ppt. is formed. | May be Al ³⁺ |
| 2. | 1 – 2 cc of the supplied salt solution is treated with dil NaOH solution drop wise and then in excess. | A white ppt. of Al(OH) ₃ is formed which dissolved in excess of the reagent. | May be Al ³⁺ |
| 3. | 1 cc of the supplied salt solution, disodium hydrogen phosphate solution is added. | A gelatinous white ppt. of AIPO ₄ is formed which is soluble in dil. HCI solution. | Al ³⁺ confirmed. |

iv. Analysis of Gr. IIIB Basic Radicals (Zn²⁺)

| Experiment | Observation | Inference |
|---|--|-----------------------------|
| 1. $1 - 2 \text{ cc}$ of the supplied salt solution is saturated with solid NH ₄ Cl followed by the addition of dil NH ₄ OH solution till alkaline. Then H ₂ S gas is passed through it. | A white ppt. is formed. | May be Zn ²⁺ |
| 1 -2 cc of the supplied salt solution is treated with potassium ferrocyanide solution drop by drop and then in excess. | A white ppt is obtained. | May be Zn ²⁺ |
| Dil. NaOH solution is added to 1 cc of the salt solution drop by drop and | A gelatinous white ppt. is formed which is soluble in excess of NaOH solution. | Zn ²⁺ confirmed. |

then in excess.

iv) Analysis of Gr. IV Basic Radicals (Ca²⁺)

| Experiment | Observation | Inference |
|---|--|-----------------------------|
| 1 – 2 cc of the supplied salt solution is saturated with solid NH₄Cl and then made alkaline with dil NH₄OH solution. Then saturated solution of ammonium carbonate [(NH₄)₂CO₃] is added. | A white ppt. of CaCO ₃ is formed. | May be Ca ²⁺ |
| The above ppt. is dissolved in a minimum quantity of dil CH₃COOH. The solution is boiled to remove CO₂ and then ammonium oxalate solution is added to it. | A white ppt. of CaC_2O_4 is formed which is soluble in dil. HCl but insoluble in CH ₃ COOH. | May be Ca ²⁺ |
| 3. Flame test is performed with the white ppt. formed above. | Brick red flame is noticed. | Ca ²⁺ confirmed. |
| iii) Analysis of Gr.V Bas | ic Radicals (NH₄⁺, Na⁺, K⁺) | |

Tests for NH₄⁺

| Experiment | Observation | Inference |
|---|--|-----------------------------|
| A small quantity of the salt is treated with soda lime and two drops of water and then the mixture is rubbed in a mortar. | A colourless gas having smell of ammonia which produced dense white fumes with a glass rod dipped in conc. NH ₄ OH. There is no change in the colour of the residue. | NH₄ ⁺ confirmed. |
| Nessler's reagent is added to 1 cc of the salt solution. | A brown ppt. is obtained. | NH₄ ⁺ confirmed. |

Tests for Mg²⁺

| Experiment | Observation | Inference |
|---|--------------------------|-----------------------------|
| 1 – 2 cc of the supplied salt solution is saturated | A white ppt. is formed. | May be Mg ²⁺ |
| with solid NH ₄ Cl followed | | |
| by the addition of dil | | |
| NH₄OH solution till | | |
| alkaline. Then dihydrogen | | |
| sodium phosphate | | |
| solution is added to it. | | |
| 2. 1 cc of the salt solution is | A blue ppt. is obtained. | Mg ²⁺ confirmed. |

| acidified with dil. HCI and | |
|-----------------------------|--|
| then treated with a few | |
| drops of magneson | |
| reagent followed by the | |
| addition of excess of dil | |
| NaOH solution. | |

Tests for Na⁺

| Experiment | Observation | Inference |
|---|-------------------------------------|----------------|
| Potassium pyroantimonate solution is added to 1 cc of the supplied salt solution. | A white crystalline ppt. is formed. | Na⁺ confirmed. |

Tests for K⁺

| Experiment | Observation | Inference |
|---|--------------------------|---------------|
| 1 cc of the salt solution is treated with two drops of cobalt nitrate solution followed by the addition of solid NaNO₃ and dil. CH₃COOH solution. | A yellow ppt. is formed. | K⁺ confirmed. |

Hence, the basic part of the supplied salt is _____ and the acid part of the salt is _____.

Thus, the salt supplied is _____.