

Class – XII	Subject: Chemistry(Practical)Term-1&2	Experiment (2021_22)
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Exp. No	Aim
	QUANTITATIVE ANALYSIS(Term-1&2)
1	Prepare 250 ml of 0.1MSolution of Oxalic Acid From Crystalline Oxalic Acid
2	Determination of Concentration/Molarity of KMnO4 Solution by Titrating it
	against a 0.1M Standard Solution of Oxalic acid
3	Determination of Concentration/Molarity of KMnO4 Solution by Titrating it
	against a Standard Solution of Ferrous ammonium sulphate
	QUALITATIVE ANALYSIS(Term-1&2)
4	To Identify the given inorganic salt[Ba(NO3)2]
5	To Identify the given inorganic salt [ZnCo3]
6	To Identify the given inorganic salt [Pb(NO3)2]
7	To Identify the given inorganic salt PbCl2
8	To Identify the given inorganic salt MgSO4
9	To Identify the given inorganic salt [BaSO4]
10	To Identify the given inorganic salt [Sr(NO3)2]
	CHROMATOGRAPHY(Term-1)
11	Separate the Coloured Components Present in the Mixture of Red and Blue Inks by
	Ascending Paper Chromatography and Find their Rf Values
12	Separate the Coloured Components Present in the Given Grass/Flower by Ascending Paper
	Chromatography and Determine their Rf Values

ORGANIC COMPOUNDS(Term-2)

15	To Identify functional group of Aldehyde (-CHO)
16	To Identify functional group of Ketone (-CO-)
17	To Identify functional group of Alcohol (-OH)
18	To Identify functional group of Carboxylic acid (-COOH)

Aim: Prepare 250 ml of M/10 Solution of Oxalic Acid From Crystalline Oxalic Acid

Theory

Molecular mass of crystalline oxalic acid
$$\begin{pmatrix} COOH \\ | & 2H_2O \\ COOH \end{pmatrix} = 126$$

Hence, for preparing 1000 ml of 1M oxalic acid, weight of oxalic acid crystals required = 126 g

:. For preparing 250 ml of 0.1M solution,

oxalic acid crystals required = $\frac{126}{1000} \times 250 \times 0.1 = 3.150$ g.

Apparatus

Watch glass, analytical balance, weight box, fractional weight box, 250 ml beaker, glass rod, 250 ml measuring flask and wash bottle.

Chemical Required

Oxalic acid crystals and distilled water.

Procedure

1. Take a watch glass, wash it with distilled water and then dry it.

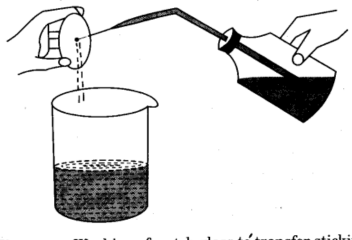
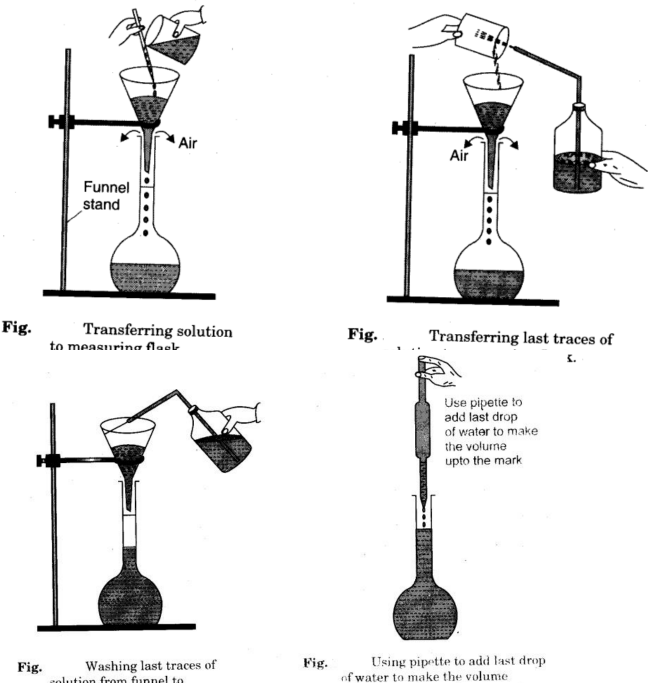


Fig. Washing of watch glass to transfer sticking particles to beaker.

2. Weigh the clean and dried watch glass accurately and record its weight in the note book.

- 3. Weigh 3.150 g oxalic acid on the watch glass accurately and record this weight in the note-book.
- 4. Transfer gently and carefully the oxalic acid from the watch glass into a clean 250 ml beaker. Wash the watch glass with distilled water with the help of a wash bottle to transfer the particles sticking to it into the beaker [Fig].
 - The volume of distilled water for this purpose should not be more than 50 ml.
- 5. Dissolve oxalic acid crystals in the beaker by gentle stirring with a clean glass rod.
- 6. When the oxalic acid in the beaker is completely dissolved, transfer carefully the entire solution from the beaker into a 250 ml measuring flask (volumetric flask) with the help of a funnel [Fig].



solution from funnel to the measuring flask.

upto the mark.

7. Wash the beaker with distilled water. Transfer the washings into the measuring flask [Fig].

- 8. Finally wash the funnel well with distilled water with the help of a wash bottle to transfer the solution sticking to the funnel into the measuring flask [Fig].
- 9. Add enough distilled water to the measuring flask carefully, up to just below the etched mark on it, with the help of a wash bottle.
- 10. Add the last few drops of distilled water with a pipette until the lower level of the meniscus just touches the mark on the measuring flask [Fig].
- 11. Stopper the measuring flask and shake gently to make the solution uniform through-out. Label it as oxalic acid solution.

Result:- 250 ml of M/10 Solution of Oxalic Acid From Crystalline Oxalic Acid is prepared.

EXPERIMENT-2

Aim:

To determine the strength of potassium permanganate by titrating it against the standard solution of 0.1M oxalic acid.

Theory:

Potassium permanganate is a strong oxidising agent and in the presence of sulfuric acid it acts as a powerful oxidising agent. In acidic medium the oxidising ability of $KMnO_4$ is represented by the following equation.

In acidic solution,

$MnO_{4^-} + 8H^{\scriptscriptstyle +} + 5e^{\scriptscriptstyle -} \rightarrow Mn^{\scriptscriptstyle 2+} + 4H_2O$

Solution containing MnO_{4^-} ions are purple in colour and the solution containing Mn^{2+} ions are colourless and hence permanganate solution is decolourised when added to a solution of a reducing agent. The moment there is an excess of potassium permanganate present the solution becomes purple. Thus, **KMnO**₄ serves as self indicator in acidic solution.

Potassium permanganate is standardized against pure oxalic acid. It involves a redox reaction. Oxalic acid is oxidised to carbon dioxide by KMnO₄, which itself gets reduced to MnSO₄. Oxalic acid reacts with potassium permanganate in the following way.

The chemical reaction at room temperature is given below.

Reduction Half reaction:- $2KMnO_4 + 3H_2SO_4 \rightarrow K_2SO_4 + 2MnSO_4 + 3H_2O + 5[O]$

Oxidation Half reaction:- $5(COOH)_2 + 5[O] \rightarrow 5H_2O + 10CO_2\uparrow$

The overall reaction takes place in the process is

 $Overall\ reaction: -\ 2KMnO_4 + 3H_2SO_4 + 5(COOH)_2 \rightarrow K_2SO_4 + 2MnSO_4 + 8H_2O + 10CO_2 \uparrow$

The ionic equation involved in the process is given below.

Reduction Half reaction:- [MnO₄⁻ + 8H⁺ + 5e⁻ \rightarrow Mn²⁺ + 4H₂O] x 2

Oxidation Half reaction:- $[C_2O_4^2 \rightarrow 2CO_2 + 2e^-] \ge 5$

Overall Ionic reaction:- $2MnO_4^- + 16H^+ + 5C_2O_4^2 \rightarrow 2Mn^{2+} + 10CO_2 + 8H_2O_3$

This titration cannot be carried out in the presence of acids like nitric acid or hydrochloric acid because itself is an oxidising agent. So hydrochloric acid chemically reacts with $KMnO_4$ solution forming chlorine which is also an oxidising agent.

Materials Required:

- 1. Oxalic acid
- 2. Potassium permanganate solution
- 3. 1.0M sulfuric acid
- 4. Chemical balance
- 5. Burette
- 6. Burette stand
- 7. Pipette
- 8. Conical flask
- 9. Funnel
- 10. Measuring flask
- 11. Weighing bottle
- 12. White tile
- 13. Burnet
- 14. Wire gauze

Apparatus Setup:

- 1. In burette $KMnO_4$ solution
- 2. In Conical flask 10ml of oxalic acid + Sulfuric acid
- 3. Indicator Self indicator (KMnO₄)
- 4. End Point Appearance of permanent pale pink colour.

Procedure:

(a) Preparation of 0.1N standard solution of oxalic acid:

The quantity of oxalic acid required for the 250ml of the solution having a normality of 0.1N can be calculated as follows.

Equivalent weight of oxalic acid = Molecular weight/No of electrons lost by one molecule

Equivalent weight of oxalic acid = 126/2 = 63

Strength = Normality x Equivalent weight

Strength = $1/10 \ge 63 = 6.3 \text{ g/l}$

For the preparation of 1 litre of N/10 oxalic acid solution amount of oxalic acid required = 6.3 g

- 1. Weigh an empty watch glass using a chemical balance.
- 2. Weigh 6.3g of oxalic acid accurately in the watch glass.
- 3. With the help of a funnel transfer the oxalic acid into the measuring flask.
- 4. Now wash the funnel with distilled water without removing the funnel from the flask.
- 5. Make the solution up to the marked point with distilled water and make sure the oxalic acid is fully dissolved.
- 6. This solution is 0.1N standard solution of oxalic acid.

(b) Titration of potassium permanganate solution against standard oxalic acid solution:

- 1. Rinse the burette with the potassium permanganate solution and fill the burette with potassium permanganate solution.
- 2. Fix the burette in the burette stand and place the white tile below the burette in order to find the end point correctly.
- 3. Pipette out 10ml of 0.1N standard oxalic acid solution in a conical flask.

- 4. Add a test tube full of sulfuric acid in order to prevent oxidation of manganese to form manganese dioxide.
- 5. Heat the mixture upto 60°C before titrating with potassium permanganate.
- 6. Note down the initial reading in the burette before starting the titration.
- 7. The hot solution is titrated against potassium permanganate solution and simultaneously swirl the solution in the flask gently.
- 8. Initially the purple colour of KMnO₄ is discharged with oxalic acid. The appearance of permanent pink colour reveals the end point.
- 9. Repeat the titration until concordant values are obtained.
- 10. Note down the upper meniscus on the burette readings. Record the reading in the observation table given below in order to calculate the molarity of KMnO₄ given.

Observation:

S.No	Volume of oxalic acid in ml	Burette Reading		Volume(V) of KMnO ₄ used V = (y-x)ml
		Initial(x)	Final(y)	

Calculations:

To calculate the strength of given KMnO4 in terms of molarity the following formula is used

$\mathbf{a}_1\mathbf{M}_1\mathbf{V}_1 = \mathbf{a}_2\mathbf{M}_2\mathbf{V}_2$

Where a₁ and a₂ are stoichiometric coefficient of oxalic acid and KMnO₄ in a balanced chemical equation.

 $a_1 = 2$

 $a_2 = 5$

Where

 M_2 and M_1 are molarities of potassium permanganate and oxalic acid solutions used in the titration.

 V_2 and V_1 are the volume of potassium permanganate and oxalic acid solutions used in the titration.

Therefore,

 $KMnO_4 = Oxalic acid$

 $5M_2V_2 = 2M_1V_1$

 $M_2 = (2M_1V_1/5M_2V_2)$

The strength of KMnO₄ is calculated by using the molarity.

Strength = Molarity x Molar mass

Results and Discussion:

- 1. Molarity of KMnO₄ is _____
- 2. The Strength of $KMnO_4$ is _____M.

Precautions:

- 1. Clean all the apparatus with distilled water before starting the experiment and then rise with the solution to be taken in them.
- 2. Rinse the pipette and burette before use.
- 3. Potassium permanganate is dark in colour, so always read the upper meniscus.
- 4. Use dilute sulfuric acid for acidifying the potassium permanganate.
- 5. Take accurate readings once it reaches the end point and don't go with average readings.
- 6. Use antiparallex card or autoparallex card while taking the burette readings.
- 7. Do not use rubber cork burette as it is can be attacked by $KMnO_4$.
- 8. The strength of the unknown solution should be taken upto two decimal places only.

EXPERIMENT-3

Aim:

To determine the strength of a given potassium permanganate solution against a standard ferrous ammonium sulfate (Mohr's salt) solution.

Theory:

Potassium permanganate is a strong oxidant in the presence of sulfuric acid. Mohr salt is a double salt forming a single crystalline structure having the formula (NH4)2. FeSO4. 6H2O. The chemical name for Mohr's salt is ferrous ammonium sulfate.

In this titration Mohr salt acts as a reducing agent and potassium permanganate acts as an oxidising agent. So, the reaction between Mohr's salt and potassium permanganate is a redox reaction. In this redox reaction, ferrous ion from Mohr's salt gets oxidised and pink coloured of manganese present in potassium permanganate, which is in the +7 oxidation state gets reduced to colourless Mn^{2+} state.

The chemical reaction and the molecular chemical equation is given below.

Reduction half reaction -

$2KMnO_4 + 3H_2SO_4 \rightarrow K_2SO_4 + 2MnSO_4 + 3H_2O + 5[O]$

Oxidation half reaction -

 $[2FeSO_4(NH_4)_2SO_4.6H_2O + H_2SO_4 + [O] \rightarrow Fe_2(SO_4)_3 + 2(NH_4)_2SO_4 + 13H_2O] \ge 5$

Overall reaction -

 $2KMnO_4 + 10FeSO_4(NH_4)_2SO_4.6H_2O + 8H_2SO_4 \rightarrow K_2SO_4 + 2MnSO_4 + 5Fe_2(SO_4)_3 + 10(NH_4)_2SO_4 + 68H_2O_4(NH_4)_2SO_4 + 68H_2O_4($

The ionic equation involved in the process is given below.

Oxidation half reaction – $[Fe^{2+} \rightarrow Fe^{3+} - e^{-}] \ge 5$

Reduction half reaction – $MnO_{4^-} + 8H^+ + 5e^- \rightarrow Mn^{2+} + 4H_2O$

Overall ionic equation – $MnO_{4^-} + 8H^+ + 5Fe^{2+} \rightarrow Mn^{2+} + 5Fe^{3+} + 4H_2O$

This titration is based upon oxidation-reduction titrations. When ferrous ammonium sulfate solution is titrated against potassium permanganate in the presence of acidic medium by sulfuric acid. Acidic medium is necessary in order to prevent precipitation of manganese oxide. Here $KMnO_4$ acts as a self indicator and this titration is called permanganate titration.

Materials Required:

- 1. Mohr's salt (ferrous ammonium sulfate)
- 2. Potassium permanganate solution
- 3. Dilute sulfuric acid
- 4. Chemical balance
- 5. Burette
- 6. Burette stand
- 7. Pipette
- 8. Conical flask
- 9. Funnel
- 10. Measuring flask
- 11. Weighing bottle
- 12. White tile
- 13. Burnet
- 14. Wire gauze
- 15.

Apparatus Setup:

- 1. In burette $KMnO_4$ solution
- 2. In Conical flask 10ml of Ferrous Ammonium Sulfate (Mohr's salt) + Sulfuric acid
- 3. Indicator Self indicator (KMnO₄)
- 4. End Point Colourless to permanent pale pink colour.

Procedure:

(a) Preparation of 0.05M standard solution of ferrous ammonium sulfate:

The quantity of Mohr's salt required for the 250ml of the solution having a normality of 0.05N can be calculated as follows.

The molar mass of mohr's salt = 392 g/mol

Strength = Normality x Equivalent weight = $(1/20) \times 392 = 19.6 \text{ g/L}$

For preparing 250ml of N/20 Mohr's salt solution, Mohr salt required = $(19.6/1000) \times 250 = 4.9 \text{ gm}$

- 1. Weigh an empty watch glass using a chemical balance.
- 2. Weigh accurately 4.9gm of Mohr's salt in a chemical balance.
- 3. With the help of a funnel transfer the Mohr's salt into the measuring flask.
- 4. Now wash the funnel with distilled water without removing the funnel from the flask.
- 5. Make the solution up to the marked point with distilled water and make sure the Mohr's salt is fully dissolved.
- 6. This solution is 0.05N standard solution of Mohr's salt.

(b) Titration of potassium permanganate solution against standard ferrous ammonium sulfate (Mohr's salt) solution:

- 1. Wash and rinse the burette and pipette with distilled water and then rinse with the corresponding solution to be filled in them.
- 2. Rinse the burette with the potassium permanganate solution and fill the burette with potassium permanganate solution.
- 3. Fix the burette in the burette stand and place the white tile below the burette in order to find the endpoint correctly.
- 4. Rinse the pipette and conical flask with standard ferrous sulfate solution.
- 5. Pipette out 10ml of 0.05N standard Mohr's salt solution into the conical flask.

- 6. Add a test tube full of sulfuric acid in order to prevent oxidation of manganese to form manganese dioxide.
- 7. Note down the initial reading in the burette before starting the titration.
- 8. Now start the titration, titrate against potassium permanganate solution and simultaneously swirl the solution in the flask gently.
- 9. Initially, the purple colour of KMnO₄ is discharged with ferrous ammonium sulfate. The appearance of a permanent pink colour reveals the endpoint.
- 10. Repeat the titration until concordant values are obtained.
- 11. Note down the upper meniscus on the burette readings.
- 12. Record the reading in the observation table given below in order to calculate the molarity of KMnO₄ given.

Observations:

S.No	Volume of ferrous ammonium sulfate (Mohr's salt) used	Burette Ro	eading	Volume(V) of KMnO₄ used V = (y-x)ml
		Initial (x)	Final(y)	

Calculations:

To calculate the strength of given KMnO4 in terms of molarity the following formula is used

$\mathbf{a}_1\mathbf{M}_1\mathbf{V}_1 = \mathbf{a}_2\mathbf{M}_2\mathbf{V}_2$

Where a_1 and a_2 are stoichiometric coefficient of **ferrous ammonium sulfate** and KMnO₄ in a balanced chemical equation.

 $a_1 = 1$

 $a_2 = 5$

Where

 M_2 and M_1 are molarities of potassium permanganate and **ferrous ammonium sulfate** solutions used in the titration.

 V_2 and V_1 are the volume of potassium permanganate and **ferrous ammonium sulfate** solutions used in the titration.

Therefore,

$KMnO_4 =$ ferrous ammonium sulfate

 $5M_2V_2 = 1M_1V_1$

 $M_2 = (1M_1V_1/5M_2V_2)$

The strength of KMnO₄ is calculated by using the molarity.

Strength = Molarity x Molar mass

Results and Discussion:

- 3. Molarity of KMnO₄ is _____
- 4. The Strength of $KMnO_4$ is _____M.

Precautions:

- 9. Clean all the apparatus with distilled water before starting the experiment and then rise with the solution to be taken in them.
- 10. Rinse the pipette and burette before use.
- 11. Potassium permanganate is dark in colour, so always read the upper meniscus.
- 12. Use dilute sulfuric acid for acidifying the potassium permanganate.
- 13. Take accurate readings once it reaches the end point and don't go with average readings.
- 14. Use antiparallex card or autoparallex card while taking the burette readings.
- 15. Do not use rubber cork burette as it is can be attacked by $KMnO_4$.
- 16. The strength of the unknown solution should be taken upto two decimal places only.

Aim: To analyse the given salt for acidic and basic radicals.

Experiment	Observations	Inference
-	Observations	Interence
 Physical examination : (a) Noted the colour of the given salt. 	White	Cu^{2+} , Fe^{2+} , Fe^{3+} , Ni^{2+} , Mn^{2+} , Co^{2+} absent.
(b) Noted the smell of the salt.	No specific odour	NH_4^+ , S^{2-} and CH_3COO^- may be absent.
2. Dry heating test		
Heated a pinch of the salt in a dry test tube and noted the fol- lowing observations :		
(a) Gas evolved	A reddish brown gas evolved which turned freshly prepared FeSO ₄ solution black.	NO_3^{-} may be present.
(b) Sublimation	No sublimate formed.	Ammonium halides, alu- minium chloride, iodide may be absent.
(c) Decrepitation	No crackling sound observed.	Lead nitrate, barium nitrate, sodium chloride, potassium chloride and potassium iodide may be absent.
(d) Fusion	Salt does not fuse.	Alkali (sodium, potassium) salts may be absent.
(e) Colour of the residue	White	Zn^{2+} , Pb^{2+} may be absent.

Experiment	Observations	Inference
5. Flame test Prepared a paste of the salt in conc. HCl and performed flame test.	Persistent grassy green flame on prolonged heating.	Ba ²⁺ present.
 6. Borax bead test Did not perform this test since the given salt was white. 7. Dil. sulphuric acid test 	_	Cu^{2+} , Ni^{2+} , Fe^{3+} , Mn^{2+} , Co^{2+} may be absent.
Treated a pinch of the salt with dil. H_2SO_4 and warmed.	No gas evolved.	CO $^{2-}$, S ²⁻ , NO -, SO $^{2-}$ may $_{2}$ $_{3}$ be absent.
8. KMnO ₄ test To a pinch of the salt added dil. H_2SO_4 warm and then a drop of KMnO ₄ solution.	Pink colour of KMnO4 was not discharged.	Cl ⁻ , Br ⁻ , I ⁻ , C O ^{2–} , Fe ²⁺ may 2 4
9. Conc. sulphuric acid test		be absent.
Heated a pinch of the salt with conc. sulphuric acid and added to it a paper pellet.	A reddish brown gas evolved which turned FeSO4 solution black.	NO_3^{-} may be present.
10. Confirmatory test for nitrate		
(a) Copper chips test. Heated a pinch of the salt with conc. sulphuric acid and a few copper chips.	Reddish brown gas evolved.	NO_3^{-} confirmed.
(b) Ring test. To 2–3 ml of the salt solution, added freshly pre- pared FeSO4 solution. Now added conc. sulphuric acid along the sides of the test tube.	A dark brown ring formed at the junction of the two liquids.	NO_3^{-} confirmed.
11. Heated a pinch of salt with conc. NaOH solution	No ammonia gas evolved.	
12. Preparation of Original Solution (O.S.)		NH4 ⁺ absent.
Shook a pinch of the salt with water.	Solution obtained	
13. To a part of the O.S. added $1-2$ mls of dilute hydrochloric acid.	No ppt. formed.	Labelled it as Original Solu tion (O.S.) Group I absent. (Pb ²⁺ absent)
14. Through a part of the above solution, passed H_2S gas.	No ppt. formed.	Group II absent
15. To the remaining solution, added a pinch of solid ammonium chloride. Boiled the solution, cooled it and added excess of am- monium hydroxide solution.	No ppt. formed.	(Pb ²⁺ , Cu ²⁺ , As ³⁺ , absent) Group III absent. (Fe ³⁺ , Al ³⁺ absent)

Experiment	Observations	Inference
16. Through a part of this solution, passed H_2S gas.	No ppt. formed.	Group IV absent. (Zn ²⁺ , Mn ²⁺ , Ni ²⁺ , Co ²⁺ , absent)
17. To the remaining ammonical solution added ammonium carbonate solution.	White ppt. formed.	Group V present. (Ca^{2+} , Ba^{2+} , Sr^{2+} may be present)
18. Confirmatory test for		
Barium		
Filtered the above white ppt. Dissolved the ppt. in hot dilute acetic acid.	Yellow ppt.	Ba ²⁺ confirmed.
(a) Pot. chromate test. To one part of the above solution, added a few drops of pot. chromate solution.(b) Flame test. Performed flame	Persistent grassy green flame on prolonged heating.	Ba ²⁺ confirmed.
test with the salt.		

Result. Acid radical: NO_3^-

Basic radical: Ba²⁺.

Experiment- 5



To analyse the given salt for acidic and basic radicals.

Experiment	Observations	Inference
1. Physical examination		
(a) Noted the colour of the given salt.	White	Cu^{2+} , Fe ³⁺ , Ni ²⁺ , Mn ²⁺ , Co ²⁺ absent.
(b) Noted the smell of the salt.	No specific odour	NH_4^+ , S^{2-} and CH_3COO^- may be absent.
2. Dry heating test Heated a pinch of the salt in a dry test tube and noted the following :		
(a) Gas evolved	A colourless, odourless gas evolved which turned lime wa- ter milky.	CO_3^{2-} may be present.
(b) Sublimation	No sublimate formed.	Ammonium halides, iodidemay be absent.
(c) Decrepitation	No crackling sound observed.	Lead nitrate, barium nitrate, sodium chloride, potassium chloride and potassium iodide may be absent.
(d) Colour of the residue	Yellow when hot and white when cold.	Zn^{2+} may be present.
3. Flame test	Green flashes seen with naked	
Prepared a paste of the salt in conc. HCl and performed flame test.	eye.	Zn ²⁺ may be present.
4. Borax bead test	_	
Did not perform this test since the given salt was white.		Cu^{2+} , Ni ²⁺ , Fe ²⁺ , Fe ³⁺ , Mn ²⁺ , Co ²⁺ may be absent.
5. Dil. Sulphuric acid test Treated a pinch of the salt with dil. H_2SO_4 and warmed.	Colourless, odourless gas evolved with brisk efferves- cence, turned lime water milky.	CO_3^{2-} present
Shook a pinch of salt with water taken in test tube.	Salt did not dissolve.	Insoluble CO_3^{2-} indicated.
6. KMnO4 test	Pink colour of KMnO4 was not	
To a pinch of the salt added dilute H_2SO_4 warm and then a drop of KMnO ₄ solution.	discharged.	Cl ⁻ , Br ⁻ , I ⁻ , Fe ²⁺ , C ₂ O_4^{2-} are absent.
7. Conc. Sulphuric acid test		
Did not perform this test because the salt reacted with dil. H_2SO_4 .	_	Cl ⁻ , Br ⁻ , l ⁻ , NO ⁻ , CH COO ⁻ , $_{3}^{3}$ C ₂ O ₄ ²⁻ are absent.
8. Confirmatory tests for carbonate		
(a) Shook a pinch of the salt with water.	Salt did not dissolve.	
(b) To the salt added dil. HCl.	Brisk effervescence with evolu- tion of colourless, odourless gas which turned lime water milky.	Insoluble carbonate indi- cated. Insoluble carbonate con- firmed
11. Heated a pinch of salt with conc. NaOH solution	No ammonia gas evolved.	firmed. NH4 ⁺ absent.

Experiment	Observations	Inference
12. Preparation of Original solution (O.S.)		
(a) Shook a pinch of the salt with water.	Insoluble	Labelled it as O.S.
(b) Shook a pinch of the salt in dil. HCl.	Clear solution obtained.	
13. As the O.S. is prepared in dil. HCl.		Group I absent. (Pb ²⁺ absent)
14. Through a part of O.S. passed H ₂ S gas.	No ppt. formed.	Group II absent (Pb ²⁺ , Hg ²⁺ , Cu ²⁺ , As ³⁺ absent).
15. To the remaining solution, added a pinch of solid ammonium chloride. Boiled the solution, cooled it and added excess of am- monium hydroxide solution.	No ppt. formed	Group III absent. (Fe ³⁺ , Al ³⁺ absent).
16. Through a part of this solution, passed H_2S gas.	Dull white ppt. formed.	Group IV present. (Zn ²⁺ present)
17. Confirmatory tests for \mathbf{Zn}^{2+} ion		
Dissolved the above dull white ppt. in dil HCl. Boiled off H ₂ S.		
Divided the solution into two parts.		
(a) To one part added NaOH solution dropwise.	White ppt. soluble in excess of NaOH.	Zn ²⁺ confirmed.
(b) To another part, added potassium ferrocyanide solution.	Bluish white ppt.	Zn ²⁺ confirmed.

Result. Acid Radical : CO_3^{2-}

Basic Radical : Zn^{2+} .

Aim:To analyse the given salt for acidic and basic radicals.

Experiment	Observations	Inference
1. Physical examination :		
(c) Noted the colour of the given salt.	White	Cu^{2+} , Fe^{2+} , Fe^{3+} , Ni^{2+} , Mn^{2+} , Co^{2+} absent.
(d) Noted the smell of the salt.	No specific odour	NH_4^+ , S^{2-} and CH_3COO^- may be absent.
2. Dry heating test		
Heated a pinch of the salt in a dry test tube and noted the fol- lowing observations :		
(f) Gas evolved	A reddish brown gas evolved which turned freshly prepared FeSO ₄ solution black.	NO_3^{-} may be present.
(g) Sublimation	No sublimate formed.	Ammonium halides, alu- minium chloride, iodide may be absent.
(h) Decrepitation	No crackling sound observed.	Lead nitrate, barium nitrate, sodium chloride, potassium chloride and potassium iodide may be absent.
(i) Fusion	Salt does not fuse.	Alkali (sodium, potassium) salts may be absent.
(j) Colour of the residue	White	Zn^{2+} , Pb^{2+} may be absent.

Experiment	Observations	Inference
11. Flame test Prepared a paste of the salt in conc. HCl and performed flame test.	Persistent grassy green flame on prolonged heating.	Ba ²⁺ present.
 12. Borax bead test Did not perform this test since the given salt was white. 12. Dil culture acid test 	—	Cu^{2+} , Ni^{2+} , Fe^{3+} , Mn^{2+} , Co^{2+} may be absent.
13. Dil. sulphuric acid test Treated a pinch of the salt with dil. H_2SO_4 and warmed.	No gas evolved.	CO $^{2-}$, S ²⁻ , NO $^{-}$, SO $^{2-}$ may $_{2}$ $_{3}$ be absent.
14. KMnO ₄ test		oc absent.
To a pinch of the salt added dil. H_2SO_4 warm and then a drop of KMnO ₄ solution.	Pink colour of KMnO ₄ was not discharged.	Cl ⁻ , Br ⁻ , I ⁻ , C O ^{2–} , Fe ²⁺ may 2 4
15. Conc. sulphuric acid test		be absent.
Heated a pinch of the salt with conc. sulphuric acid and added to it a paper pellet.	A reddish brown gas evolved which turned FeSO ₄ solution black.	NO_3^{-} may be present.
16. Confirmatory test for nitrate		
(c) Copper chips test. Heated a pinch of the salt with conc. sulphuric acid and a few copper chips.	Reddish brown gas evolved.	NO_3^- confirmed.
(d) Ring test. To 2–3 ml of the salt solution, added freshly pre- pared FeSO ₄ solution. Now added conc. sulphuric acid along the sides of the test tube.	A dark brown ring formed at the junction of the two liquids.	NO_3^{-} confirmed.
16. Heated a pinch of salt with conc. NaOH solution	No ammonia gas evolved.	
17. Preparation of Original Solution (O.S.)		NH4 ⁺ absent.
Shook a pinch of the salt with water.	Solution obtained	
18. Confirmatory test for Barium	ppt. formed.	Labelled it as Original Solu tion (O.S.) Group I absent. (Pb ²⁺ present)
Add KI in original Solution		
	Yellow ppt formed	
		Pb ²⁺ conformed

Result. Acid radical: NO_3^-

Basic radical: Pb²⁺.

Experiment- 7

To analyse the given salt for acidic and basic radicals.

Experiment	Observations	Inference
3. Physical examination		
(a) Noted the colour of the given salt.	White	Cu^{2+} , Fe^{3+} , Ni^{2+} , Mn^{2+} , Co^{2+} absent.
(b) Noted the smell of the salt.	No specific odour	NH_4^+ , S^{2-} and CH_3COO^- may be absent.
 4. Dry heating test Heated a pinch of the salt in a dry test tube and noted the following : (a) Gas evolved 	A colourless, odourless gas evolved	Cl⁻ may be present.
(b) Sublimation	No sublimate formed.	Ammonium halides, iodidemay be absent.
(c) Decrepitation	No crackling sound observed.	Lead nitrate, barium nitrate, sodium chloride, potassium chloride and potassium iodide may be absent.
(d) Colour of the residue	Yellow when hot and white when cold.	Zn^{2+} may be present.
9. Flame test		
Prepared a paste of the salt in conc. HCl and performed flame test.	White Flame observed with nakedeye.	Pb ²⁺ may be present.
10. Borax bead test		
Did not perform this test since the given salt was white.		Cu^{2+} , Ni^{2+} , Fe^{2+} , Fe^{3+} , Mn^{2+} , Co^{2+} may be absent.
11. Dil. Sulphuric acid test	Colourless, odourless gas	
Treated a pinch of the salt with dil. H_2SO_4 and warmed.	evolved	Cl [−] present
12. Conc. Sulphuric acid test		
Did not perform this test because the salt reacted with dil. H_2SO_4 .		Cl⁻, Br⁻, I⁻, NO ⁻, CH COO⁻, 3 3
13. Confirmatory tests for carbonate	—	$C_2O_4^{2-}$ are absent.
1. Silver nitrate test		
Acidify a portion of aqueous solution (or sodium carbonate extract) with dil. HNO ₃ . Boil for some time, cool and add silver nitratesolution.	A white ppt. is formed which is soluble in ammonium hydroxide.	Cl [−] is Conformed
2. Manganese dioxide test Heat a pinch of the salt with a small quantity of manganese dioxide and	Evolution of greenish yellow gas having a pungent irritating smell. It turns moist starch-	Cl [−] is Conformed

conc. H_2SO_4 .	iodide paper blue.	
11. Heated a pinch of saltwith conc. NaOH solution	No ammonia gas evolved.	
		NH ₄ ⁺ absent

Experiment	Observations	Inference
12. Preparation of Original solution (O.S.)		
(a) Shook a pinch of the salt with water.	Insoluble	Labelled it as O.S.
(b) Shook a pinch of the salt in dil. HCl.	Clear solution obtained.	
13. As the O.S. is prepared in dil. HCl.	ppt. formed.	Group I present. (Pb ²⁺ present)
17. Confirmatory tests for Pb²⁺ ion		
Add KI in original Solution	Yellow ppt formed	Pb ²⁺ conformed

Result. Acid Radical : Cl^-

Basic Radical : Pb^{2+} .

Experiment- 8

Aim To analyses the given salt for one anion and one cation present in it.

SI. No.	Experiment	Observation	Inference
1.	Noted the colour of the given salt.	White	Cu^{2+} , Fe ²⁺ , Ni ²⁺ ,Co ²⁺ , Mn ²⁺ are absent.
2.	Noted the smell of the salt.	No specific smell.	S ^{2–} , SO ^{2–} CH ₃ COO [–]
			may be absent.
3.	Heated 0.5 g of the salt in a dry test tube and noted the colour of the gas evolved and change in the colour of the residue on heating and cooling.	 (i) No gas was evolved. (ii) No particular change in colour of the residue is observed when heated and when cooled. 	(i) CO_3^{2-} may be present, NO_3^{-} , NO_2^{-} , Br ⁻ may be absent. (ii) Zn ²⁺ may be absent.
4.	Prepared a paste of the salt with conc. HCl and performed the flame test.	No distinct colour of the flame seen.	Ca^{2+} , Sr^{2+} , $Ba^{2+} Cu^{2+}$ may be absent.
5.	Borax bead test was not performed as the salt was white in colour.	_	_
6.	Treated 0.1 g of salt with 1 mLdil. H_2SO_4 and warmed.	No effervescence and evolution of vapours.	CO_{3}^{2-} , SO_{3}^{2-} , S^{2-} , NO_{2}^{-} , $CH_{3}COO^{-}$ absent.
7.	Heated 0.1 g of salt with 1 mLconc. H_2SO_4 .	No gas evolved.	Cl ⁻ , Br ⁻ , I ⁻ , NO ₃ ⁻ , C O $_{2}^{-}$ are absent.
8.	Acidified 1 mL of aqueous salt solution with conc. HNO_3 . Warmed the contents and ythere added tion 5 drops of ammonium	No yellow precipitate	PO_4^{3-} absent.

9.	Acidified water extract of the salt with dil. HCl and then added $2mL$ of $BaCl_2$ solution.	A white ppt. is obtained which is insoluble in conc. HNO_3 and conc. HCl.	SO_4^{2-} present.
10.	Heated 0.1 g of salt with 2 mLNaOH solution.	Ammonia gas is not evolved.	$\operatorname{NH}_{4}^{+}$ absent.
11.	Attempted to prepare original solution of the salt by dissolving 1g of it in 20 mL water.	Clear solution formed	Water soluble salt is present.
12.	To a small part of the above salt solution added 2 mL of dil. HCl.	No white precipitate formed.	Group–I absent.
13.	Passed H_2S gas through one portion of the solution of step 12.	No precipitate formed.	Group–II absent.
14.	Since salt is white, heating with conc. HNO_3 is not required. Added about 0.2 g of solid ammonium chloride and then added excess of ammonium hydroxide to the solution of step 12.	No precipitate formed.	Group–III absent.
15.	Passed H_2S gas through the above solution.	No precipitate formed.	Group-IV absent.
16.	Added excess of ammonium hydroxide solution to the original solution and then added 0.5 g of ammonium carbonate.	No precipitate formed.	Group–V absent.
17.	To the original solution of salt added ammonium hydroxide solution, followed by disodium hydrogen phosphate solution. Heated and scratched the sides of the test tu.	White precipitate.	Mg ²⁺ confirmed.

Result

The given salt contains:

Result. Acid Radical : SO_4^{2-}

Basic Radical : Mg^{2+} .

Experiment- 9

SI. No.	Experiment	Observation	Inference
1.	Noted the colour of the given salt.	White	Cu^{2+} , Fe^{2+} , Ni^{2+} , Co^{2+} , Mn^{2+} are absent.
2.	Noted the smell of the salt.	No specific smell.	S^{2-} , SO_{3}^{2-} CH ₃ COO ⁻ may be absent.
3.	Heated 0.5 g of the salt in a dry test tube and noted the colour of the gas evolved and change in the colour of the residue on heating and cooling.	 (i) No gas was evolved. (ii) No particular change in colour of the residue is observed when heated and when cooled. 	(iii) CO_3^{2-} may be present, NO_3^{-} , NO_2^{-} , Br^{-} may be absent. (iv) Zn^{2+} may be absent.
4.	Prepared a paste of the salt with conc. HCl and performed the flame test.	Green colour of the flame seen.	Ba ²⁺ may be present.
5.	Borax bead test was not performed as the salt was white in colour.		_
6.	Treated 0.1 g of salt with 1 mLdil. H_2SO_4 and warmed.	No effervescence and evolution of vapours.	$CO_3^{2-}, SO_3^{2-}, S^{2-}, NO_2^{-}, CH_3 COO^{-}$ absent.
7.	Heated 0.1 g of salt with 1 mLconc. H_2SO_4 .	No gas evolved.	Cl ⁻ , Br ⁻ , I ⁻ , NO ₃ ⁻ , C O ₄ ⁻ are absent.
8.	Acidified 1mL of aqueous salt solution with conc. HNO ₃ . Warmed the contents and ythere edded to for the solution of ammonium	No yellow precipitate	PO_4^{3-} absent.

9.	Acidified water extract of the salt with dil. HCl and then added $2mL$ of $BaCl_2$ solution.	A white ppt. is obtained which is insoluble in conc. HNO_3 and conc. HCl .	SO_4^{2-} present.
10.	Heated 0.1 g of salt with 2 mL NaOH solution.	Ammonia gas is not evolved.	\mathbf{NH}_{4}^{+} absent.
11.	Attempted to prepare original solution of the salt by dissolving 1g of it in 20 mL water.	Clear solution formed	Water soluble salt is present.
12.	To a small part of the above salt solution added 2 mL of dil. HCl.	No white precipitate formed.	Group–I absent.
13.	Passed H_2S gas through one portion of the solution of step 12.	No precipitate formed.	Group–II absent.
14.	Since salt is white, heating with conc. HNO_3 is not required. Added about 0.2 g of solid ammonium chloride and then added excess of ammonium hydroxide to the solution of step 12.	No precipitate formed.	Group–III absent.
15.	Passed H_2S gas through the above solution.	No precipitate formed.	Group–IV absent.
16.	Added excess of ammonium hydroxide solution to the original solution and then added 0.5 g of ammonium carbonate.	No precipitate formed.	Group–V present.
17.	Confirmatory test for Barium		
	(a) Pot. chromate test. To one part of the above solution, added a few drops of pot. chromate solution.	Yellow ppt.	Ba ²⁺ conformed
	(b) Flame test. Performed flame test with the salt.	Persistent grassy green flameon prolonged heating.	Ba ²⁺ conformed

Result

The given salt contains:

Acid Radical : SO₄^{2–}

Basic Radical : Ba^{2+} .

Aim:To analyse the given salt for acidic and basic radicals.

Experiment	Observations	Inference
1. Physical examination :		
(e) Noted the colour of the given salt.	White	Cu^{2+} , Fe^{2+} , Fe^{3+} , Ni^{2+} , Mn^{2+} , Co^{2+} absent.
(f) Noted the smell of the salt.	No specific odour	NH_4^+ , S^{2-} and CH_3COO^- may be absent.
2. Dry heating test		
Heated a pinch of the salt in a dry test tube and noted the fol- lowing observations :		
(k) Gas evolved	A reddish brown gas evolved which turned freshly prepared FeSO ₄ solution black.	NO_3^{-} may be present.
(l) Sublimation	No sublimate formed.	Ammonium halides, alu- minium chloride, iodide may be absent.
(m)Decrepitation	No crackling sound observed.	Lead nitrate, barium nitrate, sodium chloride, potassium chloride and potassium iodide may be absent.
(n) Fusion	Salt does not fuse.	Alkali (sodium, potassium) salts may be absent.
(0) Colour of the residue	White	Zn^{2+} , Pb^{2+} may be absent.

Experiment	Observations	Inference
17. Flame test Prepared a paste of the salt in conc. HCl and performed flame	Red flame on prolonged heating.	Sr ²⁺ present.
test. 18. Borax bead test Did not perform this test since	_	Cu ²⁺ , Ni ²⁺ , Fe ³⁺ , Mn ²⁺ , Co ²⁺
the given salt was white.19. Dil. sulphuric acid test		may be absent.
Treated a pinch of the saltwith dil. H_2SO_4 and warmed. 20. KMnO4 test	No gas evolved.	CO $^{2-}$, S ²⁻ , NO $^{-}$, SO $^{2-}$ may $_{2}$ $_{3}$ be absent.
To a pinch of the salt added dil. H_2SO_4 warm and then a drop of KMnO4 solution.	Pink colour of KMnO4 was not discharged.	Cl ⁻ , Br ⁻ , I ⁻ , C O ^{2–} , Fe ²⁺ may ² ⁴
21. Conc. sulphuric acid test		be absent.
Heated a pinch of the salt with conc. sulphuric acid and added to it a paper pellet.	A reddish brown gas evolved which turned FeSO4 solution black.	NO_3^{-} may be present.
22. Confirmatory test for nitrate		
(e) Copper chips test. Heated a pinch of the salt with conc. sulphuric acid and a few copper chips.	Reddish brown gas evolved.	NO_3^{-} confirmed.
(f) Ring test. To 2–3 ml of the salt solution, added freshly pre- pared FeSO ₄ solution. Now added conc. sulphuric acid along the sides of the test tube.	A dark brown ring formed at the junction of the two liquids.	NO_3^{-} confirmed.
18. Heated a pinch of salt with conc. NaOH solution	No ammonia gas evolved.	
19.Preparation of Original Solution (O.S.)		NH4 ⁺ absent.
Shook a pinch of the salt with water.	Solution obtained	
20. To a part of the O.S. added $1-2$ mls of dilute hydrochloric acid.	No ppt. formed.	Labelled it as Original Solu tion (O.S.) Group I absent. (Pb ²⁺ absent)
21. Through a part of the above solution, passed H_2S gas.	No ppt. formed.	Group II abcont
22. To the remaining solution, added a pinch of solid ammonium chloride. Boiled the solution, cooled it and added excess of am- monium hydroxide solution.	No ppt. formed.	Group II absent (Pb ²⁺ , Cu ²⁺ , As ³⁺ , absent) Group III absent. (Fe ³⁺ , Al ³⁺ absent)

Experiment	Observations	Inference
16. Through a part of this so- lution, passed H ₂ S gas.	No ppt. formed.	Group IV absent. (Zn ²⁺ , Mn ²⁺ , Ni ²⁺ , Co ²⁺ , absent)
17. To the remaining ammonical solution added ammonium carbonate solution.	White ppt. formed.	Group V present. (Ca^{2+} , Ba^{2+} , Sr^{2+} may be present)
18. Confirmatory test for		
1. Amm. sulphate test To the second part of the solu- tion, add 1 ml of amm. sulphate solution and warm.	White ppt.	Sr ²⁺ confirmed.
2. Flame test Perform the flame test with the original salt.	Crimson red flame.	Sr ²⁺ confirmed.

Result. Acid radical: NO_3^-

Basic radical: Sr²⁺.

Aim: To separate the coloured components present in the mixture of red and blue inks by ascending paper chromatography and find their R_tvalues.

APPARATUS

Gas jar, glass rod, filter paper strip (Whatman No. 1 filter paper), jar cover and a fine capillary tube.

REQUIREMENT

A mixture of red and blue inks, alcohol and distilled water.

PROCEDURE

1. Take a Whatman filter paper strip $(20 \times 2 \text{ cm})$ and draw a line with pencil above 3 cm from one end. Draw another line lengthwise from the centre of the paper as shown in Fig. 6.4.

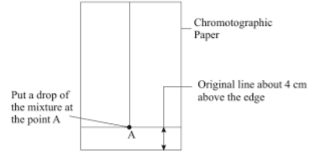


Fig. 6.4. Spotting of the mixture.

2. With the help of fine capillary tube, put a drop of the mixture of red and blue inks at the point P. Let it dry in air. Put another drop on the same spot and dry again. Repeat 2–3 times, so that the spot is rich in the mixture.

3. Suspend the filter paper vertically in a gas jar containing the solvent (eluent) with the help of a glass rod in such a way that the pencil line (and the spot) remains about 2 cm above the solvent level (50% alcohol + distilled water).

4. Cover the jar and keep it undisturbed. Notice the rising solvent along with the mix- ture of red and blue inks. After the solvent has risen about 15 cm you will notice two different spots of blue and red colours on the filter paper.

5. Take the filter paper out of the jar and mark the distance that the solvent has risen on the paper with a pencil. This is called the solvent front.

6. Dry the paper. Put pencil marks in the centre of the blue and red spots.

7. Measure the distance of the two spots from the original line and the distance of the solvent from the original line.

8. Calculate the R_f values of the blue and red inks by using the formula :

R = D is tance travelled by the blue or red ink from the point of application Distance travelled by the solvent from the original line

Substance	Distance travelled by different components	istance travelled by solvent	R _f Value
Red ink + Blue ink	A cm (Red Ink)	X cm	A/X
	B cm (Blue Ink)	X cm	B/X

OBSERVATIONS AND CALCULATIONS

PRECAUTIONS

- 1. Use good quality pencil for drawing the reference line so that the mark does not dissolve in the solvent in which the chromatography is carried out.
- 2. Always make use of a fine capillary tube.
- 3. Keep the jar undisturbed and covered during the experiment.
- 4. A spot should be small and rich in mixture.
- 5. Allow the spot to dry before putting the strip in the jar.

- 6. Keep the strip erect. Do not let it to be curled.
- 7. Do not allow the spot to dip in the solvent.

Aim: To separate the coloured components present in the given grass/flower by as- cending paper chromatography and determine their R_t values.

In this experiment, crush fresh flowers or grass in a mortar and extract the juice with acetone. Use this solution for making the spot. Proceed as in Expt. 6.1.

OBSERVATIONS AND CALCULATIONS

Colour of the spot	Distance travelled by the spot from the original line	Distance travelled by the solvent from the original line	R Value
Green (Chlorophyll)	A cm	X cm	A/X
Yellow Xanthophyll)	B cm	X cm	B/X
Red (Carotene)	C cm	X cm	C/X

To identify the functional group present in the given organic compound.

Experiment	Observations	Inference
 Test for unsaturation Dissolved 0.2 ml of organic com- pound in 2 ml CCl Then added bromine-water dropwise. Test for carboxylic group Added a pinch of NaHCO to 0.2 ml of 	Brown colour of bromine not discharged. No effervescence. No green or violet colour obtained. No effervescence.	Inference No unsaturation is present. Carboxylic group is absent. Phenolic group is absent. Alcoholic group is absent. Carbonyl group is present. May be an aldehyde or a ketone. Aldehyde is present. Amino group absent.

RESULT : - Aldehyde group (--CHO).

To identify the functional group present in the given organic compound.

Experiment	Observations	Inference
1. Test for unsaturation Dissolved 0.2 ml of organic com- pound in 2 ml CCl ₄ . Then added bromine-water dropwise.	Brown colour of bromine not discharged.	No unsaturation is present.
2. Test for carboxylic group Added a pinch of NaHCO ₃ to 0.2 ml of organic compound in a test tube.	No effervescence.	Carboxylic group is absent.
3. Test for phenolic group Added 0.2 ml of organic compound to 2–3 ml neutral FeCl ₃ solution in a test tube.	No green or violet colour obtained.	Phenolic group is absent.
4. Test for alcoholic group Added a small piece of sodium to 1 ml of the given liquid in a dry test tube.	No effervescence.	Alcoholic group is absent.
5.Test for carbonyl group Shook 0.2 ml of organic compound with 2–3 ml of 2, 3-dinitrophenyl hydrazine in a test tube.	Orange-yellow ppt. formed.	Carbonyl group is present. May be an aldehyde or a ketone.
6. Test for aldehydic group Warmed 1 ml of organic compound with 1 ml of Tollen's reagent in a test tube over a water bath.	Silver mirror formed on inner side of the test tube.	Aldehyde is present.
7.Test for amine group To a small amount of organic liq- uid in test tube, added 1 ml conc. of HCl and a few drops of CHCl ₃ . Then, added 2 ml of alc. KOH so- lution and warmed test tube.	No offensive smelling gas evolved.	Amino group absent.
8. TESTS FOR KETONES		
Place 0.5 ml of the given liquid (or 0.5 g of solid) in a clean test tube and add about 0.1 g of finely powdered m-dinitrobenzene. Now add about 1 ml of dilute sodium hydroxide solution and shake.	Appearance of violet colour which slowly fades	confirms ketonic group.

RESULT : - Ketone (-CO-)

To identify the functional group present in the given organic compound.

Experiment	Observations	Inference
4. Test for unsaturation Dissolved 0.2 ml of organic com- pound in 2 ml CCl ₄ . Then added bromine-water dropwise.	Brown colour of bromine not discharged.	No unsaturation is present.
 5. Test for carboxylic group Added a pinch of NaHCO₃ to 0.2 ml of organic compound in a test tube. 	No effervescence.	Carboxylic group is absent.
6. Test for phenolic group Added 0.2 ml of organic compound to 2–3 ml neutral FeCl ₃ solution in a test tube.	No green or violet colour obtained.	Phenolic group is absent.
6. Test for alcoholic group Added a small piece of sodium to 1 ml of the given liquid in a dry test tube.	Brisk effervescence.	Alcoholic group is present. Carbonyl group is present. May
7. Test for carbonyl group Shook 0.2 ml of organic compound with 2–3 ml of 2, 3-dinitrophenyl hydrazine in a test tube.	Orange-yellow ppt. formed.	be an aldehyde or a ketone.
8. Test for aldehydic group Warmed 1 ml of organic compound with 1 ml of Tollen's reagent in a test tube over a water bath.	No observation	Aldehyde is absent.
9. Test for amine group To a small amount of organic liq- uid in test tube, added 1 ml conc. of HCl and a few drops of CHCl ₃ . Then, added 2 ml of alc. KOH so- lution and warmed test tube.	No offensive smelling gas evolved.	Amino group absent.

RESULT : - Alcohol (-OH)

To identify the functional group present in the given organic compound.

not discharged. Brisk effervescence.	No unsaturation is present.
No green or violet colour	Carboxylic group is present. Phenolic group is absent.
No effervescence.	Alcoholic group is absent. Carbonyl group is present. May
Orange-yellow ppt. formed. No observation	be an aldehyde or a ketone.
No offensive smelling gas evolved.	Aldehyde is absent.
	Amino group absent.
	obtained. No effervescence. Drange-yellow ppt. formed. No observation No offensive smelling gas

RESULT : - Carboxylic acid (-COOH)