- Chromate CrO<sub>4</sub><sup>--</sup>. This can be tested by following methods:
- (i) To an aqueous solution, add barium chloride solution. A pale yellow precipitate insoluble in (ii) To an aqueous solution or solution in acetic acid, add silver nitrate solution. A brownish red acetic acid indicates the presence of chromate.
- (iii) Treat its acidic solution with  $H_2O_2$ . This gives deep blue solution of chromium pentoxide precipitate of silver chromate is obtained.

which fades quickly on standing.

(ii) In acidic solution, pass H<sub>2</sub>S gas. This turns the solution green. (v) Acidify the solution with dilute  $H_2SO_4$  or acetic acid and add diphenyl carbazide reagent. A

deep blue colouration is produced.

- 5. Permanganate, MnO<sub>4</sub>. All permanganates are, as a rule, soluble in water imparting purple solution. Following tests can be applied to ascertain the presence of permanganate radical. (i) To an aqueous solution, add concentrated HCl and boil. Chlorine gas is evolved which turns
- (ii) To concentrated aqueous solution, add NaOH solution and warm the mixture, A green solustarch-iodide paper blue. tion is obtained with the liberation of oxygen gas. The green solution when poured into a beaker full of water and contents acidified, purple colours of permanganate is restored.
- 6. Arsenate, AsO<sub>4</sub><sup>3-</sup> (i) To a neutral solution of arsenate, add silver nitrate solution. A brownish red precipitate insoluble in acetic acid but soluble in mineral acid and in ammonia indicates the presence of arsenate.
- (ii) To an aqueous solution, add nitric acid and ammonium molybdate. Boil the mixture. A yellow crystalline precipitate of ammonium arsenomolybdate, (NH<sub>4</sub>)<sub>3</sub>AsMo<sub>12</sub>O<sub>40</sub> indicates the presence of arsenate. This precipitate is soluble in ammonia and in alkalies. The ionic equation for the above test is written as:

bove test is written as:  

$$AsO_4^{3-} + 12MoO_4^{2-} + 3NH^{4+} + 24H^+ \longrightarrow (NH_4)_3 AsMo_{12}O_{40} \downarrow + 12H_2O_{40}$$
precipitate

(iii) Acidify the solution with large amount of concentrated HCl and to it, add KI solution. This will precipitate iodine which can be tested as usual. The ionic equation is :

AsO<sub>4</sub><sup>3-</sup> + 2H<sup>+</sup> + 2I<sup>-</sup> 
$$\implies$$
 AsO<sub>3</sub><sup>3-</sup> + I<sub>2</sub>  $\downarrow$  + H<sub>2</sub>O

2.6. Tests with sodium carbonate extract

Klarge number of acid radicals can be confirmed by using sodium carbonate extract of the mixture. This is especially useful when the mixture is less soluble in dilute acids. For example, many halides and sulphates do not readily dissolve in acids and hence they may not be tested.

When mixture is heated strongly in the minimum quantity of water with excess of Na<sub>2</sub>CO<sub>3</sub>, the double decomposition takes place i.e., the partner radicals are changed. Sodium joins with anions of mixture and carbonate attaches with basic radicals of mixture. After heating, it is mixed with water where sodium salts which contain anions of mixture dissolve (as sodium salts are soluble) and carbonates of the metals remain as insolubles. In this way, we separate the anions from the cations of the mixture after filtration.

## Preparation of Sodium Carbonate Extract

Take one part of mixture and four parts of pure Na<sub>2</sub>CO<sub>3</sub> in a porcelain dish. Add 15-20 ml distilled water and heat while stirring with the glass rod. Heat it to boiling gently for about 5-10 minutes, add water to make up the loss by evoparation and filter. The filtrate is the desired sodium carbonate extract. Use 1-2 ml of this extract for each test.

## **Precautions**

- they should be tested in the residue left by action of Na<sub>2</sub>CO<sub>3</sub> or in certain instances, in separate portion of original mixture.
- 2. During neutralization of soda extract with acids, sometimes there appears ppt. though it becomes soluble in excess of acid. It is desirable to filter off such a ppt. This is because the cations forming amphoteric oxides may partially pass into soda extract (e.g. AlO<sub>2</sub><sup>-</sup>). The copper ion forms a complex compound and also partially passes in soda extract colouring it bluish. On subsequent neutralization, all these compounds are decomposed and corresponding cations are precipitated, e.g.,

$$NaAlO_2 + CH_3COOH + H_2O \longrightarrow Al(OH)_3 \downarrow + CH_3COONa$$

Similarly, copper is precipitated as basic carbonate.

3. Before adding a particular reagent to test acid radicals, extract must be made acidic because carbonate should be decomposed otherwise confusion may arise due to precipitation of the carbonates of the metals. For example, in test for sulphate, we add  $BaCl_2$  solution to acidified sodium carbonate extract. The appearance of white ppt. indicates the presence of  $SO_4^{2-}$  radical. If sufficient acid is not added prior to the addition of  $BaCl_2$ , some undecomposed sodium carbonate will remain in solution. This will give a white ppt. of  $BaCO_3$  even in absence of  $SO_4^{2-}$ . This will obviously create a confusion.

Hence in testing various radicals by the sodium carbonate extract, acidify according to the following chart:

Radicals to be tested	Acid used for neutralization	Reagent for precipitation
Chloride	dil. HNO <sub>3</sub>	AgNO <sub>3</sub>
Bromide	"	,,
Iodide	"	37
Sulphate	dil. HCl	BaCl <sub>2</sub>
Oxalate	CH <sub>3</sub> COOH	CaCl <sub>2</sub>

(4) Do neutralization very carefully, shaking the liquid thoroughly after each addition and avoid an excess of acid otherwise certain anions (e.g., S<sup>2-</sup>, NO<sub>2</sub><sup>-</sup> etc.) may also be lost.

## 2.7. Combination Tests of Acid Radicals

Generally the following combinations are given:

- 1. Sulphide, Sulphite, Sulphate and Thiosulphate
  - 2. Nitrite and Nitrate
  - 3. Nitrite and Iodide
  - 4. Nitrite and Sulphide
- 5. Nitrate in presence of Bromide and Iodide
- 6. Chloride, bromide and Iodide
- 7. Oxalate and Fluoride
- 8. Carbonate and oxalate
- 9. Carbonate in presence of Sulphite
- 10. Sulphite and Fluoride
- 11. Phosphate, Arsenate and Arsenite

All the combinations are not essential to be tested; the radicals which respond test in preliminary examination, their combinations are to be seen and carried out.

1, Sulphide, sulphite, sulphate and thiosulphate present

Systematic analysis is required for detection of S<sup>2-</sup>, S<sub>2</sub>O<sub>3</sub><sup>2-</sup>, SO<sub>3</sub><sup>2-</sup> and SO<sub>4</sub><sup>2-</sup> ions when present together, because some of these interfere in test of others. For example, if the solution contains S<sup>2-</sup>

and  $SO_3^{2-}$ , addition of acid liberates  $H_2S$  and  $SO_2$  simultaneously and the two immediately interact.

 $SO_2 + 2H_2S \longrightarrow 3S \downarrow + 2H_2O$ 

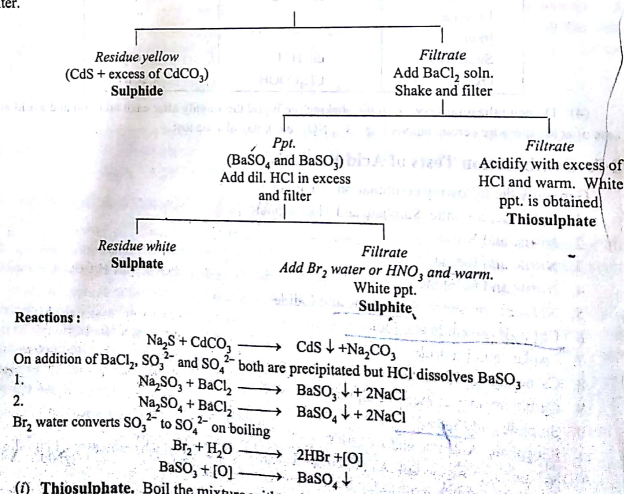
As a result of this reaction, only the gas present in excess will be detected. If this gas is SO<sub>2</sub>, addition of acid would be accompanied by the same effects (formation of  $SO_2$  and S) as would be observed if the solution contained  $S_2O_3^{2-}$ . Thiosulphate would, therefore, be wrongly found. It is obvious that a mixture of  $SO_3^{2-}$  and  $S_2O_3^{2-}$  would behave on acidification in the same way as  $S_2O_3^{2-}$  by itself

 $Na_{2}SO_{3} + 2HCl \longrightarrow 2NaCl + SO_{2} \uparrow + H_{2}O$   $Na_{2}S_{2}O_{3} + 2HCl \longrightarrow 2NaCl + SO_{2} \uparrow + S \downarrow + H_{2}O$ 

Finally in presence of S<sup>2-</sup>, the test for SO<sub>3</sub><sup>2-</sup> with acidified K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub> can not be used (H<sub>2</sub>S also reduces acidified  $K_2Cr_2O_7$  to green chromium salt). Similarly, detection of  $SO_4^{2-}$  is difficult in presence of  $S_2O_3^{2-}$  or mixture of  $S^2$  and  $SO_3^{2-}$ . In both cases, a white ppt. of sulphur is formed which being insolvable in said. being insoluble in acids, it may be mistaken for BaSO<sub>4</sub>.

The systematic analysis of mixture of  $S^{2-}$ ,  $S_2O_3^{2-}$ ,  $SO_3^{2-}$  and  $SO_4^{2-}$ , described in detail below, is based on (a) precipitation of  $S^{2-}$  by the action of CdCO<sub>3</sub>, when  $SO_3^{2-}$ ,  $SO_4^{2-}$  and  $S_2O_3^{2-}$  remain in solution; (b) precipitation of  $SO_3^{2-}$  and  $SO_4^{2-}$  by BaCl<sub>2</sub> (or SrCl<sub>2</sub>) or Sr(NO<sub>3</sub>)<sub>2</sub>, which does not precipitate  $S_2O_3^{2-}$  (a) the fact that BaSO<sub>4</sub> (as SaCO<sub>3</sub>) in the fact that BaSO<sub>4</sub> (as SaCO<sub>3</sub>) in the sact that BaSO<sub>4</sub> precipitate  $S_2O_3^{2-}$ ; (c) the fact that  $BaSO_4$  (or  $SrSO_4$ ) is almost insoluble while  $BaSO_3$  (or  $SrSO_3$ ) is soluble in dilute acids.

Take the sodium carbonate extract and add excess of freshly prepared CdCO3, shake well and filter.



(i) Thiosulphate. Boil the mixture with water and filter. Cool the filtrate and add to it a few drops of ferric chloride solution. Dark violet colour appears which fades slowly.

(a) mix + 2 m dittisory of Acid Radicals				
	Test (Procedure)	Observation		
	Add 2ml of dil H2 soy to compor mixer in a test tube.	Efferviscence with evolutions of colourless gas. (Carbonate may be present)		
	- Pass the gas into lime water	lime water turns nilty		
	-> When passfin excess -	milky colour disserpears.		
Nitrite (NO5)	Add 2ml of dil Hasoyin mix.	Reddish brown fumes in cold		
	→ Mixing dil HSOy+ fresh Starch Soln+ KI soln(fresh)	, Blue colous		
3:Sulphide		Colourless gas with smell of Lotton eggs.		
	solution, when exposed to gas evolved	Black. Confirmation by Na CO3 extract test		
Sulphite (SO <sub>3</sub> <sup>2</sup> -)	Add 2ml of dil H_Say in mixture 4 heat it	Colourless gas with Chocking smell of Sulphus		
<b>7</b> /1	dipped in Acidified dichemate solution to the gas evolved	1: 1 ter Da Dis		
PP .	but 4-5 deops of mixture with dil Hsoy in FMnDy	the Kmn Oy		
		Scanned by CamScanner		

Radical Procedure Observation white or yellowish? Add sock of Mixture in mel of dil H2 Soys heat it Thiosulphale white tue sitify appears. (5203) Confirmation By Na cos extract. Add & my Mixture in 2ml -No fumes (snells of dil H2 Soy's heat it like rinegar) - Aqueous sol of mixturet fene deaps of Neutral Fects solution , Blood red colouration ( et a Colour dissappear on addition of Acid) Chloride Add 2ml of Mixture in Conc" H. Soy in Mixture warm gently Colourless pungent gas comes out. Bringa rod dipped in Cone. NHYOH near the mouth of test tube Dense fumes Dark ged vapours → \$ 1 part mixture + 3 times of Cheonyl Chloride K2 (2207 + Com. H2504 + heats ale evolved MaoHsol" becomes then pass the vapous into dil NaOH jelloue ppt. obtained Acetate to the sol" which's is soleble in Altalihydsoxides & dil HNO3 but insoluble in chicoon e dis NH4017 Bromide Add 2ml of cone. 12.50y in Mixture & heal gently Reddish brown Vapouis. s Pall yellow PPt. sparingly soluble - taxers Normal mixture + dil HN103 + heat, filter the solution & to the in NHYOH filtrate Add Ag NO3 col

Radicals	Procedure		Observation
Iodide (I')	Add me of con nixtures mae		Violet pungent va pour evolved confirmations by Na co3 extract.
Nitsale (NOs)	Add me of come  Mixture of we  Ring test-  2ml of Naços of  Acidify with of  add V Inl of  slowly of  liquids theory  cool the test to  the stream of	extract lil Hssoy Conc. Hssoy the two thoroughly ube under	Light brown gas evolved (NOs haybe present)  A sharp brown  sing will be formed At the junction of
	then add fre feso, solo st the side of t Allowed to s-minitus.	outly from he test tense	turo layer.