Answer to problem on sodium hydroxide solution

a) The volume of 1.11 *N* hydrochloric acid solution used in titrating sodium hydroxide sample was for total alkali regardless of whether sodium carbonate is present as impurity or not; *i. e.*, the same volume of hydrochloric acid solution would be consumed if sodium carbonate was precipitated as barium carbonate (explain this fact by the related balanced chemical equations).

HCl NaOH $N_1 \times V_1 = N_2 \times V_2$ $1.11 \times 4.5 = N_2 \times 10$ $N_2 = 0.4995$ the normality of the prepared sodium hydroxide solution

b) Any solid sodium hydroxide sample taken from the original container would contain sodium carbonate as impurity by 3% w/w. Thus, you have to calculate how much sodium hydroxide was used to prepare the 10 milliliters- sample. This is calculated from the volume of hydrochloric acid used for total alkali:

1.11 *N* HCl 1 *N* HCl $N \times V = \dot{N} \times \dot{V}$ 1.11 × 4.5 = 1 × \dot{V} $\dot{V} = 4.995$ mL of 1 *N* hydrochloric acid solution would be required for total alkali (V₃)

$$wt = V_3 \times ch.factor$$

 $wt = 4.995 \times 0.04$

wt = 0.1998 g of total alkali (sodium hydroxide and the impurifying sodium carbonate) calculated as sodium hydroxide was used originally to prepare 10 mL of solution

 $\frac{3}{100} \times 0.1998 = 0.005994$ g of sodium carbonate was present in the sample used to prepare 10 mL of solution

 $wt = V_2 \times ch.factor$

 $0.005994 = V_2 \times 0.053$

 $V_2 = 0.113$ mL of 1 *N* hydrochloric acid solution was consumed by the amount of sodium carbonate present in the sample assayed

1.11 *N* HCl 1 *N* HCl $N \times V = \dot{N} \times \dot{V}$

 $1.11 \times V = 1 \times 0.113$

 $\dot{V} = 0.1 \text{ mL of } 1.11 \text{ N}$ hydrochloric acid solution would be consumed by the amount of sodium carbonate present in the sample assayed

It is important to note that the percent w/w is related to solid samples. If you were asked to calculate the percent w/v, then this is related to the prepared solution as follows:

for total alkali: $\frac{0.1998}{10} \times 100 = 1.998 \% w/v$ for sodium carbonate: $\frac{0.005994}{10} \times 100 = 0.05994 \% w/v$

Assay of Magnesium Sulfate MgSO4. 7H2O Mwt.=246.5gm/mole

- Contain not less than 99.5% MgSO₄ calculated with reference of dried substance.
- Description: colorless crystal, or white crystal powder.
- Solubility: soluble at 20°C in 1.5 parts of water, sparingly soluble in alcohol 96% i.e. practically insoluble in 96% ethanol.
- Uses: laxative agent.

Assay: Each 1ml of M/20 EDTA = 0.012325gm MgSO4.7H2O



- The technique involves titrating metal ions with a complexing agent or chelating agent (Ligand) and is commonly referred to as complexometric titration. This method represents the analytical application of a complexation reaction.
- In this type of titration we use *metal ion indicator*.
- Chemical principle:
- It is type of complexometric titration.

MgSO4 + Na2H2EDTA (MgEDTA) + 2H + Na2SO4

- Magnesium ions form relatively weak complex with (EDTA ;Ethylenediaminetetra-acetic acid).
- The stability of metal complex is pH dependent. Lower the pH of the solution, lesser would be the stability of complex (because more H⁺ ions are available to compete with the metal ions for ligand)
- $\, \ast \,$ But at $\,$ pH 10 the reaction is rapid and quantitative .

- ♦ NH3- NH4Cl buffer is used to maintain this pH.
- EDTA which is polyamine carboxylic acid is one of the most frequently used chelating agent:



EDTA form soluble, stable 1:1 chelates with metal ion.



- Secause of the low water solubility of the free acid form of EDTA, the disodium salt (Disodium EDTA) is used in preparation of std. sol.
- Indicator: Metal Indicator
- Metal indicators are used to detect the end point in complexometric titration.
- They are dyes which produce one color in the presence of metal ions and a different color when they are absent.
- The color changes are the result of displacement of the H⁺ ions by the metal.
- The metal indicator form 1:1 complex (MI) which must be weaker than the Metal-EDTA complex.

M + I _ MI

MI + EDTA - M-EDTA + I pink blue

● Chemical factor: it forms 1:1 chelate so; 1Mwt. MgSO4.7H2O = 1Mwt. EDTA 0.012325gm MgSO4.7H2O = 1ml of M/20 EDTA ● Chemical principle:

 At the onset of titration, the buffered reaction media contains the MI complex and free M:

M + I — MI

 When the titrant (EDTA) is added to the system, it will progressively chelate the free metal ions in the solution, and then a competitive reaction will take place, 1:1metal ions and EDTA, since the MI complex is weaker than M_EDTA. Finally, at the equivalent point, EDTA removes the last traces of metal ions from its colored complex with ind.(MI) to free metal ind.(I) of another color.

- Mordant black indicator: it has blue color at pH10 and most of its complexes are reddish.
- <u>Note:</u> In pH less than 6.3 and more than 11.5, the indicator and its complexes are reddish,
 - <u>so</u> it is necessary to carry out the titration in the presence of *buffer at pH 10*.



• Calculation:

V ***** ch. factor= gm wt. of MgSO4.7H₂O

wt **#** 100/10= %w/v of unknown.

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wt **#** 100/10= %w/v of unknown.

ASSAY OF MAGNESIUM HYDROXIDE IN ANTACID SUSPENSIONS (MAALOX)



- Maalox suspension contains two active ingredients, aluminium hydroxide and magnesium hydroxide. These are both medicines called antacids. They are used to reduce the acidity of the stomach's juices.
- Magnesium ion is complexed with EDTA. The solution is titrated with standard EDTA solution to the first permanent appearance of violet color.
- Mordant black is used as the indicator.



• Procedure:

- *note*: We must first dilute the Maalox suspension before assay:
- 1. Transfer 10mL unknown to 250-mL conical flask.
- 2. Add 5mL of pH-10 buffer.
- 3. Add 10ml D.W.
- 4. Add about 0.1gm of Mordant Black indicator.
- 5. Titrate with M/40 EDTA solution until the color changes from pink to violet color.

• Calculation:

• Molecular Weight of $Mg(OH)_2 = 58.3$

o mg of Mg(OH)2 Found = M (EDTA) x V (EDTA) in mL x Mwt. of Mg(OH)2 • <u>Complexometric titrations</u> use reagents containing ligands that form complexes with metal ions. The indicator is a substance that forms a colored complex with the ions, but this complex must be weaker than that formed with the reagent. The metal ion complexes preferentially with the reagent, until at the endpoint no metal-indicator complex remains, and its colour disappears.

An example is the determination of calcium and/or magnesium using EDTA (ethylenediaminetetra-acetic acid).

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ASSAY OF BOBAX

Borax <u>or</u> Sod. borate <u>or</u> Sod. Tetraborate Chemical formula Na₂B₄O₇.10H₂O Mwt.= 381.4 gm

- **Physical properties**: Borax is sod.tetraborate decahydrated, should be kept in well closed container, soluble in 20 parts of water.
- Uses; antiseptic and astringent in mouth paints.
- **Procedure**: 10mL of unknown

10mL of distilled water

2drops of methyl red indicator

Titrate with <u>N/2 HCl</u> solution, the color of sol. changes from yellow → pink-orange(1st end point).

Record the volume of N/2 HCl consumed.
Boil and cool the solution,

Add 4gm mannitol and 2drops of phenolphthalein ind. The color of sol. is pink

Now titrate with <u>N/1 NaOH</u> until the color of sol. changes from pink \rightarrow yellow \rightarrow pink(2nd end point).

Record the volume of N/1 NaOH consumed.



2nd titration

<u>N/1 NaOH</u> solution

Boil , cool and add 4gm mannitol and 2drops of ph.ph ind. The buret contains the titrant. This flask contains the solution to be titrated and the indicator.

- Chemical principle:
- Borax is a salt of weak acid with strong base.

 $Na_2B_4O_7.10H_2O + 2HCI \longrightarrow 2NaCI + 4H_3BO_3 + 5H_2O$

Soric acid(H₃BO₃) will be liberated, it is very weak acid.

Soli the solution to remove CO₂.

- Add mannitol because boric acid is weak acid to convert it into strong complex acid which is strong enough to be titrated directly with NaOH.
- Mannitol should be used in high conc. to prevent the hydrolysis of the complex.

4H₃BO₃ + 4NaOH — Mannitol > 4NaBO₂ + 8H₂O

sod. metaborate

Mannitol should be used in high conc. to prevent the hydrolysis of this complex.



mannityl boric acid

- Chemical factor: <u>home work</u>
- \geq For borax with 0.5N HCl (0.09535 gm)
- For borax with 1N NaOH (0.09535 gm)
- ➢ For sod. carbonate with 0.5N HCl (0.0265gm)
- ➢ For boric acid with 1N NaOH (0.06184gm)

Mwt. Boric acid=61.84 gm/mol Mwt. Na₂CO₃= 106gm/mol • Calculations:

we must first correct the volume of HCl & NaOH

1. If we have **<u>pure borax</u>**, the vol. of *N*/2 HCl must be equal to the vol. of 1*N* NaOH.

 $V'_{of N/2}$ HCI = $V'_{of 1N NaOH}$ (the differences up to 0.3 mL) take the average of the two volumes & calculate the % w/v of Borax.

- 2. <u>Impure borax</u> may contain Na₂CO₃ or boric acid as impurities.
- a) Mixture of Borax & Na₂CO₃

Na₂CO₃ is basic; it can react with HCl, so both borax and carbonate will react with HCl Na₂CO₃ + 2HCl \longrightarrow 2NaCl + H₂O + CO₂

- While NaOH react only with boric acid liberated from borax. Then after vol. correction;
 - V1 of N/2 HCl > V2 of 1N NaOH
- V₁- V₂= vol. of N/2 HCl react with Na₂CO₃
- Calculate the % w/v for Borax and Na₂CO₃.
- b) Mixture of Borax & boric acid
- $Na_2B_4O_7.10H_2O + 2HCI \longrightarrow 2NaCI + 4H_3BO_3 + 5H_2O$

liberated boric acid

H₃BO₃ + NaOH Mannitol NaBO₂ + 2H₂O

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So both the liberated and free boric acid react with NaOH, then

Correct the volumes and compare:

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liberated boric acid

H₃BO₃ + NaOH Mannitol NaBO₂ + 2H₂O

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So both the liberated and free boric acid react with NaOH, then

Correct the volumes and compare:

Assay of Chlorinated lime

Or bleaching powder Or Ca. Chlorohypochlorite [Ca(OCI)CI]

Assay of Chlorinated lime [Ca(OCI)CI]

- Contain not less than 30%w/w available chlorine.
- It is dull-white powder with characteristic odor, partially soluble in water and in 96% ethanol.
- Used as disinfectant.



the color of the solution changed from brown to yellow.

* End point is colorless

• Chemical principle:

Ca(OCI) CI+2CH₃COOH \longrightarrow (CH₃COO)₂Ca+ HOCI+ HCI

hypoclorous acid

 $HOCI + HCI \longrightarrow Cl_2 + H_2O$

available chlorine

The *available chlorine* is the Cl₂ which is liberated from the substance by the action of acid.

This liberated chlorine will displace an equivalent amount of I₂ from KI.

 $CI_2 + 2KI \longrightarrow 2KCI + I_2$

The liberated I₂ is titrated against sod. thiosulfate using starch mucilage as indicator.

 $I_2 + 2Na_2S_2O_3 \longrightarrow 2NaI + Na_2S_4O_6$

Chemical factor:

1ml of 0.1N Na₂S₂O₃ \equiv 0.003545gm *available chlorine*. Calculation:

Correct the sod. thiosulfate volume=V'

V' X ch. factor= gm of *available chlorine* in 10ml. (Gm/10)X100= %w/v of *available chlorine*.

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Assay of copper sulfate

Cupric sulfate CuSO4.5H2O *Mwt.=249.68 gm*

- Properties: blue, fine prism or blue crystalline powder.
- **Solubility**: soluble at 20°c in 3parts of water and 3parts of glycerol, almost insoluble in 95% alcohol.
 - It contains not less than 98.5% and not more than 101%w/w CuSO4.5H₂O.
- **Uses**: as fungicide.

Procedure: the titration should be carried in stoppered flask.

During titration the color of the solution changed from **brown** to **faint yellow** then add1ml starch and 1gm KSCN and complete titration.

≈0.1N Na2S2O3

10ml unknown _____ 30ml D.w. 1.5 gm KI 5ml acetic acid ____ The buret contains the titrant. This flask contains the solution to be titrated and the indicator.



• Notes:

- 1. KI is added to liberate I2.
- 2. Excess KI is added to dissolve I2.
- 3. HAC is added to make the medium slightly acidic.
- 4. In acidic media the liberated I₂ is adsorbed by the ppted CuI.
- 5. KSCN(pot. thiocyanate) is added near the end point.

KSCN+ CuI → CuSCN+ KI

this KI will return to

react with $CuSO_4 \rightarrow I_2$

Chemical factor:

1ml of 0.1N Na₂S₂O₃ 0.02497gm CuSO₄.5H₂O

Calculation:

- □ correct the volume of Na₂S₂O₃=V'
- □ V' X Ch. Factor= gm wt. of CuSO₄.5H₂O in 10ml
- Gm/10) X 100= %w/v of CuSO₄.5H₂O in the unknown

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Assay of Ferrous Sulfate

FeSO4.7H2O Mwt.=278gm/mole

- **Appearance**: The blue-green heptahydrate is the most common form of ferrous sulfate ,it is bluish green crystals, freely soluble in water.
- Content: Ferrous Sulfate (crystal) contains 98.0— 104.0% of ferrous sulfate heptahydrate (FeSO4· 7H2O)
- **Used** in the prevention and treatment of iron deficiency.



• Chemical principle:

10FeSO4+2KMnO4+8H2SO4

2004+5Fe2(SO4)3+K2SO4+ 8H2O

Or $MnO4 + 8 H^+ + 5Fe^{+2}$ $H^+ + 4H_2O + 5 Fe^{+3}$

- Ferrous sulfate is a *reducing agent*.
- It will be titrated against standard solution of potassium permanganate(*oxidizing agent*) in acidic media.
- *Note*: you must write the oxidation and reduction equations by each.

• Chemical factor:

<u>Derive</u> the ch. Factor from the previous chemical equation.

1ml. of 0.1N KMnO4 \equiv 0.0278gm of FeSO4

• Calculation:

Correct the volume of KMnO4 consumed=V'

Then multiply the corrected volume (V') by the Ch. Factor.

to get the weight;

V' x ch. Factor = gm of FeSO4 in the unknown.

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Assay of Hydrogen Peroxide



H2O2: Mwt.=34.02 gm/mole Different H2O2 solution: 30%,27%w/w and 6%, 3%w/v

- H2O2 is slightly more viscous than water.
- Aqueous solution of H₂O₂ colorless and odorless. Rapidly decomposes when contact with organic matter and certain metals; Cu, Fe, Mn and if allowed to stay in alkaline solution. $2H_2O_2 \rightarrow O_2 + H_2O$

- In acidic solution Fe²⁺ is oxidized to Fe³⁺ (H₂O₂ acting as an oxidizing agent):
 2 Fe²⁺ + H₂O₂ + 2H⁺ → 2 Fe³⁺ + 2H₂O
- hydrogen peroxide acts as a reducing agent in its reaction with KMnO₄ in our experiment.
 2MnO₄⁻ + 5H₂O₂ + 6H⁺ → 2Mn²⁺ + 5O₂ + 8H₂O

• Uses:

It is a *disinfectant* or *antiseptic* for treating wounds for its antimicrobial activity.

it is an effective *cleaning agent* prevents infection transmission in the hospital environment.



End point: colorless \implies *pink*

• Notes:

- 1. We make the media acidic **to**;
- Prevent decomposition of H₂O₂
- Prevent reaction of $KMnO_4 \longrightarrow MnO_2$
- Increase oxidizing power of KMnO₄
- 2. Titration should be done slowly

• Chemical principle:

 $2KMnO_4 + 5H_2 O_2 + 3H_2SO_4 \longrightarrow K_2SO_4 + 2MnSO_4 + 5O_2 + 8H_2O_4 \longrightarrow K_2SO_4 \longrightarrow K_2S$

- *You must write the oxidation and reduction equations to calculate the eqwt. for both KMnO₄ & H₂O₂.
- Chemical factor:

According to the chemical equation;

2Mwt of KMnO₄ \equiv 5Mwt of H₂O₂

then complete the derivation of the Ch. Factor.

The Chemical factor is equal to: 1ml of 0.1N KMnO₄ \equiv 0.001701gm of H₂O₂

- Calculation:
- Correct the volume of KMnO₄
 NxV = N'xV' (corrected)
- Multiply the corrected volume by the Ch. Factor. to get the **weight (wt.)** of H₂O₂
- the **%w/v** of H₂O₂ = **wt**. /10 x 100

• There is <u>2 methods</u> to calculate the concentration of hydrogen peroxide in commercial products:

1. %W/V:

10ml of concentrated sol. was diluted to 200ml with D.W. \longrightarrow Then 20ml is taken from the dil. one and titrated with approximately 0.1N KMnO₄

 conc.
 diluted

 10
 200
 x= 20X10/200=1ml of conc. sol.

 x
 20
 (original sol.)

Correct the volume of $KMnO_4 = Vol.$

Vol. x ch. Factor =wt. of H_2O_2 in <u>1ml</u>.

Then calculate %w/v

2. Volume strength of the solution:

Decomposition occurs according to this equation:

 $2 H_2O_2 \longrightarrow 2 H_2O + O_2$ <u>Example</u>: if the %w/v of H_2O_2 is 8.5%w/v *i.e.* It contains 8.5 gm of H_2O_2 in 100ml solution. Vol. <u>wt.</u> $X = 0.085 \text{ gm of } H_2O_2 \text{ in 1ml}$ 8.5 100ml Χ 1ml (from the definition) $2 H_2O_2 \longrightarrow 2 H_2O + O_2$ 2**×**34gm mwt. 22400ml $y = 28 m l of O_2$ 0.085gm y

So the *volume strength* of the sample is 28.

Assay of sodium hydroxide solution

NaOH solution

 From B.p, NaOH solution contains not less than 97.5% w/w of total alkali (as NaOH) and not more than 2.5 % w/w Na₂CO₃.

• Assay:

- > 10 ml of unknown(bulb pipette).
- > 25 ml of distilled water.
- > Add 2.5 ml of barium chloride solution.
- Titrate with 1N HCl solution using 1-2 drops of phenolphthalein ind..
- > The first end point from pink ---> colorless(turbid)
- To the turbid sol. add 5 drops of Bromophenol Blue ind. and complete titration with 1N HCl.
- > The second end point bluish violet yellowish green



Chemical principle:

- NaOH is strong base, absorbs CO₂
 2NaOH + CO₂ → Na₂CO₃ + H₂O
- both NaOH and Na₂CO₃ react with HCl
 NaOH + HCl → NaCl + H₂O
 Na₂CO₃ + 2HCl → 2NaCl + H₂O + CO₂

When we assay a sample, we do the assay for total alkalinity contributed to NaOH and Na₂CO₃.

 Barium chloride (BaCl₂) is added to precipitate all carbonate

• 1st titration:

 $NaOH + HCI \longrightarrow NaCI + H_2O$

Why HCl do not react with BaCO₃?

Why the end point is turbid?

• 2nd titration:

 $2HCI + BaCO_3 \longrightarrow BaCI_2 + H_2O + CO_2$

definition of <u>chemical factor</u>: the weight of substance that is chemically equivalent to 1ml of std. solution.

- Calculation of the *chemical factor*:
- a) From reaction of HCl with NaOH:
- 1Mwt of NaOH \equiv 1 Mwt HCl
- $1 \text{ Mwt of NaOH} \equiv 1 \text{ eqwt HCL}$
- $1 \approx 40 \text{ gm of NaOH} \equiv 1 \text{ liter of } 1N \text{ HCl}$
- $40/1000 \text{ gm NaOH} \equiv 1 \text{ml of } 1N \text{ HCl}$
- 0.04 gm of NaOH \equiv 1ml of 1N HCl of total alkalinity calculated as NaOH(chemical factor)

```
b) From reaction of 2HCl with Na<sub>2</sub>CO<sub>3</sub>
2Mwt of HCl \equiv1Mwt of BaCO<sub>3</sub> \equiv1Mwt Na<sub>2</sub>CO<sub>3</sub>
1Mwt Na<sub>2</sub>CO<sub>3</sub> \equiv 2Mwt of HCl
1Mwt Na<sub>2</sub>CO<sub>3</sub> \equiv 2 eqwt of HCl
\frac{1}{2} Mwt Na<sub>2</sub>CO<sub>3</sub> \equiv 1 eqwt of HCl
106/2 \text{ gm Na}_2\text{CO}_3 \equiv 1 \text{ liter of } 1N \text{ HCL}
53 gm Na<sub>2</sub>CO<sub>3</sub> \equiv 1liter of 1N HCL
 53/1000 \text{ gm Na}_2\text{CO}_3 \equiv 1\text{ml of } 1N \text{ HCL}
0.053 \text{ gm Na}_2\text{CO}_3 \equiv 1 \text{ml of } 1 \text{N HCL} (chemical)
     factor)
```

• Calculations :

V₁ is the of HCl consumed in the 1st titration V₂ is the of HCl consumed in the 2nd titration $V_1+V_2=V_3$ total HCl consumed. correct the V₃ according to this equation: $V \times N = V' \times N'$ Corrected V₃ x 0.04= gm wt. of total alkali Then the % w/v of total alkali Corrected V₂ x 0.053= gm wt. of Na₂CO₃ Then the % w/v of Na₂CO₃ in the unknown

Preparation & Standardization of ≈ 0.1 N Potassium Permanganate (KMnO4) solution

Redox Reaction

- Preparation of \$0.1N KMnO4 solution:
- Dissolve 3.3 gm of solid KMnO₄ in 1000ml. distilled water, in a flask.
- ➢ Heat the solution on a hotplate for 15min. at 70-80 °c.
- Cool the solution to room temp. and allow stand for 2 days in a dark place.
- Filter the solution through the glass wool or asbestos before use.

- <u>Standardization of KMnO4 solution</u>:
- Filter the prepared soln. through glass wool as said before.
- Rinse and Fill the burette with filtered KMnO₄ soln.
- Weigh accurately about 200mg sod. oxalate (Na₂C₂O₄), previously dried at 110 °c to constant weight.
- Dissolve it with 100ml of D.W. in a conical flask. Add 7ml. of conc. H₂SO₄.
- Heat the soln. to about 70°c.
- Titrate against KMnO₄ soln. until a permanent pale pink color appear (persist for 15 sec.).
- Record the volume of KMnO₄ consumed.

• **OXIDATION- REDUCTION REACTIONS:**

Methods of analysis involve a change in valence of the reacting substances, thus there is always a transfer of electrons.

Electrons lost by reducing agent are gained by oxidizing agent. The oxidizing agent will be reduced; gain of electrons and then decreasing in oxidation number, but the reducing agent will be oxidized; losing of electrons and then increasing in oxidation number. We can calculate the eqwt. of the reactants: Mwt.

Eq. wt =_____

no. of e-gained or lost by each molecule

Permanganate ion (MnO₄⁻) acts as an oxidizing agent in acidic media, being reduced to Mn⁺⁺:
 ⁺⁷/_{MnO4} + 8H⁺ + 5⁻e — Mn⁺² + 4H₂O

 \Box Oxalate ion (C₂O₄⁼)acts as reducing agent:

C₂O₄⁼ → 2CO₂ + 2e⁻
- KMnO₄ is widely used in volumetric analysis as a standard and oxidizing agent because:
- 1. It is strong oxidizing agent.
- The intense color of KMnO₄ solution is sufficient to signal the end point in most titrations thus it is used as self indicator.
- On the other hand,
- KMnO₄ is a strong oxidizing agent, then it will oxidize any organic matter present in D.W., so heating for 15min. is to accelerate the oxidation of these organic matter.

- 2) KMnO₄ soln. is allowed to stand for 2days to ensure the completion of the decomposition reaction.
- 3) It is filtered through asbestos to remove all traces of manganese dioxide (MnO₂).
- 4) KMnO₄ is unstable in the presence of direct sunlight, organic matter ex. Filter paper.
- 5) H₂SO₄ is added since KMnO₄ is better oxidizing agent in acidic media, H₂SO₄ is added to keep [H⁺]ion conc. constant through out the titration.
- 6) Heat to about 70 °c because the oxidation of sod. oxalate is rapid enough if the temp. above 60 °c.

- Sod. oxalate, not oxalic acid, is used because:
- 1. The salt can be obtained in very pure condition.
- 2. Free from water of crystallization.
- 3. It can be dried to 130 °c without decomposition.
- 4. Stable to air.

The chemical equation for the std. of pot. Permanganate:

 $5Na_2C_2O_4 + 2KMnO_4 + 8H_2SO_4 \longrightarrow 2MnSO_4 + 5Na_2SO_4 + K_2SO_4 + 8H_2O_4 + 10CO_2$

Calculations

$\frac{\text{KMnO4}}{\text{N X V}} = \frac{\text{Na}_2\text{C}_2\text{O}_4}{\text{WT}} \times 1000$

eqwt

67gm/mole

The preparation and standardization of 0.1N sodium thiosulfate Na2S2O3.5H2O

Preparation:

Dissolve about 26gm sod. thiosulfate and 200mg sod. carbonate in 1000ml of recently boiled and cooled D.W.

Standardization:

- Put into the <u>stoppered flask</u> the followings;
- 0.1–0.2 gm of potassium dichromate (K2Cr2O7) 100ml of D.W.
- 3gm potassium iodide (KI)
- 2gm sod. Bicarbonate(NaHCO3)
- 5ml conc. HCl
- Close the flask with the stopper and swirl to mix.
- Then keep the flask in dark place for 10min.

- Rinse the stopper and inner walls of the flask with D.W.
- Titrate the liberated I₂ with sod. thiosulfate the color change from brown to faint yellow.
- Add 1ml of starch indicator the color of the solution change to dark blue.
- Complete titration until the color change to green-blue.



End point: dark blue \longrightarrow green-blue

Notes for preparation:

Why do we use recently boiled and cooled water?
 a. to destroy sulfur bacteria
 b. to expel CO2
 Na2S2O3 +2H2CO3 → 2NaHCO3+H2S2O3
 H2S2O3 → H2SO3 + S↓
 thiosulfuric acid
 sulfar
 SO2 +H2O

Na2CO3 is added to the solution as preservative

Chemical principle of standardization:

- Pot. Dichromate is an oxidizing agent K2Cr2O7 +6KI +14HCI 3I2+2CrCI3+8KCI+7H2O
- The liberated I2 is titrated against Na2S2O3 2Na2S2O3+ I2 Na2S4O6 +2NaI

sod. tetrathionate

This indirect procedure is known as *iodometry*.

Notes for standardization:

- We use stoppered flask to prevent the loss of I₂.
- 2. KI is added to liberate I₂.
- 3. Excess KI is added, Why?
 I[−]+ I₂ → I₃[−] tri-iodide ion (soluble form of iodine)
- 4. The standardization should be carried in acidic media and not in alkaline media: because alkaline media I₂ will react with OH⁻ to give:
 - $I_2 + OH^- \rightarrow HI + IO^-$ Hypoiodite ion
 - $3 \text{ IO}^- \longrightarrow \text{ IO}_3^- + 2\text{ I}^-$

These ions will oxidize the thiosulfate to: $IO^- + IO_3^- + S_2O_3^- \longrightarrow SO_4^-$

thiosulfate

sulfate

excess

atmospheric O₂



Sod. bicarbonate is added to react with formed HI: NaHCO₃+ HI \longrightarrow NaI+ H₂O+ CO₂ \nearrow

- 6. We put the flask in dark place for 10min.?
- 7. Starch ind. is added at the end (when the color of the solution change to yellowish green) or we can said the ind. is added when the iodine concentration is low because β -amylase which the soluble form of starch react :

 β -amylase+I₂+I⁻ blue colour complex

the color change is reversible, the color being discharged (change to colorless) when iodine is reduced by Na₂S₂O₃.

This reversibility is reduced when the iodine conc. is high for this reason starch ind. should be added until most of I₂ has been reduced by Na₂S₂O₃.

8. The end point is a pale green and not colorless why?

Calculation:

Na2S2O3 $K_2Cr_2O_7$ N x V = (Wt/eqwt) x1000

eqwt = 49

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 - $3 \text{ IO}^- \longrightarrow \text{ IO}_3^- + 2\text{ I}^-$

These ions will oxidize the thiosulfate to: $IO^- + IO_3^- + S_2O_3^- \longrightarrow SO_4^-$

thiosulfate

sulfate

excess

atmospheric O₂



Sod. bicarbonate is added to react with formed HI: NaHCO₃+ HI \longrightarrow NaI+ H₂O+ CO₂ \nearrow

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eqwt = 49

The preparation and assay of Aqueous Iodine Solution (Lugol's Solution)

- It contains 5% w/v *lodine* (limits 4.75– 5.25% w/v) and 10% w/v *Pot. lodide* (limits 9.5– 10.5% w/v).
- Used in pre-operative treatment of thyrotoxicosis.
- <u>Preparation</u>:
 - l2 50gm
 - KI 100gm

purified water sufficient to 1000ml *Dissolve KI in a little amount of water then add the I2 and shake well until the I2 is completely dissolved, then add sufficient water to 1000ml using volumetric flask.



b) Assay of KI,

* titrate with M/20 KIO3 (Pot. lodate) until dark brown sol. change to pale brown, add 1ml of <u>amaranth</u> sol.; the color of sol. Change to red and then continue tit. Until the color of sol. change to yellow -orange

10ml D.W



From the no. of ml. of M/20 KIO3 required subtract ¼ no. of ml of N/10 Na2S2O3 required in the assay of I2

each 1ml of the remainder is \equiv 0.0166gm of Kl

Chemical principle for I2 assay:

I2 will oxidize chemically equivalent amount of sod. thiosulfate.

 $I_2 + 2Na_2S_2O_3 \longrightarrow 2NaI + Na_2S_4O_6$

 <u>Chemical principle for KI assay</u>: KIO₃ (pot. lodate)is fairly strong oxidizing agent, it can quantitatively react with both iodide I⁻ and iodine I₂.

*If the conc. of HCl not exceed 1N the reaction: $KIO_3 + 5KI + 6HCI \implies 6KCI + 3I_2 + 3H_2O$

- *In the presence of conc. HCl exceeding 4N, I2 produced by the previous reaction is further oxidized to iodine monochloride (ICl): KIO₃+ 2I₂+ 6HCl → KCl+ 5ICl+ 3H₂O
- * <u>so</u> the overall reaction between KIO₃ & KI in the presence of high conc. HCL >4N can be expressed by:
 - $KIO_3 + 2KI + 6HCI \longrightarrow 3KCI + 3ICI + 3H_2O$
 - The presence of high conc. HCl is required to decrease :
 - $HCI + ICI \longrightarrow ICI_2^- + H^+$

stable cpd

- In this experiment we use amaranth indicator, it is a dye, dyes which are destroyed or changed in the presence of oxidizing titrant can be use as indicator.
- We can not use starch as indicator because starch_ iodide complex can not be formed in high acid conc. used in the iodate titration.

• Chemical factor. <u>Homework</u>

Each1ml of 0.1N Na₂S₂O₃ \equiv 0.01269gm of lodine Each1ml of M/20 KlO₃ \equiv 0.0166gm of Kl Each1ml of M/20 KlO₃ \equiv 0.02538gm of iodine

Calculation:

For I2: <u>original sol.</u>
 25
 X
 100
 X= 2.5ml of original sol.
 10

V' x ch.fact. 0.01269x 100/2.5 = %w/v of iodine in orig. sol. V' the corrected volume of sod. thiosulfate.

For KI: original sol. 25 X 25 X 5X = 1.25ml of original sol.

 $V \text{KIO3 M/20} = V \text{KIO3 M/20 (used} - \frac{1/4}{1/4} V \text{Na2S2O3}$ (req. for KI) in the total assay) 0.1N

Vol. of KIO3 (req. for KI)x 0.0166 x 100/1.25= %w/v of KI

- It is preferred to use iodate (KIO3) of known M rather than of known N, it depends on the nature of reaction.
- KIO3 sol. used in the titration of sol. containing both iodide and iodine.
- The equivalent of KIO3 in its reaction with KI differs from its equivalent when react with iodine.

The preparation and assay of Aqueous Iodine Solution (Lugol's Solution)

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