Kinetics of Oxidation of Schiff Base by Chloramine –T Catalysed by Mixture of Os (Viii) and Ru (Iii) – Effect of Osmium Tetraoxide and Ruthenium Chloride

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ABSTRACT : The kinetic study of simple and substituted anils ware studied under various conditions viz., solvent ,oxidants , electrolyte , catalysts and mixture of catalysts and temperature; simple and substituted anils , pyridinium Chloro Chromate(PCC) ,Pyridinium Dichromate (PDC) were prepared in the laboratory and kinetic study were carried out using Pyridinium dichromate , substrate , perchloric acid and osmium tetraoxide . The graphs were drawn at all relevant places and results were obtained in full satisfaction.

Keywords: diverse nature , chloramine [CAT], sulphonamide , oxidants , water – bath , evaporating , stirring ,

1.INTRODUCTION :

Among oxo derivatives of variable valence metals , Chromium compound play the most important role , in oxidative reaction .A number of Chromium reagents are readily available . Almost every oxidisable fuctional group may undergo Chromium oxidation .Chromium (VI) containing reagents include Chromium acid , dichromate ion , Chromyl chloride , chromyl acetate , t - butyl Chromate , Chromylnitrate and Co – ordination complexes of Chromium trioxide .¹⁻²

Chromium (VI oxidations are usually performed under acidic condition . Co – solvents like (Jone's reagent) benzene, methylenechloride (Two Phase system) are often added in order to deal with water insoluble organic complexes . The oxidation rate is generally high under acid catalysed conditions, however the low pH of the reaction medium and the presence of water favouring hydrolytic condition exclude the use of this class of reagents for the oxidation of molecules containing acid sensitive groups.

Polymer supported Chromium (VI) reagents have also been developed. These reagents offer the advantage of reducing the work of procedure to more filtration. Recently some neutral or almost neutral Chromium (VI) reagents have been developed to effect oxidation under mild conditions. Pyridinium Chlorochromate (PCC) introduced by Covey et al³ is widely used in the oxidation of alcohols. Pyridinium flourochromate has a less pronounced acidity and is an effective agent for the oxidation of Plycyclic organic substitutes . PFC was developed by Bhatachargee and coworker .⁴

In 1986, Narayanan and Balasuramaniam⁵ introduced Pyridinium bromochromate. This is an efficient oxidant for alcohols and a brominating agent as well .Firouzahadi and Coworkers used Zinc chromates. This is a very useful reagent for the oxidation of variety of organic compounds , including alcohols , oximes , olefins and aromatic hydrocarbons.

They have also shown chromium peroxide complexes as versatile, mild and efficient oxidants in organic synthesis.Chlorotrimethyl Silone Imidazolium dichromate , 2,2' - Bipyridyl chlorochromate (BPCC) and the corresponding chromate (BPC) have been proposed as oxidants for the oxidation of the hydroxyl group to thr carbonyl group.4 – (dimethyl amino) – pyridinium chlorochromate and chromyl nitrate have been shown to be efficient and mild oxidant in aprotic media.Naphthyridium chlorochromate Pvrazinium chlorochromate tetrabutvl ammonium chlorochromate , dimethylpyrazole chromate, tetrabutyl ammonium dichromate, Pyridine complex of oxodiperoxochromium and the dipyridine complex of chromium trioxide are some of the Chromium (VI) oxidants introduced in the last twenty years.Quinolinium Chlorochromate was used by Bhavani et al to oxidize methyl phenyl sulphoxides.Quinolinium dichromate was introduced by Balasubramanian and Prathiba⁶ as an effective oxidant under non - aqueous conditions.

Pyridium Chlorochromate (PCC) is irreplaceable in the generation of functional groups in highly unsaturated carbinols and pyrols . yet mildly acidic character of Pyridinium chlorochromate precludes its use with acid sensitive substrates or products. Pyridium Chlorochromate oxidation has been investigated under various condition on different substrates .Despite the introduction of numerous oxidants based on chromium (VI) not much work has been done to investigate the mechanism of oxidation in

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Oxidation of Anils

Several oxidation of anils have been studied :

1.2.1.Effect of varying PCC concentration

The reaction was investigated by varying the concentration at constant substrate concentration .Th reaction was found to be first order with respect to PCC [Table -1.2]

Table -1.2

 $\begin{array}{ll} [Anil] &= 1.5 \ x \ 10^{-2} \ M \\ [H_2C_2O_4] &= 5.0 \ x \ 10^{-2} \ M \end{array}$ ACOH :H2O = 70:30 (V/V)Temp. = 35^{0} c $[HClO_4] = 3.87 \times 10^{-3} M$

$[PCC] \times 10^3$	Kobs. X 10 ⁴ s ⁻¹
1.2	1.932
1.5	1.921
2.00	1.811

1.2.2.Effect of varying the concentration of Anil

The reaction was carried out under pseudo - first order conditions. In this study, the concentration of PCC, Oxalic acid, Perchloric acid and percentage of oxalic acid were kept constant and the concentration of the alone varied. The reaction was found to be first order with respect to anil. [Table -1.3] Table -1.3

 $[PCC] = 1.5 \times 10^{-3} M \qquad ACOH : H2O = 70:30 (V/V)$ $[H_2C_2O_4] = 5.0 \times 10^{-2} M$ Temp. = $35^{\circ} c$ $[\text{HClO4}] = 3.87 \text{ x } 10^{-3} \text{ M}$

[ANIL] x 10^2	Kobs. X 10 ⁴ s ⁻¹
1.00	1.08
1.50	1.923
2.50	2.785
3.00	3.512

1.2.3.Effect of varying solvent Composition

The reaction was studied by the different composition of acetic acid under constant [Reactants]. It was found that as the percentage of acetic acid increased, the rate decreased. [Table -1.4]

Table - 1.4

 $[ANIL] = 1.5 \times 10^{-3} M$ $[\text{HClO}_4] = 3.87 \text{ x } 10^{-3} \text{ M}$ $[H_2C_2O_4] = 5.0 \times 10^{-2} M$ Temp. = 35[°] c

% AcOH	Kobs. X 10 ⁴ s ⁻¹
40	3.921
50	3.207
60	3.256
70	1.932

It was found that the rate deccreases with increase of percentage of acetic acid.

1.2.4Effect of varying Oxalic acid

The reaction was carried out under by varying , the concentration of , Oxalic acid and by keeping other [reactans] constant. . The result indicate that as the concentration of oxalic acid increased , the rate constant also increased.

[Table – 1.5]

Table – 1.5

$$[ANIL] = 1.5 \times 10^{-2} M ACOH :H2O = 70:30 (V/V)$$
$$[HClO_4] = 3.87 \times 10^{-3} M Temp. = 35^{0} c$$

$[H_2C_2O_4] \ge 10^4$	Kobs. X 10 ⁴ s ⁻¹
5.0	1.932
7.5	2.872
10.0	3.871
12.0	5.965

1.2.5.Effect of varying Perchloric acid

The reaction was studied by varying , the concentration of perchloric acid and by keeping other [reactans] constant. . The result indicate that as the concentration of Perchloric acid increased , the rate constant also increased.

[Table -1.6]

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Table -1.6
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[ANIL] = 1.5 \times 10^{-2} M ACOH :H2O = 70:30 (V/V)
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$[H_2C_2O_4] = 5.0 \times 10^{-2} M$	Temp. = $35 \degree c$
$[\text{HClO}_4] \ge 10^4$	Kobs. X 10^4 s ⁻¹
3.90	1.932
5.80	2.785
7.70	2.952
9.70	4.609

1.2.6.Effect of varying Sodium Perchlorate

To follow the Primary Salt effect, the reaction was studied with varying concentration of Sodium Perchlorate by keeping other [reactants] constant. It was found that as the concentration of Sodium Perchlorate increased, the rate constant also increased. This shows that the participation of ions which are similar in their sign or dipole –ion interaction in the rate determining step. [Table 1.7]

Table – 1.7

 $\begin{array}{ll} [\text{ANIL}] &= 1.5 \ x \ 10^{-2} \ \text{M} & \text{ACOH} : \text{H2O} \\ &= 70:30 \ (\text{V/V}) \\ & [\text{H}_2\text{C}_2\text{O}_4] \\ &= 5.0 \ x \ 10^{-2} \ \text{M} & [\text{HClO}_4] = 3.87 \ x \ 10^{-3} \ \text{M} \\ & [\text{PCC}] &= 1.2 \ X \ 10^{-3} \ \text{M} & \text{Temp.} = 35^{-0} \ \text{c} \end{array}$

Table 2.5

[NaClO4] X 10 ³	K _{obs} x 10 ⁴ S ⁻¹
0.00	1.932
2.50	3.548
5.00	4.657
7.50	5.086
10.00	5.799

1.2.7.Effect of MnSO4

The reaction rate decreases tremendously with the increase in the concentration of MnSO4 [Table - 1.8] This may be due to the formation of Cr(VI) in the rate determining step. Table - 1.8

 $\begin{array}{ll} [\text{ANIL}] &= 1.5 \text{ X } 10^{-2} \text{ M} \\ [\text{H}_2\text{C}_2\text{O}_4] &= 5.0 \text{ x } 10^{-2} \text{ M} \\ [\text{ HClO4}] &= 3.87 \text{ X } 10^{-3} \text{ M} \end{array} \begin{array}{l} \text{AcOH} : \text{H2O} &= 70 : 30 (\text{V/V}) \\ [\text{ PCC}] &= 1.2 \text{ X } 10^{-3} \text{ M} \\ \text{Temp.} &= 35^{-0} \text{C} \end{array}$

[MnSO4] X 10 ⁴	$K_{obs} \ge 10^4 \text{ S}^{-1}$
0.00	1.932
1.00	1.053
2.00	0.723
3.00	0.575

1.2.8 Effect of temperature

The rate of oxidation of some *meta* and *para* substituted anils had been studied at four different temperatures viz., 35° c , 40° c, 45° c and 50° c . It was observed , as we expected ,that the rate increases very much with increase of temperature . [Table – 1.9]

Table – 1.9

 $\begin{array}{ll} [\text{Substrate}] &= 1.5 \ x \ 10^{-2} \ \text{M} & \text{ACOH} : \text{H2O} \\ [\text{H}_2\text{C}_2\text{O}_4] &= 1.0 \ x \ 10^{-4} \ \text{M} & [\text{PCC}] \\ [\text{HCIO}_4] &= 4.0 \ x \ 10^{-3} \ \text{M} & \text{Temp.} = 35^{-0} \ \text{c} \end{array}$

S.No.	Substituents	Kobs. X 10 ²	s ⁻¹		
		35^{0}	40^{0}	45^{0}	50^{0}
2	Н	1.932	3.335	4.127	6.735
3	P-CH ₃	3.91	6.123	8.315	10.371
4	P – O CH3	6.22	9.132	12.612	16.431
5	p- Cl	1.023	2.425	3.671	4.923
6	m - Cl	0.705	1.412	2.213	3.824

2. Kinetics of Chloraminometric oxidation

Coull and co –workers were thr first to investigate the kinetics and mechanism of decomposition of H2O2 by CAT in the presence of HCl.The reaction was found to be first orders each of [H2O2] and [CAT [, but inverse first order dependence on the concentration of RNH2.

Oxidation of kinetics of Alpha – hydroxyl acids¹ with CAT in alikaline medium in the presence of Os (VIII) as catalyst was found a follow first order each in [CAT] AND Os (VIII) and was independent of substrate concentrations. Os (VIII) catalysed oxidation of acetone and ethyl methyl ketone by CAT was reported³.

3.EXPERIMENTAL

2.1

3.1.Preparation of Anils General method of preparation of Anils

Ph

$$C = N - C6H4 X$$

H
Where $X = H$, p - CH3, p - OCH3, p - Cl. p - Br, P - COOH,
p - NO2, m - CH3, m - Cl, m- NO2

The anils were prepared 87a by refluxing equimolar quantities of benzaldehyde and aniline or substituted anilines in alcohol for about 2 to 3 hrs. The resulting solution was cooled and poured into the cold water. The precipitated anil was filtered, washed, dried and recrystallised from alcohol. The purity of anils were checked by determining their melting points (Table 2.1).

3.2.Preparation of Chloramine – T (CAT)

a) Preparation of Pure p- tolenesulphonamide

The procedure used was that of Vogel.⁷⁹ Ten grams of p- tolenesulphonylchloride and 20 g of ammonium carbonate were ground in a mortar to a fine powder. The mixture was heated in a an evaporating dish on a water bath for 1 - 2 hrs. and the mixture was stirred frequently with glass rod. After cooling the mixture was extracted with a little cold water to remove excess ammonium salts. The crude p- tolenesulphonylchloride was recrystallised from boiling water and dried at 100^{0} C. The pure product melted at

138⁰ C.

b) Preparation of Dichloramine – T (DCT)

This was prepared by the method of Krausand crede . p- tolenesulphonamide was dissolved in ten parts of 1 : 10 caustic soda and diluted with twenty parts of water . Chlorine gas was then passed into the solution until a voluminous white precipitate of dichloramine – T was formed .This was collected on a Buckner funnel , thoroughly washed twice with 5 - 8 parts of water and finally with enough 10% alcohol to make a thin paste . The dilute alcohol washing was done very quickly and the substance was separated with air of a vacuum filter . It was then dried in a vacuum drier dichloramine –T was obtained , and its melting point is $82 - 83^0$ c.

C) Preparation of Chloramine - T (CAT)

Forty five ml of 10% sodium hydroxide solution was heated to a temperature of about 80^{0} and to which added 6 g of dichloramine – T in small quantities , strring the mixture gently after each addition until a clear solution was obtained. When the addition was completed , the hot solution was filtered and then allowed to cool spontaneously . The crystals were filtered with suction , washed with little brine solution , and dried in a desiccators over anhydrous calcium chloride . It was recrystallised from water ; m.p. $175 - 179^{0}$ C.

3.3.Reagents

Analar samples are sodium perchlorate ,Sulphuric acid , Perchloric acid , Sodium chloride , Osmium tetroxide were used. Ruthenium chloride was dissolved in water containing dil. HCl and the solution was standardized by the method of Houriuchi and Ichijo 13

4.Kinetic measuments

The reaction was carried out in 60 % aqueous acid (V/V) . Following solutions of desired concentrations were prepared .

- 1. Anil in Acetic acid
- 2. Chloramine T in water
- 3. Sodium perchlorate in water
- 4. Sodium Chloride in water
- 5. Ruthenium chloride in water containing dilHCl
- 6. Osmium tetroxide in dil.Sodium hydroxide .

The solutions were thrmostated at least for one hour before the run. The kinetics was carried out by mixing the substrate and oxidant in a appropriate solvent mixture. The reaction was carried out under pseudo first order conditions , taking always the substrate in excess.

The progress of the reaction was monitored by withdrawing aliquots from the reaction mixture at regular intervals of time and estimating the unreacted CAT by iodometric method.

Product analysis

Under kinetic conditions , the reactants ratio anil : CAT (1 : 10) were mixed and allowed to react for about 2 days. The product was extracted with chloroform . the solvent was removed at reduced pressure after drying over anhydrous Na₂SO₄. The products were separated , after identifying by TLC , into individual components by Column Chromatography . Benzaldehyde was confirmed by its semicarbazone (m.p.221^o c , lit. 222^o c) and 2,4 – DNP (m.p.236^o c , lit. 239^o c) derivatives.

Nitrobenzene was identified by recording IR spectrum and exhibits three bands at 1626, 1500, and 1019 cm^{-1} attributed to stretching frequency of C-N. One sharp band at 1452 cm^{-1} due to stretching vibratin of N-O and band at 530 cm^{-1} due to ring deformation and CNO bending vibration.

5. Stoichiometry

The stoichiometry of the reaction was determined by studying the reaction under the conditions of CAT >> anil and it was found to be 1:1

6.RESULTS AND DISCUSSINS 6.1.Effect of varying Os (VIII)

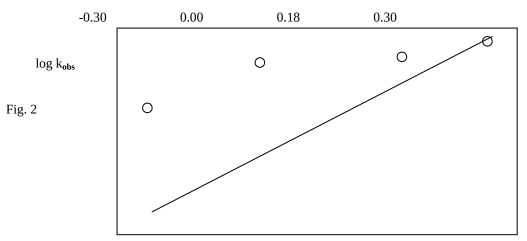
The reaction was carried out by varying the concentration of Os (VIII) and by keeping othe [reactants] constant. The results indicate that as the concentration of Os (VIII) increased, the rate constant also increased. the plot of log Os (VIII) Vsa log kobs. Gave a straight line of slope 0.29 [Table 5.4. Fig.2] Table -5.4

 $[CAT] = 1.0 \times 10^{-3} M$ $[Ru(III)] = 1.0 \times 10^{-5} M$

[Anil] =
$$1.0 \times 10^{-2} M$$

AcOH : $H2O = 60 : 40 (V/V)$
Temp. = $40^{0} c$

[Os(VIII)] x 10 ⁵ M	$K_{obs} X 10^4 s^{-1}$
0.5	2.516
1.0	2.997
1.5	3.495
2.0	3.753



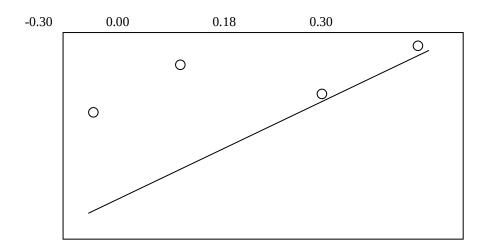
6.2.Effect of varying Ru(III) concentration

The reaction was followed with varying concentration of Ruthenium trichloride and by keeping other [reactants] constant. It was found that as the concentration of Ruthenium trichloride increased, the rate constant also increased. The plot of log [Ru(III)] against logKobs. gave a straight line of slope of 0.34 (Table 5.5; Fig. 5)

Table – 5.5

$[CAT] = 1.0 \times 10^{-3} M$	[Anil] = $1.0 \times 10^{-2} M$
$[OsO_4] = 1.0 \ge 10^{-5} M$	AcOH : H2O = 60 : 40 (V/V)
	Temp. = 40° c
1(III) v 10 ⁵ M	$K = X 10^4 \mathrm{s}^{-1}$

$[Ru(III)] \times 10^{3} M$	K_{obs} . X 10 ⁴ s ⁻¹
0.5	2.516
1.0	2.997
1.5	3.495
2.0	3.753



log k_{obs}

Fig. 3

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