Electron Transfer Reactions: a Treatise

V. Jagannadham

Department of Chemistry, Osmania University, Hyderabad, 500007, India

Abstract

In the introduction the concept of oxidation-reduction is well defined. In the discussion a systematic classification of electron transfer reactions is made based on the nature of the redox partners. There were total of eight sets of redox reactions under two main headings four sets for each class of outer sphere electron transfer reactions, and inner sphere electron transfer reactions: containing (i) Inorganic oxidant and inorganic substrate, (ii) Inorganic oxidant and organic substrate, (iii) Organic oxidant and organic substrate, and (iv) Organic oxidant and inorganic substrate. At least one example for each set of redox partners is given and well explained. At the end an example of atom transfer reaction, inner sphere synchronous three and four electron transfer reactions are included.

Keywords

Electron Transfer Reactions, Inner Sphere Electron Transfer Reactions, Outer Sphere Electron Transfer Reactions

1. Introduction

First one must understand the meaning of oxidation and reduction before going to define the concept. The word redox reaction envisages transfer of electron(s), hydrogen atom, hydride ion, oxygen atom, and chlorine atom or chlorinium ion between two redox partners. The word oxidation (of course it necessarily involves concurrent reduction) has been defined in many ways. The most prominent definitions, however, refer to addition of oxygen, removal of hydrogen, or transfer of electron(s), of which the last one is the most favored one at present time. There are several thousands of research publications in literature on different kinds of redox processes. Only some representative publications related to this review were presented and discussed. To understand the nature of the mechanism of a redox reaction it is necessary to find out whether an atom or group or electron transfer has occurred, which atoms are transferred or how many electrons are transferred and what the transition states of all steps are like. For obtaining such information kinetic measurements are useful [1-6]. Acceptance of one electron by an oxidant is equivalent to acceptance of a hydrogen atom and acceptance of two electrons is equivalent to acceptance of a hydride ion. In general most of the redox reactions can be classified into two broad categories following the scheme of Wiberg [7].

1. Direct electron transfer reactions.
2. Atom, group or ion transfer reactions.

1.1. Direct Electron Transfer Reactions

Electron transfer reactions may be classified as one equivalent or two equivalent reactions depending on the number electrons being transferred during the oxidation. This kind of classification was first given by Kirk and Brown [1] in 1928. The oxidants like Mn(III), Ce(IV), Fe(III), Cu(III), Ag(II), Co(III), Ni(III) etc act exclusively as one electron abstractors, while Pb(IV), Ru(III), Ag(III), IO4−, Cl2, N-halo compounds, hypohalous acids, peroxo-anions etc are two-equivalent oxidants. Some of the oxidants like Mn(VII), Cr(VI), Pt(IV), Tl(III), V(V) etc act as both one and two equivalent oxidants depending on the conditions employed. In some special cases Cr(VI) and Mn(VII) acted as three and four electron oxidants respectively.

It is generally observed that oxidations by two-equivalent oxidants proceed much faster than one-equivalent oxidants, because during two-equivalent oxidations, no high-energy free radicals are formed and there is overall twice the free energy change. For one-equivalent oxidants the rate is affected by the addition of vinyl monomers or mercuric chloride or in the presence of oxygen, but not in the case of two-equivalent oxidations. These two different reaction pathways can also be distinguished from product analysis and stoichiometry of certain reactions. Direct electron transfer is more likely with simple one-electron oxidants like Ce(IV), Co(III) and Fe(III). But with other one or two-electron oxidants either hydrogen atom or hydride ion abstraction takes place. Oxidation of hydrazine or sulfite has been generally used to distinguish between one and two-electron oxidants [8, 9] on the basis of the products formed. One-equivalent oxidants tend to convert sulfite to dithionate and hydrazine to ammonia and nitrogen according to the following reaction mechanisms: (a and b, eqns. 1 - 5)

\[
\begin{align*}
\text{(1)} & \quad \text{M}^{n+} + \text{SO}_3^{2-} \rightarrow \text{M}^{(n-1)+} + \text{SO}_3^{5-} \\
\text{(2)} & \quad 2\text{SO}_3^{5-} \rightarrow \text{S}_2\text{O}_6^{2-} \quad \text{(dithionate)}
\end{align*}
\]

\[
\begin{align*}
\text{(3)} & \quad \text{M}^{n+} + \text{N}_2\text{H}_4 \rightarrow \text{M}^{(n-1)+} + \bullet \text{N}_2\text{H}_3 + \text{H}^+
\end{align*}
\]
The two equivalent oxidants have been found to react in the following manner: (c and d, eqns. 6 - 11)

c) \( M^{n+} + SO_3^{2-} \rightarrow M^{(n-2)+} + SO_3 \) (6)
\[ SO_3 + H_2O \rightarrow SO_2^{−} + 2H^+ \] (7)

d) \( M^{n+} + N_2H_4 \rightarrow M^{(n-2)+} + N_2 + 2H^+ \) (8)
\[ 2N_2H_2 \rightarrow N_2 + N_2H_4 \] (9)
\[ N_2H_4 \rightarrow NH_3 + HN_3 \] (10)

The redox reactions involving metal ions can also be classified based on whether the transfer of electrons involves ionic sphere or coordination sphere of the metal ion as
1. Outer-sphere electron transfer reactions
2. Inner-sphere electron transfer reactions

1.1.1. Outer-Sphere Electron Transfer Reactions

In reactions of this type the metal ion retains its full coordination shell and there is a direct electron transfer from the reductant to the oxidant. The electron given by the reducing agent must be transferred from the primary bond system of one complex to that of the other. The essential feature of this mechanism is that there is no transfer of ligands between the reactants. Kinetically the rate of reaction is faster than the rate of substitution of ligands, or the ligand exchange or displacement is slower than electron transfer. Typical examples of this type are: (eqns. 12 and 13)

\[ \text{Fe}(CN)_{5}^{2−} + 3\text{I}_2 \rightarrow \text{Fe}(CN)_{5}^{−} + 3\text{I}_2 \] (12)
\[ \text{Ru}(NH_3)_5^{2+} + 3\text{Ru} (ND)(OAc) \rightarrow \text{Ru}(NH_3)_5^{3+} + 3\text{Ru} (ND)(OAc) \] (13)

The evidence for this type of mechanism is usually given by a rate law corresponding to an activated complex containing all the ligands in the coordination sphere of both the metal ions. (eqn. 14)

\[ \text{rate} = k[\text{Fe}(CN)_{5}^{2−}] [\text{I}_2] \] (14)

Thus the oxidation of organic compounds proceeds by the outer sphere mechanism, if the oxidant is substitution inert.

1.1.2. Inner-Sphere Electron Transfer Reactions

The inner sphere mechanism may be seen, if one of the reactants is labile. In reactions of this class, electron transfers are preceded by the substitution of coordination sphere of one of the ions, with the formation of bridged intermediate in which the two reactants are linked by a common ligand. In this case, ligand displacement is faster than electron transfer process. A typical example of this type is given in eqn. 15

\[ Cr^{3+} + (NH_3)_5Co^{3+} + X + 5H^+ \rightarrow Cr^{3+} + X + Co^{2+} + 5NH_3 \] (15)

Where \( X = H_2O, OH^-, Cl^-, OAc^- \) etc,

Electron transfer takes place through the bridged intermediate activated complex. Eqn. 16 to yield the above products. The Cr(II) and Co(II) ions are both labile and the aquapenta-amino cobalt(II) ion which is formed as the product, loses amine ligands rapidly.

\[ [(NH_3)_5Co^{3+} X \cdots \cdots \cdots Cr^{2+} (H_2O)_3] \] (16)

In the oxidation of \( \text{Ca(NO)}_{5}^{2−} \) by \( \text{Fe}(CN)_{5}^{2−} \), where both metal ions are substitution inert, a product believed to have a bridged structure. Eqn. 17

\[ [(CN)_3\text{Co}^{3+} \cdots \cdots \cdots \text{CNFe}^{2+} (CN)_3] \] (17)

has actually been isolated. An example of this type was the oxidation of DMSO by Ce(IV) [10], which involves a reversible rapid complex formation between them with a kinetic proof yielding the final products in the rate determining step, for which the rate law was given by the equation 18

\[ \frac{−d[Ce(IV)]}{dt} = \frac{k_1[Ce(IV)][DMSO]}{1 + k_2[DMSO]} \] (18)

This implies that the complex was formed between DMSO and Ce(IV) before the electron was transferred from the substrate to the oxidant. Oxidation by metal ions that proceeds via an inner-sphere electron transfer mechanism, when the substrate acts as a ligand or contains donor groups such as an oxo or −OH or −COOH that readily form complexes with metal ions.

Inner-sphere electron transfer mechanisms have also been shown for two redox electron reactions involving metal complexes, an example of this type is the Pt(II)-Pt(IV) exchange reaction [11].

However the oxidizing agents like Cr(VI) [12] and Mn(VII) [13] have also shown a synchronous three electron and four electron redox change, when more than one reductant was used. This type of redox process was first termed as “Co-oxidation” by Hasan and Roček [12] in 1972 in Cr(VI)-propane-2-ol-oxalic acid system. A similar study was reported in 1986 using Mn(VII) wherein a synchronous four electron transfer took place in Mn(VII)-propane-2-ol-lactic acid system [13].

Apart from the above metal ion - metal ion and metal ion - organic molecule reactions there were several reports on redox reactions wherein both the redox partners are organic molecules, which will be discussed in detail in the latter sections.

1.2. Atom, Group or Ion Transfer Reactions

These reactions include
1. Hydrogen, oxygen, chlorine atom or group reactions
2. Hydrogen atom abstraction reactions
3. Hydride ion abstraction reaction
4. Proton abstraction from a precursor-ester or complex.
5. Displacement mechanisms.
6. Addition-elimination mechanisms.

1.2.1. Hydrogen, Oxygen, Chlorine Atom or Group Transfer Reactions

When redox reactions take place in aqueous medium the transfer of atom or group is assumed. For example Fe^{2+} ion may act as a reducing agent by transferring a hydrogen atom from its hydration shell to a substrate eqn.19

\[ \text{Fe}(H_2O)_3^{2+} + R^- \rightarrow \text{Fe}(H_2O)_3^+ OH^{2+} + R - H \] (19)
In many redox reactions of oxyanions, using $^{18}$O labelling, for example has showed oxygen atom transfer. A well-known oxygen atom transfer reaction was hypochlorite oxidation of nitrite ion. Taube [14] proposed the following mechanism for this reaction eqn. 20

$$NO_2^- + ^{18}OCl^- \rightarrow [O_2N\cdots\cdots^{18}O\cdots\cdotsCl]^\text{+} \rightarrow O_2N^{18}O^- + Cl^- \quad (20)$$

The activated complex in this case cleaves to produce nitrate and chloride ions. With CrCl$_2^+$, FeCl$_2^+$, AuCl$_4^-$ and [Co(NH$_3$)$_5$Cl]$^2+$ as oxidizing agents the transfer of chlorine atom has been shown to occur similarly [15]. By using Co(NH$_3$)$_5$X as an oxidant, the transfer of X has been demonstrated, for X = NCS$^-$, N$_3^-$, PO$_4^{3-}$, CH$_3$COO$^-$, C$_2$O$_4^{2-}$ and SO$_4^{2-}$ [16].

1.2.2. Hydrogen Atom Abstraction Reactions

In all homolytic cleavages of C-H bonds of organic substrates, abstraction of hydrogen is energetically more favored. Free radical substitution of alkanes (e.g. CH$_4$) with chlorine in the presence of light or heat, one of the propagation steps involves the abstraction of hydrogen, eqn. 21

$$CH_4 + Cl^- \rightarrow \text{CH}_2^- + HCl \quad (21)$$

Hydrogen atom abstraction occurs most readily from $\alpha$-position to un-dissociated groups, particularly at allylic hydrogens because of resonance stabilization of the resulting free radical, eqn. 22.

$$>C-C-H \rightarrow >C-C^-< \leftrightarrow >C^-C=C< \quad (22)$$

Similarly benzylic hydrogens are also readily removed in the case of benzaldehyde, toluene and tetralin which all produce carbon centered radicals, stabilized by resonance interactions with aromatic ring (Scheme 1).

If both aliphatic and aromatic hydrogens are present in a molecule, aliphatic hydrogens are preferentially abstracted, because of the higher dissociation energy of aromatic C-H bond than that of aliphatic C-H bond. These kinds of hydrogen atom abstraction reactions are very frequently observed [17]. Hydrogen atom abstraction can also occur from $\alpha$-position of alcohols [18] with sufficiently reactive reagents like hydrogen atom or hydroxyl radical eqn. 23 In the oxidation of aldehydes by Cr(VI), Roček and Ng [19] proposed the transfer of a hydrogen atom from the aldehyde carbon to the oxidant.

$$CH_3CH_2OH + OH^* (H^+) \rightarrow CH_3^*CHOH + H_2O(H_2) \quad (23)$$

1.2.3. Hydride Ion Abstraction Reaction

When C-H bond cleaves heterolytically, the hydride ion abstraction can usually occur. Hydride ion abstractions have been proposed in explaining the mechanism of large number of organic reactions such as Meerwein Ponnendorf Verley reduction [20], the Cannizaro reaction [21], the Tischenko reaction [22], the Leukart reaction [23], the Sommelet conversion of primary alkyl halides to aldehydes [24, 25], the transition metal ion catalyzed oxidation of organic compounds [26] and many reactions in which a carboxylation abstracts a hydride ion intermolecularly or intramolecularly belong to this category [27, 28].

As an example the base catalyzed disproportion of aldehydes lacking an $\alpha$-hydrogen atom (Cannizaro reaction) can be represented as shown in Scheme 2:

![Scheme 2. Base catalyzed disproportion of aldehyde](image)

When the Cannizzaro reaction is conducted with ArCDO (deutero aldehyde), the alcohol was found to contain two deutero atoms, thereby establishing that the sequence involves a direct transfer of hydride ion from one aldehyde molecule to the other. Hydride ion transfer mechanisms have also been proposed in the oxidation of benzyl alcohol by acid KMnO$_4$ [29], bromine [30] and also in the oxidation of propane-2-ol by Ru(VIII) [31].

An oxidation mechanism involving either a hydride ion abstraction or a hydrogen atom abstraction is based on a logical deduction of indirect evidence. If the hydride ion is removed in the rate determining step, then a negative $\rho$ value of high magnitude and a fairly large kinetic isotope effect can usually be observed. On the other hand hydrogen atom abstraction reactions generally exhibit moderate isotope effects and $\rho$ values in the range from −0.5 to −1.5 [32, 33].

![Scheme 3. Proton abstraction from a precursor-ester or complex](image)
A number of oxidations are involved in the formation of an ester intermediate (usually of an inorganic acid) [34], and then the cleavage of this intermediate by proton loss to any available base. An example of this mechanism can be seen in the oxidation of aromatic aldehydes by permanganate (Scheme 3) [35].

The formation of intermediate ester was supported by the presence of $^{18}$O in the acid when KMn$^{18}$O$_4$ was used.

The oxidation of glycols by Pb(IV) acetate [36], and periodic acid [37], also followed this pattern, but the positive leaving group is carbon instead of hydrogen. It is seen that whenever proton loss occurs, electron-withdrawing groups accelerate the reaction rate. For instance, Ru(III) catalyzed substrate and then part of it is lost Scheme 5:

$$\text{organic molecules} \rightarrow \text{oxidizing agent} \rightarrow \text{oxidation product}$$

There are some reactions [18, 41] that follow this pattern of addition of an oxidizing agent and the loss of part of the agent, usually in a different oxidation state.

When an oxidizing agent enters the reaction in its oxidized state – 1 and the $\text{OH}^-$ departs with its oxygen in the – 2 state, so it is reduced and the substrate is oxidized. There are some reactions [18, 41] that follow this pattern of addition of an oxidizing agent and the loss of part of the agent, usually in a different oxidation state.

A mechanism is the actual process by which a reaction takes place - which bonds are broken, in what order, how many steps are involved, the relative rate of each step, etc. In order to state a mechanism completely: one should have to specify the positions of all atoms, including those in solvent molecules, and the energy of the system, at every point in the process. A proposed mechanism must explain all the experimental facts available. It is always subject to change as new experimental facts are discovered. The usual course is that the gross features of a mechanism are the first to be known and then increasing attention is paid to finer details. The tendency is always to probe more deeply, to get more detailed descriptions.

Although for most reactions gross mechanisms can be written today with a good degree of assurance, no mechanism is known completely. There is much about the fine details, which is still puzzling, and for some reactions even the gross mechanism is not yet clear. The problems involved are difficult because there are so many variables. Many examples are known where reactions proceed by different mechanisms under different conditions. In some cases there are several proposed mechanisms, each of which completely explains all the data.

The original observation that the results of studies on several redox reactions lend themselves to simple generalizations was satisfying and provided confidence that the mechanisms for more complex reactions might also be understood. However, the exceptions to these generalizations show that it is dangerous to assume that even simple reaction mechanisms can be easily understood. These exceptions have the potential to provide new insight into the mechanism for several redox reactions by carrying new experiments.

As said in the above paragraphs, a good Physical Organic Chemist is always eager in finding the way to reach a specific goal for the problem that he has in his mind. In this course of time, this review would help for writing gross mechanisms of several synthetically hitherto known important reactions and paves the way to get an insight into finer details.

A criticism which is anticipated from the readers of this review is about the inclusion of the first two sections of discussion on inner and outer-sphere electron transfer reactions between inorganic redox partners. One may ask where the organic chemistry or physical-organic chemistry here is. This criticism may spring from different points of view where the lines should be drawn which divide and bound the review. Some topics like these first two sections are treated as essential core material in this review. The title itself of the review warrants inclusion of such topics, because they involve all the essential features of kinetics like stoichiometry of the reaction, and effect of [H$^+$], ionic strength, dielectric constant, temperature, [catalyst] and [complexing agent] on rates. They will be very useful to start with for a beginning kineticist.

2. Scope of the Review

The chemical mechanisms of several oxidation-reduction reactions have been well studied and are continued to be studied. The investigations on the kinetics and mechanism of these reactions have played an important role in the development of Physical Organic Chemistry.

3. Discussion
3.1. Outer-Sphere Electron Transfer Reaction between an Inorganic Oxidant and Inorganic Substrates

Establishment of the nature of the reactive species of the Ni(III) ion - Kinetics of oxidation of iodide and thiocyanate ions by nickel oxyhydroxide in aqueous dilute sulfuric acid medium [42]:

Solid NiO(OH) was prepared by the oxidation of Ni(II) ions by bromine in aqueous alkali. The reaction of this NiO(OH) with iodide and thiocyanate ions in aqueous H2SO4 medium was overall second order, being first order each in [oxidant] and [reductant]. The rate law at constant [H+] was found to be as given in eqn. 24:

\[ \text{Rate} = k [\text{Ni(III)}][X] \]  
(24)

Where k is second order rate constant and X = iodide or thiocyanate ion.

The consumption of Ni(III) was determined in solutions containing excess of [Ni(III)] (2 mM) over [iodide] (1 mM) with different [H+]. Assuming that all iodide was oxidized completely, \( \frac{d[Ni(III)]}{dt} \) was found to be 1 in the range 0.25 to 1.0 M of \( \text{H}^+ \). It was therefore concluded that the overall reaction could be as follows: eqn. 25:

\[ 2\text{Ni(III)} + 2F^- \rightarrow 2\text{Ni(II)} + I_2 \]  
(25)

The rates of oxidation were affected by added \( \text{H}^+ \) at constant ionic strength. The order in [H+] was found to be two, suggesting that Ni(OH)2+ may be the reactive species as given in the equation 26.

\[ \text{Ni}(O)\text{OH} + 2\text{H}^+ \rightarrow \text{Ni(OH)}_{2}^+ + \text{H}_2\text{O} \]  
(26)

The rates of oxidation were affected by added SO42- ions at constant [H+] and the experimental data obeyed Debye-Hückel equation: eqn. 27:

\[ \log k = \log k_0 + 1.01Z_aZ_b\sqrt{\mu} \]  
(27)

indicating that the rate determining step to involve an ion-ion. From the value (- 1.90) of the slope of the linear plot (r = 0.987) of log k vs. \( \sqrt{\mu} \) it was concluded that Ni(OH)2+ was assumed to be the reactive species of Ni(III), being iodide to be as I. A reaction between two species that proceeds via a polar transition state should be sensitive to variation of solvent polarity. If the reaction was of the same sequence as shown below (eqn. 28) should have a solvent effect on rates.

\[ \Gamma^- + \text{Ni(OH)}_{2}^+ \rightarrow \text{Ni(OH)Ni•••••I}^+ \]  
(28)

In fact the rates were found to be dependent on solvent polarity and obeyed the eqn. 29.

\[ \log k = \log k_o - \frac{NZ_aZ_be^2}{RTD\Delta r} \]  
(29)

Here \( r_c \) is the distance of closest approach of the reactants in the transition state. Kinetic evidence for no complex formation between Ni(OH)2+ and either iodide or thiocyanate ions suggests that the reaction was an outer-sphere electron-transfer process. The distance of closest approach for iodide ion is smaller (2.97 Å) than that for thiocyanate (3.60 Å), which is reasonable to say that the later is bigger in size and difficult to approach the oxidant in the transition state. The high negative \( \Delta S^\theta \) (-132 J. K-1. mol-1) in the reaction of thiocyanate was due to HCN produced as the end product which is a source of \( \text{H}^+ \) to be solvated, and that causes freezing of solvent, whereas the positive \( \Delta S^\theta \) (23.0 J. K-1. mol-1) for the reaction of iodide is due to oxidation of solva(hydra)ted I to give I2 and resulting in the generation of free solvent molecules from frozen shell around I. Hence, based on the above arguments the mechanism can be depicted as shown in scheme 6.

\[ \text{NiO(OH)} + 2\text{H}^+ \rightarrow \text{Ni(OH)}_{2}^+ + \text{H}_2\text{O} \]

\[ \text{Ru(III)} \rightarrow \text{Ru(V)} + \text{Ru(III)} \]

\[ \text{Ni(OH)}_{2}^+ + \text{I}^- \rightarrow \text{Ni(OH)} + \text{I}^+ \]

\[ \text{I}^+ + \text{I}^- \rightarrow \text{I}_2 \]

Scheme 6. Reaction between Ni(III) and iodide ion

Apart from this, effect of ionic strength on the Ru(III) catalyzed decomposition of Ni(III) was studied in aqueous sulfuric acid, at varying ionic strengths was overall second order, first order each in [Ru(III)] and [Ni(III)]. Effect of [H+] showed that two moles of acid was required to generate Ni(OH)2+ from Ni(O)OH. The slope of the linear plot according to Debye-Hückel equation was 5.7. As RuCl3 exists in aqueous acid medium as Ru3+ it may be concluded that the reactive species of Ni(III) as Ni(OH)2+ [43]. Hence in this study the effect of ionic strength on Ni(III) decomposition provides yet additional evidence for the participation of Ni(OH)2+ as the reactive species in aqueous acid medium. Based on the foregoing discussion the mechanism of Ru(III)-catalyzed decomposition of Ni(III) could be depicted as shown Scheme 7:

\[ \text{Ni(OH)}_{2}^+ + \text{Ru(III)} \rightarrow \text{Ru(IV)} + \text{Ni(OH)}^+ \]

\[ 2\text{Ru(IV)} + \text{H}_2\text{O} \rightarrow 2\text{Ru(III)} + 2\text{H}^+ + \frac{1}{2}\text{O}_2 \text{or} \]

\[ 2\text{Ru(IV)} \rightarrow \text{Ru(V)} + \text{Ru(III)} \]

\[ \text{Ru(V)} + \text{H}_2\text{O} \rightarrow \text{Ru(III)} + 2\text{H}^+ + \frac{1}{2}\text{O}_2 \]

Scheme 7. Self decomposition Ni(III) ion in presence of Ru(III)

3.2. Inner-Sphere Electron Transfer Reaction between Inorganic Oxidant and Inorganic Substrates

One-electron reduction of Ni(III) by hydroxylamine and hydrazine via addition/elimination - an example of an inorganic inner sphere electron transfer reaction [44]:

The rates of oxidations in aqueous solution of NH2OH and N2H4 by Ni(III) ion were measured iodometrically in the presence of different \( \text{Ni(O)}\text{OH} \) and \( \text{N}_2\text{H}_4 \) to give I2 and resulting in the generation of free solvent molecules from frozen shell around I. Hence, based on the above arguments the mechanism can be depicted as shown in scheme 8.

\[ \text{Ni(O)}\text{OH} + 2\text{H}^+ \rightarrow \text{Ni(O)}\text{OH}^+ + \text{H}_2\text{O} \]

\[ \text{Ni(O)}\text{OH}^+ + \text{Ni(III)} \rightarrow \text{Ni(IV)} + \text{Ni(II)} \]

\[ \text{Ni(IV)} + \text{Ni(II)} \rightarrow \text{Ni(III)} \]

\[ \text{Ni(III)} + \text{Ni(II)} \rightarrow \text{Ni(IV)} + \text{Ni(II)} \]

Scheme 8. Self decomposition Ni(III) ion in presence of Ru(III)
rates were acid-catalyzed with a total third order in [H\(^+\)] accounting for Ni(OH)\(^{2+}\) and 'NH\(_2\)OH were the redox partners in the rate determining step. And the rates were marginally affected by added SO\(_4\)\(^{2-}\) ions. The substituent effects of OH and NH\(_2\) on rates and activation parameters were discussed as follows: The rate of oxidation for NH\(_2\)OH was greater than that of H\(_2\)NNH\(_2\) based on the fact that only removal of one of the hydrogens as H\(^+\) from NH\(_2\) and forming a free radical of the type 'NHOH. The presence of free radicals was tested by positive polymerization of acryl amide. The non-involvement of the O-H bond breakage or removal of one of the electron of lone pair on nitrogen by the oxidant was ruled out because the first fact should support the non-reactivity of hydrazine since it does not contain any O-H bond and the second fact should support the higher reactivity of hydrazine since the +I effect of NH\(_2\) > than that of OH. There was no effect of temperature on K, the formation constant. However the rate constant for the slow step, the k increased with increase in temperature. The very high negative \(\Delta S\) values (~ - 135 J (mol. K\(^{-1}\)) were in good agreement with the liberation of a proton in the rate-determining step, which may be hydrated resulting in the freezing of solvent. The stiochiometry of the reaction was found to be oxidant : reductant = 2 : 1. Based on the above experimental observations a suitable mechanism was proposed as shown in scheme 8. The first two steps account for a total third order in [H\(^+\)], which was in fact observed experimentally.

\[
\text{Ni(OH)OH} + 2H^+ \rightarrow \text{Ni(OH)}^{2+} + H_2O
\]
\[
\text{NH}_2\text{OH} + H^+ \rightarrow \text{NH}_2\text{OH}^+
\]
\[
\text{Ni(OH)OH}_2^{2+} + \text{NH}_2\text{OH} \equiv \text{Ni(NH}_2\text{OH)}^{2+} + H_2O
\]
\[
\text{Ni(NH}_2\text{OH)}^{2+} \text{slow} \rightarrow \text{Ni}^{2+} + \text{NHOH} + H^+
\]
\[
\text{NHOH} + \text{Ni(OH)}^{2+} \text{fast} \rightarrow \text{HNO} + \text{Ni}^{2+} + H_2O
\]
\[
2\text{HNO} \rightarrow \text{N}_2O + H_2O
\]

Scheme 8. Reaction between NH\(_2\)OH and Ni(III)

The kinetic evidence for the formation of the complex [Ni(NH\(_2\)OH\(^{2+}\)] was found from the linear plot of \(1/kvs.1/\text{[NH}_2\text{OH]}\) with a positive intercept on y-axis.

3.3. Outer-Sphere Electron Transfer Reaction between an Inorganic Oxidant and Organic Substrates

3.3.1. Chromium(VI) ion

Cr(VI) oxidation of benzaldehydes [45], benzaldoximes [46], aliphatic esters [47,48], toluenes [49], anisoles [50,51], dimethyl sulfoxide [52] and phenols [53] was carried out in various media. All these reactions obeyed total second order kinetics. Identification of products, discussion of mechanism and application of Hammett and Taft equations were some of the salient features of these reactions.

3.3.1.1. Benzaldehydes [45]

A search for resonance effects in the kinetics of chromic acid oxidation of a series of benzaldehydes was made: The kinetics of oxidation of a series of eight meta and para-substituted benzenalddehydes by chromic acid has been re-investigated in acetic acid-water mixture with a view on the search for resonance effects of para-substituents. The overall order of the reaction was two, first order in each reactant at constant acidity. The products of oxidation were the corresponding carboxylic acids. Hammett’s reaction constants (\(\rho_m\)) and (\(\rho_p\)) for meta and para-substituents were calculated separately as 0.851 and 1.23 respectively. The reason for the higher \(\rho_p\) value than \(\rho_m\) was due to the direct resonance interaction of para-substituents with the reaction center. The individual contributions of inductive and resonance effects were calculated for para-substituents. The para-substituents did not fit in to the correlation of meta-substituents, instead they correlated well with \(\sigma_{p} + \sigma_{I}^{\prime}\). Based on these results, we proposed a mechanism as shown in scheme 9, which shows a heterolytic cleavage of C-H bond, and which is different from that proposed by Wiberg and Szeimies assuming the homolytic cleavage of aldehydic C-H bond for which one could expect a very low \(\rho\) value:

\[
\text{PhCH=NOH} + \text{Cr(VI)} \rightarrow \text{PhCH(ON)} + \text{Cr(IV)}
\]

\[
\text{PhCH(ON)} \rightarrow \text{PhCHO} + \text{HNO}
\]

Scheme 9. Reaction between aldehyde and Cr(VI)

3.3.1.2. Benzaldoximes [46]

Kinetics of chromic acid oxidation of benzaldoxime in acetic acid-water mixtures: Kinetics of oxidation of a series of six benzaldoximes by chromic acid in 30% (vol./vol.) acetic acid-water mixture was measured at constant acidity and ionic strength. The reaction was over-all second order, being first-order in each reactant. The rates of oxidation were not affected by the addition of salts like Na\(_2\)SO\(_4\) and NaHSO\(_4\).
The reaction was acid catalyzed with an order in \([H^+]\) to be four and the rate of oxidation increased with increase in percentage of acetic acid. Hammett's equation was applicable with a reaction constant \(\rho = -2.00\) and a correlation coefficient \(r = 0.998\) indicating an electron deficient center in the transition state. The Arrhenius plots for four benzoaldehydes met at 325 K, which was well above the experimental temperature range indicating the reaction series was considered to be enthalpy-controlled process. Formation of a very unstable cyclic chromate ester in a fast step and subsequent decomposition of this in a slow step had been suggested as a probable mechanism as shown in scheme 10:

3.3.1.3. Aliphatic Esters [47]

Kinetics and mechanism of oxidation of some aliphatic esters by chromic acid in the presence and absence of oxalic acid in acetic acid-water medium: The oxidation kinetics of aliphatic esters by chromic acid was examined in aqueous acetic acid in the presence and absence of oxalic acid. In the absence of oxalic acid the reaction was overall second order, being first order in each reactant. It was acid catalyzed and the reaction rates were independent of added salts and it was dependent upon solvent polarity. In the presence of oxalic acid the rate of oxidation increased appreciably. The products of oxidation in the absence of oxalic acid were acetic acid and the corresponding aldehydes (Me2CO in the case of Me2CHOAc). The formation of free radicals was observed only in the presence of oxalic acid. There could be a possibility of ester hydrolysis before the oxidation could take place. The kinetic study was done in 20% aqueous acetic acid in presence of 3.0 M sulfuric acid. Under these conditions the observed rate constant for the hydrolysis of methyl acetate was found to be 7.88 X 10^{-4} min^{-1}. Since this was much lower than the observed rate of oxidation of methyl acetate by Cr(VI) (1.0 X 10^{-2} min^{-1}), a direct oxidation of the ester had been envisaged. Activation parameters were calculated and from the isokinetic plot it was concluded that the reaction sequence to be entropy controlled one. A suitable mechanism suggested in the absence of oxalic acid (in the presence of oxalic acid will be discussed later under the heading of direct three electron transfer reactions) with the formation of an initial very unstable complex in the fast step which later gives acetic acid and the corresponding carbonyl compound as the products (scheme 11). The other steps were essentially similar to those mentioned above.

\[
\text{CH}_3\text{COOCH}_2\text{R} + \text{HCrO}_4^- \rightarrow \text{complex} \\
\text{complex} \xrightarrow{\text{slow}(k)} \text{CH}_3\text{COOH} + \text{RCHO} + \text{Cr(IV)} \\
\text{Cr(IV)} + \text{Cr(VI)} \xrightarrow{\text{fast}} 2\text{Cr(V)}
\]

or

\[
2\text{Cr(IV)} \xrightarrow{\text{fast}} \text{Cr(V)} + \text{Cr(III)} \\
\text{CH}_3\text{COOCH}_2\text{R} + \text{Cr(IV)} \xrightarrow{\text{H}_2\text{CrO}_4\text{slow}(k)} \text{CH}_3\text{COOH} + \text{RCHO} + 2\text{H}^+ + \text{Cr(III)}
\]

Scheme 11. Reaction between ester and Cr(VI)

3.3.1.4. Application of Taft Equation [48]

Taft four parameter equation as a mechanistic tool in chromic acid oxidation of aliphatic esters: The Taft four parameter equation was used to correlate the kinetic data in the chromic acid oxidation of CH3COOCHRR1 (R = H, Me; R1 = H, Me, Et, Me2CH, Pr). Both polar and steric effects were operative in this reaction. The mechanism involved attack of a positively charged species at the C-H bond of CHRR1. The Taft reaction constants \(\rho^*\) and \(\sigma^*\) were found to be -7.14 and 2.72 respectively indicating the electron withdrawing and more bulky groups favoring the reaction.

3.3.1.5. Applicability of the Taft four parameter equation [49]

Kinetic study of chromic acid oxidation of eight o-substituted toluenes in acetic acid-water mixtures: The kinetics of the Cr(VI) oxidation of o-RC6H4Me (R = MeO, EtO, Me, F, Cl, Br, I, NO2), to the corresponding o-RC6H4CHO in 80% aqueous acetic acid and 1 M HClO4 had poor linear free energy correlation with either \(\rho^*\) or \(E_a\). A four parameter Taft linear free energy relationship, \(\log k/k_0 = \rho^*\sigma^* + \delta E_a\) was found to be applicable satisfactorily. The polar (P) and steric (S) effects were quantitatively separated. And the polar and steric interaction energies were evaluated by the use of the equations \(P_{\text{polar}} = -2.303 \rho^*\sigma^* RT\) and \(S_{\text{steric}} = -2.303 \delta E_a RT\). The total polar and steric interaction energies (P + S) were well correlated with the \(\Delta G^*\).

Scheme 12. Reaction between anisole and Cr(VI)

3.3.1.6. 1,10-Phenanthroline Catalyzed and Un-Catalyzed Oxidation of Anisole by Chromic Acid In Acetic Acid-Water Mixture [50]

The hydroxylation of anisole, to give o - and p -
HOC₆H₄OMe, by H₂CrO₄ in aqueous acetic acid-H₂SO₄ was accelerated by the presence of 1,10-phenanthroline as catalyst. The rate-limiting step involved the attack of chromium-phenanthroline complex on anisole. Under otherwise similar conditions there was no reaction between 1,10-phenanthroline and Cr(VI). The mechanism in the absence of 1,10-phenanthroline is shown in scheme 12.

An LFER for the rates of the title reaction, which was first order in each of substrate and oxidizing agent at constant H⁺ concentration and ionic strength, showed that the site of attack was the C atom ortho to the MeO group. The reaction mechanism was suggested as shown in scheme 14.

3.3.1.8. Kinetics and Mechanism of Oxidation of Dimethyl Sulfoxide by Chromic Acid in Aqueous Acid Medium [52]

The title reaction showed first order dependence each on [oxidant] and [substrate]. The reaction was acid-catalyzed and showed fourth order dependence in [H⁺]. The rate was affected to a small extent by added sulfate ions and was dependent upon solvent polarity. Added Mn(II) retards the rate while the rate was enhanced if 1,10-phenanthroline was also present along with Mn(II). The rate retarding effect of Mn(II) is expected if Mn(II) ions catalyze the disproportion of the intermediate valency states of Cr(VI) i.e. Cr(IV) or Cr(V). However added 1,10-phenanthroline in the presence of Mn(II) ions catalyzed the rate of oxidation. This might be due to the fact that the added Mn(II) is removed as its complex and the disproportion of the Cr(IV) or Cr(V) was not diminished hence the higher rate of oxidation. This was an additional evidence in Cr(VI) oxidations to show the formation of Cr(IV) and Cr(V) as intermediates. The sole reaction product was identified as dimethyl sulfone. Based on the above discussion, the mechanism could be written as shown in scheme 15 with an initial formation of the complex in a fast step and subsequent decomposition of the complex in the rate-determining step. The remaining steps being similar to those as mentioned in the section 3.3.1.2 above for benzaldoximes.

3.3.1.9. Kinetics and Mechanism of Oxidation of Phenols by Chromic Acid in acetic Acid-Water Mixture [53]

The title reactions were first order each in [chromic acid] and [phenol]. The phenols having electron-donating substituents in the benzene ring accelerated the rates and vice versa indicating an electron deficient center with a Hammett
3.3.2. Nickel(III) ion

Reactions of Ni(III) ion with aliphatic primary [54] and secondary alcohols [55] and glycols [56] were studied. The total order of the reaction was found to be two. The oxidation process of primary and secondary alcohols went through the abstraction of a α-hydrogen. A synchronous carbon-carbon and oxygen-hydrogen bond cleavage was shown to take place in the case of glycols based on the kinetic and entropic evidences. The products of oxidation were the corresponding carbonyl compounds.

3.3.2.1. One-electron Reduction of Ni(III) Ion by Aliphatic Primary Alcohols in Aqueous H2SO4 Medium. A Kinetic and Mechanistic Approach [54]

The rates of reactions of a series of seven aliphatic alcohols in aqueous solution with Ni(III) ion were studied isomerically at constant acidity and ionic strength under pseudo-first-order conditions. The overall order of the reaction was two: first order each in [oxidant] and [reductant]. The rate of reduction of Ni(III) was not affected by added salts like (NH4)2SO4 and K2SO4. The reaction was acid catalyzed at constant ionic strength with an order of two in [H+]. The rate of reduction also increased with increase in percentage of acetic acid. The plot of log k vs. 1/D gave a straight line with a positive slope indicating that one of the reactants is a positive ion i.e. Ni(OH)2+. Taft equation was applicable with a reaction constant of ρ* = -1.41 and correlation coefficient r = 0.91. A mechanism involving an outer-sphere electron transfer from alcohol to Ni(III) ion was proposed. The mechanism of oxidation of both primary and secondary alcohols can be depicted as shown in scheme 16.

3.3.2.2. One-Electron Oxidation of Aliphatic Secondary Alcohols by Ni(III) Ion in Aqueous Sulfuric Acid-Acetic Acid Medium: a Kinetic and Mechanistic Study [55]

Oxidation of a series of aliphatic secondary alcohols MeCHROH (R = Me, Et, Pr, Bu, hexyl) by Ni(III) in aqueous acetic acid and H2SO4 medium at constant acidity and ionic strength exhibited overall second-order kinetics. The rates increased with increasing [H+] at constant ionic strength. The Taft equation gave ρ* = -1.00 at 303 K. A mechanism involving an outer sphere electron transfer from alcohol to Ni(III) ion was proposed. The mechanism of oxidation of both primary and secondary alcohols can be depicted as shown in scheme 17.

3.3.2.3. A Synchronous C-C and O-H Bond Cleavage of Some 1,2-Diols by Ni(III) Ion in Sulfuric Acid Medium: a Kinetic, Entropic and Mechanistic Approach [56]

The kinetics of oxidation of HOCHRCHR1OH (R = R1 = H, Me; R = H, R1 = Me, Et) by Ni(III) ion was examined in aqueous H2SO4. The rates were affected by acidity and dielectric constant but not by added SO42- ions. The reaction was overall second order, first order in each reaction component, and proceeded via a synchronous C-C and O-H bond cleavage, which was confirmed by product analysis, kinetic parameters and large negative entropies of activation. Ni(OH)2+ was shown to be the reactive species from the application of Amis equation to the rate data at different solvent compositions. From the product analysis, stoichiometry, presence of free radicals and foregoing kinetic results the mechanism was given as shown in scheme 18 taking ethylene glycol as a typical example:

Further, one can also propose an alternate mechanism for the oxidation of glycols by Ni(III) as given shown scheme 19:

Both the mechanisms given above tend to give the same products irrespective of the mode of the redox process. The only difference in both the mechanisms was that one involves production of a proton and the other a carbocation i.e. CH3OH with the same free radical i.e. CH2OH and Ni(II) as the additional products in the rate determining step. Among the two mechanisms, I am of the opinion that the first mechanism appeared to be the most likely one based on the large negative entropies of activation ( - 115 to - 170 J. mol⁻¹ K⁻¹) observed which originate from the dehydration of proton produced in the rate determining step resulting in the freezing of solvent molecules, where as the positive ion produced as shown in the second mechanism was a carbocation which
could not be hydrated. Hence one should get more positive entropies of activation. Therefore the author certainly was of the view that the oxidation process of glycols by Ni(III) involved a synchronous cleavage of C-C and O-H bonds in the rate determining step.

3.3.3. Thallium(III) ion

Oxidation of acetophenone-semicarbazones, acetone—semicarbazones [57], the ruthenium(III) adducts of ethanamines [58], aliphatic alcohols [59] and mannitol [60] were studied with Tl(III) ion. All these reactions followed total second order kinetics. The pathway of semicarbazones took place via a dimer of the hydroxy semicarbazone that was first formed in the rate-determining step by a \( \alpha \)-hydrogen abstraction followed by reaction with water. There was no reaction of Tl(III) ion with ethanamines, alcohols and mannitol in the absence of Ru(III). Here Ru(III) was used as a catalyst.

3.3.3.1. Kinetics of Oxidation of Semicarbazones by Thallium(III) [57]

Oxidation of semicarbazones of acetone, methyl ethyl ketone, acetophenone and 4'-methylacetophenone by Tl (III) in aqueous acetic acid containing \( H_2SO_4 \) and \( Cl^- \) was examined. The reaction followed a total second order. The rate of oxidation decreased with increasing \( [H^+] \) and with increasing \( [Cl^-] \). These two observations could be interpreted with the following (eqns. 31 and 32) two equilibriums assuming \( Tl(OAc)_3^- \) as the reactive species.

\[
Tl(OAc)_3^- + H^+ \rightleftharpoons Tl(OAc)_2^{2+} + HOAc \tag{31}
\]

\[
Tl(OAc)_2^{2+} + Cl^- \rightleftharpoons Tl(OAc)_2Cl \tag{32}
\]

The rate increased with decreasing percentage of acetic acid. A plot \( \log k \) vs.1/D was linear with a positive slope indicating that one of the redox partners was a positive ion i.e. \( Tl(OAc)_2^{2+} \). The product was observed to be the corresponding carbonyl compound. Induced polymerization was not observed when acryl amide was added, suggesting that no free radicals were involved in the reaction. The mechanism suggested could be depicted as shown scheme 20:

![Scheme 20. Reaction of semicarbozones with Tl(III)](image)

3.3.3.2. Kinetics of Oxidation of Ethanolamines by Thallium(III): Ruthenium(III) Catalysis [58]

Kinetics of oxidation of monoethanolamine, diethanola-
mine and triethanolamine by Tl(III) with Ru(III) as catalyst has been studied in acetic acid-water mixture. The reaction was first order in \([Tl(III)]\) and fractional order in \([substrate]\) and \([Ru(III)]\). The reaction ceased even in lower concentrations of \( H_2SO_4 \) and NaCl. The rate decreased with increase in percentage of acetic acid. This revealed that the reactive species of thallium in aqueous acetic acid was supposed to be \( Tl(OAc)_3^- \). A mechanism involving formation of an adduct between the catalyst and the substrate in a fast step which reacts later with the oxidant in a slow step to give products has been proposed. Thermodynamic and activation parameters for formation of the adduct and for its reaction with Tl(III) have been determined respectively. The negative \( \Delta S \) values may be attributed to the more rigid nature of the adduct formed, which also accounted for the decrease in number of molecules on going from the reactants to the adduct formation stage. The negative \( \Delta S^* \) values were in agreement with formation of an \( H^+ \) in the rate determining step, which could be solvated hence freezing the movement of free molecules. The non-involvement of the free radical intermediates was confirmed by negative polymerization test. The products of the reaction were found to be HCHO and NH₃.

3.3.3.3. Kinetics of Oxidation of Aliphatic Alcohols by Thallium(III): Ruthenium(III) Catalysis [59]

Kinetics of oxidation of aliphatic alcohols by Tl(III) with Ru(III) as catalyst were studied in acetic acid. The reaction was first order in \([Tl(III)]\) and fractional order in \([substrate]\) and \([Ru(III)]\). The reaction ceased even in lower concentrations of \( H_2SO_4 \) and NaCl. The rate decreased with increasing percentage of acetic acid, indicating \( [Tl(OAc)_3^-] \) was the reactive species. The noninvolvement of free radical intermediates was confirmed by a negative polymerization test. Mechanisms involving adduct formation between the catalyst and substrate in a fast step and the oxidation of this adduct by the oxidant in the rate determining step to form the products had been proposed. The product of the reaction was the corresponding carbonyl compound.

3.3.3.4. Kinetics and Mechanism of Oxidation of Mannitol by Thallium(III): Ruthenium(III) Catalysis [60]

The rates of oxidation of D-mannitol by Tl(III) with Ru(III) as catalyst had been measured in acetic acid medium. The reaction was first order in \([Tl(III)]\), fractional order each in \([substrate]\) and \([Ru(III)]\). The reaction ceased even in lower concentrations of \( H_2SO_4 \) and NaCl. The rate decreased with increase in percentage of acetic acid. This revealed that the reactive species of thallium in aqueous acetic acid was supposed to be \( Tl(OAc)_3^- \). The non-involvement of the free radical intermediates was confirmed by negative polymerization test. Mechanisms involving adduct formation between the catalyst and substrate in a fast step and the oxidation of this adduct by the oxidant in the rate determining step to form the products had been proposed. The product of the reaction was D-mannose.
The rate law (eqn. 33) and the mechanism (scheme 21) which could explain all the studies of above three redox processes are given below:

The rate law:

\[
rate = \frac{kk'[\text{III}][\text{substrate}][\text{Ru(III)}]}{1 + K[\text{substrate}] + K[\text{Ru(III)}]}
\]

(33)

And the mechanism is shown in Scheme 21:

\[
\begin{align*}
\text{Ru(III)} + \text{HO-CH}_2\text{R} & \rightleftharpoons (\text{adduct})^+ + \text{Ru(III)} \\
(\text{adduct})^+ + \text{TI(III)} & \rightarrow \text{Ru(III)} \\
(\text{adduct})^+ & \rightarrow \text{H-O-C-H-R} \\
\text{H-O-C-H-R} & \rightarrow \text{CHR} + \text{H}^+
\end{align*}
\]

\[k_{\text{cat}} = \frac{k_{\text{cat}}[\text{III}][\text{substrate}][\text{Ru(III)}]}{1 + K[\text{substrate}] + K[\text{Ru(III)}]}
\]

Scheme 21

3.4. Inner-Sphere Electron Transfer Reaction between an Inorganic Oxidant and Organic Substrates

3.4.1. Cerium(IV) ion: Silver(I) Catalyzed and Un-catalyzed Oxidation of Dimethyl Sulfoxide by Ceric Nitrate in Nitric Acid Medium [61]

The un-catalyzed oxidation of DMSO by Ce(IV) involves complex formation in HNO\textsubscript{3} medium which yields dimethyl sulfone as the final product. The evidence for the complex formation between Cr(IV) and DMSO was obtained from kinetic observations and the absorption spectra of both Ce(IV) and Ce(IV)-DMSO complex which showed a shift in \(\lambda_{\text{max}}\) from 340 nm to 360 nm. The orders in [Ce(IV)] and [DMSO] were one and fractional respectively. The intra molecular electron transfer in the complex resulted in the liberation of DMSO\textsuperscript{2+} radicals which in subsequent steps convert in to sulfone. The formation of free radicals was confirmed by acrylonitrile polymerization. With Ag(I) as catalyst, an Ag(I)-DMSO adduct was proposed. This adduct reacts with Ce(IV) bimolecularly to give a Ag(II)-DMSO adduct which yields the products. The evidence for the formation of Ag(II) as the intermediate was obtained from kinetic features and spectral studies. When bipyridyl added to the reaction mixture a brown colored complex of Ag(II) with a characteristic maximum at 454 nm was obtained supporting the formation of Ag(II)-DMSO adduct as the intermediate in the catalyzed process. The kinetic evidence for the formation of intermediate complexes both in the catalyzed and un-catalyzed processes was the linear double reciprocal plot obtained when \(1/k_{\text{obs}}\) was plotted against \(1/\text{substrate}\) or \(1/[\text{catalyst}]\). The rates exhibited fractional order in [Ag(I)] as well as [DMSO] and first order in [Ce(IV)]. The other kinetic features observed in the catalyzed process were same as that observed in the absence of Ag(I). From the effect of [H\textsuperscript{+}] on rates it was concluded that the neutral species of cerium i.e. Ce(NO\textsubscript{3})\textsubscript{4} was the reactive one in the catalyzed and un-catalyzed reactions. In the catalyzed redox process, there could be a possibility of competition between Ag(I) and Ce(IV) for DMSO to give Ag(I)-DMSO adduct or Ce(IV)-DMSO complex. Analysis of the kinetic data from double reciprocal plots the formation constants \(K\) (in the absence of catalyst) and \(K_{\text{cat}}\) (in the presence of catalyst) were found to be 9.0 mol\textsuperscript{-1} and 30.3 mol\textsuperscript{-1}. The value of 9.0 mol\textsuperscript{-1} for \(K\) (the formation constant of Ce(IV)-DMSO complex) in the absence of catalyst is far less than the value of 30.3 mol\textsuperscript{-1} for \(K_{\text{cat}}\) (the formation constant of Ag(I)-DMSO adduct) in the presence of catalyst. This clearly indicated that the Ag(I)-DMSO adduct was exclusively formed in the presence of Ag(I). Based on these results the mechanisms suggested were as follows (Schemes 22 and 23):

Uncatalyzed:

\[
\begin{align*}
\text{DMSO + Ce(IV)} & \rightarrow \text{complex} \\
\text{complex} & \rightarrow \text{DMSO}^* + \text{Ce(III)} \\
\text{DMSO}^* + \text{Ce(IV)} & \rightarrow \text{DMSO}_2 + \text{Ce(III)} + 2\text{H}^+
\end{align*}
\]

Scheme 22. Reaction of DMSO with Ce(IV)

The rate law from this mechanism came out to be (eqn. 34):

\[
rate = \frac{kK[\text{IV}][\text{DMSO}]}{1 + K[\text{DMSO}]} \tag{34}
\]

Catalyzed:

\[
\begin{align*}
\text{DMSO + Ag(I)} & \rightarrow \text{complex} \\
(\text{adduct})^+ + \text{Ce(IV)} & \rightarrow \text{Ru(III)} \\
(\text{adduct})^+ & \rightarrow \text{H-O-C-H-R} \\
\text{H-O-C-H-R} & \rightarrow \text{CHR} + \text{H}^+
\end{align*}
\]

\[k_{\text{cat}} = \frac{k_{\text{cat}}[\text{IV}][\text{DMSO}]}{1 + K[\text{DMSO}]} \tag{35}
\]

The activation energy determined for catalyzed process was 83.1 kJ mol\textsuperscript{-1} smaller than the one for un-catalyzed process i.e. 89.2 kJ mol\textsuperscript{-1} indicating a moderate catalysis by Ag(I) in the oxidation of DMSO by Ce(IV).

3.4.2. Nickel(III) ion

Oxidation of benzaldehydes [62], \(\alpha\)-hydroxy acids [63] and ethanolamines [64] has been studied. In all the three systems studied, the rate law was found to be as shown in eqn. 36

\[
rate = \frac{kk'[\text{III}][\text{substrate}]}{1 + K[\text{substrate}]} \tag{36}
\]

where \(K\) is the formation constant of the complex between the substrate and Ni(III) and \(k\) is the first order rate constant for the decomposition of the complex to yield the products. Benzoic acids from benzaldehydes and the corresponding carbonyl compounds from the other two systems were the products. Applicability of the Hammett equation in benzaldehyde oxidation and discussions on the contribution of activation parameters in the oxidation of ethanolamines were some of the salient features enumerated.
3.4.2.1. Kinetics and Mechanism of Oxidation of Benzaldehydes by Nickel(III) Ion in Acetic Acid-Water Mixtures [62]

The rates of oxidation of a series of benzaldehydes (BA) by Ni(III) ion were measured in acetic acid-water mixtures at constant acidity [H$_2$SO$_4$] and ionic strength. The reaction was fractional order in [BA] and first order in [Ni(III)]. The reaction was acid catalyzed, with an order of two in [H$^+$], and was unaffected by added Na$_2$SO$_4$. Also the rates were sensitive to solvent polarity. Hammett’s equation was applicable with a reaction constant $\rho$ = 1.92 and correlation coefficient $r$ = 0.978. Formation of an intermediate in a fast step and subsequent decomposition of this in a slow step by a homolytic cleavage of aldehydic C-H bond was proposed as a probable mechanism. The presence of free radical intermediates was confirmed by acryl amide polymerization test. From the iso-kinetic plot the reaction series was considered to be entropy controlled one. The product was identified as benzoic acid from its melting point. The mechanism was as given in scheme 24.

\[
\begin{align*}
C_6H_5CHO + Ni(OH)^{2+} & \xrightleftharpoons{K^{(fast)}} \xrightarrow{k \ (slow)} C_6H_5C=O + Ni^{(II)} + H_2O \\
C_6H_5C=O + Ni(OH)^{2+} & \rightarrow C_6H_5COOH + Ni^{(II)}
\end{align*}
\]

Scheme 24. Reaction between benzaldehydes and Ni(III)

3.4.2.2. Oxidation of Lactic Acid (LA) and Mandelic Acid (MA) by Nickel(III) Ion in Sulfuric Acid Medium Via Addition/Elimination [63]

The oxidation of lactic acid and mandelic acid by Ni(III) ion was studied in aqueous acid medium. A rapid reversible formation of a complex between the oxidant and reductant, which later decomposes in a slow step by an inner-sphere electron transfer process, was observed. Acetaldehyde, benzaldehyde and CO$_2$ were detected as the products. The oxidation path proceeds via a C-C bond cleavage. An increase in the acidity and a decrease in the solvent polarity increased the rates. The formation of free radicals was identified by a positive acryl amide polymerization test. The formation constants and unimolecular decomposition rate constants for the complexes were separated. From these kinetic observations the mechanism suggested was as given in scheme 25 taking mandelic acid as a typical example:

\[
C_6H_5C(OH)CO_2H + Ni^{(III)} \xrightarrow{K \ (adduct)} C_6H_5C=O + Ni(OH)^{2+} + H^{+}
\]

Scheme 25. Reaction between mandelic acid and Ni(III)

The rate law came out to be as shown in eqn. 37:

\[
rate = \frac{kk[Ni(III)][MA]}{1+K[MA]} \tag{37}
\]

The more negative $\Delta S^*$ values (-35 to -56 J. (K mol.$^{-1}$)$^{-1}$) for the first step were an indication of formation of a rigid complex. Again the more negative $\Delta S^*$ values (-30 to -40 J. (mol. K)$^{-1}$)$^{-1}$ was in support of the production of H$^+$ in the rate determining step.

3.4.2.3. Reactivity of Ethanolamines with Nickel(III) in Aqueous Solution: Control by Enthalpy and Entropy of Activation [64].

Ethanolamine, diethanolamine, and triethanolamine react with Ni(III) in aqueous solution via an inner-sphere electron transfer mechanism. The orders in [substrate] and [oxidant] were fractional and one, respectively, suggesting the formation of a complex ($K$-path) between the substrate and the oxidant, which decomposes ($k$-path) in a rate-determining step. Thermodynamic and activation parameters of $K$ and $k$ paths were determined and discussed. Solvent immobilization due to generation of H$^+$ in the rate-determining step leads to the negative activation entropies. The rates increased with increasing [H$^+$]. The mechanism suggested was as shown in scheme 26:

\[
\begin{align*}
C_2H_5(OH)CH_2NH_3^+ + Ni(OH)^{2+} & \rightleftharpoons Ni(H_2NCCH_2OH)^{3+} + H_2O \\
complex & \rightarrow CH_2O + H^+ + *CH_2NH_2 + Ni^{(II)} \\
*CH_2NH_2 + Ni^{(III)} & \rightarrow CH_2O + NH_3^+ + Ni^{(II)}
\end{align*}
\]

Scheme 26. Reaction between ethanalamine and Ni(III)

An alternate mechanism could also be suggested which involves the formation of a carboxylation as shown in the case of Ni(III) and 1,2-diols [15]. But the mechanism suggested above was the most likely one based on the similar grounds offered for Ni(III) and 1,2-diols reaction. The rate law also came out to be similar to that observed for Ni(III) and 1,2-diols reaction. The $p^*$ values for the formation of the complex (on $K$: -0.92) and for the decomposition of the complex (on $k$: -1.7) showed that the decomposition step was more sensitive to effect of substituents. From the application of iso-kinetic relationship both on the formation and decomposition of the complex, it was observed that the formation step was supposed to be enthalpy controlled and the decomposition step to be entropy controlled. The product formaldehyde was identified by chromatropic acid test and the ammonium ion by Nessler’s reagent test.

3.5. Outer-sphere Electron Transfer Reaction between Organic Oxidants and Inorganic Substrates: Oxidants are the Nitrobenzenes

One electron reduction of nitrobenzenes by carbon dioxide radical anion (CO$_2^-$) and hydrogen atom (H$^+$) [65]: During the course of characterization of nitrobenzene radical anions (RA$^-$), nitrobenzenes were subject to the reaction of CO$_2^{**}$. The yields of RA$^-$ were characterized based on the thiocyanate (G(OH) + G(H) = 6.6) and methyl chloride (G($e^-_{aq}$) =
3.0)) dosimetries. Reactions of aqueous carbon dioxide radical anion (CO$_2^-$) with p-substituted nitrobenzenes were studied by pulse radiolysis attached with time resolved uv absorption technique. CO$_2^-$ radicals were produced by the reaction of formate ion with primary radicals (OH) generated from water radiolysis in N$_2$O saturated aqueous solutions. At pH 7 these radicals were quantitatively oxidized by nitrobenzenes to yield carbon dioxide and nitrobenzene radical anion (RA$^-$) with second order rate constants (~ 2 X 10$^8$ M$^{-1}$ s$^{-1}$) close to little lower than the diffusion controlled limit (5 X 10$^9$ M$^{-1}$ s$^{-1}$) for almost all the twelve nitrobenzenes studied except the one with an electron donating substituent, the 4-methoxy nitrobenzene which reacted with a rate constant of 1.2 X10$^8$ M$^{-1}$ s$^{-1}$. The yield of RA$^-$ was found to be quantitative when studied spectroscopically compared to that obtained due to direct reaction of $e_{aq}$ with nitrobenzene. And similarly H$^+$ (produced from radiolysis of water) atoms were oxidized to hydrogen ions whose yields were found to be quantitative when compared to the CH$_3$Cl dosimetry. The following is the reaction scheme 27.

![Reaction scheme 27](image)

Scheme 27. Reaction of carbon dioxide radical/hydrogen radical with nitrobenzene

3.6. Do Nitrobenzenes with Electron-Donating Substituents React with CO$_2^-$ via Inner-Sphere Electron Transfer Reaction?

One electron reduction of nitrobenzenes attached with electron-donating substituents by carbon dioxide radical anion (CO$_2^-$) [65]: The investigations focused on this heading were not studied in detail in this direction in 1984 when this article was published in JACS [65] except for the determination of the optical properties like the extinction coefficients of nitrobenzene radical anions. While writing this review a gap was figured out for a reaction that involves an electron-donating substituent, the 4-methoxy nitrobenzene which reacted with a rate constant of 1.2 X10$^8$ M$^{-1}$ s$^{-1}$. The yield of RA$^-$ was found to be quantitative when studied spectroscopically compared to that obtained due to direct reaction of $e_{aq}$ with nitrobenzene. And similarly H$^+$ (produced from radiolysis of water) atoms were oxidized to hydrogen ions whose yields were found to be quantitative when compared to the CH$_3$Cl dosimetry. The following is the reaction scheme 27.

![Reaction scheme 27](image)

Scheme 27. Reaction of carbon dioxide radical/hydrogen radical with nitrobenzene

3.7. Inner-Sphere Electron Transfer Reaction between Organic Oxidants and Organic Substrates: Oxidants are the Nitrobenzenes.

3.7.1. One-Electron Reduction of Nitrobenzenes by α-Hydroxymethyl Radical via Addition/Elimination.

An Example of an Organic Inner-Sphere Electron-Transfer Reaction [65]

Reactions of aqueous α-hydroxymethyl radicals $^*$CH$_2$OH with p-substituted nitrobenzenes were investigated by using product analysis, in-situ-radiolysis-ESR spectroscopy, pulse radiolysis methods and also time resolved optical and conductance detection techniques. $^*$CH$_2$OH was produced by H abstraction from methanol by the OH (and H) radicals generated by irradiation of N$_2$O-saturated aqueous solutions containing 0.1-2.0 M methanol. On addition of 0.1 to 1.0 mM para-substituted nitrobenzenes to the solutions, at pH 3-6 nitroxide type radicals were produced, as observed by in-situ-radiolysis-ESR experiments. These radicals were formed by addition of $^*$CH$_2$OH to the nitro group as shown in scheme 29. Nitrobenzene radical anions were not detected.

![Reaction scheme 29](image)

Scheme 29. Reaction between α-hydroxymethyl radical and nitrobenzene

Experiments were also carried out using optical detection. The time dependent transient spectra of nitroxide type radicals showed $\lambda_{max}$ at 280-310 nm. From the optical detection system the bimolecular rate constants were determined for the formation of these radicals. They range from 10$^7$ to 10$^9$ (M $^{-1}$ s$^{-1}$) depending on the R of the nitrobenzene in the para position. Further the non-formation of ions in this reaction was tested using time resolved conductance technique. 4-Nitrobenzenediazonium tetrafluoroborate (R = N$_2$O$_4$) was the only nitrobenzene to give ions from the reaction of $^*$CH$_2$OH. This reaction did not need any base to get the final
products. The quantity of formaldehyde formed per \( ^*\text{CH}_2\text{OH} \) was quantitative indicating that this particular nitrobenzene with strong electron withdrawing group (\( N_\text{2}^* \); Hammett’s \( \sigma \) value is 1.93 the known substituent with highest \( \sigma \) value) at para position could oxidize \( ^*\text{CH}_2\text{OH} \) by addition/elimination to give the final products stochiometrically.

At \( \text{pH} > 7 \) the \( \alpha \)-hydroxymethyl radicals were oxidized quantitatively to yield formaldehyde and \( \text{H}^+ \), and the nitrobenzenes are reduced to the radical anions. The mechanism of this redox reaction depends strongly on the substituents on both the reactants. In the case of \( ^*\text{CH}_2\text{OH} \) radical, the reaction proceeds via addition to the nitro group to produce an alkoxynitroxyl radical; this undergoes only an \( \text{OH}^- \) catalyzed heterolysis to give \( \text{HCHO} \) and the nitrobenzene radical anion with any nitrobenzene as shown in scheme 30:

\[
\begin{align*}
\text{Nitroxide type radical} & \xrightarrow{+ \text{OH}^-} \text{HCHO} + \text{nitrobenzene radical anion} \\
\text{Scheme 30. Reaction between } \alpha \text{-hydroxymethyl radical and nitrobenzene in presence of } \text{OH}^-
\end{align*}
\]

The nitrobenzene radical anions were also characterized by time dependent optical absorption spectra which showed strong absorptions at 300-360 nm and 420-560 nm depending on the nitrobenzene. Between \( \text{pH} \text{ 9-10} \) the rate constant for the formation of the radical anion was dependent of [nitrobenzene] and the second order rate constants were similar to those measured at \( \text{pH} \leq 6 \) for the formation of the nitroxide radicals indicating that the addition step was rate determining. From this observation and from the rate constants for the formation of un-ionized nitro radical the rate constant for uni-molecular heterolysis of ionized nitroxide radical was calculated to be \( \geq 5 \times 10^5 \text{ s}^{-1} \).

At \( \text{pH} \geq 10 \) the rate constants for the formation of radical anion increased with increase in \( \text{pH} \) up to 11.5 indicating that this was due to the direct electron transfer reaction of ionized hydroxymethyl radical (\( ^*\text{CH}_2\text{O}^- \) (\( pK_a = 10.7 \)) which was a stronger electron donor than the un-ionized hydroxymethyl radical (\( ^*\text{CH}_2\text{OH} \)) with nitrobenzene (eqn. 38):

\[
\text{RC}_\alpha\text{H}_2\text{NO}_2 + ^*\text{CH}_2\text{O}^- \rightarrow \text{RC}_\alpha\text{H}_2\text{NO}_2^- + \text{CH}_2\text{O} \tag{38}
\]

The rate constants for this electron transfer reaction were close to diffusion controlled limit (\( k = 3 \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \)) and were essentially independent of the nature of the substituent in nitrobenzene ring.

### 3.7.2. Reaction of 6-yl Radicals of Uracil, Thymine, and Cytosine and Their Nucleosides and Nucleotides with Nitrobenzenes via Addition to Give Nitroxide Radicals. Hydroxide ion-catalyzed Nitroxide Heterolysis [66]

The 6-yl radicals produced by addition of \( \text{OH} \) to C(5) of the C(5)/C(6) double bond of naturally occurring pyrimidine bases, nucleosides, and nucleotides, or those formed by H-abstraction from C(6) of 5,6-dihydropyrimidines, react with para-substituted nitrobenzenes by addition to yield nitroxide-type radicals which were characterized by ESR and optical detection techniques. In basic solution the nitroxide radicals deprotonate: with the radicals derived from the free bases deprotonation occurs at N(1), with those from the nucleosides and nucleotides of uracil deprotonation at N(3) is observed. The ionized 6-yl radicals react with the nitrobenzenes also by addition with rate constants considerably higher than those for the case of the neutral radicals. The neutral nitroxide radicals react with \( \text{OH}^- \) to give the nitroxide radical anion which was able to undergo a unimolecular heterolysis reaction and results in the formation of the nitrobenzene radical anion. In the case of the nitroxide radicals having N(1)-H (the radicals derived from the free bases), the \( \text{OH}^- \) catalysis of nitroxide heterolysis proceeds via deprotonation at N(1) and the unimolecular rates are comparatively high. On substitution of N(1) by a Me or (deoxy)ribosyl(phosphate) group the site of \( \text{OH}^- \) attack is changed to N(3)-H (with the uracils) or to N(4)-H (with the cytosines). With radicals from N(1)-alkylated pyrimidines the rates of heterolysis of the nitrobenzene adducts are considerably lower than if N(1) carries a proton. The addition/\( \text{OH}^- \) catalyzed elimination sequence of interaction of pyrimidin-6-yl radicals and nitrobenzenes results in the ultimate transfer of an electron from the pyrimidine radical to the nitrobenzene and was therefore an example of a one-electron redox reaction. The course of events was shown in scheme 31 where 4-NAP indicates 4-nitroacetophenone.

\[
\begin{align*}
\text{Scheme 31. Reaction events of 4-NAP and pyrimidin-6-yl radical}
\end{align*}
\]

\[
\begin{align*}
\text{Scheme 32. Reaction events of } \alpha \text{-hydroxy ethyl radical with nitrobenzene}
\end{align*}
\]
3.7.3. One-Electron Reduction of Nitrobenzenes by α-Hydroxy Ethyl Radical via Electron Transfer and Addition/Elimination: Effect of Substituents on Rates and Activation Parameters for Unimolecular Heterolysis of Nitroxyl-Type Tetrahedral Intermediates [65]

In the reaction of HOCH2CH3 with nitrobenzenes both addition and electron transfer took place, the fraction of electron transfer increasing with the electron-withdrawing power of the substituent in the nitrobenzene. Occurrence of both these reaction paths was successfully studied by time dependent absorption spectroscopy. The transient spectra clearly showed that the adducts absorbed at 280-300 nm and the radical anions at 330-350 nm. The nitroxyl-type adducts underwent a spontaneous uni-molecular heterolysis \((k_s)\) to give CH3CHO, \(H^+\) and nitrobenzene radical anion. The rate constants \(k_s\) (from \(<10^2\) to \(5 \times 10^3 \text{ s}^{-1}\)) for this heterolysis increased with the electron-withdrawing strength of the substituent if it was on the benzene, and decrease if the substituent was on the CH3 carbon of the nitroxyl. The heterolysis was characterized by low (5-10 kcal/mol) activation enthalpies and strongly negative (-5 to -25 e.u.) activation entropies, which originated from hydration of a proton in the transition state. From the effect on the activation parameters exerted by substituents on the electron-acceptor and -donor parts of the nitroxy radical, the heterolysis proceeded via a push-pull mechanism, and was entropy controlled. Both Hammett and Taft equations obeyed for heterolysis reaction and yielded \(\rho (=1.4)\) when substituents were changed in nitrobenzene ring and \(\rho^* (= -1.15)\) when substituents were changed in the alcohol moiety. From the isokinetic plot (\(\beta = 255 \text{ K}\)), these reactions were supposed to be entropy controlled one. The reaction is shown in scheme 32.

The formation of acetaldehyde was quantitative from GC analysis. In the ESR spectra the formation of nitroxy type radicals could not be detected as in the case of \(^{1} \text{H} \text{CH}_2 \text{OH}\). However by the use of time dependent optical absorption and conductivity detection techniques both the nitroxy type radicals and radical anions were detected. With increase in pH the \(k_s\) was found to increase according to the equation \(k_{\text{hbd}} = k_s + k_c [\text{OH}^-]\). At pH > 10.5 the rates of formation radical anions became independent of pH but dependent on [nitrobenzene] and the simultaneous biphasic formation of nitroxy type radicals and radical anions became monophasic with very high rates, which clearly showed that the ionized hydroxy ethyl radical i.e. \(\text{CH}_2\text{CHO}^+ (pK = 11.3)\) reacted by direct electron transfer.

The reaction path of \(k_s\) took place via a synchronous O-H and C-O bond breaking and via a solvent assisted deprotonation as shown in scheme 33:

This was supported by studying the kinetic isotope effect on \(k_s\). The \(k^{\text{H}} / k^{\text{D}} = 2.2\) at 20°C using 4-nitrobenzonitrile with \(\text{CH}_2\text{CHOH}\) clearly showed that the above observation showing simultaneous breaking of N-O and C-O bonds in the transition state.

3.7.4. One-Electron Reduction of Nitrobenzenes by \(^{1}\text{OH}\) and \(H^+\) Radical Adducts of 6-Methyluracil and 6-Methylisocytosine via Electron Transfer and Addition/Elimination: Effect of Substituents on Rates and Activation Parameters for Unimolecular Heterolysis of Nitroxyl-Type Tetrahedral Intermediates [67]

The radicals formed by OH radical addition to C-5 of 6-methyluracil or of 6-methylisocytosine, i.e., 5-hydroxy-5,6-dihydro-6-methyluracil-6-yl or 5-hydroxy-5,6-dihydro-6-methylisocytosin-6-yl, or that produced by H addition to C-5 of 6-methyluracil (or by H abstraction from C-6 of 5,6-dihydro-6-methyluracil), i.e., 5,6 - dihydro-6 - methyluracil -6-yl, react in aqueous solution with para-substituted nitrobenzenes to give both nitrobenzene radical anions and nitroxy-type radicals with rate constants that vary from \(\approx 7 \times 10^7\) to \((2-5) \times 10^9 \text{ M}^{-1} \text{s}^{-1}\), depending on the pyrimidine radical and on the nitrobenzene. The nitroxy radicals undergo a spontaneous unimolecular heterolysis \((k_s)\) to yield (additional) nitrobenzene radical anion and oxidized pyrimidine with rate constants of \(10^3\) to \(5 \times 10^5 \text{ s}^{-1}\), depending on the structure of the pyrimidine and of the nitrobenzene. This reaction is characterized by activation energies of 30-40 kJ mol\(^{-1}\) and by activation entropies of -7 to -89 J mol\(^{-1}\) K\(^{-1}\) (entropy control). The addition/elimination sequence constitutes a case of inner-sphere electron transfer reaction. The rate constants for the heterolysis reaction are a measure of the reducing power of 5,6-dihydro-6-methylpyrimidin-6-yl radicals. On this basis, the cytotoxic radicals are better reductants than the corresponding uracil radicals, and the radicals derived by H atom addition to pyrimidines are stronger reductants than those formed by OH radical addition. The experimental observations for studying these reactions were essentially similar to that studied with hydroxyl ethyl radicals and nitrobenzenes (Scheme 34).

3.8. Outer-Sphere Electron Transfer Reaction between Organic Oxidants and Organic Substrate: Oxidants are the Nitrobenzenes
In the oxidation of α-hydroxy isopropyl radical (CH₃)₂COH by nitrobenzenes [65], the nitroxyl radical of the type formed in the case of α-CH₂OH could not be detected and all nitrobenzenes oxidize (CH₃)₂COH by simple outer sphere electron transfer. The yield of acetone in this reaction was 86% and the remaining 14% was accounted for β-radicals, which has no reducing properties. Hence α-hydroxy isopropyl radicals were oxidized quantitatively. However a much weaker oxidant like 4-nitro-N,N-dimethyl aniline could not oxidize this radical by electron transfer, instead the course of the reaction was just similar to the events observed with that of α-CH₂OH. As the substituents in nitrobenzene become more and more electron withdrawing the reaction turned out to be addition/elimination and finally becomes pure electron transfer.

With R = H, both addition/elimination took place similar to that observed with CH₃C(OH)₂H. But if R = Me only addition and then OH⁻ catalyzed heterolysis took place and this situation was just similar to that observed with methanol radical using any nitrobenzene. Therefore 4-nitro-N,N-dimethyl aniline was the only nitrobenzene which behaved like much weaker oxidant to oxidize (CH₃)₂COH (Scheme 35).

In the oxidation of α-hydroxy isopropyl radical (CH₃)₂COH by nitrobenzenes [65], the nitroxyl radical of the type formed in the case of α-CH₂OH could not be detected and all nitrobenzenes oxidize (CH₃)₂COH by simple outer sphere electron transfer. The yield of acetone in this reaction was 86% and the remaining 14% was accounted for β-radicals, which has no reducing properties. Hence α-hydroxy isopropyl radicals were oxidized quantitatively. However a much weaker oxidant like 4-nitro-N,N-dimethyl aniline could not oxidize this radical by electron transfer, instead the course of the reaction was just similar to the events observed with that of α-CH₂OH. As the substituents in nitrobenzene become more and more electron withdrawing the reaction turned out to be addition/elimination and finally becomes pure electron transfer.

With R = H, both addition/elimination took place similar to that observed with CH₃C(OH)₂H. But if R = Me only addition and then OH⁻ catalyzed heterolysis took place and this situation was just similar to that observed with methanol radical using any nitrobenzene. Therefore 4-nitro-N,N-dimethyl aniline was the only nitrobenzene which behaved like much weaker oxidant to oxidize (CH₃)₂COH (Scheme 35).

3.9. Oxidations by N-Bromoacetamide (NBA)

3.9.1. Kinetics and mechanism of oxidation of dimethyl sulfoxide by N-bromoacetamide (NBA): Outer-sphere oxygen atom transfer reaction [69]

The title oxidation was carried out at constant [H⁺] and ionic strength. The order in [NBA] was unity. Pseudo first-order rate constants were calculated for various [DMSO]. The order in [DMSO] was found to be zero. An analysis of the rate dependence on [H⁺] at constant ionic strength revealed that reactive species was found to be H₂OBr⁺, formed in the rate-determining step by addition of H₂O⁺ to HOBr. The rates were negligibly affected by added salts and dependent on solvent polarity. The sole product of the reaction was identified as dimethyl sulfone. The activation entropy for this reaction was compared with that of a NBA-alcohol reaction structurally related to DMSO. In the oxidation of iso-propanol (structurally related to DMSO) by NBA the activation entropy was found to be - 98.0 J (K mol⁻¹). This was explained by the production of H⁺ in the rate determining step by an α-H abstraction from the alcohol and this could be solvated extensively resulting in the freezing of free solvent molecules. But in the present reaction the small controlled rate constants were not due to large activation enthalpies but to very negative activation entropies [-10 to -80 J (mol K)⁻¹]. In fact, the enthalpies (2 to 16 kJ mol⁻¹) were in most cases below those for diffusion in water. The results were explained in terms of formation of a charge-transfer complex between the reactants and an ionic type transition state that results from electron transfer between the reactants as shown in scheme 36.

Solvent immobilization caused to the ionic transition-state structures, leads to the negative activation entropies. The two different types of final products, i.e., nitroxyl and radical anion, are derived from the ion-pair-type transition state by collapse (giving addition/nitroxyl) or by separation by solvent (yielding electron-transfer/radical anion). In the hypothetical case as shown below, that the electron transfer was taking place followed by or concerted with proton transfer and (full) hydration of the proton, the electron would be trapped on the nitrobenzene for good, since hydration of the proton was irreversible due to enormous exothermicity of this reaction. In other words proton transfer would “freeze”, i.e. finalize electron transfer. (See scheme 37).

3.9.2. Oxidations by N-Bromoacetamide (NBA): Outer-sphere oxygen atom transfer reaction [69]

The title oxidation was carried out at constant [H⁺] and ionic strength. The order in [NBA] was unity. Pseudo first-order rate constants were calculated for various [DMSO]. The order in [DMSO] was found to be zero. An analysis of the rate dependence on [H⁺] at constant ionic strength revealed that the reactive species was found to be H₂OBr⁺, formed in the rate-determining step by addition of H₂O⁺ to HOBr. The rates were negligibly affected by added salts and dependent on solvent polarity. The sole product of the reaction was identified as dimethyl sulfone. The activation entropy for this reaction was compared with that of a NBA-alcohol reaction structurally related to DMSO. In the oxidation of iso-propanol (structurally related to DMSO) by NBA the activation entropy was found to be - 98.0 J (K mol⁻¹). This was explained by the production of H⁺ in the rate determining step by an α-H abstraction from the alcohol and this could be solvated extensively resulting in the freezing of free solvent molecules. But in the present reaction the small controlled rate constants were not due to large activation enthalpies but to very negative activation entropies [-10 to -80 J (mol K)⁻¹]. In fact, the enthalpies (2 to 16 kJ mol⁻¹) were in most cases below those for diffusion in water. The results were explained in terms of formation of a charge-transfer complex between the reactants and an ionic type transition state that results from electron transfer between the reactants as shown in scheme 36.

Solvent immobilization caused to the ionic transition-state structures, leads to the negative activation entropies. The two different types of final products, i.e., nitroxyl and radical anion, are derived from the ion-pair-type transition state by collapse (giving addition/nitroxyl) or by separation by solvent (yielding electron-transfer/radical anion). In the hypothetical case as shown below, that the electron transfer was taking place followed by or concerted with proton transfer and (full) hydration of the proton, the electron would be trapped on the nitrobenzene for good, since hydration of the proton was irreversible due to enormous exothermicity of this reaction. In other words proton transfer would “freeze”, i.e. finalize electron transfer. (See scheme 37).

3.9.1. Kinetics and mechanism of oxidation of dimethyl sulfoxide by N-bromoacetamide (NBA): Outer-sphere oxygen atom transfer reaction [69]

The title oxidation was carried out at constant [H⁺] and ionic strength. The order in [NBA] was unity. Pseudo first-order rate constants were calculated for various [DMSO]. The order in [DMSO] was found to be zero. An analysis of the rate dependence on [H⁺] at constant ionic strength revealed that the reactive species was found to be H₂OBr⁺, formed in the rate-determining step by addition of H₂O⁺ to HOBr. The rates were negligibly affected by added salts and dependent on solvent polarity. The sole product of the reaction was identified as dimethyl sulfone. The activation entropy for this reaction was compared with that of a NBA-alcohol reaction structurally related to DMSO. In the oxidation of iso-propanol (structurally related to DMSO) by NBA the activation entropy was found to be - 98.0 J (K mol⁻¹). This was explained by the production of H⁺ in the rate determining step by an α-H abstraction from the alcohol and this could be solvated extensively resulting in the freezing of free solvent molecules. But in the present reaction the small...
activation entropy of only -12 J (K mol)$^{-1}$ was an indication that there was no such proton production in the rate determining step instead a proton was consumed liberating a free water molecule which could in fact lead to a positive entropy of activation. Based on the above arguments the mechanism suggested was as shown in scheme 38:

$$\text{CH}_3\text{CONHBr} + \text{H}_2\text{O} \xrightleftharpoons[k]{K} \text{CH}_3\text{CONH}_2 + \text{HOBr}$$

$$\text{HOBr} + \text{H}_2\text{O}^+ \xrightarrow[k]{\text{slow}} \text{H}_2\text{OBr}^+ + \text{H}_2\text{O}$$

$$\text{H}_2\text{OBr}^+ + \text{DMSO} \xrightarrow{\text{slow}} \text{DMSO}_2 + \text{HBr} + \text{H}^+$$

Scheme 38. Reaction between DMSO and NBA

From the above mechanism the reaction followed the rate equation (eqn. 39):

$$\text{rate} = \frac{kk[NBA][\text{H}_2\text{O}^+]}{K + [\text{AA}]}$$

(39)

where AA stands for acetamide. From the plot of $1/$rate vs. [AA] the $k$ and $K$ were separated on which all the effects were performed.

3.9.2. Kinetics and Mechanism of Oxidation of Benzaldehydes by N-Bromoacetamide in Acetic Acid-Water Mixture [70]

The title oxidations in 40% acetic acid - water mixture were overall second-order, first-order in each [reactant]. The reactions were acid-catalyzed and the rate was not affected by the addition of $\text{Na}_2\text{SO}_4$. A plot of log $k$ vs. Brown's $\sigma^+$ gave $\rho^+ = -1.675$ $(r = 0.97)$ indicating an electron deficient center in transition state. However, p-nitro benzaldehyde deviated from the Hammett plot, its rate constant was found to be higher than expected. This might be due to the high degree of hydration of this compound compared to the other benzaldehydes. From the application of iso-kinetic relationship the reaction series was considered to be entropy controlled one. A probable mechanism involving formation of a cyclic intermediate in a fast step and subsequent decomposition in a slow step by a hydride transfer route was proposed as shown scheme 39:

$$\text{CH}_3\text{CONHBr} + \text{H}_2\text{O} \xrightarrow[K]{\text{fast}} \text{CH}_3\text{CONH}_2 + \text{HOBr}$$

$$\text{PhCH(OH)}_2 + \text{HOBr} \xrightarrow[k]{\text{slow}} \text{PhCOOH} + \text{H}_2\text{O} + \text{HBr}$$

Scheme 39. Reaction between benzaldehyde and NBA

3.9.3. Kinetics and Mechanism of Uncatalyzed and Ruthenium(III) Catalyzed Oxidation of Mandelic Acid by N-Bromoacetamide in Aqueous Sulfuric Acid Medium [71]

Rates of uncatalyzed and Ru(III)-catalyzed oxidation of mandelic acid (MA) by N-bromoacetamide (NBA) were measured in aqueous sulfuric acid. The overall order of the uncatalyzed reaction was one in each [reactant], i.e., NBA and MA. Under catalyzed conditions the order in [NBA] is unity, but that in [MA] and [Ru(III)] were fractional. The rates increase with increase in $[\text{H}^+]$ and decrease with increase in $[\text{acetamide}]$. The catalyzed redox process involves formation of an adduct between the catalyst and MA in a rapid reversible step, which later reacts with NBA bi-molecularly in a slow step to give rise to products. Benzaldehyde was identified as the product from the 2,4-DNP test. Activation energy for catalyzed process (64.3 kJ mol$^{-1}$) was less than the un-catalyzed reaction (77.3 kJ mol$^{-1}$) indicating a good catalytic effect. The value of formation constant $(K = 26.7$ lit. mol$^{-1}$) between Ru(III) and MA was much higher than the bimolecular rate constant $(1.67$ lit mol$^{-1}$ min$^{-1}$) in the un-catalyzed process, which indicated that Ru(III) interacts exclusively with MA in the formation of the adduct. This also got further support from the high negative entropy change (-59.0 J mol$^{-1}$ K$^{-1}$) for the formation of the adduct. From the above experimental facts the mechanisms for both the processes were as shown in scheme 40:

$$\text{MA} + \text{Ru(III)} \xrightarrow[K]{\text{slow}} \text{Ru}^{3+} + \text{H}_2\text{OBr}^+ + \text{Br}^- + \text{H}_2\text{O}$$

$$\text{Ru}^{3+} + \text{C}_6\text{H}_5\text{CHO} + \text{CO}_2 + \text{HBr}$$

Scheme 40. Ru(III) catalyzed reaction between mandelic acid and NBA

From the mechanism the rate law came out to be (eqn. 40):

$$\text{rate} = \frac{kk[NBA][\text{H}^+][\text{MA}]}{K + [\text{AA}]}$$

(40)

where AA stood for acetamide. The possibility of further reaction of benzaldehyde product with oxidant was ruled out since the reaction conditions were under pseudo-first order. Hence the concentration of benzaldehyde produced would be very small.

The mechanism under catalyzed conditions ran to be (Scheme 41):

$$\text{MA} + \text{Ru(III)} \xrightarrow[K]{\text{slow}} \text{(adduct)}^{3+} + \text{H}_2\text{OBr}^+ \xrightarrow[k]{\text{slow}} \text{(adduct)}^{5+} + \text{Br}^- + \text{H}_2\text{O}$$

$$\text{(adduct)}^{5+} \xrightarrow[\text{fast}]{\text{Ru}(\text{III})} \text{C}_6\text{H}_5\text{CHO} + \text{CO}_2 + \text{Ru(III)} + 2\text{H}^+$$

Scheme 41. Reaction of mandeliciacid with NBA in presence of Ru(III)
From the above mechanism the rate came out to be:
\[
rate = \frac{k_{cat} K[\text{H}_2\text{OBr}^-][\text{MA}][\text{Ru(III)}]}{1 + K'[\text{MA}] + K'[\text{Ru(III)}]}
\]
(41)

3.9.4. Kinetics and Mechanism of Oxidation of Aromatic Amines by N-Bromoacetamide [72]

Kinetics of oxidation of anilines by N-bromoacetamide in methanol-water (50:50 vol./vol.) mixture had been investigated in the presence of mercuric acetate and excess acetamide. The total order of the reaction was two. Electron-releasing substituents in the aromatic ring accelerate the reaction rates and electron-withdrawing substituents retarded them. The value of the reaction constant (\(\rho\)) obtained from the Hammett plot (correlation coefficient = 0.998) was -0.785. The product of the reaction was identified as the azobenzene. A mechanism shown in scheme 42 involving an electrophilic attack of the oxidant on the amino group had been proposed to explain the observed results.

\[
\text{CH}_3\text{CONHBr} + \text{H}_2\text{O} \xrightarrow{K} \text{CH}_3\text{CONH}_2 + \text{HOBr}
\]

\[
\text{HOBr} + C_6H_5NH_2 \xrightarrow{\text{slow}} C_6H_5NH^+ + H_2O + Br^- \\
C_6H_5NH^+ + C_6H_5NH_2 \xrightarrow{\text{fast}} C_6H_5\tilde{N}H_2 - NHC_6H_5 \\
C_6H_5\tilde{N}H_2 - NHC_6H_5 \xrightarrow{\text{fast}} C_6H_5NHNHC_6H_5 \\
C_6H_5NHNHC_6H_5 \xrightarrow{\text{fast}} C_6H_5N = NC_6H_5 + H_2O + HBr
\]

Scheme 42. Reaction of anilines with NBA

The very high negative \(\Delta S^\circ\) values (-74 to -234 J mol\(^{-1}\) K\(^{-1}\)) were explained due to formation of an ionic transition state (Scheme 43) with an extensive charge separation leading to the transfer of either two electrons or hydride ion in the rate determining step as shown below: the transition state was formed from two neutral molecules, viz. the substrate and HOBr.

\[
\text{CH}_3\text{CONH}_2 + \text{HOBr} \xrightarrow{\text{fast}} \text{HOBr} + \text{CH}_3\text{CONH}_2
\]

Scheme 43

The Brønsted plot was fairly linear with a \(\beta\) value of 0.186 indicating that there was extensive bond formation between aniline and HOBr in the transition state. The positive value of Brønsted coefficient and the negative value of Hammett’s reaction constant point out that the rate increased with the increase in basicity of the approaching nucleophile. From the iso-kinetic plot the reaction series was considered to be entropy controlled one.

3.9.5. Kinetics and Mechanism of Bromination of Phenols by N-Bromoacetamide: Effect of Substituents on Decomposition of Intermediate Adducts [73]

The title reactions were studied in 40% aqueous methanol. A mechanism involving an intermediate six-membered cyclic transition state between phenol and HOBr (produced in situ by hydrolysis of N-bromoacetamide) was postulated for the title reaction as shown in scheme 44:

\[
\text{CH}_3\text{CONHBr} + \text{H}_2\text{O} \xrightarrow{K} \text{CH}_3\text{CONH}_2 + \text{HOBr}
\]

\[
\text{HOBr} + \text{Ph} \xrightarrow{K} \text{PhBr}
\]

Scheme 44. Reaction of phenols with NBA

From the above mechanism the rate law (eqn. 42) came out to be:
\[
rate = \frac{kK_k[\text{NBA}][\text{phenol}]}{(K_k + [\text{AA}])[1 + K[\text{phenol}]]}
\]
(42)

The rate showed that the order in [NBA] was unity and that in [phenol] was fractional. The rate showed an inverse dependence on [AA]. The product of the reaction was identified as o-bromophenol. A Hammett plot for \(k\) gave \(\rho = -0.60\) (\(r = 0.986\)) taking \(\sigma\) values of \(m\)-substituents as para and of \(p\)-substituents as meta assuming ortho carbon as the reaction center. From the iso-kinetic plots the reaction series was assumed to be entropy controlled one.

3.9.6. Kinetics and Mechanism of Bromination of Anisoles by N-Bromoacetamide in 50% Aqueous Acetonitrile [74]

The rates of bromination of anisoles with HOBr produced in situ by the hydrolysis of N-bromoacetamide (NBA) have been measured in 50% aqueous acetonitrile in the presence of mercuric acetate and acetamide. The formations of an intermediate cyclic adduct between HOBr and anisole which decomposes in a slow step to yield the products has been proposed to explain the observed results. The thermodynamic parameters for the hydrolysis of NBA and adduct formation steps have been evaluated. The activation parameters for the first order decomposition of the adduct have also been calculated. The corresponding o-bromoanisoles have been identified as major components in the products. The kinetic trends, rate law and the mechanism were similar.
3.9.7. Kinetics and Mechanism of \textit{in situ} Bromohydration of Cinnamic Acids by N-Bromoacetamide [75]

Bromohydroxylation of cinnamic acids (CA) by N-bromoacetamide (NBA) was studied in aqueous methanol containing Hg(OAc)$_2$ and acetamide. The influence of substituents on the decomposition rate constant of the adduct was investigated. The corresponding bromohydrins were identified as the reaction products. The mechanism suggested was as given in scheme 45:

\[
\text{CH}_3\text{CONHBr} + \text{H}_2\text{O} \rightleftharpoons K \text{CH}_3\text{CONH}^+ \text{HOBr} \\
\text{Ph} \cdot \text{CH} = \text{CH} \cdot \text{COOH} + \text{HOBr} \rightleftharpoons K \text{Ph} \cdot \text{CH} = \text{CH} \cdot \text{COOH} \\
\text{O} \cdot \text{Br}^+ \\
\text{H} \text{adduct} \\
\text{Ph} \cdot \text{CH} = \text{CH} \cdot \text{COOH} \xrightarrow{k_{\text{slow}}} \text{Ph} \cdot \text{CH} = \text{CH} \cdot \text{COOH} \\
\text{O} \cdot \text{Br}^+ \\
\text{H} \text{adduct}
\]

\textbf{Scheme 45.} Reaction of cinnamic acids with NBA

From the above mechanism the rate law (eqn. 43) came out to be:

\[
\text{rate} = \frac{kKK_{[\text{NBA}][\text{CA}]}(K_{\text{h}} + [\text{AA}][1 + K[\text{CA}])}{(K_{\text{h}} + [\text{AA}])} \tag{43}
\]

The values of $K$ and $K_{\text{h}}$ did not show any systematic variation on changing the substituents in the benzene ring. But Hammett equation was applicable on $k$, the first order rate constant for the decomposition of the adduct with $\rho$ value of -1.36 indicating an electrophilic addition of halogen to olefinic double bond. Also the Hammett’s $\rho$ decreased with temperature indicating that the decomposition of the adduct was assumed to be an entropy controlled process and this was reflected from the iso-kinetic plot.

3.9.8. Solvent Effects on Kinetics of Oxidation of Dimethyl Sulfoxide by N-Bromoacetamide. An Example of Inner Sphere Oxygen Atom Transfer Reaction [76]

Rates of oxidation of DMSO by N-bromoacetamide (NBA) in neutral aqueous solution were measured in the presence of acetamide. The order in [DMSO] was one below 0.01 mol dm$^{-3}$, fractional at [DMSO] 0.01-0.5 mol dm$^{-3}$, and zero at [DMSO] > 0.5 mol dm$^{-3}$. Different rate laws were operative under these three conditions, though HOBr was the effective oxidizing species in all the cases. The influence of the variation of solvent composition on the reaction rate was studied by using MeOH-H$_2$O of various compositions. Thermodynamic parameters for NBA hydrolysis and adduct formation between HOBr and DMSO were evaluated. The activation parameters for the first-order decomposition of the adduct were also calculated for solvent compositions of 0-80% vol. aqueous MeOH. The free energies of activation increase with the MeOH content of the medium, whereas the enthalpies and entropies of activation show a more complex, but partially compensating behaviour. The mechanism suggested was given below in scheme 46 for the fractional order condition with DMSO:

\[
\text{CH}_3\text{CONHBr} + \text{H}_2\text{O} \rightleftharpoons K \text{CH}_3\text{CONH}^+ \text{HOBr} \\
\text{CH}_3\text{CONHBr} + \text{HOBr} \rightleftharpoons K \text{CH}_3\text{CONH}^+ \text{HOBr} \\
\text{CH}_3\text{CONHBr} + \text{DMSO} \rightleftharpoons K \text{CH}_3\text{CONH}^+ \text{HOBr}
\]

\textbf{Scheme 46.} Reaction of DMSO with NBA

From the above mechanism the rate law (eqn. 44) came out to be:

\[
\text{rate} = \frac{kKK_{[\text{NBA}][\text{DMSO}]}(K_{\text{h}} + [\text{AA}][1 + K[\text{DMSO}])}{(K_{\text{h}} + [\text{AA}])} \tag{44}
\]

This rate equation was operative when [DMSO] was 0.01 to 0.5 mol dm$^{-3}$, below 0.01 mol dm$^{-3}$ the reaction was total second order, first order in each [reactant] and above 0.5 mol dm$^{-3}$ it was first order, being zero order in [DMSO]. Under high [DMSO] the zero order dependence of rate on [DMSO] and first order dependence on [HOBr] tempted us to speculate a rate determining polarization of the O-Br bond in HOBr to give a species of the type HO$^-$•••••Br$^+$. Because in the rate-determining step DMSO had to do nothing, except participating with HOBr in a fast step due to nucleophilic interaction of oxygen of HOBr with electrophilic sulfur to generate some kind of highly unstable transition state which may later decompose to give products. Will really a rate determining polarization of any chemical species be possible? Even today this is a question to readers. Therefore we did not go into the details of this concept while we published this article in Indian Journal of Chemistry.

Effect of solvent polarity: Attempts have been made to analyze medium effects on various kinetic constants i.e. $K_{\text{h}}$, $K$ and $k$. There were abrupt changes in $K_{\text{h}}$ and $K$ at the mole fraction of 0.4 of methanol. As the mole fraction of methanol increases there was about 3 to 4 times sudden increase in $K_{\text{h}}$ which suggests that there might be greater stabilization of possible intermediate (scheme 47) formed during the hydrolysis of NBA which was due to enhanced H-bonded interaction in media of higher methanol content.

\textbf{Scheme 47.} Interaction of water with NBA
On the contrary the formation constant $K$ showed a sharp decrease of about 2.5 to 3 times at mole fraction of methanol of 0.4, which could be attributed to the destabilization of the adduct formed in the second step of the reaction due to enhanced hydrophobic interaction between the methyl group of methanol and methyl groups of DMSO. The rate determining step involved an intramolecular oxygen atom transfer between DMSO and HOBr parts of the adduct. As the content of methanol increased, the rate of decomposition of the adduct decreased steadily. This could be due to the enhanced H-bonded interactions at the oxygen site of the dipolar adduct, there by making the intramolecular oxygen atom transfer difficult. Hence a decrease in the values of $k$ was observed.

### 3.10. Oxidations by HOBr

Survey of literature revealed that hypobromous acid (HOBr) is generated by hydrolysis of N-bromo compounds such as N-bromoacetamide (NBA), N-bromosuccinamidine (NBS), and N-bromobenzamide (NBB), which later reacts with substrates. We have recently identified that HOBr was found to be easily generated in situ from NaBrO$_3$ in the presence of an appropriate reducing agent such as NaHSO$_3$, which was successfully used, in the synthesis of several organic compounds [77]. And we are the first to under take the kinetic investigations using this reagent. It was very easy to handle this reagent in aqueous solution as compared to that of the aqueous solutions of several N-halo compounds. Since the aqueous solutions of N-halo compounds are not stable, every time they have to be prepared afresh when necessary instead it was very easy, less time consuming and economical to handle the sodium bromate-sodium bisulfite reagent for generating HOBr. In view of the above advantages of sodium bromate-sodium bisulfite reagent over N-halo compounds, this reagent has been selected and used in the kinetic investigations of the reactions with DMSO [78], cinnamic acids [79], benzylamines [80], phenols [81] and benzaldehydes [82]. All these reactions followed the same mechanism that there was no reaction step involving the acetamide transfer difficult. Hence a decrease in the values of $k$ was observed.

#### 3.10.1. Kinetics and Mechanism of Oxidation of Dimethyl Sulfoxide by Sodium Bromate-Sodium Bisulfite Reagent in Aqueous Medium [78]

Rates of oxidation of DMSO by HOBr produced in situ from sodium bromate -sodium bisulfite reagent were studied iodometrically in aqueous medium. The order in [DMSO] is one when [DMSO] < 0.01 mol dm$^{-3}$, fractional when [DMSO] is 0.01-0.5 mol dm$^{-3}$ and zero when (DMSO) > 0.5 mol dm$^{-3}$. Different rate laws were operative under these three conditions though HOBr was the effective oxidizing species in all the cases. A mechanism involving an intermediate four-membered cyclic transition state between DMSO and HOBr with a formation constant ($K$), which decomposes in a slow step with a rate constant ($k$) was proposed. Thermodynamic parameters for the formation of adduct and activation parameters for the first-order decomposition of the adduct were evaluated. Activation parameters were also determined while the orders in [DMSO] were unity and zero. And all these were discussed. The reaction product was identified as di-methyl sulfone (DMSO$_2$).

#### 3.10.2. Kinetics and Mechanism of in situ Bromohydroxylation of Cinnamic Acids by Sodium Bromate-Sodium Bisulfite Reagent in Aqueous Acetonitrile Medium [79]

Rates of bromohydroxylation of 4-substituted cinnamic acids with HOBr, produced in situ from sodium bromate-sodium bisulfite reagent have been studied in aqueous acetonitrile medium iodometrically. Thermodynamic parameters for the formation of the adduct and the activation parameters for the first order decomposition of the adduct have been evaluated in the temperature range and discussed. The influence of the substituents on the first order decomposition rate constant ($k/s^{-1}$) of the adduct has been investigated. The corresponding bromohydrids of cinnamic acids have been identified as the reaction products.

#### 3.10.3. Kinetics and Mechanism of Oxidation of Benzylamines by Sodium Bromate-Sodium Bisulfite Reagent in Aqueous Medium [80]

Kinetics of oxidation of benzylamines (BA) by HOBr produced in situ from sodium bromate-sodium bisulfite reagent has been studied in aqueous medium at constant pH = 9.2 iodometrically. The reaction was overall second order and first order in each reactant i.e., HOBr and BA. HOBr has been postulated as effective oxidizing species. The activation parameters for the second order rate constant ($k/dm^3 mol^{-1} s^{-1}$) have been evaluated and discussed and they indicated the reaction series was entropy controlled. The influence of the substituents has been investigated. The Hammett p value was found to be −0.64. A mechanism (scheme 48) involving transfer of a hydride ion from the methylene group of benzylamine to oxidant HOBr in the rate-determining step has been proposed. The sole reaction products have been identified as corresponding benzaldehydes and NH$_3$.

[Scheme 48. Reaction of benzylamines with HOBr]
3.10.4. Kinetics and Mechanism of Bromination of Phenols by Sodium Bromate-Sodium Bisulfite Reagent in Water -Acetonitrile Mixture [81]

Kinetics of bromination of phenols with HOBr produced in situ from sodium bromate-sodium bisulfite reagent has been studied in water-acetonitrile mixture iodometrically. The order of the reaction was found to be unity in [HOBr] and fractional in [phenol]. HOBr has been established as the effective brominating species. A mechanism involving a six-membered cyclic adduct between phenol and HOBr, which decomposes in a slow step has been proposed. Thermodynamic parameters for the adduct formation and the activation parameters for the first order adduct decomposition have been evaluated and discussed. The influence of the substituents on the first order decomposition rate constant (k/s⁻¹) of the adduct has been discussed. The sole reaction product has been identified as the corresponding o-bromophenol.

3.10.5. Kinetics and Mechanism of Oxidation of Benzaldehydes by Sodium Bromate-Sodium Bisulphite Reagent in Aqueous Acetonitrile Medium [82]

Kinetics of oxidation of substituted benzaldehydes (BH) by HOBr produced in situ from sodium bromate-sodium bisulphite reagent has been studied iodometrically in aqueous acetonitrile (50:50 v/v) medium at constant acidity. The reaction was overall second order and first order in each reactant. The reaction was acid catalyzed and the rate was not affected by the addition of Na₂SO₄. Formation a cyclic intermediate in a slow step and subsequent decomposition of this in a fast step by a hydride ion transfer route were suggested as a probable mechanism. Activation parameters have been evaluated and discussed. The influence of the substituents on the oxidation reaction rates and rate constants has been investigated. The sole reaction product has been identified as the corresponding benzoic acid.

3.11. Oxidations by Peroxo Compounds: Two Peroxo Anions, Peroxodiphosphate [83] and Peroxomonsulfuric acid [84]

3.11.1. Kinetics and Mechanism of Ruthenium(III) Catalyzed Oxidation of Lactic Acid by Peroxodiphosphate [83]

The kinetics of Ru(III)-catalyzed oxidation of lactic acid (LA) by peroxydiphosphate (PP) was studied in aqueous acid medium. An analysis of the rate dependence on [H⁺] revealed that the active oxidizing species in the oxidation could be H₂PO₄⁻. The Ru(III) catalysis of (PP-La) reaction was explained in terms of a 1:1 complex formation between Ru(III) and LA which later reacts with PP bimolecularly to give acetaldehyde and carbon dioxide as the products. The mechanism suggested was as given in scheme 49:

From the above mechanism the rate law (eqn. 45) derived was:

\[ \frac{-d[PP]}{dt} = kK[PP][LA][Ru(III)] } \]

which explained all the kinetic features.

\[ \text{PMS}^- + \text{Ru}^{III} \rightarrow \text{PMS}^2^- \]

\[ \text{PMS}^2^- + \text{LA} \rightarrow \text{PMS}^- + \text{Ru}^{III} + \text{H}^+ + \text{CO}_2 \]

Scheme 49. Reaction peroxydiphosphate with lactic acid

3.11.2. Reactivities of Mono and Di-Anions of Peroxomonsulfuric acid Towards Benzaldehydes: A Kinetic and Mechanistic Study [84]

The rates of reactions of different species of peroxomonsulfate (PMS) with benzaldehydes RC₆H₄CHO (R = H, 4-NO₂, 3-NO₂, 4-Me, and 4-MeO) have been measured iodometrically in CH₃CN-water in the presence of phosphate buffers. The reaction was overall second order and the rate increased with the increase in pH and was very fast and almost instantaneous at pH > 8. This could be due to the slow accumulation of more reactive PMS²⁻ (see below). Hence any reaction beyond this pH could not be conducted. Neutral peroxomonsulfuric acid reacts very slowly with benzaldehydes. The locus of the Hammett plot had a minimum near the point for benzaldehyde showing that there was a change in the rate-determining step. A mechanism involving addition/elimination between PMS and benzaldehyde near neutral pH has been proposed as shown in Scheme 50:

Scheme 50. Reaction of benzaldehyde with peroxy mono sulfonic acid

It was evident from the mechanism and the non-linear Hammett plot that the first reversible addition of PMS to the carbonyl was accelerated by electron-withdrawing substituents in the phenyl ring since the nucelophilicity of carbonyl carbon increases. This was followed by the elimination of SO₄²⁻ from the adduct, a reaction which could be accelerated due to presence of electron-donating substituents in the aromatic nucleus. The balance between these two factors could result in a minimum or a maximum in the overall rate. In the present investigation the minimum rate was observed for benzaldehyde. PMS could exist in two anionic forms, viz. PMS⁻ (HOOSO₃H ⇌ HOOSO₃⁻ + H⁺), due to dissociation of sulfuric acid proton with a pKₐ of ~3.00) and PMS²⁻ (HOOSO₃⁻ ⇌ `OOSO₃⁻ + H⁺), due to dissociation of peroxyde proton with a pKₐ of 9.4) depending on the pH in aqueous solution. Since PMS was inert, both PMS⁻ and PMS²⁻ could react with BA as shown in scheme 51:

\[ \text{BA} + \text{PMS}^- \rightarrow \text{products} \]

\[ \text{BA} + \text{PMS}^2^- \rightarrow \text{products} \]

Scheme 51. Reactions of anions of PMS with benzaldehyde
Hence the rate law (eqn. 46) could be:

$$\text{rate} = k_1 [BA][PMS^-] + k_2 [BA][PMS^{2-}]$$  \hspace{1cm} (46)

This could be rearranged to eqn. 47:

$$\text{rate} = k_1 [BA][PMS^-] + \frac{K_h k_2 [BA][PMS^-]}{[H^+]}$$  \hspace{1cm} (47)

where $K$ is the second dissociation constant of PMS. From the rate dependencies on $H^+$ at constant [BA], $k_2$ the second order rate constants for the reaction of PMS$^{2-}$ could be obtained. The values of $k_2$ did not show any regular trend with the substituents in the benzaldehyde ring but they were found to be 3 to 100 orders of magnitude higher than $k_1$ again depending on the substituent in the benzaldehyde ring. And this was not surprising because PMS$^{2-}$ was known to be more reactive than PMS$^-$ since PMS$^{2-}$ with two negative charges on it was more nucleophilic to be added to electrophilic carbon of carbonyl group of benzaldehyde than PMS with one negative charge. The very high negative entropy $\Delta S^\circ$ values (-130 to -275 J K$^{-1}$ mol$^{-1}$) were in accordance with the formation of the addition complex.

3.12. Inner-Sphere Synchronous Three Electron Transfer Reactions

A review has been published on this aspect [85] along with some original publications of the work [86, 87], and [88]. In literature there were several examples on Cr(VI) oxidations. An interesting feature of this reaction was that Cr(VI) oxidation of a mixture of a hydroxy acid or di-carboxylic acid and 2-propanol proceeds much faster than that of either of the two substrates alone. A termolecular complex was supposed to decompose in a rate-determining step [86]. Free radicals were detected as intermediates by polymerization technique. Acetone from 2-propanol and CO$_2$ from acid were the products. The individual oxidation proceeds by two-electron oxidation and the mixed system proceeds by three-electron oxidation and this is now known as “co-oxidation”. A study of mixed substrate oxidation of an alcohol and an aldehyde showed that the redox process took place in a single step three-electron oxidation [87]. Another important aspect of this kind of study was that a substrate having three functional groups of same kind or different could also proceed in a single step three-electron oxidation [88].

3.12.1. Co-Oxidation of Iso-Propanol and Lactic Acid in the Presence of Chromic Acid: A Case of Three-Electron Oxidation Study [86]

The rate of oxidation of (CH$_3$)$_2$CHOH by chromic acid was increased considerably in the presence of lactic acid. The rates were acid catalyzed and dependent on the polarity of solvent but independent of added salts. A mechanism in which formation of a 1:1:1 termolecular complex followed by its decomposition in a single 3-electron redox reaction was suggested. However, the acceleration in rate was not observed when 1,2-glycols were used in place of LA indicating the absence of “co-oxidation”. This might be due to the fact that 1,2-glycols are not good complexing agents like the one, hydroxy acid or a di-carboxylic acid. From the products and the kinetic trends the mechanism was suggested as given in scheme 52:

**Scheme 52.** Reaction of 2-propanol with Cr(VI) in presence of lactic acid

3.12.2. Three-Electron Oxidation: Chromic Acid Oxidation of a Mixed Substrate System of an Alcohol and Aldehyde [87]

The kinetics of oxidation of CH$_3$CH$_2$CHO (I) and (CH$_3$)$_2$CHCHOH (II) and their mixture by chromic acid was examined in aqueous acetic acid. A first-order dependence on [Cr(VI)] was observed. First-order kinetics in [I] or [II] alone were also found; fractional orders were found for I and II when their mixture was oxidized. The rates were independent of added salts and changes in solvent polarity. A mechanism was proposed involving neutral molecules and formation of a di-ester (shown in scheme 53) of chromic acid with aldehyde and alcohol in two successive equilibrium steps. The di-ester then decomposes in a slow step to give the products. Equilibrium constants and the rate constants for the uni-molecular decomposition of the di-ester were evaluated.

**Scheme 53.** Reaction of mixture of an alcohol and an aldehyde with Cr(VI)

Other steps were similar to those mentioned in “co-oxidation”. However distinguishing between these two rate determining steps was not possible. May be one could do the distinguishing by the study of isotope labelling.

3.12.3. Kinetics and Mechanism of Oxidation of Glycerol by Chromic Acid in Sulfuric Acid Medium [88]

The title reaction was overall second order, first order in each [reactant]. The reaction was acid-catalyzed and the rates
were independent of added salts, but dependent upon solvent polarity. The oxidation products were HCHO and CHOCHO, formed via a free radical mechanism in direct three electron change step. The oxidation of glycerol was compared with that of 1,2-diol and 1-propanol. From the change in activation entropy, it was found that the large negative value for glycerol oxidation (-164 J K⁻¹ mol⁻¹); but for 1,2-diol and 1-propanol oxidation they were only -98.0 and -90.0 respectively) could lead to suggest the formation of a rigid binary complex (Scheme 54) in the rate determining step which latter gives the products. Here glycerol was supposed to act as a tridentate ligand with its three OH groups.

**Scheme 54.** Decomposition of the binary complex of Cr(VI)

Other steps being again similar as shown earlier for mixed substrate oxidation and co-oxidation.

3.13. Inner-Sphere Synchronous Four Electron Transfer Reaction: A Rapid Reaction of Manganese(VII) with Two Component Substrate Systems Containing 2-Propanol and some Bifunctional Compounds: A Kinetic Study

The rates of the permanganate ion oxidation of some bifunctional compounds such as hydroxy acids and 1,2-glycols in the presence of 2-propanol (IPA) were higher than the rates of oxidation of either of those compounds or IPA alone under identical conditions. The orders in [substrate] and rates of oxidation of either of those compounds or IPA alone (Mn(II)), unlike in the Cr(VI) co-oxidation, mixed oxidation and glycerol oxidation where the end product of Cr(VI) was Cr(III) which is a stable oxidation state of chromium. Therefore the possibility of further oxidation of acetone and acetaldehyde by Mn(III) could be possible in which case the kinetics would have been much complicated. But this possibility was ruled out on the basis of the fact that Mn(III) oxidation of organic substrates needs high mineral acid concentrations say 2 to 3 M, where as with Mn(VII), four electron oxidation was carried out only in presence of 0.40 M sulfuric acid in which case the oxidation by Mn(III) was supposed to be negligible.

4. Conclusions

Organic chemistry and inorganic chemistry investigated by the methods and techniques of Physics and Physical Chemistry have been particularly fruitful areas of study. And although physical chemists have always investigated the properties and reactions of organic compounds, the discipline that today we recognize as Physical Organic Chemistry really began when organic chemists themselves became seriously interested in the relationship between molecular structure and chemical properties, and in how, at the molecular level the reactions of organic compounds take place. This review in my opinion may lead the readers to offer to the students at an early stage the important distinction between experimental facts and reaction mechanisms.

Structures of reactants and products are usually the principal features of a chemical reaction to be established. Equilibrium constants for reversible processes are also normally determinable, and a third experimental aspect of a reaction which may be quantified is its rate (or velocity). Molecular structures and reaction parameters can be expressed numerically with associated degrees of precision which are usually taken as measures of accuracy. In contrast a mechanism - a view of how reactants become products - constitutes a set of deductions derived by theory and intuition from the experimental results. A mechanism is not, therefore, a primary experimental feature in the way that a molecular structure, an equilibrium constant, or a rate constant is, and we can never express it in fine detail and with complete confidence.

The general organic chemistry text book may not describe completely how reactions take place, why one reactant is more reactive than the other under the same experimental conditions. It is aimed in this review, however, is not to demonstrate in detail, but how experimental results could lead towards a mechanistic understanding of a particular reaction. This may not be possible over a wide subject area. But indeed it is the aim of the author to illustrate in a single progressive account the application of principles by which this knowledge is used to interpret the experimental results with in the general context of organic chemistry. This attempt could lead to the idea of interfacing between organic and physical chemistry may be seen as an amalgamation.

**Dedication:** This review is dedicated to Prof. Dr. Steen Steenken (Retd) of the then Max-Planck-Institute for Radia-
Acknowledgement: It is my pleasure to owe my heartfelt gratitude and sincere thanks to Dr. B. K. Mishra, M.Sc., Ph.D., D.Sc., Professor and Head, Department of Chemistry, Sambalpur University, Jyoti Vihar, Orissa-768 019, India and to Professor G. B. Behera, M.Sc., Ph.D., D.Sc., (Retd.), Department of Chemistry, Sambalpur University, without these two good personalities, my review would have not come into existence for which I was dreaming since a long time.

I should not forget to thank Professor Dr. D. Schulte-Frohlinde, (Retd, now living in USA) the then Director, Max-Planck-Institute for Radiation Chemistry, Muelheim a.d. Ruhr, Germany for having invited me as a Post-Doctoral Fellow during 1980-88 and attaching me to a giant Physical Organic Chemist Professor Dr. Steen Steenken

I take this opportunity to convey my sincere gratitude to Professor Dr. Steen Steenken (Retd), Max-Planck-Institute for Bioinorganic Chemistry (The then Max-Planck-Institute for Radiation Chemistry), Muelheim a. d. Ruhr, Germany.

I also thank Professor John P. Richard and Professor Tina L. Ames (then at the Department of Chemistry, University of Kentucky, Lexington, Kentucky) Department of Chemistry, State University of New York at Buffalo, Buffalo, U.S.A. for their good wishes.

I thank all my PhD students Dr. P. Musala Reddy, Dr. T. Rayapa Reddy, Dr. S. Venkateswarlu, Dr. S. Venkateswar Rao and Dr. T. Satyanarayana Reddy, Readers, Department of Chemistry, Sardar Patel college, Secunderabad from where I also started earning my bread and butter through teaching chemistry during 1978-1983, Dr. (Mrs.) P. R. Sharadamani, Reader, Department of Chemistry, Kasturba Gandhi College for women, Secunderabad, Dr. Yogyaraj, Lecturer, Department of Chemistry, N. B. Science College, Hyderabad, Dr. G. Krishna Reddy, Postgraduate Teacher in Chemistry, then at Kendriya Vidyalaya, Kanchanbagh, Hyderabad, Dr. R. Sanjeev, as a DST (New Delhi) funded JRF, Dr. J. Viroopakshappa, Reader, Department of Chemistry, Tara Government Degree College, Sangareddy (Andhra Pradesh) were instrumental in contributing a lot to my research program since last 37 years.

Thanks are also due to Professor Masaaki Mishima and Professor Yuho Tsuno, Department of Chemistry, Faculty of Science, Kyushu University, Fukuoka, Japan, Dr. Yong Gu Lee, a Post-Doctoral Fellow from the Department of Chemistry, Kangwon National University, Chuncheon, Korea, and Dr. Douglas J. Rice, then a Graduate Student in Dr. John’s laboratory, in the Department of Chemistry, University of Kentucky, Lexington, Kentucky who contributed their share to my research program.

I express my thanks to my wife Vijaya, my son Suresh, and my daughter Sharada for their patience and uncomplaining tolerance during my entire research activity.

Finally I wish to record my indebtedness to my father Akkaiah (Late) and to my mother Savithramma who gave me the birth to exist in this world in my present position.

I have written this review with utmost care not to involve any grammatical mistakes taking the help of my friends, who teach English, but some mistakes will inevitably remain; these are my own responsibility.

REFERENCES

[1] R. E. Kirk and A.W. Brown., J. Am. Chem. Soc., 50, 337 (1928)
[2] F. A. Cotton and G. Wilkinson, Advanced inorganic chemistry., 3rd ed. Wiley Eastern Pvt. Ltd., (1976) 674.
[3] W. G. Movins and R. G. Link., J. Am. Chem. Soc., 91, 5394 (1969)
[4] J. W. Baker and T. G. Heggs and R. F. W. Bader., J. Chem. Soc., 616 (1955)
[5] C. G. Swain, R. A. Wiles and R. F. W. Bader., J. Am. Chem. Soc., 83, 1945 (1961)
[6] C. G. Swain, R. F. W. Bader, R. M. Esteve, Jr. and R.N. Griffin., J. Am. Chem. Soc., 83, (1951) 61
[7] K. B. Wiberg, Surv. Prog. Chem., 1, 211 (1963)
[8] P. Saffir and H.Taube., J. Am. Chem. Soc., 82, 13 (1960)
[9] R. T. M. Fraser and H.Taube., J. Am. Chem. Soc., 82, 4152 (1960)
[10] S. Venkateswar Rao and V. Jagannadharnad., React. Kinet. Catal. Lett., 27, 239 (1985)
[11] F. Basolo, M. L. Morris and R. G. Pearson, Discuss. Faraday Soc., 29, 80 (1960)
[12] F. Hasan and J. Roček, J. Am. Chem. Soc., 94, 3181 (1972)
[13] M. Anand Rao, B. Sethuram, T. Navaneeth Rao and V. Jagannadham., Oxidin. Commsns., 9, 247 (1986)
[14] H. Taube, Can. J. Chem., 37, 129 (1959)
[15] (a) H. Taube, H. Meyers and R. L. Rich., J. Am. Chem. Soc., 75, 4118 (1953), (b) H. Taube and H. Meyers., J. Am. Chem. Soc., 76, 2103 (1954)
[16] H. Taube, J. Am. Chem. Soc., 77, 4481 (1955)
[17] R. Stewart, Oxidation Mechanisms: Application to Organic Chemistry, W. A. Benjamin Inc., 66, (1964)
[18] V. Jagannadham and S. Steenken, J. Am. Chem. Soc., 106, 6542 (1984)
[19] J. Roček and C.G. Ng., J. Am. Chem. Soc., 95, 1522, 2840 (1974)
[20] R. B. Woodward, N. L. Wendler and F. J. Brutschy, J. Am. Chem. Soc., 67, 1425 (1945)
[21] L. P. Hammett., Physical Organic Chemistry, P. 350, McGraw-Hill Book Co., Inc., New York, NY, 1940
[22] E. Pfeil. Chem. Ber., 84, 229 (1951)
[23] C. B. Pollard and D. C. Young, J. Org. Chem., 16, 661 (1951)
[73] S. Venkateshwarlu, and V. Jagannadham, Indian J. Chem., Sect. A, 27A, 314 (1988)

[74] P. R. Sharadamani, and V. Jagannadham, Natl. Acad. Sci. Lett. (India), 14, 331 (1991)

[75] P. R. Sharadamani, V. Jagannadham, and V. Venkat Rao, Indian J. Chem., Sect. A, 30A, 514 (1991)

[76] P. R. Sharadamani, and V. Jagannadham, Indian J. Chem., Sect. A, 30A, 518 (1991)

[77] a) H. Ohta et al. J. Org. Chem., 54, 1668 (1989); b) H. Masuda et al. J. Org. Chem., 59 5550 (1994); c) H. Ohta et al. Chem. Lett., 773 (1990); d) K. Takase, et al. Chem. Lett., 871 (1995)

[78] J. Viroopakshappa and V. Jagannadham, Indian Acad. of Sci., 114, 149 (2002)

[79] J Viroopakshappa and V Jagannadham, Indian J. Chem., 42B, 374 (2004)

[80] J Viroopakshappa, Ph.D. thesis, Osmania University, Hyderabad, India, 2003

[81] J Viroopakshappa and V Jagannadham, Ind. J. Chem., 43A, 532 (2004)

[82] J Viroopakshappa and V Jagannadham Oxid. Commns. 27, 623, 2004

[83] V. Jagannadham, M. Anand Rao, B. Sethuram and T. Navaneeth Rao, Oxid. Commns. 8, 41, (1985/86)

[84] Ahmad Naseeruddin, A. S. R. Swamy, and V. Jagannadham, Indian J. Chem., Sect. A 26A, 427 (1987)

[85] V. Jagannadham, Indian J. Chem. Sci., 2, 21 (1988)

[86] V. Jagannadham, M. Anand Rao, B. Sethuram, and T. Navaneeth Rao, Oxid. Commun. 8, 31 (1986/1985)

[87] P. Musala Reddy, V. Jagannadham, B. Sethuram, and T. Navaneeth Rao, React. Kinet. Catal. Lett. 21, 437 (1982)

[88] P. Musala Reddy, V. Jagannadham, B. Sethuram, and T. Navaneeth Rao, Indian J. Chem., Sect. A, 21A, 608 (1982)

[89] M. Anand Rao, B. Sethuram, T. Navaneeth Rao, and V. Jagannadham, Oxid. Commun. 9, 247 (1986)