Micellar Catalysis on 1,10-Phenanthroline Promoted Chromic Acid Oxidation of Propanol in Aqueous Media

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(Received December 29, 2011; Accepted December 30, 2011)

Key words: Propanol, Chromic acid, Phenanthroline, Micellar catalyst

The catalysis of reaction by organized media is of increasing interest. In these systems, aggregates act as nano or microreactors, compartmentalizing and concentrating or diluting reagents and thereby altering the observed rate of chemical reactions. Surfactant systems are well known for their high capacity for reactant solubilization in spite of their type: organic, inorganic, polar and non-polar. The green chemistry aspect of using water as environmentally and user-friendly solvent for chemical reactions has sparked much interest with an impressive number. A micellar system appears to be homogeneous since these aggregates are of colloidal size; however in reality the absorbed reagents are in a microheterogeneous two phase system. The catalytic efficiency will be governed by the affinity of the reactivity of the bound reagent molecules. Oxidation is a key reaction for different organic synthesis. Chromic acid is a widely used oxidizing agent. 1, 10-phenanthroline is well known to act as a promoter for Cr(VI) oxidation of organic substances. Bipyridine (bpy) and picolinic acid are also used as good promoter for chromic acid oxidation but here the work is only concentrated on the promoter phenanthroline. Under the kinetically reaction conditions [propanol]_T >> [Cr(VI)]_T (subscript T stands for the total concentration), and the alcohol upon oxidation yields propanal which was confirmed by the production of 2,4-dinitrophenylhydrazone derivatives. The overall stoichiometry of the reaction may be represented as:

\[ 3\text{CH}_3\text{CH}_2\text{CH}_2\text{OH} + 2\text{HCrO}_4^- + 8\text{H}^+ \rightarrow 2\text{Cr(III)} + 3\text{CH}_3\text{CH}_2\text{CHO} + 8\text{H}_2\text{O} \]

The rate of disappearance of Cr(VI) shows the first order kinetics on [Cr(VI)]_T. The pseudo first-order rate constants \( k_{obs} \) have been determined from plots of \( \ln([\text{Cr(VI)}]_0-kt) \). The formation of a Cr(III)-phen complex (identified spectroscopically) indicates that phen undergoes complexation with the higher oxidation states (which are labile) of chromium. Due to the inertness of Cr (III), phen does not bind the Cr(III) produced after the completion of Cr(VI). Here it is quite reasonable to accept that Cr(VI)-phen complex formed at the pre-equilibrium step which is the active oxidant for this reaction. The observed rate law and the cyclic transition state reduction of Cr(VI) to Cr(IV) occur through H+ transfer (rate determining step) is supported by our previous work on the oxidation of propanol to propanal.

Scheme 2 is drawn on the basis that the reaction follows first order dependency on [propanol], [Cr(VI)], [Phen], and [H+] (Not shown in the paper).

The colours of the final solutions in the absence of promoter and in the presence of the promoter are far different indicating different types of species. The observed final colour of the solution in the absence of phen under the experimental condition is pale blue (\( \lambda_{max} = 413 \text{ nm and 586 nm} \)) and the corresponding transitions are 586 nm for \( ^4\text{A}_{2g}(\text{F}) \rightarrow ^4\text{T}_{1g}(\text{F}) \) and 413 nm for \( ^4\text{A}_{2g}(\text{F}) \rightarrow ^2\text{T}_{2g}(\text{F}) \) of Cr(III) species while the final colour of the solution of phen-promoted reaction under the same kinetic condition is pale

\[ \text{CH}_3\text{CH}_2\text{CH}_2\text{OH} + \text{HCrO}_4^- + \text{H}^+ \rightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{O}^- + \text{Cr}^{3+} + \text{H}_2\text{O} \]

1 + \text{H}^+ \rightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{O}^- + \text{CrO}_3^- + \text{OH}^- \]

Scheme 1. Cr(VI) oxidation of propanol in the absence of 1, 10-phenanthroline.
violet ($\lambda_{\text{max}}=546$ nm, for $^4A_{2g}(F) \rightarrow ^4T_{2g}(F)$ of Cr(III) species). The spectra of the final solution without promoter reaction and pure chromic sulphate solution in aqueous sulphuric acid media are identical. This is consistent with the final Cr(III) species which is simply Cr(III)-species for the unpromoted reaction while for the promoted reaction (phen); the final Cr(III)-species is a different species, which is Cr(III)-promoter complex. In the phen promoted reaction path, there is a hypsochromic (blue) shift (Fig. 1) for the peak due to the transition $^4A_{2g}(F) \rightarrow ^4T_{2g}(F)$ compared to the final solution without promoter path. Phenanthroline contain two strong field donor site i.e., heteroatomic shift. The peak due to the transition $^4A_{2g}(F) \rightarrow ^4T_{1g}(P)$ simply explains the charge-transfer band in the Cr(III)-promoter complex (Fig. 1). The band around 265 nm in Cr(III)-aqueous species arises as a shoulder due to $^4A_{2g}(F) \rightarrow ^4T_{1g}(P)$ transition and it is of high energy charge-transfer band i.e., in ultraviolet region. In Cr(III)-promoter complex, the
appearance of the charge-transfer band at much lower energy i.e., in visible range for the proposed Cr(III)-promoter complex is quite reasonable because of the favoured metal to ligand charge-transfer. The vacant π* M.O. of the phen switches the metal to ligand electron-transfer. The rate of the in phen promoted path is excessively enhanced by the charge-transfer band (metal to ligand) at lower energy. Wavelength of this corresponding charge-transfer band is shifted to a lower value than the wavelength of the corresponding band in unpromoted path.

The scanned spectrum indicates the gradual disappearance of Cr(VI) species and appearance of Cr(III) with an isobestic point at λ = 524 nm for the unpromoted (Fig. 2a) and phen promoted (isobestic point at λ = 518 nm) (Fig. 2b) reaction.

The following scanned spectrum is observed for SDS (in presence of promoter phen) catalyzed chromic acid oxidation of propanol with an isobestic point at λ = 520 nm (Fig. 3).

Triton X-100 with promoter (phen) gives the similar type of scanned spectrum for the disappearance of Cr(VI) species and appearance of Cr(III) species with an isobestic point at λ = 516 nm (Fig. 4).

CPC with promoter (phen) gives the similar type of scanned spectrum for the disappearance of Cr(VI) species and appearance of Cr(III) species with an isobestic point at λ = 522 nm (Fig. 5).

In unpromoted path (Scheme 1) the partitioning of neutral ester in all kind of micellar phase is possible. Partitioning of proton is maximum in SDS due to electrostatic attraction less in CPC due to electrostatic repulsion. Rate is maximum in SDS, minimum in CPC and TX-100 has effect in between them (Table 1).

The active oxidant participate is Cr(VI)-phen complex (supported by Fig. 6) reacts with the substrate (Scheme 2)
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to form a ternary complex which experience a redox decomposition in a rate determining step giving rise to organic product. As Cr(VI)-phen is a giant molecule electrostatic attraction is not operating smoothly. For promoted reaction rate is maximum in TX-100 followed by SDS (Table 2). Most probably the large hydrophobic core of TX-100 (greater than SDS) allows maximum number of neutral substrate. In case of unpromoted reaction small proton easily penetrated in SDS micelle due to electrostatic attraction, so rate is highest in SDS. But in promoted reaction maximum number of neutral substrate in TX-100 hydrophobic core predominates over electrostatic attraction by positively charged micellar head group of SDS to active oxidant Cr(VI)-phen.

**Table 2.** Reaction completion time in presence of micellar catalyst and in presence and absence of promoter

<table>
<thead>
<tr>
<th>Micellar catalyst [mol dm⁻³]</th>
<th>T_completion [hrs]</th>
</tr>
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<tbody>
<tr>
<td>None</td>
<td>1.45</td>
</tr>
<tr>
<td>TX-100</td>
<td>2×10⁻², 3×10⁻², 4×10⁻², 5×10⁻²</td>
</tr>
<tr>
<td>SDS</td>
<td>1×10⁻², 2×10⁻², 3×10⁻², 4×10⁻²</td>
</tr>
<tr>
<td>CPC</td>
<td>2×10⁻², 4×10⁻², 6×10⁻², 8×10⁻²</td>
</tr>
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[Cr(VI)]_T = 5×10⁻⁴ mol dm⁻³, [H₂SO₄]_T = 0.5 mol dm⁻³, [propanol]_T = 75×10⁻⁴ mol dm⁻³, [phen]_T = 50×10⁻⁴ mol dm⁻³, Temp = 30°C.

**EXPERIMENTAL**

1, 10-phenanthroline (AR, Merck), propanol (AR, Qualigens), K₂Cr₂O₇ (AR, BDH), sodium dodecyl sulphate (SDS) (AR, SRL), TX-100 (AR, SRL) and all other chemicals used were of highest degree of purity available commercially. All the solutions were prepared in double distilled water. Solutions of the oxidant and reaction mixtures containing the known quantities of the substrate(s) (i.e. propanol) and promoter (1,10-phenanthroline) under the kinetic conditions [propanol] >> [Cr(VI)] and acid and other necessary chemicals were separately thermostatted (±0°C). The reaction was initiated by the requisite amounts of the oxidant with the reaction mixture. Progress of the reaction is monitored by following the rate of disappearance of Cr(VI). The concentration of Cr(VI) at different time intervals was measured by a titrimetric quenching technique using excess of standard Mohr’s solution and unreacted Fe(II) was estimated by a standard Ce(IV) solution using ferroin indicator. The pseudo first order rate constants were calculated from the slopes of the plot of log [Cr(VI)] versus time t, which were linear at least for three half lives. The scanned spectra, spectrum after completion of the reaction and other spectra were recorded with a UV-VIS spectrophotometer [UV-1800 and UV-VIS NIR-3600 (SHIMADZU)]. Quartz cuvettes of path length 1 cm were used. Under the experimental conditions, the possibility of decomposition of the surfactants by Cr(VI) was investigated and the rate of decomposition in this path was kinetically negligible.

Acknowledgement. Thanks to University of Burdwan and CSIR, New Delhi for offering the opportunity to write
and for providing financial help in the form of a project and fellowship.

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