Catalytic Studies of Complexes of Organic Compounds
Part-4: Synthesis, Characterization, Catalytic activity of Cd(II) Complex of Chiral Schiff base

C. J. PATIL* and S. B. SALVE

Organic Research Laboratory, Smt. G. G. Khadse College, Muktainagar-425306, MS, India.
*Corresponding author E-mails: drcjpatil@yahoo.com

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ABSTRACT

Chiral Schiff base ligand from 3,5-Diiodo-salicylaldehyde with a chiral amine (1S,2S)- (+)-1,2-Diaminocyclohexane is synthesized, and its Cd (II) complex was synthesized. These were analysed by the physical constant, TLC, Colour, UV-Vis, FTIR and 1H NMR method. Also, efforts were made to study the catalytic activity of Cd(II) chiral Schiff base complex. The oxidation of benzyl alcohol was used as model reaction using acetonitrile as solvent. The present reaction system was heterogeneous system of catalysis.

Keywords: (1S,2S)-(+)-1,2-Diaminocyclohexane, O-N Donor Chiral Schiff base, Cd(II) complex, Benzyl alcohol, Oxidation and Catalysis.

INTRODUCTION

Oxidation of alcohol to the corresponding carbonyl compound (aldehyde, ketone or acid) is the key step in many organic synthetic methods. The catalytic oxidation of alcohol to aldehyde or ketone has been developed employing aqueous hydrogen peroxide (H2O2) in presence of Cd complex as catalyst is a safe process. In particular, benzyl alcohol (primary aromatic) was oxidized selectively to their corresponding aldehyde can be optimised in the present conditions. The oxidative transformation of primary alcoholic compounds to the aldehydes remains very problematic due to the oxidation of primary alcohols the other possible product (uncontrolled oxidation process) is the analogous carboxylic acids1-4. Last but not the least, tandem oxidation may be achieved without the need to isolate the any intermediate or change in solvent. Schiff bases and their metal complexes are of prime importance in the field of synthetic chemistry. Literature reports that these complexes are studied in the view of organic intermediates, metal complexation5 also voltammetric6 reduction and oxidation processes. The potential of chiral schiff base complex as catalyst towards oxidations, ring closer, epoxidation, hydrogenations, polymerizations and various coupling reactions7-8. Chiral Schiff bases are the reported as a catalyst viz. in asymmetric cyclopropanation9 and also in varied organic reactions.
The oxidative transformation of primary and secondary alcohols selectively into the respective aldehydes and ketones without forming undesired product is the most important reaction for both industrial applications and academic interest\textsuperscript{10-11}, still many processes of oxidation in use are non-environmentally friendly.

Varied reagents \textsuperscript{4,12-13} \(O_2\), \(H_2O_2\) \textsuperscript{14} and \(H_2O_2–TPAP\) doped ormosils\textsuperscript{15} were used for oxidative catalysis of alcohol to equivalent aldehydes and ketones. A phase transfer catalyst nBu\(_4\)NBr in aq. CH\(_2\)Cl\(_2\), sodium periodate in presence of Mn(TNP) Cl was used for the transformation of arylalkane and cycloalkane into the related alcohols and ketones at room temperature as reported\textsuperscript{16a} in the literature. Meanwhile a cheap protocol was reported\textsuperscript{16b} using \(H_2O\) as efficient solvent for the oxidation of aromatic alcohol and amines with \(H_2O_2\) (34\%) by heteropolyoxometalates. Also amechanistic approach was reported\textsuperscript{16c} for the oxidation of alcohol with \(H_2O_2\) (34\%) using simple heteropoly acids (\(H_3PW_{12}O_{40}\) and \(H_3PMo_{12}O_{40}\)) as water tolerant catalyst. Recently, in the literature \(WO_3\)ZnO/Fe\(_2\)O\(_3\) was prepared and was tested as nanophoto catalyst for the oxidation of benzylic alcohol and hence to form 2-substituted benzimidazoles by reacting with \(\alpha\)-Phenylendiamine\textsuperscript{16d}. An ecofriendly, simple and efficient method is described for the oxidation of some alcohols with 34\% hydrogen peroxide, catalysed by \(H_3PW_{12}O_{40}\) as simplest class of heteropolyoxometallates\textsuperscript{16e}.

Ru(III) complexes of the type [RuX\(_2\)(EPh\(_2\))\(_2\)(L)] (where \(X = Cl\) or Br; \(E = P\) or As; \(L\) = monobasic bidentate Schiff base ligands) have been synthesized and characterized. The catalytic efficiency of one of the ruthenium complex was determined in the case of oxidation of primary and secondary alcohols into their corresponding aldehydes and ketones in the presence of NMO (N-methylmorpholine-N-oxide) as co-oxidant\textsuperscript{17}. An enantiopure GO (galactose oxidase) enzyme model has been used synthesized from readily available (R)–binam and Cu(OTf)\(_2\)(TEMPO = 2,2,6,6–Tetramethyl–piperidin–1–oxyl) and has been effectively used as an efficient chiral catalyst for the oxidative kinetic resolution of secondary alcohols.\textsuperscript{18}

The dilute \(H_2O_2\) is an oxidant of ideal choice. It is cheap, readily available and gives water as the only by product. Many systems using aqueous \(H_2O_2\) as oxidant and Ligand-based catalysts under catalytic conditions have been reported by J. F. Larrow \textit{et al.},\textsuperscript{19} Also literature contains synthesis of ligand from (1R,2R)-(−)-1,2-Diaminocyclohexane and their complexes, However, literature reveals that there is scare or no publication on use of Cd complexes of chiral Schiff bases, an asymmetric catalyst, for the oxidative transformation of benzyl alcohol. As per the literature survey done by us there is no literature available on heterogenous catalytic oxidative transformation of alcohols by these synthesized complex N,N’-(S,S)-1,2-Cyclohexylenebis-(3,5-diodosalicylidene-iminato) cadmium (III) chloride, complex using N,O-donors along with \(H_2O_2\) as oxidant.

In continuation of our reported studies\textsuperscript{14} of complex of organic compound with respect to behaviour, here we have reported the synthesis of chiral Schiff base from 3,5-Diiodo-salicylaldehyde with chiral cyclodiamine, viz. (1S,2S)-(−)-1,2-Cyclohexylenebis-(3,5-diiodosalicylidene-iminato) cadmium (III) chloride, complex using N,O-donors along with \(H_2O_2\) as oxidant.

\textbf{MATERIAL AND METHODS}

The chemicals used were of synthesis grades, for the present work. TLC plates of aluminium material of Merck make (silica gel 60 F254) were used with TLC grade solvents for monitoring the progress and reaction completion. Iodine chamber was used to highlight the starting and final products spot identification in TLC. The estimation of elemental halogen and cadmium in complex was estimated by reported methods\textsuperscript{20-21}. FTIR spectral determination were made in 4000-600 cm\(^{-1}\) frequency on a Bruker Spectrum 2000 FT-IR spectrophotometer at SAIF, Kochi for the ligands while FTIR of complexes were recorded on Perkin FT-IR Spectrophotometer, at IISER, Bhopal in the in the range 4000-450 nm. The other instruments used for analysis are mentioned in earlier reports\textsuperscript{14}. 
**EXPERIMENTAL**

**Synthesis of Ligand**

The ligand, L was synthesized as per previously reported work procedure\(^{14}\). TLC using mobile phase Ethyl Acetate: n-Hexane 2.5:0.5 was performed, to monitor the completion of reaction. The yellow-brown coloured crude product was filtered, dried and recrystallized from ethanol, dried and kept in vacuum desiccator containing calcium carbonate as desiccant. On further drying the yellow-brown pure product was obtained, m.p. = 123-126°C.

\[
\text{(S,S)-N,N'-Bis(3,5-diiodo-salicylidene)-1,2-cyclohexanediamine, L}
\]

**Synthesis of Cd(II) complex of Ligand**

The cadmium complex was prepared according to reported work\(^{14,22}\). To the solution of (S,S)-N,N'-Bis-(3,5-diiodosalicylidene)-1,2-diaminocyclohexane, (50 mmol) in alcohol (40 mL) in 2 proportion added a solution of 1 proportion of Cadmium (II) chloride monohydrate (CdCl\(_2\).H\(_2\)O), (25 mmol) in aqueous alcohol (25 mL). Bulk reaction was reflux under stirring 2-3 h till a homogeneous TLC is monitored in 2.5:0.5 Ethyl Acetate: n-Hexane as mobile phase. The crude brown material formed was filtered and recrystallized from ethanol. It was dried and kept in vacuum desiccator.

**RESULTS AND DISCUSSION**

Ligand (S,S)-N,N'-Bis-(3,5-diiodosalicylidene)-1,2-diaminocyclohexane and the Cd(II) complex was synthesized in two steps. Firstly, the selected 3,5-Diiodosalicylaldehyde and the Chiral amine were condensed to get (S,S)-N,N-Bis-(3,5-diiodosalicylidene)-1,2-diaminocyclohexane. In the second step, cadmium (II) chloride monohydrate (CdCl\(_2\).H\(_2\)O) was used to prepare its Cd(II) complex i.e. \([(\text{DiiodoSalicylo})\text{Cd (II)Cl}]\). As mentioned in our reported work\(^{14}\), reports that 2nd mole may react separately after the 1st mole reacted or both moles reacts in the same step. The route of synthesis of ligand and the Cd(II) complex is depicted respectively in Scheme 1 and Scheme 2.

**Table 1:** The M. F., F. wt., colour and the percentage yield of Chiral Schiff bases, L and its Complex, CdL

| Code | M. F. | F. wt. (g/mol.) | Colour      | %Yield |
|------|-------|----------------|-------------|--------|
| L    | \(C_{20}H_{18}O_2N_4\) | 825.96         | Brown       | 80.3   |
| CdL  | \(C_{20}H_{18}O_2N_4Cl\) | 971.82         | Light brown to yellow | 89.3   |

Synthesized chiral Schiff base and its Cd(II) complex were analysed by colour, TLC, m.p., elemental (C, H, N, Cl and Cd) and spectral data viz. UV-Viz, FTIR and \(^1\)H NMR data (400 MHz, DMSO) results. The results are as displayed below.
The elemental analysis data of the ligand and complex are satisfied in agreement for proposed molecular formulas. UV-Vis spectra for the Ligand, L and its Complex were recorded using ethanol as solvent. The obtained values of elemental analysis for the schiff base ligand and its metal chelates are consistent with the calculated values. The complexation of chiral Schiiff base with cadmium (III) ion showed UV-Vis differences in $\pi \rightarrow \pi^*$ transition was shifted from 258 nm to 265 nm, and for the $n \rightarrow \pi^*$ transition was changed from 343 nm to 347 nm. The UV-Vis spectra for ligand, L before and after the complex, CdL formation is depicted in Figure 1.

The ligand exhibited a band around 2930 cm$^{-1}$ weak band due to intramolecular H-bond between H of –OH group and N of C=N group, which is in agreement with earlier reports. The stretching vibrations of phenolic OH and phenolic C=O in the free ligand appear at 2930 and 1210 cm$^{-1}$ respectively and the stretching bands of phenolic C=O have been found at higher values (1245 cm$^{-1}$) in Cd chelate confirming the coordination through phenolic oxygen (M=O). In the FTIR spectrum of ligand, the absorption frequency at 1625 cm$^{-1}$ appeared because of Schiff base which on chelation, it is shifted to lower wave numbers upto 15 cm$^{-1}$ in metal chelates indicating the participation of azomethine nitrogen in coordination with central metal ion (M-N) which is due to the reduction of double bond character in C=N bond. The complexation of Schiff base with cadmium ion understood with significant differences in FTIR frequency for $>\text{C=O}$ and $>\text{C=N}$ groups.

The notice of absence of phenolic
absorption band in the FTIR spectrum of ligand after the complex formation with cadmium ion was an evidenced for the complexation with the chiral Schiff base. The appearance of new band in FTIR spectrum as 550 cm\(^{-1}\) indicates complexation. The frequency of C=\(\text{N}\) shifted to lower cm\(^{-1}\) and disappearance of band due to –OH indicates complexation and deprotonation respectively. These results are in concurrence to earlier findings\(^{14}\).

The coordination of N of the Schiff base nitrogen is indicated by appearance of a newer metal–ligand weak frequency absorption at 490 cm\(^{-1}\) because of \(\nu_\text{Cd-N}\)\(^{29-30}\). The extra agreement of formation of complex has been provided due to band at 550 cm\(^{-1}\) with weak-intensity, which is attributed to \(\nu_\text{Cd-O}\) of phenolic part of Schiff base in the complex\(^{30}\). The FTIR spectral characteristics frequencies are in concurrence with that reported for the similar compounds as reported\(^{29-30}\).

The \(^1\)H NMR (in DMSO) of Chiral Schiff bases, L and its complex, CdL are depicted in Figure 3A and Figure 3B respectively.

\[ \text{L (C}_{29}\text{H}_{19}\text{O}_{3}N_{2}I_{2}) \]: Brown powder, (995 mg, 80.3 \%), m.p. = 123-126\(^\circ\)C; R\(_f\) = 0.91 (hexane:ethyl acetate, 80:20); Anal. Calc. for [C\(_{29}\)H\(_{19}\)O\(_{3}\)N\(_{2}\)I\(_{2}\)]; FW 825.96; C 29.08; H 2.2; N 3.39\%; Found: C 29.10; H 2.19; N 3.42 \%; FTIR (in cm\(^{-1}\)) : 2915 (w), \(\nu_{\text{CH}}\): 1625, \(\nu_{\text{CD}}\): 1440, \(\nu_{\text{C=O}}\): 1400, \(\nu_{\text{C=N}}\): 1270, \(\nu_{\text{C=O}}\): 620, \(\nu_{\text{C=O}}\): 550 \(\nu_{\text{C=O}}\) and 495 \(\nu_{\text{C=O}}\) \(^1\)H NMR data (400 MHz, DMSO): (\(\delta\) in ppm, assigned to, proton number) \(\delta = 1.25-1.65\) (m, 8H (4 x CH\(_2\))) 2 x C-CH\(_2\)-CH\(_2\)-CH-N (proton \(\gamma\) to CH=N) and 2 x C-CH\(_2\)-CH\(_2\)-CH-N (proton \(\beta\) to CH=N) in cyclohexane ring; \(\delta = 2.55\) (m, 2H) -CH\(_2\)-CH-N- (proton \(\beta\) to CH=N) in cyclohexane ring; \(\delta = 2.60\) (m, 2H) -CH\(_2\)-CH-N- (proton \(\beta\) to CH=N) in cyclohexane ring; \(\delta = 8.33\) (s, 2H x CH=N protons); \(\delta = 8.08-8.31\) (d, 4 Ar-H protons); \(\delta = 9.88\) (bs, 2H phenolic H on different rings).

\[ \text{Fig. 3B. } ^1\text{H NMR of the Chiral Schiff base complex, CdL (after complexation with Cd)} \]

\(\text{LCd(C}_{29}\text{H}_{19}\text{O}_{3}N_{2}I_{2}\text{CdCl})\): Light brown yellow powder, (434 mg, 89.3 \%), m.p. = 236-239 (decompose)\(^a\)C; R\(_f\) = complex at baseline (hexane: ethyl acetate, 80:20); Anal. Calc. for [C\(_{29}\)H\(_{19}\)O\(_{3}\)N\(_{2}\)I\(_{2}\)CdCl]; FW 971.82; C 24.72; H 1.66; N 2.88; Cl 3.65; Cd 11.57 \%; Found: C 24.71; H 1.65; N 2.87; Cl 3.66; Cd 11.55 \%. FTIR (in cm\(^{-1}\)) : 1615, assigned to \(\nu_{\text{Cd-N}}\); 1440, \(\nu_{\text{Cd-Cd}}\); 1270, \(\nu_{\text{C-C}}\); 620, \(\nu_{\text{C=O}}\) and 495 \(\nu_{\text{C=O}}\) \(^1\)H NMR data (400 MHz, DMSO): (\(\delta\) in ppm, assigned to, proton number) \(\delta = 1.25-1.65\) (m, 8H (4 x CH\(_2\))) 2 x C-CH\(_2\)-CH\(_2\)-CH-N (proton \(\gamma\) to CH=N) and 2 x C-CH\(_2\)-CH\(_2\)-CH-N (proton \(\beta\) to CH=N) in cyclohexane ring; \(\delta = 2.55\) (m, 2H) -CH\(_2\)-CH-N- (proton \(\beta\) to CH=N) in cyclohexane ring; \(\delta = 8.01\) (s, 2H CH=N protons); \(\delta = 7.72-8.00\) (d, 4 Ar-H protons); Ar-H).

The ligand under study shows a singlet at 9.88 \(\delta\)value indicate the phenolic proton and a singlet at 8.33 shows >CH=N- proton, \(\delta\) multiplet signal at 8.08-8.31 \(\delta\) is assigned to aromatic \(\delta\) proton. The methylene protons are absorbed as bunch of peaks in the range 1.35-1.75 \(\delta\) values (proton \(\beta\)- and \(\gamma\)- to CH=N) of cyclohexane ring, the 2.60 \(\delta\) multiplet for (proton \(\alpha\)- to CH=N) cyclohexane ring.

The Cd(II) complex shows a weak singlet at 8.01 \(\delta\) shows >CH=N- protons indicating involvement of N of azomethine in complexation with cadmium, multiplet signal at 7.72-8.00 \(\delta\) is assigned to aromatic protons. The methylene protons are absorbed as bunch of peaks in the range 1.25-1.65 \(\delta\) values of cyclohexane ring, the 2.55 \(\delta\) multiplet for cyclohexane ring. A very weak peak at 9.88 \(\delta\) value indicated...
The absence of phenolic protons hence confirms the complexation with O. We have reported\textsuperscript{14} the similar findings in earlier work and are also in concurrence with the results of Wen, Z. et al.,\textsuperscript{31}

The Cd (II) complex shows values of the molar conductivity ($
\Lambda_m$) at the concentrations of $10^{-3}$ M in dichloromethane is 19.1 S cm$^{-1}$ mol$^{-1}$, which indicates that the present Cd(II) complex is of slight polar nature because of electrolytic behavior of halogen (chlorine). According to the molar conductance values, the Schiff base complex have 1:1 metal:ligand ratio, as 1:1; consequently one chloride ion is outside the coordination sphere in the complex. Hence the chloride must be coordinated to the metal center\textsuperscript{14,32}.

The results of oxidation catalysis of benzyl alcohol by using H$_2$O$_2$ as oxidant and the cadmium complex of chiral Schiff base as catalyst in acetonitrile to form benzaldehyde are summarized in Table 2.

**Table 2: Screening conditions for oxidation* of primary alcohols**

| Entry | Alcohol | Product | Time (h) | Conversion (%) | Ref. |
|-------|---------|---------|----------|---------------|------|
| 1     | C$_6$H$_5$CH$_2$OH | C$_6$H$_5$CHO | 08       | 79.00         | 2    |
| 2     | C$_6$H$_5$CH$_2$OH | C$_6$H$_5$CHO | 04       | 75.02$^a$     | 14   |
| 3     | C$_6$H$_5$CH$_2$OH | C$_6$H$_5$CHO | 03       | 88.00$^b$     | 17   |
| 4     | C$_6$H$_5$CH$_2$OH | C$_6$H$_5$CHO | 22       | 71.00         | 18   |
| 5     | C$_6$H$_5$CH$_2$OH | C$_6$H$_5$CHO | 04       | 51.33$^c$     | This work$^{30}$ |

*Reaction was performed under open-air conditions fitted with a condenser; $^a$Cr (III)L complex, isolation purpose multiple size batch was conducted, alde % determined by HPLC and comparing with authentic samples; $^b$achiral [RuCl$_2$(AsPh$_3$)$_2$]$_2$ (OHA-ampy)]/NMO and $^c$chiral catalyst Cd (II)L complex used here.

The product of oxidation catalysis obtained was dried over sodium sulphate and was concentrated in microflask under vacuum (carefully) and resulting liquid product was purified by eluting the reaction mass using hexane:ethyl acetate on silica gel column to obtain the benzaldehyde as a colourless liquid showed FTIR (in cm$^{-1}$): 1605 (s), assigned to $\nu$C=O$^1$; 1525, $\nu$C=C$_{(ar)}$; 1222 $\nu$C=O$^2$; 2710 $\nu$C-H$_{(ar)}$; 2860, $\nu$CHO$^3$ $^1$H NMR data (400 MHz, DMSO): ($\delta$ in ppm, assigned to, proton number) $\delta = 10.3$ (s, 1H) -CHO; $\delta 6.52$-7.75 (m, 5 Ar-H). Similar report$^{14}$ was made with a chiral Schiff base from 3,5-Dibromosalicylaldehyde for the oxidative transformation of benzyl alcohol to benzaldehyde.

The Table 2 indicated a comparison of the present oxidation catalysis with the results available in literature which indicated that, time wise it is less time than required by other Heterogeneous catalysis and it matches with the conversion of the benzaldehyde formation\textsuperscript{2,14,17-18}. In the present work, % conversion of oxidation of benzyl alcohol to benzaldehyde by H$_2$O$_2$ oxidation in presence of catalytic CdL, using solvent acetonitrile is 51.33% which is slightly less as compared to that with the Cr(II) catalyst from chiral Schiff base with dibromo substituent which is small in size, and hence it is more efficient catalyst. As because the size of iodine makes the ligand more bulkier than when it is bromo substituent the rate of oxidation is less than the Cr(III) L catalyst recently reported\textsuperscript{14}.

Literature showed that the many ruthenium catalytic reactions were reported with a oxidants of wide variety viz., hydrogen peroxide\textsuperscript{33}, molecular oxygen\textsuperscript{15}, aqueous H$_2$O$_2$\textsuperscript{16,18}, NaIO$_4$\textsuperscript{33}, chloramine-T\textsuperscript{34}, iodosylbenzene\textsuperscript{35} and benzoquinone\textsuperscript{36}, where as N-methylmorpholine-N-oxide (NMO) as oxidant is scarcely available in literature. Oxidation of alcohols by complex, (g6-p-cyemene)ruthenium (II) 2-(naphthylazo) phenolate using N,O-donors by use of NMO\textsuperscript{37}.

**Possible mechanism for the Benzyl alcohol to Benzaldehyde oxidation reaction**

A possible route of oxidation mechanism based on the above discussed data can be proposed as below,

\begin{center}
\includegraphics[width=\textwidth]{scheme3.png}
\end{center}

Scheme 3. The mechanistic route proposed for oxidative transformation of the benzyl alcohol in presence of aq. H$_2$O$_2$ (aqueous) solution catalyzed by chiral Schiff base Cd (II) complex.
It is marked, presence of coordinated chiral ligand in Cd(II) complex (Scheme 3) may form Cd(II) intermediate-I within the reaction because of the reaction of oxidant with the complex catalyst as reported \(^{38-42}\). Specifically, the detection of intermediate Cd(II) species from reaction intermediate-II has occurred by monitoring the successive spectral observation of complex in the oxidation process (above Eq.-1). Strength of peculiar absorption band maximum is changed little during transformation, which is dependent on oxidative addition. It is because of formation of positive species in the oxidative transformation to oxide intermediate-II via intermediate-I in coordination of oxidant to the central metal ion \(^{43,44}\). Several reports have been found in literature that the oxo-cadmium (II) complex is probably an intermediate-III in the oxidative transformation process of organic reaction catalyzed by Cd (II) complex \(^{39-42}\).

Thus, Cd (II)--O is formed through the oxygen transfer from oxidant molecule to give potential intermediate-III. The modification in the colour of the reaction at the start of the oxidative catalytic reaction of the solution from light yellow to slight brown in is in support for oxygen atom transfer from \(\text{H}_2\text{O}_2\) (oxident) to the central metal ion (Cd(II)) to oxo-cadmium (II) intermediate-III. When an aqueous \(\text{H}_2\text{O}_2\) was mixed to solution, colour turned to brown and marked the change is because of the formation of Cd(II)--O moiety, which explains reaction of peroxydo function of Cd(II)-oxidant and the presence of strong transition (charge transfer) in intermediate-III. The active oxo-cadmium (II) intermediate could oxidatively transfer benzyl alcohol (substrate) through the complex with benzyl alcohol to potential oxo-cadmium (II) intermediate IV to regenerate oxidized catalyst complex (intermediate-I) and yield chemo selective product (i.e., benzaldehyde).

Conceptually, above sequence can be written in the form of Scheme 4 as below...

Typical mechanism of catalytic reaction for Cd\(^{2+}\)-moiety is discussed here. But, on the basis of colour change through catalytic system of reaction and depending on mechanisms assigned for reactions of similar nature\(^{39-42,45}\), one could predict that important process is oxo-cadmium (II) moiety presence in catalytic transformation through alcohols oxidation, as depicted in Scheme 3, which is scarely discussed.

**CONCLUSION**

The chiral Schiff base was synthesized from chiral amine and its mononuclear Cd(II) complex was synthesized. The prepared Cd(II) complex was characterized by different analytical and spectral methods. Cd(II)-complex was tested as catalysts for one pot conversion of benzyl alcohol to aldehyde by the oxidation by use of \(\text{H}_2\text{O}_2\) in mild conditions. The possible oxidized products of benzyl alcohol were benzaldehyde and or an impurity, benzoic acid. It is heterogeneous system of catalysis. In the visible region, complex exhibited a peculiar absorption band responsible for the metal-to-ligand charge transfer (MLCT). The prepared CdL is a promising catalyst in oxidation of alcohol using hydrogen peroxide as terminal oxidant. The main benefit of this catalytic system are the easy synthesis of ligand and its Cd complex and also modification of ligand is easy process. In present work, % conversion (oxidation) benzyl alcohol to benzaldehyde by \(\text{H}_2\text{O}_2\) oxidation in presence of catalytic CdL, using solvent acetonitrile is less than the Cr(III)L catalyst with comparatively less bulkier ligand (substituent) recently reported\(^1\). This work leads to the development of new class of catalyst system for oxidative transformation of benzyl alcohol to benzaldehyde and related work is now underway.

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**Conflicts of Interest**

The authors declare no conflict of interest.
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