COORDINATION COMPOUNDS OF Cu (II) AND Ni (II) WITH SCHIFF BASES DERIVED FROM FORMYLMENTHONE AND AROMATIC AMINES

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Abstract: The Schiff bases obtained from formylmenthone and o-, m-, p-toluidine behave as bidentate ligands with O and N donor atoms in Ni(II) and Cu(II) complexes by the type NiL₂ and CuL₂(H₂O)₂. The donor atoms and the possible geometry for the complexes were assigned by means of chemical, thermodifferential analyses and electronic, EPR and IR spectra.

Keywords: formylmenthone, complexes, synthesis, thermodifferential analyses, EPR, IR

1. Introduction

Terpenes represent the largest class of secondary plant metabolites, being used for antimicrobial activity. Terpenes act on gram-positive bacteria, less than gram-negative bacteria. [1] Monoterpenoids with nitrogen atoms, generally oxymes, or amines have various biological properties, such as anticancer [2], antimicrobial [3], antiviral [4].

A series of Schiff bases of menthone with aromatic amines has been synthesized and were evaluated for anticonvulsant activity. [5, 6]

Our previous paper [7] has reported the preparation and characterization of the coordination compounds of some "3d" metals with Schiff bases derived from 3-formylmenthone and aromatic amines. The position of the methyl substituent on the benzene ring has determined the type of the complex and monodentate or bidentate coordination of the ligands. It seemed desirable to investigate the coordination ability of the Schiff bases derived from 2-formylmenthone and o-, m-, p-toluidine on the same "3d" metals, for purposes of comparison.

2. Materials and Methods

2.1. Materials

Cu(CH₃COO)₂·H₂O and Ni(CH₃COO)₂·4H₂O p.a. Merck were used.

The syntheses were carried out using reagents and solvents of the highest quality of analytical reagent grade and were used without further purification. Distilled water was used in all experiments.

2.2. Characterization Techniques
The electronic diffuse reflectance spectra within 300-1100 nm range were obtained with VSU-2P Zeiss-Jena Spectrophotometer, using MgO as a standard. EPR spectra were recorded at room temperature on polycrystalline powders, on ART-5-IFA Spectrograph.

The klystron frequency was 9060 MHz and modulation of the magnetic field 100 KHz. The EPR spectral parameters were calculated against a Mn (II) standard.

Thermodifferential analyses were carried out with a Paulik-Paulik-Erdey Derivatograph Q 1500-D MOM. Conditions of measurements: temperature range up to 1000°C, heating program: 10 degree/min, sensitivity DTA= 1/10, m$_4$=0.0180 g, S=20 and m$_5$ = 0.0372 g, S=50; atmosphere over sample air.

IR spectra (KBr pellets) were measured on a BIO-RAD FTS-135 Spectrometer.

2.3. Synthesis

2.3.1. General synthesis of ligands

0.01 mol formylmenthone was dissolved in 200 mL ethylic alcohol. A solution of toluidine dissolved in 40 mL ethyl alcohol is added in hot drop in the reaction mixture, after which it is maintained at the room temperature for 50 hours.

2.3.1.1. Synthesis of 6-isopropyl-3-methyl-2-{(2'-methylphenylamino) methylen]cyclohexan-1-one (ortho-L) (3a)

Compound 3a was synthesized according to general procedure of the ligands. The crystals were filtered off, washed with water and dried in air; 1.39 g Yield (51%); red-brown crystals; m.p.= 135°C; IR (KBr, cm$^{-1}$): 3200-3320 (m, ν$\text{O-H}$), 1700 (sh, ν$\text{C=O}$), 1550, 1590, 1630 (vs, ν$\text{C=N}$).

2.3.1.2. Synthesis of 6-isopropyl-3-methyl-2-{(3'-methylphenylamino) methylen]cyclohexan-1-one (metha-L) (3b)

Compound 3b was synthesized according to general procedure of the ligands. The crystals were filtered off, washed with water and dried in air; 1.17 g Yield (43%); brick-red crystals; m.p.= 105°C; IR (KBr, cm$^{-1}$): 3310-3450 (m, ν$\text{O-H}$), 1700 (vw, ν$\text{C=O}$), 1560, 1620, 1640 (s, ν$\text{C=N}$).

2.3.1.3. Synthesis of 6-isopropyl-3-methyl-2-{(2'-methylphenylamino) methylen]cyclohexan-1-one (para-L) (3c)

Compound 3c was synthesized according to general procedure of the ligands. The crystals were filtered off, washed with water and dried in air; 1.47 g Yield (54%); red-brown crystals; m.p.= 110°C; IR (KBr, cm$^{-1}$): 3300-3400 (m, ν$\text{O-H}$), 1700 (vw, ν$\text{C=O}$), 1520, 1580, 1630 (s, ν$\text{C=N}$).

2.3.2. General synthesis of the complexes

The complexes were prepared by mixing warm aqueous methanol solution (50%) of metal acetate (1 mmol) and ligands (2 mmols). The resulting precipitates were filtered and washed with aqueous methanol solution (50%) and dried at room temperature. The metal content was obtained gravimetrically.

2.3.2.1. Synthesis of Cu (ortho-L)$_2$.H$_2$O (5a)

Compound 5a was synthesized according to general procedure of the complexes.
After the cooling at the room temperature, the pale green crystals were filtered off, washed with water and dried in air; 0.64 g Yield: (20%). IR (KBr, cm⁻¹): 3423 (s, νOH), 1558 (vs, νC=N), 1615(vs, νC=N), 1111 (w, H₂O coord.). Elemental analysis data calculated for Cu (ortho-L):(H₂O): Cu 9.94%, H₂O 5.63%. Found: Cu 10.37%, H₂O 7.03%.

2.3.2.2. Synthesis of Cu (metha-L):(H₂O): (5b)

Compound 5b was synthesized according to general procedure of the complexes.

After the cooling at the room temperature, the blue crystals were filtered off, washed with water and dried in air; 0.54 g Yield: (17%). IR (KBr, cm⁻¹): 3395 (m, νOH), 1557 (s, νC=N), 1610(vs, νC=N), 1094 (w, H₂O coord.). Elemental analysis data calculated for Cu (metha-L):(H₂O): Cu 9.94%, H₂O 5.63%. Found: Cu 9.64%, H₂O 6.80%.

2.3.2.3. Synthesis of Cu (para-L):(H₂O): (5c)

Compound 5c was synthesized according to general procedure of the complexes.

After the cooling at the room temperature, the gray crystals were filtered off, washed with water and dried in air; 0.76 g Yield: (24%). IR (KBr, cm⁻¹): 3399 (m, νOH), 1514 (vs, νC=N), 1545 (s, νC=N), 1613 (s, νC=N), 1111 (w, H₂O coord.). Elemental analysis data calculated for Cu (metha-L):(H₂O): Cu 9.94%, H₂O 5.63%. Found: Cu 10.09%, H₂O 6.91%.

2.3.2.4. Synthesis of Ni (metha-L):(H₂O): (5d)

Compound 5d was synthesized according to general procedure of the complexes.

After the cooling at the room temperature, the light blue crystals were filtered off, washed with water and dried in air; 0.44 g Yield: (14%). IR (KBr, cm⁻¹): 3398 (m, νOH), 1514 (vs, νC=N), 1543 (s, νC=N), 1510 (s, νC=N), 1635 (s, νC=N). Elemental analysis data calculated for Ni (metha-L):(H₂O): Ni 9.80%, H₂O 5.63%. Found: Ni 10.09%.

2.3.2.4. Synthesis of Ni (para-L):(H₂O): (5e)

Compound 5e was synthesized according to general procedure of the complexes.

After the cooling at the room temperature, the cream crystals were filtered off, washed with water and dried in air; 0.67 g Yield: (21%). IR (KBr, cm⁻¹): 3422 (m, νOH), 1510 (m, νC=N), 1543 (m, νC=N), 1558 (m, νC=N). Elemental analysis data calculated for Ni (metha-L):(H₂O): Ni 9.80%, Found: Ni 10.50%.

3. Results and Discussion

3.1. Synthesis

Five new ligands able to generate complexes have been synthesized (Scheme 1).
Scheme 1. Synthesis of Schiff bases. Reagents and conditions: CH₃OH, warm (i)

The presence of the >C=O and >C=N - groups respect to each other favours keto-enolic tautomerism (Scheme 2). This tautomerism has been attributed to an intramolecular hydrogen bond and might explain their chelating ability. NMR studies have shown that such Schiff bases with the carbonyl and azomethine groups exist in solution as the enolic tautomer (4a-c) and that the tautomer distribution was solvent dependent (Scheme 2) [8, 9].

Scheme 2. Tautomeric forms.

3.2. Elemental Analyses

The elemental analyses are consistent with their formulation as anhydrous, monomeric Schiff bases chelates CuL₂ and NiL₂) or hydrated, monomeric complexes with water molecules either as coordinated or as crystalline water. All the metal chelates are colored, readily soluble in organic solvents (chloroform, acetone, methanol), but sparingly soluble in water.

3.3. Thermodifferential Analysis

The thermodifferential analysis are consistent with their formulation as anhydrous, monomeric Schiff bases chelates. Curves for compounds Cu(para-L)₂.(H₂O)₂ and Ni(para-L)₂ are shown in detail in Figures 1a and 1b.
Figure 1a. Thermodiffferential curve of Cu(para-L):2H₂O
For compound Cu(para-L)$_2$.H$_2$O (Figure 1a) the mass loss observed within 60-215°C range in the TG curve corresponds to the loss of two water molecules per molecule of each copper compound. The TG curve shows that the water molecules are liberated in two steps (weight loss at 170°C, found: 2.98%, calcd., for H$_2$O: 2.88%; weight loss at 215°C, found: 7.03%, calcd., for 2H$_2$O: 5.76%). Hence, the two water molecules are present as coordinated water, in Cu(II) complexes.

The results indicate that compound Ni(para-L)$_2$.2H$_2$O (Figure 1b) is stable up to 150°C, but rapidly loses ligands at 565°C (weight loss at 565°C, found: 88.86%, calcd., for 2L: 90.14%).

3.4. Electronic diffuse reflectance

The electronic diffuse reflectance spectra of the complexes are given in Figures 2 and 3.

The spectra of the Ni (II) complexes (5d and 5e, Figures 2) are like and are consistent with tetrahedral Ni (II) complexes [10].
Figure 2. Electronic diffuse reflectance spectra of Ni(meta-L)2 blue and Ni(para-L)2 red

Figure 3. Electronic diffuse reflectance spectra of Cu(ortho-L)2 green; Cu(meta-L)2 blue and Cu(para-L)2 red
The spectra present the band ν₃ (660 nm) assignable to the ³T₁(P)→³T₁ transition weak band (760 nm, compound Ni (meta-L)₂) or the shoulder (760 nm, compound Ni (para-L)₂) are assigned as spin forbidden transitions to components of 1D levels [10]. The band ν₁ assignable ³T₂→³T₁ transition (near 500 nm) is covered by a broad and intense band due to the ligands (500 nm). It is present in the spectra of the copper complexes, also. The spectra of the copper complexes (Cu(ortho-L)₂·(H₂O)₂, Cu(meta-L)₂·(H₂O)₂, Cu(para-L)₂·(H₂O)₂) in Figure 3 are similar and present a band at 800 nm that could be assigned to a d-d transition associated with a distorted octahedron [11-13].

3.5. EPR spectra

EPR spectra of the three copper complexes recorded at room temperature on polycrystalline samples present a similar intense EPR signal characteristic monomeric species of Cu (II) ion with a third order anisotropy for the factor g resulting from distortion of octahedral geometry (Figure 4).

This anisotropy is compatible with Cu (II) ion in a compressed rhombic-octahedral geometry [12, 13] with R>1 and supports the electronic spectra. In a three g value spectrum with g₁ < g₂ < g₃, the value of R = (g² - g₁)/(g₃ - g₂) may be significant: if R > 1, a predominant dz² ground state is present and would be consistent with compressed axial or rhombic symmetry with slight misalignment of the axes. If R < 1, a predominant dx²-y² state is present [12, 13].

Figure 4. EPR spectra and SPR spectral parameters of Cu(ortho-L)₂ green; Cu(meta-L)₂ blue and Cu(para-L)₂ red
Complex  |  $g_1$  |  $g_2$  |  $g_3$
--- | --- | --- | ---
$	ext{Cu(ortho-L)}_2$ | 1.9804 | 2.0439 | 2.4586
$	ext{Cu(meta-L)}_2$ | 1.9804 | 2.0605 | 2.4586
$	ext{Cu(para-L)}_2$ | 1.9959 | 2.0605 | 2.4466

3.6. IR spectra

The keto-enolic tautomeration is supported by presence of the bands due to $\nu$O-H and $\nu$C=N. The band due to $\nu$C=O (1700 cm$^{-1}$) occurs as a very weak band or a shoulder. The band due to $\nu$O-H occurs as a broad band (3200-3600 cm$^{-1}$ range) with two unresolved peaks. The band due to $\nu$C=N occurs as a broad band (1500-1600 cm$^{-1}$ range) with three unresolved peaks.

A comparison of the position of the bands in spectra of the complexes with their position in the IR spectra of free ligands shows changes of the bands due to $\nu$O-H and $\nu$C=N. Upon coordination, the stretching frequencies, $\nu$C=N, are shifted to lower values and stretching frequencies $\nu$O-H are shifted to higher values. These changes are generally noticed upon coordination of the Schiff bases containing an N and O donor atoms, by the both donor atoms [14]. The new band near 1111 cm$^{-1}$ occurring in the IR spectra of the three copper complexes only could be assigned to the coordinated water molecules in agreement with Fujita [15] and is be consistent with the results of the thermodifferential analysis.

4. Results

On the basis of elemental, differential analyses and spectral measurements, we conclude that Cu (II) ion is hexacoordinated in a compressed rhombic geometry, while Ni (II) ion is tetracoordinated in a tetrahedral geometry.

The ligands acted bidentately with both O and N donor atoms by deprotonation of the OH group making evident the participation of the ligands in the enolic tautomeric form.

Cu (II) coordinates by atom donors, N and O in a plane and by shorter bonds to axial water molecules in a compressed rhombic geometry (Figure 5).

Figure 5. Structural formula proposed for the Cu (II) complexes
The tetrahedral environment of the Ni (II) ion suggested by electronic spectra is obtained by both donor atoms, N and O.

We may conclude that the metallic ion determines the type of the new complexes.

The arguments for the structure of the new chelates were fully supported by the spectral data (electronic, EPR and IR spectra).

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