New nickel (II) complexes with benzilbis (semicarbazone) and dithiocarbamate ligands

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Abstract

The preparation of new nickel (II) complexes with mixed ligands including benzilbis- (semicarbazone) - SCH₂ and one of dithiocarbamate ligands (ammonium pyrrolidenedithio- carbamate - Pyrdtc or ammonium indolinedithiocarbamate - Indtc) have been carried out in both neutral and basic medium. The resulted complexes have been characterized using elemental analysis by atomic absorption spectroscopy and by precipitation method, molar conductance measurements, refraction measurements, magnetic susceptibilities, infrared and electronic spectra. The study suggested that the above ligands formed neutral complexes having general formula [Ni(SCH₂)(L)X] or [Ni₂(SCH₂)₂(L)₂X_n] in neutral medium and [Ni(SCH)(L)] or [Ni₂(SCH)₂(L)₂] in basic medium (where SCH=deprotonated (-1H) from SCH₂ ligand; L= the ligand Pyrdtc or Indtc; X= Cl⁻ or NO₃⁻ or CH₃COO⁻ or SO₄⁻², n= 1 or 2). Hexacoordinated mononuclear (monomer) complexes or dimeric complexes have been investigated by this study, having octahedral geometries.

Keywords: Mixed ligands, Benzilbis(semicarbazone), Dithiocarbamate, Nickel(II) complexes. Available online at http://www.vetmedmosul.org/ijvs

الخلاصة

حضرت معقدات جديدة للنيكل (II) مع مزيج من ليكندي بنزل بس (سميكاربازون) – SCH₂ وأحد ليكندات الثايوكارباميت (امونيوم بايرولدين ثنائي ثايوكارباميت – Pyrdtc او امونيوم اندولين ثنائي ثايوكارباميت – Pyrds Indtc). شخصت المعقدات الناتجة باستخدام التحليل العنصري بجهاز طيف الامتصاص الذري والطريقة الترسيبية وقياس التوصيلية الكهربائية ومعامل الانكسار والمغناطيسية وأطياف الاشعة تحت الحمراء والالكترونية. واستنتج ان الليكندات تعطي معقدات متعادلة ذات صيغ عامة [Ni(SCH₂)(L)] أو منه الاشعة تحت الحمراء والالكترونية. واستنتج ان الليكندات تعطي معقدات متعادلة ذات صيغ عامة (Ni(SCH₂)(L)] أو منه بروتون، LSCH]في الوسط المتعادل و (Ni(SCH)(L)] أو (SCH₂(SCH)₂(L)] في الوسط القاعدي (حيث SCH = الليكند SCH منه بروتون، LSCH اليكند عائمات الدراسة ان المعقدات المعقدات متعادلة ذات صيغ عامة SCH₂ المعقدات الاليكند تحلي

Introduction

Nickel complexes occurred in several nickel-containing enzymes have been proposed to be involved in catalytic reaction (1).

A good deal of work has been reported on the preparation and structural investigation of semicarbazone

and their complexes (2,3). This is due partially to their capability of acting as multidentate, NO, NNO and ONNO donors with the formation of either mono- or bi- or poly-nuclear complexes (4,5). In addition to their interesting ligational properties, semi carbazones and their complexes have important biological applications (6).

Dithiocarbamate formed an impor- tant class of biologically active ligands (7). An extremely large number of dithiocar bamate complexes with transition and non transition metal ions have been known (8,9).

There has been growing interest in the formation of mixed ligands chelates involveing ligands containing different functional groups and transition metals of different oxidation states which can formed chelates with ligands containing different donation sites (10,11). On the other hand, coordination compounds with mixed ligands are of considerable impor tance in the field of metalloenzymes and other biological activities (12). Hence a large body of the coordination chemistry of mixed ligands with transition and non-transition metal ions have been reported recently (13,14). Due to the importance of semicarbazone ligands, we took a modest part in the chemistry of mixed ligands containing semicarbazones and their complexes, and some articles have been published so for on their coordination chemistry with transition and non-transition metal ions (15.16)

In the present work, nickel (II) complexes with mixed ligands {benzilbis- (semicarbazone) - SCH_2 and ammonium pyrrolidenedithiocarbamate - Pyrdtc or ammonium indolenedithiocarbamate - Indtc} have been prepared and characterized physicochemically. The structures of the ligands are shown in Figure (1).

Materials and methods

Starting Materials

All the chemicals used were supplied from Fluka, BDH and Aldrich.

Table 1: Amounts of ligands and medium of reaction.

Preparative methods

Semicarbazone ligands have been prepared according to literature method (17). Dithiocarbamate ligands have been prepared according to literature method (18).

A general procedure has been adopted for the preparation of the complexes in neutral and basic medium.

In neutral medium

A solution of 0.5 gm $(2x10^{-3} \text{ mole})$ of NiCl₂.6H₂O in 5 ml water has been added to the solution of gm SCH₂ $(2X10^{-3})$ and $2X10^{-3}$ mole of one of the dithiocarba- mate ligands in a small amount of ethanol and water with heating until a clear solution has been resulted (the amounts in grams were listed in Table 1). The mixtures have been refluxed for three hours followed by evaporation to half their volumes then cooled. The products were separated by filteration, washed with petroleum ether and dried. The complexes of the other salts Ni(CH₃COO)₂.4H₂O or (Ni(NO₃)₂.6H₂O or NiSO₄.7H₂O) have been prepared by applying same procedure.



Figure (1): Structures of the ligands.

No.	Complexes	Wt of SCH ₂	Wt of Pydtc or Indtc	Wt of salt	Medium
1	[Ni(SCH ₂)(Pyrdtc)Cl]	0.682	0.345	0.5	Neutral
2	$[Ni_2(SCH)_2(Pyrdtc)_2]$	0.682	0.345	0.5	Basic
3	[Ni(SCH ₂)(Indtc)Cl]	0.682	0.446	0.5	Neutral
4	[Ni(SCH)(Indtc)]	0.682	0.446	0.5	Basic
5	[Ni(SCH ₂)(Pyrdtc)NO ₃]	0.558	0.282	0.5	Neutral
6	$[Ni_2(SCH)_2(Pyrdtc)_2]$	0.558	0.282	0.5	Basic
7	[Ni(SCH ₂)(Indtc)NO ₃]	0.558	0.365	0.5	Neutral
8	$[Ni_2(SCH)_2(Indtc)_2]$	0.558	0.365	0.5	Basic
9	[Ni ₂ (SCH ₂) ₂ (Pyrdtc) ₂ SO ₄]	0.577	0.292	0.5	Neutral
10	[Ni(SCH)(Pyrdtc)]	0.577	0.292	0.5	Basic
11	[Ni ₂ (SCH ₂) ₂ (Indtc) ₂ SO ₄]	0.577	0.378	0.5	Neutral
12	[Ni(SCH)(Indtc)]	0.577	0.378	0.5	Basic
13	$[Ni_2(SCH_2)_2(Pyrdtc)_2(Ac)_2]$	0.652	0.330	0.5	Neutral
14	[Ni(SCH)(Pyrdtc)]	0.652	0.330	0.5	Basic
15	[Ni(SCH ₂)(Indtc)Ac]	0.652	0.426	0.5	Neutral
16	[Ni(SCH)(Indtc)]	0.652	0.426	0.5	Basic

In basic medium

The complexes have been prepared by applying the same amounts used for neutral medium, and after mixing the metal salts with the ligands and heating, sodium hydroxide solution (1M) has been added until pH of the solutions have been adjusted to about 9-10. The mixtures have been allowed to stand, then the products were filtered off and washed with petroleum ether and dried.

Analytical and physical measurements

Nickel contents have been deter-mined by applying precipitation method after the decomposition of the complexes with concentrated nitric acid and also determined by atomic absorption spectro-scopy (19). Refraction measurements have been carried out with Atago Illumina-tion, Atago Co-LTD, Japan using 10⁻³ M dimethylformamide solution. Conductivity measurements have been carried out with an electrolytic conductivity measuring set LF-42 (apparatus A) and Multiline f/SET-2WTW Wissenschaf Tecchnische Werkst- attem 82362 Weiheim (apparatus B) using 10^{-3} M dimethylformamide solution at 25 °C. The infrared spectra has been recorded on FT-IR Bruker type Tensor 27 in the range 400-4000 and 200-4000 cm⁻¹ using KBr pellets. Electronic spectra has been recorded on Shimadzu UV-1650 PC UV-Vissible Spectrophotometer for 10⁻³ M solutions of the ligands and their complexes in dimethylformamide at 25 °C, using a

Table 2: Analytical and physical properties of the complexes.

1cm cell. Magnetic suscepibility of the complexes have been measured by Bruker-BM6.

Results and discussion

The reaction of nickel(II) salts, benzilbis (semicarbazone) and dithiocarba- mate ligands in 1:1:1 and 2:2:2 molar ratio in both neutral and basic medium may be represented by the following equations:

$$\begin{split} \text{NiX}_{2}.\text{mH}_{2}\text{O} + \text{SCH}_{2} + \text{LNH}_{4} &\to [\text{Ni}(\text{SCH}_{2}) \ (\text{L})\text{X}] + \text{NH}_{4}\text{X} \\ + \text{mH}_{2}\text{O} \\ 2\text{NiX}_{2}.\text{mH}_{2}\text{O} + 2\text{SCH}_{2} + 2\text{LNH}_{4} &\to [\text{Ni}_{2}(\text{SCH}_{2})_{2}(\text{L})_{2}\text{X}_{2}] + \\ 2\text{NH}_{4}\text{X} + 2\text{mH}_{2}\text{O} \\ 2\text{NiSO}_{4}.7\text{H}_{2}\text{O} + 2\text{SCH}_{2} + 2\text{LNH}_{4} &\to [\text{Ni}_{2}(\text{SCH}_{2})_{2}(\text{L})_{2}\text{SO}_{4}] + \\ (\text{NH}_{4})_{2}\text{SO}_{4} + 14\text{H}_{2}\text{O} \\ \text{NiX}_{2}.\text{mH}_{2}\text{O} + \text{SCH}_{2} + \text{LNH}_{4} + \text{NaOH} \to [\text{Ni}(\text{SCH})(\text{L})] + \\ \text{NH}_{4}\text{X} + \text{NaX} + (\text{m}+1)\text{H}_{2}\text{O} \\ 2\text{NiX}_{2}.\text{mH}_{2}\text{O} + 2\text{SCH}_{2} + 2\text{LNH}_{4} + 2\text{NaOH} \to \\ [\text{Ni}_{2}(\text{SCH})_{2}(\text{L})_{2}] + 2\text{NH}_{4}\text{X} + 2\text{NaX} + (2\text{m}+2)\text{H}_{2}\text{O} \\ \\ \text{NiSO}_{4}.7\text{H}_{2}\text{O} + \text{SCH}_{2} + \text{LNH}_{4} + \text{NaOH} \to [\text{Ni}(\text{SCH})(\text{L})] + \\ \text{NaNH}_{4}\text{SO}_{4} + 8\text{H}_{2}\text{O} \\ (\text{where } \text{X} = \text{CI}^{-} \text{ or } \text{NO}_{3}^{-} \text{ or } \text{CH}_{3}\text{COO}^{-}; \text{LNH}_{4} = \text{Pyrdtc or } \\ \text{Indtc ligands; SCH} = \text{deprotonated } \text{SCH}_{2} \text{ ligand; m} = 4 \text{ or } 6). \end{split}$$

Color	M.P or d °c	M		**M	Ni% Calc. (Obs.)	Theo M Wt	Ref $x 10^{-3}$.
101		app.A	app.B	iviell.	1176 Cule. (0000)		ner are .
hite	212				_	324.38	-3.010
le yellow	150*	_	_	_	—	164.17	-3.010
le yellow	216	_	_	_	—	212.21	+0.005
le green	190*	04.36	04.36	2.600	10.395 (11.640)	0564.76	-4.015
ellowish green	280*	07.99	07.99	1.190	11.112 (11.050)	1056.52	-4.015
ellowish orange	165*	14.54	13.81	2.411	09.581 (09.571)	0612.80	-4.015
ellowish green	210*	22.53	21.80	2.343	10.189 (10.180)	0576.23	-4.015
le green	170*	19.62	18.90	2.392	08.788 (08.781)	0591.26	-5.020
le green	280*	14.54	14.54	1.031	11.112 (11.710)	1056.52	-4.015
ellowish green	235*	10.90	11.63	2.880	09.183 (09.140)	0639.30	-2.005
ellowish orange	210*	27.62	29.07	1.855	10.188 (09.710)	1152.46	-5.020
ige	205*	18.17	08.72	2.707	09.387 (09.389)	1154.47	-4.015
ige	220*	05.81	02.91	2.611	11.112 (09.910)	0528.26	-3.010
ige	260*	12.36	13.81	1.982	09.581 (09.571)	1250.55	-4.015
ige	225*	05.09	05.81	2.542	10.189 (10.180)	0576.23	-3.010
le green	200*	13.08	06.54	1.812	09.980 (10.440)	1176.52	-3.010
ellowish green	220*	10.90	03.27	2.049	11.112 (09.950)	0528.26	+4.015
ige	250*	03.63	04.51	2.982	09.226 (08.420)	0636.30	-4.015
ige	205*	09.47	09.47	2.669	10.189 (10.175)	0576.23	-4.015
	hite le yellow le yellow le green ellowish green ellowish green le green le green ellowish green ellowish green ge ge ge ge le green ellowish green ge ge ge ge ge ge ge	olorM.P or d $^{\circ}$ chite212le yellow150*le yellow216le green190*ellowish green280*ellowish orange165*ellowish green210*le green280*ellowish green235*ellowish orange210*ge220*ge260*ge220*ge20*ge250*ge205*	IdeaM.P or d $^{\circ}$ Capp.Ahite212—le yellow150*—le yellow216—le green190*04.36Ellowish green280*07.99Ellowish green216*14.54ellowish green210*22.53le green170*19.62le green280*14.54ellowish green235*10.90Ellowish green235*10.90Ellowish green220*05.81ge220*05.81ge225*05.09le green200*13.08Ellowish green220*03.63ge250*03.63ge250*09.47	IndextM.P or d ${}^{\circ}$ CIndextIndextapp.Aapp.Bhite212le yellow150*le yellow216le green190*04.3604.3604.3610wish green280*280*07.9907.9907.9910wish green210*22.5321.80le green10*19.6218.90le green280*14.5414.54ellowish green235*10.9011.63ellowish orange210*27.6229.07ge205*200*13.0806.54ellowish green200*13.0806.54ellowish green220*05.8102.91ge20*10.9003.27ge250*03.6304.51ge250*09.4709.47	IndextM.P or d 0 CMapp.Aapp.B**Meff.hite212————le yellow150*———le yellow216———le green190*04.3604.362.600ellowish green280*07.9907.991.190ellowish orange165*14.5413.812.411ellowish green210*22.5321.802.343le green170*19.6218.902.392le green280*14.5414.541.031ellowish green235*10.9011.632.880ellowish orange210*27.6229.071.855ge205*18.1708.722.707ge220*05.8102.912.611ge260*12.3613.811.982ge225*05.0905.812.542le green200*13.0806.541.812ellowish green220*03.6304.512.982ge250*03.6304.512.982ge205*09.4709.472.669	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c c c c c c c c c c c c c c c c c c c $

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 $\Lambda_{M:}$ Molar conductivities in Ω^{-1} cm² mol⁻¹; ** $\mu_{eff:}$ Magnetic moment in Bohr Magneton; d=decomposition points; app= apparatus.

The resulted complexes were color-ed solid, moderately soluble in ethanol, soluble in dimethylformamide and dimethylsulfoxide. Elemental analyses revealed that the complexes have the compositions $[Ni(SCH)(L)_nX_n]$ and $[Ni_n(SCH)_n(L)_n]$. The molar conductivities $(^{M})$ of the comp-lexes in 10⁻³ M dimethylformamide are determined, the values shown in Table-2 approached those expected for non electrolytes (20). The magnetic moments of the complexes calculated from the corrected magnetic susceptibilities determ-ined at room temperature are shown in Table-2. The μ_{eff} values reported for the complexes (1.031-1.982 and 2.049-2.982 B.M.) supported the monomeric or dimeric structures indicating octahedral geometry around the metal ions (21). Refraction measurement (Table 2) of the complexes were found to be either positive or negative values (22). The molar refraction is an additive and constitutive property. It has been used to construct the correct structure of the compound (23).

The infrared spectra of semicarba- zone ligands (Table 3) showed a strong band at 1688 cm^{-1} attributed to the C=O group (2,16,24) shifted towards a lower frequency on coordination in neutral medium, indicating the formation of a chelation between the oxygen of the C=O group and the metal ion (2,24). Mean- while, in basic medium, this band was disappeared in the complexes and a new band has been observed at 1217-1251 cm⁻¹ due to C-O group, thereby establishing coordination of the ligand through the enolic oxygen atom (3,24). The next strong band at 1575 cm⁻¹ attributed to C=N group (24) shifted towards a lower frequency on coordination due to the decrease of the bond order as a result of metal nitrogen bond formation (3,24). The appearance of strong bands at 1630-1635 cm⁻¹ and 1580-1585 cm⁻¹ attributed to $v_{(C=N)} + v_{(O-C=N)}$ and azine chromophore $v_{(C=N-N=C)}$, respect- ively, supported the formation of enolic structure in basic medium (2,3,24). The other band at 3473 cm⁻¹ due to NH group remained unaltered in the complexes prepared in neutral medium indicating that there is no coordination through the NH group (3,24). Whereas in basic medium, this band was disappeared due to the enolic form. The other bands at 1450.00 cm⁻¹ attributed to δ_{NH2} (2,24) remained unaltered on complexation indicating that there is no coordination through this group and the metal ion (2,3,24).

The infrared spectra of dithiocar- bamate ligands (Table 3) showed band at 1322-1325 cm⁻¹ assigned to C=S group shifted towards a lower frequency on coordination (24,25). The other band observed at 1212-1217 cm⁻¹ due to C-S group shifted towards a lower frequency on complexation indicating the formation of a chelation between the sulfur atom and the metal ion (24,25).

On the other hand the spectra of the complexes 5 and 7 showed bands at 1450-1486 cm⁻¹, 1355-1371 cm⁻¹ and 946 cm⁻¹ due to $v_{s(NO3)}$, $v_{as(NO3)}$ and v_{NO} (v_2 , v_1 , v_5), respectively (24). The difference between v_1 . v_5 equal to 115 cm⁻¹ which

supported the bonding of nitrato group as mono- dentate ligand through the oxygen atom. The spectra of the complexes 1 and 3 showed bands at 290-300 cm⁻¹ due to v_{M-Cl} . The spectra of the complexes 9 and 11 showed bands at 1125-1180 cm⁻¹, 1068-1077 cm⁻¹, 1005-1025 cm⁻¹ and 946 cm⁻¹ due to v_{SO4} indicating the monodentate coordination of sulfate group to the metal ion^[28]. The spectra of the complexes 13 and 15 showed two bands at 1535-1540 cm⁻¹ and 1365-1370 cm⁻¹ attributed to $v_{coo(s)}$ and $v_{coo(as)}$, respectively. The presence of thes two bands supported the coordination of the acetate group to the metal ion through one oxygen atom {i.e. the acetate group act as monodentate ligand} (24). Moreover the spectra of all the complexes showed new bands at 450-500 cm⁻¹, 417-420 and 525-579 cm⁻¹ due to v_{M-N} v_{M-S} and v_{M-O} , respectively (24,25). The presence of these bands supported the formation of the complexes under inves-tigation.

The electronic spectra of the Ni (II) complexes (Table 4) showed absorption bands at 7212.72-9608.28 cm⁻¹{ v_1 calcu- lated (26)}, 15625.00-18867.92 cm⁻¹ (v_2) and 21097.04-27247.95 cm⁻¹ (v_3) attributed to the three transitions ${}^{3}A_{2g}(F) \rightarrow {}^{3}T_{2g}(F)$, ${}^{3}A_{2g}(F) \rightarrow {}^{3}T_{1g}(F)$ and ${}^{3}A_{2}g(F) \rightarrow {}^{3}T_{1}g(P)$, respectively, which were expected for d⁸ system having high spin octahedral geometries (26). The ligand field parameter B {Racah parameter (interelectronic repul-sion)}, the crystal field parameter Dq and the ligand field splitting energy (10Dq) have been calculated (26). The values of β {nephelauxetic ratio (β =B'/B)} has been determined. The apparent value of B' was always smaller than that of the free ion (B of free nickel ion=1084). The phenomenon is known as the nephelauxetic effect and was attributed to delocalization of the metal electrons over molecular orbitals that emcompass not only the metal but the ligands as well. As a result of this delocalization or (cloud expanding) the average interelectronic repulsion was reduced and B' was smaller. The values of β values of the complexes were between 0.791-1.001 indicating the covalent character of the bond concerned. The values of Dq/B indicated that the complexes had octahedral geometries of all the complexes (26).

Conclusions

This work in fact is a continuation of our studies including mixed ligand comp-lexes. In this work some observations have been achieved that lead to establish the SCH₂ ligand acted as tetradentate or tridentate chelating ligands joint to nickel (II) ion through the nitrogen and oxygen atoms. This ligand acted as neutral molecule in the complexes prepared in neutral medium, whereas in basic medium it acted as anion (-I, or -2) due to the deprotonation process that took place in enol form of this ligand. The ligands Pyrdtc and Indtc acted as anion (-1) joint to metal ion through sulfur atom, they acted as either monodentate or bidentate chelating ligands. Chloride ion, nitrate ion, acetate ion and sulfate ion acted as mono-dentate ligands in the complexes prepared in neutral medium. Nickel (II) ion

had probably hexa-coordinated, leading to high spin octahedral geometry (Figure 2).

Table 3: IR spectra of complexes (values in cm⁻¹).

No.	$\upsilon_{C=N}$	$v_{C=S}$	$\upsilon_{C\text{-}S}$	$\upsilon_{C=O}$	υ_{C-O}	υ_{OH}	$\upsilon_{M\text{-}N}$	υ_{M-S}	υ_{M-O}	Other bands
SCH ₂	1575	—	_	1688	_	3213	_	_	_	—
Pyrdtc	_	1325	1211		_	_		_	_	_
Indtc		1322	1217							_
1	1509	1164, 1002	946	1565			500	417	550	υ _{M-Cl} 290
2	1510	1060. 1010	945		1217	3200	450	418	550	_
3	1510	1078, 1024	933.	1560			454	420	549	υ _{M-Cl} 300
4	1509	1138, 1025	934		1218	3250	454	420	538	_
5	1509	1164, 1025	911	1561			450	417	526	υ _{NO} 1450, 1335, 946
6	1508	1074, 998	945		1250	3220	454	417	540	_
7	1509	1155, 1020	934	1561			454	418	540	υ _{NO} 1486, 1371
8	1509	1079, 1020	934		1250	3200	450	420	530	
9	1509	1163, 1025	914	1560			456	420	525	υSO ₄ 1180, 1077, 1025, 946
10	1510	1074	945		1250	3220	450	418	543	
11	1510	1079, 1020	999	1565			450	420	540	υSO ₄ 1125, 1068, 1005
12	1510	1104, 1015	915		1250	3200	454	420	540	
13	1510	1163, 1020	1000	1559			456	418	579	υ _{Ac} 1370, 1540
14	1509	1075, 1026	945		1251	3286	451	420	537	
15	1514	1079, 1020	910	1565			500	420	550	υ _{Ac} 1365, 1535
16	510	137, 025	933		1218	3200	454	420	539	

Table 4: Electronic spectral data of some complexes.

No.	$v_1 \text{ cm}^{-1}$	$v_2 \mathrm{cm}^{-1}$	$v_3 \text{ cm}^{-1}$	C.T cm ⁻¹	В	10Dq	Dq/B	β	C.F.S.E
1	7226.27	15723.27	20833.33	30674.84	903.28	7226.30	0.80	0.833	8671.524
2	7212.72	15723.27	21739.13	31055.90	901.59	7212.72	0.80	0.832	8655.264
3	9187.31	16077.17	22935.77	29940.11	1080.9	9187.31	0.85	0.997	11024.77
4	8286.04	18867.92	22935.77	29940.11	920.67	8286.04	0.90	0.849	9943.248
5	9225.02	15625.00	26595.74	30120.48	1085.3	9225.02	0.85	1.001	11070.24
6	9608.29	15723.27	26595.74	31055.90	1011.4	9608.29	0.90	0.933	11529.95
7	7609.74	15723.27	22935.77	30120.48	951.22	7609.74	0.80	0.878	30120.48
8	7609.74	16025.64	22935.77	29857.90	951.22	760.974	0.80	0.878	9131.688
9	8110.32	15723.27	21097.04	31250.00	1013.8	8110.32	0.80	0.935	9732.384
10	7286.99	15723.27	26595.74	31250.00	857.29	7286.99	0.85	0.791	8744.388
11	9187.28	17141.83	22935.77	29940.11	1080.9	9187.28	0.85	0.997	11024.73
12	7955.52	15772.87	22935.77	33783.78	935.99	7955.52	0.85	0.864	9546.624
13	7837.82	15723.27	27247.95	31055.90	922.10	7837.82	0.85	0.851	9405.379
14	9608.29	15723.27	26595.74	31055.90	1011.4	9608.29	0.90	0.933	11529.95
15	9187.28	17141.83	22935.77	30120.48	1080.9	9187.28	0.85	0.997	11024.73
16	8286.04	18867.92	22935.77	30120.48	920.67	8286.04	0.90	0.849	9943.248





Where $Z = Cl^{\circ}$, NO_{3}° , $CH_{3}COO^{\circ}$; R =

Figure (2): Proposed structures of the complexes.

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