# Studies on Some Transition Metal Complexes with LIX:84 

Ali, Rustom<br>University of Rajshahi

http://rulrepository.ru.ac.bd/handle/123456789/880
Copyright to the University of Rajshahi. All rights reserved. Downloaded from RUCL Institutional Repository.

# STUDIES ON SOME TRANSITION METAL COMPLEXES WITH LIX84 



MDissertation<br>Submitted to The University of Raishahi in Partial Fulfilment of The Requirements for The Degrec of J.haster of Philosophy in Chemistry

> By

## MD. RUSTOM ALI

B.Sc. (HONS) M.Sc.

RAJSHAHI
2007

PHYSICAL RESEARCH
LABORATORIES DEPARTMENT OF CHEMISTRY RAJSHAHI UNIVERSITY RAJSHAHI, BANGLADESH

## STUDIES ON SOME TRANSITION METAL COMPLEXES WITH LIX84


oithissertation



> BY
MD. RUSTOM ALI
B.Sc. (HONS) MLSc.

RAJSHAIII
2007

PHYSICAL RESEARCH
LABORATORIES
DEPARTMENT OF CHEMISTRY
RAJSHAFII UNIVERSITY
RAJSHAHI, BANGLADESH

## Declaration

I hereby declare that the entire work now submitted as a thesis to the Department of Chemistry, University of Rajshahi, towards the fulfillment for the degree of Master of Philosophy is the result of my won investigation and was carried out under the supervision of Prof Dr. Md. Shamsul Islam, Department of Chemistry, University of Rajshahi, Rajshahi. The thesis has not been submitted elsewhere for any other degree.

(Rustom Ali)
M. Phil. Fellow

# Department of chemistry <br> University of Rajshahi 

## Declaration Certificate

This is to certify that Mr. M. Rustom Ali, M.Sc. the M.Phil research fellow of the Department of Chemistry, Rajshahi University carried out his research work under my supervision in the research laboratories in the same department for his Dissertation submitted for the Degree of Master of Philosophy. This research involving the study of complexing behaviour of the three metal ions with the extractant LIX84, other physical and microbiological properties is an original work for the extension of knowledge in the fields of solvent extraction and organometallic Chemistry.During the progress of his research, he gave a successful seminar on his thesis material and passed the assigned theoretical courses successfully in this department.

(Prof. Dr. M.Shamsul Islam)
Supervisor

EXAMMHEXS
COPY

PART-I
Page
ACKNOWLEDGEMENT ..... I
ABSTRACT ..... II
LIST OF TABLES ..... III
LIST OF FIGURES ..... IV
LIST OF ABBREVIATIONS ..... V

## CHAPTER-1

## GENERAL INTRODUCTION

1.1 Transition Metal Chelates and their Stability ..... 2
1.2 Solubilities of Metal Chelates ..... 6
1.3 Electrolytic Properties of Complexes ..... 8
1.4 Magnetic Properties of Transition Metal Complexes ..... 8
1.5 IR Spectral Data of Complex Compounds ..... 11
1.6 UV-VIS Spectra of Complex Compounds ..... 12
1.7 LIX 84 as Extractant/Ligand ..... 14

## CHAPTER-2

## LITERATURE REVIEWS

2.1 Literature Reviews 17
2.2 Scope of the Present Investigation 25

## CHAPTER-3

## PRESENT AIM AND PREPARATION

Page
3.1 Aim of the Present Investigation: ..... 28
3.2 Preparation of the Complexes ..... 28
CHAPTER-4
EXPERIMENTAL TECHNIQUES
4.1 Weighing ..... 33
4.2 Melting Point Measurements ..... 33
4.3 Conductivity Measurements ..... 33
4.4 Magnetic Measurements ..... 34
4.5 Infrared Spectra ..... 35
4.6 Electronic Spectra ..... 35
4.7 Determination of Molecular Weight ..... 37
CHAPTER-5
RESULTS AND DISCUSSION
5.1 Molecular Weight ..... 39
5.2 Molar Conductances ..... 39
5.3 Magnetic Moments ..... 39
5.4 Percent Solubilities ..... 41
5.5 Infrared Spectra ..... 41
5.6 Electronic Spectra ..... 42

## LIST OF STRUCTURE OF COMPLEXES

Page
i) $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ ..... 44
ii) $[\mathrm{Ag}(\mathrm{L})]$ ..... 45
iii) $\left[\mathrm{Cd}(\mathrm{L})_{2} \cdot \mathrm{HL}\right]$ ..... 46

## LIST OF TABLES

Table-1: Color, yield, decomposition temperature and molar conductance of the complexes. ..... 47
Table-2: Data for solubility of the complexes. ..... 48
Table-3: Percentage (\%) determination of the solubility
of complexes in various organic solvents $\left(\mathrm{gdm}^{-3}\right)$. ..... 49
Table-4: Detailed magnetic data of the complexes ( $\mathrm{T}=303 \mathrm{~K}$ ). ..... 50
Table-5: Observed frequencies and assignments for LIX84. ..... 51
Table-6: Infrared Absorption Maxima in $\mathrm{cm}^{-1}$
of LIX-84 and M-LIX84 complexes. ..... 52
Table-7: Electronic spectral data of the prepared complexes with assignment. ..... 53
Table-8: Electronic spectral data of LIX84 ..... 54
Table-9: Electronic spectral data of the prepared complex $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ ..... 55
Table-10: Electronic spectral data of the prepared complex $[\operatorname{Ag}(\mathrm{L})]$ ..... 56
Table-11: Electronic spectral data of the prepared complex [Cd $\left.(\mathrm{L})_{2} . \mathrm{HL}\right]$ ..... 57

## LIST OF FIGURES

Page
Fig. 1. Infrared spectrum of LIX 84 in KBr cell. ..... 57
Fig. 2. Infrared spectrum of $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ in KBr cell. ..... 58
Fig. 3. Infrared spectrum of $[\mathrm{Ag}(\mathrm{L})]$ in KBr cell ..... 59
Fig. 4. Infrared spectrum of $\left[\mathrm{Cd}(\mathrm{L})_{2}\right.$. HL$]$ in KBr cell ..... 60
Fig. 5. Electronic spectrum of LIX 84 (in cychlohexane). ..... 61
Fig. 6. Electronic spectrum of $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ (in cychlohexane). ..... 62
Fig. 7. Electronic spectrum of $[\mathrm{Ag}(\mathrm{L})]$ (in cychlohexane). ..... 63
Fig. 8. Electronic spectrum of $\left[\mathrm{Cd}(\mathrm{L})_{2} . \mathrm{HL}\right]$ (in cychlohexane). ..... 64
REFERENCES ..... 66-69
APPENDIX ..... 98-100

## LIST OF ABBREVIATIONS

1. cm Centimeter
2. $\mathrm{cm}^{-1}$ Per centimeter
3. Fig. Figure
4. i.e. That is
5. K Degree Kelvin
6. m.p. Melting point
7. mL Milliliter
8. No. Number
9. nm Nanometer
10. \% Per cent
11. $\wedge \quad$ Conductance
12. $\lambda$ Wave length
13. $\chi_{g}$ Mass susceptibility
14. $\chi_{\mathrm{m}}$ Molar susceptibility
15. g Gramme
16. uv Utra-violet
17. IR Infrared
18. $v$ Absorption maximum frequency

PART-II

## CHAPTER-6

ANTIBACTERIAL ACTIVITY TESTING

## LIST OF TABLES

## Table 6.1: Complexes abbreviation for antibacterial activity. <br> 77

Table 6.2: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-roxy-5-nonyl acetophenone, oxime against Escherichia coli.78

Table 6.3: Antibacterial activity of the complexes 1,2,3 and ligand 2-hydroxy-5-nonylacetophenone against Shigella dyscnteriae.79

Table 6.4: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5-nonylacetophenone against Pseudomonas aeruginosa. 80
Table 6.5: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5- nonylacetophenone against. Bacillus subtilis 81
Table 6.6: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5- against nonylacetophenone Sarcina lutea. 82
Table 6.7: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5- nonylacetophenone against Staphylococcus aureus. 83

## LIST OF FIGURES

Fig-6.1 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Escherichia coli.84

Fig-6.2 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Shigella dysenteriae. 85

Fig-6.3 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Pseudomonas aeruginosa.
Fig-6.4 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Bacillus subtilis.87

Fig-6.5 Photographic representation of zone of inhibition of the complexes 1, 2, 3 and the standard compound kanamycin against Sarcina lutea.
Fig-6.6 Photographic representation of zone of inhibition of the complexes $1,2,3$ and the standard compound kanamycin against Staphylococcus aureus.

## ABSTRACT

The research work presented in this thesis deals with the preparation and characterization of some new complexes of the second transition metal ions $\left(\mathrm{Zr}^{4+}, \mathrm{Ag}^{+}\right.$and $\left.\mathrm{Cd}^{2+}\right)$ with the new solvent extraction reagent, LIX84 (i.e.2-hydroxy-5- nonylacetophenone oxime) as a ligand. For their synthesis, the prepared aqueous solutions of the metal chlorides (for Cd (II), nitrate) are allowed to react with an n-hexane solution of LIX84 (containing KOH ) in different ratios. The prepared complexes have been characterized by conductivity and magnetic measurements, infrared spectra and electronic spectral studies. Their percent solubilities in various organic solvents and decomposition temperatures have been measured. The molecular weight of the complexes have also been determined by the cryoscopic method.

The prepared complexes have compositions;

1. $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$
2. $[\operatorname{Ag}(L)]$
3. $\left[\mathrm{Cd}(\mathrm{L})_{2} \cdot H L\right]$
where, $L$ denotes a LIX84 anion.
The complex 1 is assumed octahedral and the complex 2 is linear but the complex 3 has tetrahedral structure based on experimental data. IR spectra of the complexes 1,2 and 3 indicate coordination of the ligand to the metal ion through its N and O atoms. In these complexes LIX84 acts as a uninegative bidentate ligand. From magnetic measurements, the complex $\mathbf{1}$ is found to be binuclear paramagnetic, whereas complexes 2 and 3 are diamagnetic. All the complexes are found to be insoluble in polar solvents viz, water, ethanol and acetone; but sparingly soluble in other organic solvents. Measured values of the molecular weights and magnetic moments for the complexes indicate that $\mathbf{1}$ is binuclear compound. Moreover, all the complexes have been observed to behave as non-electrolytes in DMSO or cyclohexane. Probable structures for the complexes have been suggested.

Further, the microbiological study of LIX84 and its $\mathrm{Zr}, \mathrm{Ag}, \mathrm{Cd}$, complexes has been carried out. Studies of the antibacterial activity of the prepared complexes showed them to behave as good inhibitors on mycelial growth against selected standard and the ligand has been studied for six pathogenia bacteria viz.,

## 1. Escherichira coll (Gram negative)

2. Shgiella desenteria (Gram negative)
3. Pesudomonus areuginosa (Gram negative)
4. Bacillus subtilis (Gram positive)
5. Sarcina lutea (Gram positive)
6. Staphylococcus aureus (Gram positive)
and also investigated the minimum inhibitory concentration of the complexes 2 and 3. All the results obtained is compared with the standard compound, Kanamycine.

The results of this investigation reveal that the complexes are more or less active compared to the ligand (2-hyroxy-5-nonylacetophenone oxime).

Among the complexes, complexes 2 and 3 show the excellent result.

## GENERAL INTRODUCTION

## INTRODUCTION:

Transition metals are those elements, which usually have partly, field $d$ or $f$-shells in any of their commonly occurring oxidation states ${ }^{1}$. Because of their variable valence, they form a large number of complexes with various types of ligands with a few exceptions. These compounds are colored in one if not all oxidation states, and at least a few of them are paramagnetic.

However, a few transition metal complexes with 2-hydroxy-5nonylacetophenone oxime (LIX 84) are reported ${ }^{2}$.

This compound has recently been introduced as a solvent extraction reagent by Hinkle Corporation, U.S.A. Being a 2- hydroxy-5nonylaceto phenone it is expected that LIX 84 may form water insoluble complexes with various metal cations in slightly acidic solutions under controlled pH conditions. This reagent has been used in hydrometallurgy for the extraction a few very useful metals viz. copper from aqueous ammonia solutions using the method of solvent extraction (SX). But the chemistry of the extraction processes as well as the elucidation of the structures of the extracted complexes has not yet been studied. Complexing behaviour of LIX 84 with other useful metal ions are lacking.

Some first transition metal complexes of LIX 54 have been prepared and characterized, but these await publication. However, studies with heavier
transition metals are also lacking therefore, the preparation and studies on isolated complexes of LIX 84 with such metals appear interesting from those points of view. The preparation and characterization of some complexes of Ag ( I ), Cd ( II ), Zr ( IV ) with LIX 84 have been presented in this dissertation, the reagent acting as a uninegative bidentate ligand. The knowledge acquired will be helpful for different metal extraction processes with LIX 84 using the solvent extraction method.

### 1.1 TRANSITION METAL CHELATES AND THEIR STABILIY:

A metal complex may be defined as a compound containing a central metal ion (cation) to which are attached oppositely charged ions or neutral molecules whose number usually exceeds the number corresponding to the oxidation number of the metal ion. The ions or neutral molecules which are attached (coordinated) to the central metal ion are called ligands.

Transition metal ions form stable chelate complexes with high molecular ligands containing more than one containing more than one coordinating centers, Nitrogen-sulpher, nitrogen-oxygen-sulpher or phosphorous-oxygen donor ligands are known to form very important complexes with transition metals. Such metal chelates are formed when metal cations (Lewis acids) react with large chelating anions (Lewis base). Usually, the chelating ligands destroy the hydration sphere of the metal ions in their aqueous solution forming a stable molecular species.

Most of organic ligands being weak acids of long carbon chains ( $>6 \mathrm{C}$ ) are generally soluble in organic solvents. Usually, they contain at least two coordinating functional groups on either sides of 2 or 3 atoms. One
group of the chelating molecule is an acidic OH . NH or SH each containing at least one electronegative atom $\mathrm{O}, \mathrm{N}$ or S having basic character that possesses lone pair of electrons. The other is one uncharged basic functional group or atom capable of coordinating by its donor properties to the metal. Such chelating ligands function as bases by loss of a proton from the acidic group for an equivalent of metal, and thus produce reactive anionic ligands the chelating reagents remaining as almost undissociated neutral molecules (without solvation or coordination) in solution in an inert (non polar) organic solvent having no donor atoms capable of coordination to metal ions. In principle they do not undergo dissociation or association itself or with the metal chelates in the organic phase; but they dissociate ionically in the aqueous phase or at the interface of the two phases. Equilibrium studies have much contribution to the understanding of the various factors that influence metal complex formation in aqueous solutions.

The factors affecting the formation and stability of metal chelates include ${ }^{3}$.
i) The basic strength of the chelating agent: the higher the basicity of the chelating group (higher pKa value), the greater is the stability of the metal chelate formed. Nevertheless, a more acidic chelating agent with a lower pKa value, will be more useful for the extraction of metals from acidic aqueous solutions.
ii) The electronegativity of the bonding atoms of the basic group in the chelating agent: atoms of lower electronegativity (increasing acidity of the group) tend to form stronger covalent
bonds and there by increase the stability of metal chelates. For example, nitrogen and sulpher, have better donor properties than oxygen, as their electronegativity increases in that order.
iii) Size and number of chelate rings formed; from Bayer strain theory, five or six membered rings are most stable (minimum strain). All other factors being equal, a five membered ring chelate will be somewhat more stable than its six membered ring analog. Apart from any intrinsic advantage of ring formation, chelate stability also increases with the number of fings that are formed possibly owing to increasing number of say, water mole cules that are displaced from that metal coordination sphere by one molecule of the polyfunctional reagent.
iv) The metal ion characteristics such as acidity i.e., ionic potential (i.e., charge/ionic, radius ratio) or charge ${ }^{2} /$ ionic ratio, electronegativity, coordination number and coordinating ability; as the ration increases the stability of the metal complex usually rises, provided the metal ion has available bonding or bitals for the electron pairs it accepts. The nature of the orbitals affects both the bond strength and the stereochemical arrangement of the complex ${ }^{4}$. Moreover, complexes of the more electronegative metals ( e.g. $\mathrm{Zr}, \mathrm{Ag}, \mathrm{Cd}$ ) tend to be more stable.
v) Resonance and steric effects: the stability of the chelate structure is enhanced by contribution of resonance structures of the chelate ring ${ }^{5}$ But the stability decreases because of steric
hindrance to chelate formation caused by the bulky substitution groups ${ }^{6}$; this hindrance is, however, at a minimum in tetrahedral grouping of the reagent molecules about the metal ion.

The majority of coordination compounds contain nitrogen, oxygen, sulpher or a halogen as the bonding atom. Some metals which prefer oxygen over nitrogen are Mg (II), Ca (II), $\mathrm{Si}(\mathrm{II}), \mathrm{Sn}(\mathrm{II}), \mathrm{U}$ (VI ), Fe ( III ) are Be (II). Some of which prefer nitrogen donors over those of oxygen are $\mathrm{Cu}(\mathrm{I}), \mathrm{Cu}(\mathrm{II})$ , $\mathrm{Ag}(\mathrm{I}) \mathrm{Cd}(\mathrm{II}), \mathrm{Hg}(\mathrm{II}), \mathrm{V}$ (II) $\mathrm{Co}(\mathrm{III})$ and $\mathrm{Ni}(\mathrm{II}), \mathrm{Cr}(\mathrm{III}), \mathrm{Fe}(\mathrm{II}), \mathrm{Pt}(\mathrm{IV}), \mathrm{Zn}(\mathrm{II})$ , Zr (IV), V) seem to coordinate equally well either with oxygen or nitrogen Except for $\mathrm{Cu}(\mathrm{I}), \mathrm{Ag}(\mathrm{I}) \mathrm{Au}(\mathrm{I})$ and $\mathrm{Hg}(\mathrm{II})$, oxygen donors are clearly superior to those of sulpher. With certain large ions of low charge (eg) $\mathrm{Cu}(\mathrm{I}) \mathrm{Ag}(\mathrm{I})$ $\mathrm{Hg}(\mathrm{I})$ the maximum stability occurs with sulpher containing ligands. However $d^{8}$ or $d^{10}$ ions have little enderncy to become positively charged and are therefore, able to ccept electrons from ligands Such ions are, $\mathrm{Hg}(\mathrm{II}), \mathrm{Ag}(\mathrm{I}), \mathrm{Pd}(\mathrm{II})$ and $\mathrm{Pt}(\mathrm{II})$ which form more stable complexes with ligands of high polarizability, and the order is $\mathrm{P}>\mathrm{S}>\mathrm{N}>0$.

Generaly donor atoms which gave large ligand field spliting from their strongest complexes with metal ions are particularly sensitive to ligand field stabilization whereas donor atoms producing small ligand fields tend to form relatively more stable complexes with cations which are insensitive to ligand field stabilization. Moreover, the difference in instability between complexes which produce small and large ligand fields will be mich greater for metal ions which are particularly sensitive to ligand field stabilization than those which are not. Metal chelates with least basic i.e. weakest donors dimerize to the greatest extent ${ }^{7}$.

### 1.2 SOLUBILITY OF METAL CHELATES:

Metal chelates, being essentially neutral covalent compound, are far less soluble in water than in organic solvent (dilute). Rendering the chelate molecule larger and bulker, and more hydrophobic, its coordination properties can be altered. Factors causing low affinity of the metal chelate to aqueous are zero of low charge, large sizw, non polar nature, absence of electronegative atom at the surface and a highly ordered water structure, which also cause a high affinity for the organic phase neutral nonhydrated metal chelates are usually formed by many polyvalent metal cations whose coordination number is twice the charge; but some bivalent ions $\left(\mathrm{Cd}^{2+}\right.$, $\mathrm{Zr}^{4+}$ ) having a coordination number higher than four may retain one or more water molecules in their first coordination sphere ${ }^{8}$.

The chelates of heavy metals are in many cases soluble in organic solvents. Some of them, however, are soluble in highly polar solvents, e.g. alcohol and carbon terta chlorice $\left(\mathrm{CC1}_{4}\right)$, while others are soluble only in typically non polar solvents e.g. $\mathrm{CH}_{2} \mathrm{C1}_{2}$ or $\mathrm{C}_{6} \mathrm{H}_{6}$. Therefore, systematic compariso of the effects of various solvents on their structures of electronic polarity so that they can even serve as appfoximate indicators of solvent polarity.The solutions thus obtained sometimes exhibit a marvellous variety of colours according to the change in solvent states can often be carried out only over a limited range of solvent several different kinds of spectral charges are involved in these colour changes and those observed for the $\mathrm{Cd}(I I)$ chelates are quite different from the $\mathrm{Zr}(I I)$ chelates.

The nature of a solvent may also influence the reactivity of chemical compounds. The different reactivity of a solute X in two solvents, $\mathbf{1}$ and $\mathbf{2}$ may be characterized by standard free energy of transfer, $\Delta G^{\text {tr. }}$; and with its
help the equilibrium constants of many reactions. e.g., $A+B \leftrightarrows C$ in the solvents, 1 and 2 may be calculated.

$$
-\operatorname{RT} 1 n K_{1} / K_{2}=\Delta G^{\text {tr. }}(\mathbf{C})-\Delta G^{\text {tr }}(\mathbf{A})-\Delta G^{\operatorname{tr}}(\mathbf{B})
$$

Systematic investigations ${ }^{9-12}$. using various non-aqueous solvents have led to the conclusion that this $\Delta G^{\text {tr }}$ is almost exclusively determined by short range solute-solvent interactions. In order to express these short range interactions recourse has been taken to empirical solvent parameters such as Gutmann's donor and acceptor numbers (DN \& AN), Kosower's Z, Zimoroth- Reichardt's $\mathrm{E}_{T}$ values etc.

Sone and Fukuda ${ }^{13}$ have studied the electronic spectra of the solutions of some complexes, wach of which is soluble in a good number of solvents. They found that the colours of the solutions and the $v_{\text {max }}$ of the $d$-d bands are highly sensitive to changes in solvent polarity. For instance, mixed Cu (II)chelate with tetramethylethylenediamine and aacetylacetonate ion has a bluish or violet colour in aqueous solution. The colour becomes more and more reddish with the decrease in solvent polarity expressed in DN ; and in solvents with very lox DN values, it is nearly red. On the other hand, in solvents of increasing polarity (high DN values) the complex assumes inereasingly bluish colour and in $\left(\mathrm{CCl}_{4}\right)$ it is blue. The parameter, donor number (DN) describes the nucleophilic behaviour and the acceptor number (AN) clectrophilic behaviour of the solvent.

Sone and Fukuda ${ }^{13}$ who worked with square planar complexes argue that in solvents of increasing polarity, the complexes are ionized and the solvent molecules enter into the coordination sphere. Theoriginal square
planar structure gradually tends towards the octahedral form and as a result, the d-d transition shifts towards higher wave lengths, that is lower $\mathrm{v}_{\text {mas }}$ values.

### 1.3 ELECTROLYTIC PROPERTIES OF COMPLEXES:

Conductance measurements can also be used to give useful information about the total number of ions (Complex, simple or both) yielded by a complex in solution and charge type on each ion. Electrolytic or non-electrolytic nature of the complex can be inferred from the experimental conductance data. Conductance values wall ascertain both the type and nature of electrolytes ${ }^{14,15}$.

The state of the anionic species i.e., whether the anions are in bridge, inside or out side the coordination sphere, can also be predicted from the conductance which may thus help to determine the structures of the coordination complexes.

### 1.4 MAGNETIC PROPERTIES OF TRANSITION METAL COMPLEXES:

The fundamental magnetic properties (either paramagnetism or diamagnetism) of coordination compounds or complexes are used on the basis of measurements of their magnetic susceptibilities. From the experimental data, different possible electronic configurations can be assigned to them, which may give the clear information as well as stereochemistry of the complexes and even nature of bonding present there in ${ }^{16,1}$.

Generally transition metal compounds show two types of magnetic properties, viz., paramagnetism and diamagnetism. A substance which is paramagnetic is attracted into a magnetic field with a force propotional to the field strength times the field gradient. Paramagnetism of a substance consisting of atoms, ions or molecules is generally caused by the presence of unpaired electrons. The greater the number of unpaired electrons, The greater will be the paramagnetism shown by the substance. The substance having no field than that in vacuum is called a diamagnetic substance and the property due to which it shows this behaviour is called diamagnetism.

Paramagnetic substances are characterized by the positive values of their magnetic susceptibilities. A paramagnetic substance increases the flux due to the applied field. Diamagnetic substances when brought under the influence of magnetic field, a repelled. These substances are characterized by negative values of magnetic susceptibilities. The diamagnetic character is an intrinsic phenomena, but the magnitude of the diamagnetic effect is small.

The ions of the first transition series starting with scandium ( $\mathrm{Sc}^{3+}$ ) ion and ending with $\operatorname{zinc}\left(\mathrm{Zn}^{2+}\right)$ ion, second transition series starting with yttrium $\left(\mathrm{Y}^{3+}\right)$ ion and ending with cadmium $\left(\mathrm{Cd}^{2+}\right)$ ion, and the third transition series starting with lanthanum $\left(\mathrm{La}^{3+}\right)$ ion and ending with mercury $\left(\mathrm{Hg}^{+2}\right)$ ion have progressively zero to ten electrons in the $3 \mathrm{~d}, 4 \mathrm{~d}$ and 5 d orbitals respectively. Their compounds may show paramagnetism or diamagnetism depending upon their oxidation states i.e., the presence of odd or even number of electrons in the ligand/crystal field. The greater the erystal field, the greater is the tendency of the electrons to be come paired up. If the crystal field is not so sufficient then the Hund's rule of maximum multiplicity is obeyed.

Hund's rule states that electrons tend to go separate orbitals with parallel spins provided the situation is energetically reasonable. The configuration with maximum number of unpaired electrons is of the high spin type and with minimum number of unparied electrons, the low spin type. For example $\mathrm{Fe}^{2+}$ which is $\mathrm{d}^{6}$ system in presence of octahedral field of high spin type shows paramagnetism corresponding to two unpaired electrons. But in presence of square planar field, it becomes low spin type and diamagnetic ${ }^{16}$.

The magnetism of second and third row electrons compared to that of the first row, is of much greater frequency with which diamagnetic occurs, and the paramagnetic moments are usually much lower. This is partly due to the fact that the complexes are invariably of the spin paired type ${ }^{15}$, $\mathrm{The} \mathrm{Ag}^{+}$ is of course, diamagnetic ${ }^{17}$. The possible divalence of silver and of gold in $\mathrm{Cs}_{2} \mathrm{Ag} \mathrm{AuCl} 1_{6}$ and in $\mathrm{Cs}_{2} \mathrm{Au}_{2} \mathrm{Cl}_{6}$ respectively has been examined by Elliot ${ }^{18}$. Both the compounds are diamagnetic. Zirconium and halfnium apparently form no appreciable paramagnetic compounds. The useful valence of four is diamagnetic. The mercurous ion is diamagnetic because it always forms a double ion. $\mathrm{Hg}^{+2}$. The mercury ion has no unpaired electrons and is diamagnetic ${ }^{17}$.

Generally transition metal Compounds are coloured. The Colour and magnetic properties of transition metals are related to the presence of unpaired electrons in the system. Colour is not only due to upaired electrons but also due to other factors as well as electron transitions in presence of litght. Magnetic moments are extensively used to the study of complex compounds. From the evaluated magnetic moment, one can find out the
number of unpaired electrons present in the system and thus gets an idea of its possible configuration and hence the structure.

### 1.5 IR SPECTRAL DATA OF COMPLEX COMPOUND:

All organic and inorganic species (except homonuclear diatomic molecules) absorb infrared radiation. IR spectroscopy has wide use in inorganic chemistry mainly coordination chemistry and proved to be a rich and convenient source of information concerning to the characteristic functional groups as well as structure and bonding. The condition of a reaction can be characterized from the initial IR and the product IR spectra. The field of vibrational spectroscopy of coordination chemistry has been expanded by a variety of research technich leading to many new and interesting result ${ }^{19}$. It is be coming more and more popular with ever increasing rapidly day by day. Therefore, the great utility of the vibrational spectroscopy in deducing structural and bonding information is noteworthy.

Since every different type of bond has a different natural frequency of vibration, and since the same type of bond in two different compounds is in a slightly different environment, no tow molecules of different structure will have exactly the same infrared absorption pattern or infrared spectrum. The presence or absence of a particular functional group in a molecule or the type of coordination in a compound can be assigned. IR spectrum can be used for compounds as finger prints can be used for human beings.

Penland et $\mathrm{al}^{20}$ first studied the infrared spectra or urea complexes to determine whether coordination occurs through nitrogen or oxygen. If coordination occurs through nitrogen, an increase of the $\nu(\mathrm{C}=0)$ stretching

Rajshabi University Library
Documentation Section
Document No Io...29....
Date $26 \cdot 4.08$
frequency with a decrease in the $\mathrm{v}(\mathrm{N}-\mathrm{H})$ stretching frequency results. On the other hand, if oxygen atom participates in coordination, a decrease in the $\mathrm{v}(\mathrm{C}=0)$ stretching frequency results without any appreciable change on the $\mathrm{v}(\mathrm{N}-\mathrm{H})$ stretching frequency ${ }^{21-22}$. The metal nitrogen stretching frequencies usually lie in the range $480-310 \mathrm{~cm}^{-1} 23-24$. The bands at $400-200 \mathrm{~cm}^{-1}$ are attributed due to $\mathrm{v}(\mathrm{M}-\mathrm{C} 1)$ stretching frequency ${ }^{19}$. Thus IR spectroscopy offers a variety of information about the complex compounds.

### 1.6 UV- VIS. SPECTRA OF COMPLEX COMPOUND:

The ultraviolet and visible spectroscopy is a simplex but powerful tool, which gives information on the geometries of complex molecules. In a typical electronic transition by metal complexes, the observed spectrum, in general, consists of a series of crystal field bands which are in the visible region and depends on the donor atom of the ligand and on the metal ion.

The crystal field transitions are of two types. One is the intens spinallowed transition and another is the lower intensity spin-forbidden transitions, which appear as shoulders on the spin allowed transition. The ultraviolet spectrum is complicated and consists of electronic transitions between the ligands and the metal (charge transfer) and also transitions within the ligand itself which are usually $\pi-\pi^{*}, n-\pi^{*}$ otransitions. The bands in the electronic spectra represent' different vibration transitions according to the electronic charge in rotational energy superposde on the electronic and vibrational changes.

The ions of the transition metals give rise to a great rangeof colours which vary in hue and intensity from compound to compound and with oxidation state. This has long been known to be one of their more
characteristic features. These colours arise from the absorption of the visible spectrum by the ions and their associated ligand group. The electronic dipole mechanism is the only of importance for the absorption fo light by the complex ions. when radiation of frequency equivalent to the energy separation between two levels specified by the functions $\Psi_{1}$ and $\Psi_{2}$ is applied this mechanism may be absorbe. The intensity of a band is best measured in terms of a quantity known as the ossillaror strength $f ; f$ is the area under the band when extinction co-efficient, $\varepsilon$, is plotted against frequency v .

If the terms of transition metal complexes correspond to a single energy level that absorption bands in their spectra should be exceedingly narrow. It has been that 10 Dq depends upon the metal-ligand separation in a bery critical manner. The energy differences between the highest and lowest states of a term (The overall multiplet width) is some simple multiple of the spin orbital coupling coupling is very large, the fine structure has been observed in solutions of their complexes.

The intense bands that are exhibited by the ligand around 345 nm and 303 nm are due to $n-\pi^{*}$ and $n-\pi^{*}$ respectively ${ }^{25}$. For example, electronic spectra of all complexes show three bands about $400 \mathrm{~nm}, 290 \mathrm{~nm}$ and 300 nm which may be assigned to the transitions respectively-

$$
\begin{aligned}
& { }^{4} T_{2 g}(P) \longrightarrow{ }^{4} T_{2 \mathrm{~g}}(F) \\
& { }^{4} T_{1 \mathrm{~g}}(P) \longrightarrow{ }^{4} T_{1 \mathrm{~g}}(F)
\end{aligned}
$$

characteristic of octahedral geometry ${ }^{26.27}$. By this way, UV-VIS spectroscopy plays an important part in the detection and structure determination of complex compounds.

However, there are many other technique such as Dipole moment, Xray diffraction, NMR (nuclear magnetic resonance), ESR (electron spin resonance) etc. which also furnish Valuable information on the structure of the complex compounds.

### 1.7 LIX-84 AS EXTRACTANT/LIGAND:

The high molecular weight hydroxyl amines and their suitable derivatives, which are being widely used as solvent extracting reagents all over the world in hydrometallurgy are known as LIX reagents. However, various $\alpha$ or $\beta$-hydroxy oximes (or phenolic oximes) and their derivatives Bdiketones and quinoline sulphonamides are included in this class of metal extractants.

Although the popular LIX reagemts are $\beta$-hydroxy oxime but LIX 84 is a 2-hydroxy-5-nonylacetophonone oxime. It is a low viscosity (sp.gravity O.90) water insoluble 2-hydroxy -5-nonylacetophenone oxime having a molecular weight of 277 with the structure given below.


LIX-84 is a hydrogen ion donor and hence it forms a uninegative ion which acts as a 1 igand informing the complexes with metal ions. Like other chelating metal extractants, it operates on a similar hydrogen ion cycle as,
$[\mathrm{nRH}]_{\text {org }}+\left[\mathrm{M}^{\mathrm{n}+}\right]_{\mathrm{aq}} \leftrightharpoons\left[\mathrm{R}_{\mathrm{n}} \mathrm{M}\right]_{\text {org }}+\left[\mathrm{nH}^{+]}\right]_{\mathrm{aq}}$ lacking in one- $\alpha$-hydrogen, the LIX 84 complexes of transition metals are not expected to undergo polymerisation in the solid state. The $1: 3$ dicarbonyl compound, LIX 84 can be synthesized condensation (aldehydes or ketones).

Although LIX 84 has been used as an extractant for copper (II) from aqueous ammoniacal solutions but on report on the structure of its extracted complex with Cu (II) or any other metal ion is available.

Some observed extraction properties ${ }^{28}$ of LIX 84 are outlined below.
(I) LIX 84 can be used neat without further kerosene dilution and still exhibit excellent phase separation properties. Phase separation is very rapid even at $100 \%$ concentrations.
(II) It does not load any free ammonia nor a copper-ammine complex.
(III) LIX 84 has improved selectivity for Zr (IV) over Ag ( I ) and Cd (II).
(IV) It can be stripped of its complex with an aqueous solution having very low acid concentration and for this only one stripping stage is necessary.
(V) Extraction kinetics of LIX84 are very rapid (10 min) giving almost complete extraction. Moreover, LIX 84 has been found to be completely miscible in kerosene, ethanol and other usual organic and inorganic solvents except methanol. Its metal complexes are insoluble in ethanol and water.
CHAPTER-2

### 2.1 LITERATURE REVIEWS:

The survey of literature reveals that very little work have been done on the LIX ${ }^{\text {® }} 84$ reagent. However, much work of extraction and isolation of the extracted complexes of the transition metal ions has been made with other chelating LIX. reagents, their mixtures and the kelex reagents have been published The following summarizes the main reports on the use of LIX 84 reagent and other LIX reagents for different metal ions.

Law, Henrey H., Tierney, vicki ${ }^{29}$, studied, "Direct recycling of copper from etching to platting. "In the manufacture of printed circuit boards, the copper dissolved during etching can be used to replace the Cu needed for plating using a liq. ion-exchange agent LIX 84 Extraction and stripping studies demonstrated the feasibility. The quality of the etching was not affected by the presence of the ion-exchange agent Good-quality deposits were plated from the stripping solution with a current efficiency of $100 \%$
M.S. Islam and M.A.Salam ${ }^{30}$ have prepared and characterized some complexes of first transition metal ions, $\mathrm{Ti}(\mathrm{III}), \mathrm{V}(\mathrm{II}), \mathrm{Cr}(\mathrm{III}), \mathrm{Mn}(\mathrm{II}), \mathrm{Fe}(\mathrm{II})$, Fe (III), Co (II), $\mathrm{Ni}(\mathrm{II}), \mathrm{Cu}(\mathrm{II}), \mathrm{Cu}(\mathrm{I})$ and $\mathrm{Zn}(\mathrm{II})$ with LIX 54 (phenyl alkyl beta-diketone). Their characterization has been made by elemental analyses; conductivity and magnetic measurements, infrared and electronic spectral studies. The solubilities of the prepared complexes are investigated in various organic solvents. The decomposition temperatures are also measured. All these data have not yet been published

By the survey of existing literature it is evident that this reagent is related to the various types of the treatment of liver diseases. It can be used as a dental artifact. Being an ingredient of a composition of sintered
porcelain product, LLX 84 may be also used as a dental material also LLX 84 reagent seems to be used in diabetes mellitus, and tumor growth stimulation. It is related to the transdermal pharmaceuticals to promote absorption. This reagent could be used for the prevention of treatment of gastritis.

The LIX 84 reagent has brain uptake property. So it acts as an agent of blood brain barrier. $\beta$-Adrenergic blockade restores glucose's anti ketogenic activity after exercise in carbohydrate- depleted athletes. Early T cell differentiated chronic mycloyed leukemia blast crisis with rearrangement of the break point cluster region but not of the T cell receptor B-chain genes.

This LIX 84 reagent may have industrial importance, too. There is a correlation between the oximes sensory and gas- chromatographic measurements on grape fruit juice volatiles. It' may have relation with stored -croude-oil sludge components identified. LIX 84 reagent can be used for some chromatographic investigation. It may act as an organic Pollutant. Oxime may have in cretaceous black shales. It also has some bio degradation in waste water treatment, The LIX 84 reagent acts as dental linear contg, It has also air pollutants effect. The transition metal complexes of LIX 84 reagent have some special type of physical and chemical properties. Some have coffee roasted aroma, some of them have cooked beef aroma; marasmius oreades fresh and dried mushroom aroma, pharmaceutical transdermal gels can be prepared by the metal complexes of the reagent. Most of the LLX84 complexes of transition metal elements may be used as corrosion inhibitors. These possess an excellent antirust and antifoaming properties. Cobait (II) loads and strips very quickly but cobalt (III) does not
load on LIX84 ${ }^{31,32}$ for the extraction of copper from ammonia leach solutions. LIX 64 N and other extractants with similar organic structures, have certain disadvantage ${ }^{33,}$
(a) They are limited in copper loading capacity such that high organic to aqueous flow ratios are required with large solvent extraction plants and associated capital investment costs:
(b) Some reagents will co-extract up to one mole of $\mathrm{NH}_{3}$ per mole of copper which results in excessive loaded organic washing problem and subsequent wash effluent treatment, or disposal; and
(c) Other cheating extractants loads $\mathrm{H}_{2} \mathrm{SO}_{4}$ in the stripping circuit which would also demand another washing stage. However, LIX84 does not include the disadvantages cited above, but does offer the following positive advantages;
(I) LIX84 can be used neat without further kerosene dilution and still exhibit excellent phase separation properties. Phase separation is very rapid even at $100 \%$ concentrations.
(II) Used neat, LEX84 can extract up to $40 \mathrm{gdm}-{ }^{3} \mathrm{cu}$ from an aqueous solution containing $30 \mathrm{gdm}^{-3} \mathrm{NH} 3$ thus eliminating or reducing the need for aqueous recycles in the extraction circuit.
(III) LIX84 does not load any free ammonia, nor a copper-ammine complex.
(IV) LLX84 has improved selectivity for copper over Cd and Zr .
(V) LIX84 can stripped with as 25 gdm. $\mathrm{H}_{2} \mathrm{SO}_{4}$. At greater acid strengths, 80 or $100 \mathrm{gdm}^{-3} \mathrm{H}_{2} \mathrm{SO}_{4}$ only one stripping stage is necessary.
(VI) Extraction kinetics in very rapid rapid and sssentially complete extraction is obtained in 60 second (90\%).

Thus LIX84 is a relatively weak copper extractnt which will load up to $(4.75-4.80 / \mathrm{ICu}) \mathrm{Cu}$ and can be used with no further dilution. Because it is a weak copper extractant, LIX84 can be striped with an aqueous solution whose equilibrium acid concentration is low but whose equilibrium copper concentration is high. LIX84 has been reported to be completely miscible in the solvents, acetone, toluene, isopropyl alcohol, ethanol (absolute), isodecanal, methylene chloride, cyclohexane, hexane and Kermac 470B (highly purified kerosene) to the extent $25 \mathrm{w} / \mathrm{v} \%$ However, it is almost insoluble in methanol.
D.P. Graddon and E.C. Watton have reported some adducts of copper (II) $\beta$-diketone chelates with heterocyclic bases ${ }^{33}$. Equilibrium constants have been determined at room temperature for the $1: 1$-addition of heterocyclic bases to copper (II) $\beta$-diketone chelates in three solvents, Irrespective of changes in the other two variables, the stability of the adducts increases in the order:
(a) 2:6: diethyl-pyridine<2:methyl-pyridine $<4$ : methl-pyridine.
(b) acetylacetonate<benzoylacetonate<ethyl-acetoacetate.
(c) chloroform<acetone<benzene.

Variations of stability of the adducts due to changing bases are attributed in part to changes in donor power and in part to steric effects. Variations from one copper compound to another are related to the residual positive on the copper atoms of the chelates and solvent affects are thought to be due mainly to changes in the salvation energies of the bases.

The addition compounds have been isolated by crystallization of the $\beta$ diketonates from pyridine or 4-methyl pyridine. Infrared and vesible region absorption spectra of five-co-rodinated structure for all of these compounds, indicative the formation of $1: 2$ adducts of bis (ethyl acetoacetato) copper (II) having the additional molecule of heterocylic bade free in the crystal lattice. The magnetic properties of these adducts are normal,the moment of the copper atom be $1.81-1.85 \beta . \mathrm{M}$
P.A.O' Hara, M.P.Bohrer ${ }^{34}$ worked on supported liquid membranes for copper transport. A carrier medicated liquid membrane process to printed wiring board shop as studied. Cu is transported from the alk. etching solution to the acid plation bath through a membrane consisting of a solution of a $\beta$-diketone in kerosene. A cu flux of $3.2 \mathrm{~g} / \mathrm{cm}^{2}$ was obrtained with a $50 \%$ solution $\beta$-diketone (LIX 54-100) supported by a thin ( $25 \mu \mathrm{~m}$ ) microporous polypropylene film. The effect of process variables inclding aq. stirring rate, reactant concentrations and membrane thickness indicates that diffusion of the Cu carrier complex across the membrane is the permeation rate-controlling process at sufficiently high stirring rates and Cu feed concentration. The equilibrium distribution relation between the feed solution and the membrane is nonlinear for carrier concentrations $>40 \%$. Independent measurements using. well-characterized, track etched membranes qual. verified the inverse relation between concn. and diffusivety of the Cu-carrier complex, but are not accurate enough to quart. support the simple membrane diffusion model. The selectivity of the membrane for Cu is $>100$ with respect to $\mathrm{Ni}, \mathrm{Zn}$ and Sn .

The preparation and characterization of several complexes of first transition metal ions ( $\mathrm{V}, \mathrm{Cr}, \mathrm{Mn}, \mathrm{Co}, \mathrm{Ni}$ and Zn ) with LLX 34 was made by M.S. Islam and M.A. Islam ${ }^{30}$. They characterized the prepared complexes by comductivity, soluvility, infrared spectra, electronic spectroscopic studies and elemental analysis.
M.S.lslam et al ${ }^{37}$ have prepared and characterized several new complexes of $\mathrm{Cu}(\mathrm{II}), \mathrm{Zr}(\mathrm{IV}), \mathrm{Ag}(\mathrm{I}), \mathrm{Cd}(\mathrm{II})$ and $\mathrm{Hg}(\mathrm{II})$ with- 8 Alkyl sulfonamido-quinoline (LLX 34; Abbreviated as HL ) and bis $-(2,4,4-$ trimethylpentyl) phosphonic acid (cyanes 272; HL). Their characterizatioon was made by elemental analysis, conductivity and magnetic measurements, infrared spectra and electronic studies. The solubilities of the complexes in various. solvent and decomposition temperatures have also been measured. The stereochemistry of the complexes has been discussed on the basis of the experimental data.

The extraction of Cr (VI) from its aqueous acidic sulfate solution by Cyanex-272 in kerosene has been investigated ${ }^{38}$ as a function of the aqueous phase acidity, extract ant concentration and temperatures: spectrophotometric technique was employed for the estimation of Cr (VI). The extraction ratio increases with increasing concentration of the extractant, aqueous phase acidity and temperatures. Chromium(VI) extraction by cyanex-272 is moderately slow requiring about 30 attain equilibrium cyanex-272 forms a $1: 1$ solvated complex with the metal ion in its dilute solution. From temperature dependence data, the apparent enthalpy change for the extraction reaction has been calculated to be $\mathrm{AH}=23.65 \mathrm{Kj}$ ) $\mathrm{mol}^{-1}$ The probable composition of the complex of $\mathrm{Cr}(\mathrm{VI})$ with Cyanex-272
at high loading has also been suggested from $\mathbb{R}$ spectra, chromium(VI) phosphorus ratio and molecular weight data. The mechanism of extraction at high loading is different from that at low loading.

The distribution of Ni (II) between benzene solutions of LIX 34 its aqueous acidic sulphate as well as ammoniacal sulphate solutions has been in estimated as a function of the extractable metal ion, sulphate acetate hydrogon ion and ammonia concentration. of the aqueous phase, extractant concentration,temperature and the nature of the diluents ${ }^{29}$ ratio has been found to increase with increasing concentrtion of the extractant, acetate and temperature of extraction, and with decreasing acidity and sulphate concentration. Nickel extraction has been found to be quantitative in ammoniacal sulphate medium but very poor in acidic sulphate medium. LIX 34 forms a $1: 1$ complex with the metal ion ${ }^{19}$ in its dilute solution. From the temperature dependence data, the apparent enthalpy change for the extraction reaction has been calculated to be $\Delta \mathrm{H}=3.43 \mathrm{kj} \mathrm{mol}^{-1}$. The extraction of $\mathrm{NI}(I I)$ depends on the nature of the diluent.s Toluene may be used as diluent in place of benzene.

The solvent extractive separation of Co (II) from Ni (II) in acidic sulphate-acetate medium with 8 -alkarylsulphonamidoquinoline ( $\mathrm{LI}<34$ ) has been investigated spectrophotometrically ${ }^{40}$. The effects of operaing variable such as contact time, temperature, concentrations of LIX34, metal ions; and anions $\left(\mathrm{SO}_{4}=, \mathrm{Ac}\right)$ on the extraction and separation behaviors have been studied. Experimental data suggest that the separation of Co (II) from $\mathrm{Ni}(\mathrm{II})$-is possible with short contact times of 2 minutes under setected condition.

The probable structures of the metal chalets of $\mathrm{Co}(\mathrm{II}), \mathrm{Ni}(\mathrm{II})$ and $\mathrm{Cu}(\mathrm{II})$ with 8-alkarylsulfonamido-quinoline (LIX34 or HIR) have been suggested by the sulphur-metal atom ratio (S/M), nitrogen-metal atom ratio ( $\mathrm{N} / \mathrm{M}$ ), molecular weight, magnetic moment and infrared spectral data of the complexes ${ }^{41}$. As per data both Ni (II) and Cu (II) form 1:2 solid octahedral cheaters; but Co (II) may form semi-solid solvated square planar as well as octahedral complex.

The partition of $\mathrm{Co}(\mathrm{II})$ between its aqueous acidic sulphate-acetate solutions of LDX 34 in benzene has been investigated as function of the extractable metal ion, $\mathrm{H}^{+}$; sulphate, acetate concentrations of the aqueous phase, extractant concentration, temperature and the nature of the diluents ${ }^{41}$. The extraction ratio incerases with increasing of the extractant, acetate and temperature and with. decreasing acidity and sulphate concentration. Cobalt (II) extraction has been found to be quantitative and moderately fast ${ }^{42}$. A spectrophotometric technique was employed for the estimation of Co(II). LIX 34 form a $1: 1$ complex with the metal ion in its dilute solution. From the temperature dependene data, the apparent enthalpy change for the extraction reaction has been calculated to be the extraction of Co (II) depends on the nature of the diluent. n-hexane may be used as diluent in place of benzene.

Some studies on complexes with chelatiing agents other than LIX reagent are mentioned below. the synthesis and characterization of some metal chelates of Ti (III), V (II) $, \mathrm{Mn}(\mathrm{II}), \mathrm{Fe}$ (III), $\mathrm{Co}(\mathrm{II}), \mathrm{Ni}(\mathrm{II})$ and Zn (II) with bis 1,4,4-trimethyl-pentyl) phosphinic acid are reported by M.S Islam and M.K.H Talukdar ${ }^{43}$. They characterized the complexes by elemental analysis,
conductivity and magnetic measurements and infrared and electronic spectral studies. The solubilities of the complexes in various solvents are also measured.

### 2.2 SCOPE OF THE PRESENT INVESTIGATION:

Complexes of transition metals with different known legends are of great importance, because many of them are efficient stoichiometric and catalytic reagents. These complexes have been extensively used in the therapeutic, analytical, biological and industrial fields and in co-ordination chemistry. As such, it is essential to develop new chelating agents for both therapeutic and analytical usages, and to understand the roles of transition metal ions in various properties extraction reagents ${ }^{44,45}$, by solvent extraction method are of recent interest. These complexes (usually) cheaters) displaying distinctive co-ordination chemistry ${ }^{46}$ and characteristic physical properties, may have industrial as well as biological significance. But complexes of heavier transition metals are rare. The extraction of common metals viz. Zr (IV), Ag (I) Cd (II), with various reagents has been studied frequently and their separation, especially from waste materials, low grade ores and sea nodule. The chemistry and mechanism of their extraction also has been the subjects of intense research. However, no report on such studies on complexes of lighter and heavier transition metal ions with the new extracting reagent LIX84 is available.

The transition metal complexes with legends are interesting from theoretical standpoint to broaden the present state of knowledge. With the rapid advancement of science and technology, the easily available sources of the useful transition metal ions are going to be exhausted and so substitute
sources for them will be sought for, as is now the case of cobalt. As a result, the now un important ores (with too much contaminants) will certainly become increasingly important, for which usual methods of extraction do not work. Therefore, the study of completing properties of lirst transition series of metal ions with LIX84 will be of much value for extracting the metal ions from the sea nodules, low grade ores and the metal wastes employing the solvent extraction method economically.

Under the circumstances, it is thought appropriate, in view of the various importances of the transition metal complexes, to investigate complexing behaviour of LIX84 as a uninegative bidentate ligand. As such a programme is, undertaken to synthesize several new complexes of L[X84 with both lighter and heaviour transition metal ions, and characterize them by some physical methods of chemical analysis in order to get information about their compositions and structures as well as some other physical properties. The derived will be helpful for solvent extraction of these metal ions.

### 3.1 AIM OF THE PRESENT INVESTIGATION:

The main aim or objective of this investigation is the preparation of some new second and third transition metal ions complexes with LIX84 acting as a ligand, their characterization based on some physical measurements and also solution behaviour. The Zr (IV), Ag (I) and Cd (II) ions are selected for their greater stabilities than their other oxidation states. The prepared complexes are proposed to be characterized on the basis of determination of molecular weight, IR and electronic spectra, magnetic and conductance measurements along with the melting point measurement.

Since LIX84 is a chelating extractant, the solubility of the chelate that is formed in different solvents in an important factor in the efficiency of solvent extraction. The study of the solubilities and electronic spectral of the prepared complexes in different solvents is also proposed to be made in order to throw further light on the structure of the complexes in solution.

The cadmium is in reletively low abundance in Nature (of the order of $10^{-6}$ of the earth's crust from Cd ), but has long been known because it is easily obtained from it's ores.

Cadmium minerals are scare, but as a result of its chemical similarity to zinc occurs by isomophous replacement in almost all zinc ores. This element directly reacts with halogens and with nonmetals sush as $\mathrm{S}, \mathrm{Se}$ and Pb and it Cadnium forms many alloys,s such as has being of technial importance. The oxide ( CdO ) is formed on burning the metals in air or by pyrolysis of the carbonate or nitrates; thus of oxide smokes can be obtained by combustion of the alkyls, those of cadmium being excendingly toxic.

Aquous solutions of cadmium halides appear, superficially to be in completely dissociated, that is, to be weak electrolytes. The formation constants
for halides and other ligands(oxime) differ widely and the main feature is that they are many orders of magnitude smaller than those for $\mathrm{Hg}^{2+}$. The cadmium ones are most stable and the ion $\left[\mathrm{CdCl}_{5}\right]^{3-}$ can also be formed $\mathrm{Cd}^{2+}$ is bound more strongly to $\mathrm{Cl}, \mathrm{S}$ and oxime ligands.Cadmium ${ }^{113} \mathrm{Cd} \mathrm{nmr}$ has provided useful biological systems but shifts are very sensitive to concentration and counter ions.

Again, silver and gold are widely distributed in nature. They occurs as metals and also in numerous sulfides ores usually accompanined by sulfides of $\mathrm{Fe}, \mathrm{Cu}, \mathrm{Ni}$ etc. The main sources are Africa and the USSR.Silver also occur as horn silver( AgCl ). After floaing or other concentration processes, the crucial chemical steps are cyanide loaching and zinc precipitation.

Silver is normally puriffied by electrolysis. It is a white,yellow metal and malleable metal with the highest known electrical and thermal conductivity.

However it is chemically less reactive than copper, except towared sulfur and hydrogen sulfide which rapidly blacken silver surfaces silver atoms that does not aggregate but are trappein a macrocyclic poly- nitrogen ligand are obtained by photochemicals or etectrochemical reductions of the $\mathrm{Ag}^{+}$complex. There is a great variety of silver complexes which exist in solution or in the solid state.Since the most stable $\mathrm{Ag}^{+}$complexes have the linear structure $\mathrm{L}-\mathrm{Ag}-\mathrm{L}^{+}$, chelating ligands cannot form such simple ions, hence tend to give polynuclear complex ions. Slver chloride can be obtained as rather tough sheets that are transparent over much of the infrared region and have been used for cell materials. Moreover, silver chloride and bromide are light sensitively studied because of their importance in photography.

On the other hand, zirconium occurs widely over the earth's crust but is not as very concentrated deposit. The major minerals are baddeleyite, a form of
$\mathrm{ZrO}_{2}$, and zircon $\left(\mathrm{ZrSiO}_{4}\right)$. The chemical similarity, of zirconium and hafinium are well examplified in their geochemistry for hafinium is found in nature in all zirconium minerals in the range of fractions of a percent of the zirconium content. Separation of the two elements are extremely difficult, even more so than for adjacent lanthanides, but it can now be acoomplished in salts factory by ion-exchange or solvent extraction fractonation methods.

However, zirconium metal is hard and corrosion resistant, resembling stanless steel in appearance. It will burn in air at high temperatures, reacting with a anumber of compounds called"Zirconates" may be made by combining oxides, hydroxides,nitrates,oxime and so on, of other metals with similar zirconium compounds and fring the mixtures at 1000 to $2500{ }^{\circ}$. These, like their titanimm analogue,are mived metal oxides,there are no discrete zroonate ions known.

Adition of hydroxide to zirconium(iv) solutions causes the precipitation of white gelatineous $\mathrm{ZrO}_{2} \cdot \mathrm{nH}_{2} \mathrm{O}$, where the content is variable; no true hydroside exists. On strong heating, this hydrated oxide gives hard, white, iinsoluble $\mathrm{ZrO}_{2}$. This has an extreamly high meling point $\left(2700^{\circ} \mathrm{C}\right.$ ), exceptional resitance to attack by both acids and allaalis, and good macanical properties;it is used for crucibles and furnace cores.Zrconium dioside in its monoclinic(bad deleyite)form.

Aquous chemistry of zirconium aqua ions of low and mediun valence state are not in general well defined or important for any of the heavier transition element. In Bangladesh zirconium is available in the soil of Cox's Bazar, the study for elemantal analys, quality and quantity of zirconium will provide to investigate zirconium metal in the soil of Cox's Bazar.

Therefore, this study of complaxing(extracting) beahaviour of three above mentioned three metals $\mathrm{Ag}, \mathrm{Cd}$ and Zr will be helpful for their solvent extraction.

### 3.2 PREPARATION OF THE COMPLEXES:

## (I). Preparation of $\left[\left(\mathrm{ZrO} \mathrm{O}_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}_{2}\right]\right.\right.$ complex:

LIX84 ( $2.84 \mathrm{~g} ; 0.015 \mathrm{~mol}$ ) was dissolved in n - haxane ( 40 mL ) in a beaker and $\mathrm{KOH}(0.56 \mathrm{~g}$; O .01 mol ) was dissolved in water $(20 \mathrm{~mL})$ in another beaker. These two solutions were mixed together in a 250 mL beaker. The mixture was stirred, heated on a water-bath for two tours and the resulting solution allowed to stand at room temperature for several hours.

A solution of $\mathrm{Zr}\left(\mathrm{NO}_{3}\right)_{4} \cdot 6 \mathrm{H}_{2} \mathrm{O}(2.236 \mathrm{~g} ; 0.005 \mathrm{~mol})$ in water $(50 \mathrm{~mL})$ was then added slowly to the above solution of the ligand. The resulting immiscble mixture was stirred well with the help of a magnetic stirrer and warmed on the water-bath. The deep coppy colour precipitate appeared was filtered off, washed several times with alcohol and water, and finally dried in vacuo over anhydrous $\mathrm{CaCl}_{2}$.

## (II). Preparation of $[\mathrm{Ag}(\mathrm{L})]$ complex:

An aquos solution $(20 \mathrm{~mL})$ of $\mathrm{KOH}(0.28 \mathrm{~g} ; 0.005 \mathrm{~mol})$ was added solution $(30 \mathrm{ml})$ of LIX84 $(1.23 \mathrm{~g}, 0.005 \mathrm{~mol})$ in n -hexane in a 250 mL beakr. The mixture was stirred well and heated on a water-bath for 2 hours to reduce it to half its values. This solution was then allowed to stand at room temperature for several hours. In another beaker silver nitrate ( $0.849 \mathrm{~g} ; 00005 \mathrm{~mol}$ ) was dissolved in distilled water ( 40 mL ) by heat and was added to the above solution. This resulting immiscible mixture was stirred and the black precipitate found was collected by filtration, washed several times with alcohol and water. and finally dried in vacuo over anhydrous $\mathrm{CaCl}_{2}$.

## (ID). Preparation of $\left[\mathrm{Cl}(\mathrm{L})_{2} \mathrm{HL}\right]$ complex:

A solution of KOH ( $0.56 \mathrm{~g} ; 0.015 \mathrm{~mol}$ ) in was n -hexane solvent $(40 \mathrm{~mL})$ into a 250 mL beaker. This mixture was stirred and heated on a water-bath for one hour. It was then allowed to stand at room temperature for several hours and to this was added a solution of $\mathrm{CdCl}_{2} \cdot 2 \frac{1}{2} \mathrm{H}_{2} \mathrm{O}(1.0066 \mathrm{~g} ; 0.005 \mathrm{~mol})$ in water ( 30 mL ). The resulting mixture was immiscible and stired well with a glass rod at room temperature. The white precipitates formed was filtered off, washod well with ethanol, and distilled water (to remove KCl ). The product was air dried and finally dehydrated in a vacuo desicator over anhydrous $\mathrm{CaCl}_{2}$.


## EXPERIMENTAL TECHNIQUES

### 4.1 WEIGHING:

The weighing operation was performed with the help of a METTLER TOLEDO AB204 electronic balance.

### 4.2 MELTING POINT MEASUREMENT:

The melting or decomposition temperatures of all the prepared metalLIX84 complexes were observed with an AZ 6312 model Electrothermal Melting Point apparatus. It was, however, not possible to measure the melting points beyond $360^{\circ} \mathrm{C}$.

### 4.3 CONDUCTIVITY MEASUREMENTS:

The conductivity cell was cleansed several times with distilled water, rinsed with acetone and finally allowed to dry in air.

The conductance measurements of $10^{-3} \mathrm{M}$ solution of the complexes in benzene $\left(\mathrm{C}_{6} \mathrm{H}_{6}\right)$ or $\left(\mathrm{C}_{6} \mathrm{H}_{12}\right)$ were made at room temperature using a type CG 857 No. 71798 SCHOT GERATE GmbH Germany digital conductivity meter and a dip-type cell with platinized electrodes. The cell was calibrated using .001 N potassium chloride solution at the experimental temperature.

The molar conductivities were ( $\Lambda$ ) calculated using following equation;

$$
\begin{equation*}
\Lambda=1000 / \mathrm{c} \times \text { cell constant } \times \text { experimental conductivity. } \tag{i}
\end{equation*}
$$

where, $c$ is the concentration of complex solution in the $\mathrm{mol} \mathrm{dm}^{-3}$. The experimental conductivity was obtained by subtracting the conductance of $\left.) \mathrm{C}_{6} \mathrm{H}_{6}\right)$ or $\left(\mathrm{C}_{6} \mathrm{H}_{12}\right)$ from that of solution of solution of each complex in $\left(\mathrm{C}_{6} \mathrm{H}_{6}\right)$ or $\left(\mathrm{C}_{6} \mathrm{H}_{12}\right)$

### 4.4 MAGNETIC MEASUREMENTS:

Magnetic susceptibilities of the solid complexes were measured with the help of the SHER WOOD SCLENTIFIC Magnetic Susceptibitity Balance (Gouy method). The balance was calibrated using $\mathrm{HgCo}(\mathrm{SCN})_{4}$ and the calibration was checked using a sealed off sample of $\mathrm{MnCl}_{2}$ solution calculation of the corresponding mass susceptibilities were done using the equation ${ }^{47}$.

$$
\begin{equation*}
\chi_{g}=2.08 . l .\left(\mathbb{R}-\mathbb{R}_{0}\right) / 10^{9} . \mathrm{m} \tag{ii}
\end{equation*}
$$

where,
$\mathrm{m}=$ mass of sample in grammes
$1=$ length of the sample in centimeters
$R=$ observed reading for tube plus sample
and $\mathrm{Ro}=$ observed reading for the empty tube.
The calibration constant $\left(C_{B}\right), 2.086$ was evaluated using the relations ${ }^{47}, C=C_{B} / 10^{9}$ and $\mathrm{C}_{\mathrm{B}}=\mathrm{C}_{\text {tube }} /(\mathrm{R}-\mathrm{Ro})$ where C is the constant of proportionality related to the calibration constant of the balance. Then effective magnetic moments (spin only values, $\mu_{\text {eff }}$ ) were calculated and the number of unpaired electrons ( $n$ ) in the metal ions of the following
relation ${ }^{47}$. $\mu_{\text {eff }} /$ number of metal ions in a molecule of the complexes $=$ $\sqrt{n(n+2)}$

## Experimental Procedure:

(I) The zero knob of the magnetic susceptibility balance was turned until numerical display showed zero ( O 0 ) and calibration sample, HgCo $(\mathrm{SCN})_{4}$ was inserted into sample holder. It was then allowed to settle to give the numerical display ${ }^{48}$.
(II) Reading was recorded and calibration constant $C_{B}$ was calculated from the equation,
$\mathrm{C}_{\mathrm{B}}=\mathrm{C}_{\text {tube }} /(\mathrm{R}-\mathrm{Ro})$.

## Operation of the balance:

(a) The range knob was turned to the display reads (o00). The zero was adjusted on each side.
(b) The zero knob was adjusted until the display reads (ooo). The zero was adjusted on each side.
(c) An empty tube of known weight was placed into the tube guide and the reading Ro was taken
(d) The sample was packed, and the sample mass in grammes and the sample length, 1 in cm were noted. the packed sample tube was placed into the lube guide and the reading $R$ was taken. From the data, $\chi_{g}$ was calculated.

### 4.5 INTRARED SPECTRA:

The $\mathbb{R}$ spectra of all the solid metal-LLX84 complexes were run on a FTR Shimadzu Japan Model No-8400 FOURIER TRANSFORM INFRARED SPECTROPHOTOMLTER (as KBr pellets) in the range 400 to $4000 \mathrm{~cm}^{-1}$ However, the spectrum of liquid LIX884 was obtained directly as thin film in the same range of frequency ${ }^{49-51}$. Another set of $\mathbb{R}$ spectra were not also run on an infrared spectrophotometer,

Procedure: A requisite amount of complex was put in a mortar and thoroughly powdered with potassium bromide. This mixture was then transferred in a mini disc holder, and a KBr disc was made by hand press. This was then mounted in the sample cavity of the spectrophotometer ${ }^{52}$ and its spectrum was taken. On the other hand, one drop of LIX84 (liquid) was placed into a rectangular cell made of NaCl its $\mathbb{R}$ spectrum was then taken directly as thin film. From the spectra of complexes important vibration band frequencies of different groups were found out ${ }^{19}$.

### 4.6 ELECTRONIC SPECTRA:

Electronic spectra of the complexes were run on a Lambda-19 UV/VIS/NIR spectrohotometer (PERK) NELMERAMERICA NO. All the spectra were taken for dilute solutions of the complexes in yclohexane in the wave length of 200 to 800 nm . The electronic spectra of LIXS4 was also taken and it was compared with those of the complexes in order to find out the possible d-d transitions and charged transfer bands ${ }^{5 \cdot-55}$.

### 4.7 DETERMINATION OF MOLECULAR WEIGHT:

The cryoscopic method using a Beckmann apparatus has been used for determination of the molecular weights of the extracted complexes of $\mathrm{Zr}(\mathrm{IV}), \mathrm{Ag}$ (I) and $\mathrm{Cd}(\mathrm{II})$ with LIX84. This method gives more or less accurate when the molecular weight is small but discrepancies arise when the molecular weight is high. The accuracy obtainable with the Beckmann apparatus in stated to be methine approximately $5 \%$ of the molecular weight. The mojor source of error in the determination is the problem of super cooling of sample solution Pure solvents have little tendency to super cool, but their solutions are affected to a much larger extent; for this reason, the differential between freezing tube and cooling bath of the apparatus should be maintained at $1.5^{\circ}$ to $1.0^{\circ}$ for precies work. The determination of molecular weight by observation of the depression of the freezing point induced in a pure solvent by a known amount of the substance under investigation is based on the following equation; $\quad M=k_{f} \frac{1000 w}{G \Delta T_{1}}$ where,
$\mathrm{K}_{\mathrm{f}}=$ molar depression constant for solvent=5.12 for Benzene .
$\Delta T_{f}=$ experimentally observed depression of the freezing point of the solvent.
$\mathrm{w}=$ weight of the added solute taken in g .
$G=$ weight of the solvent taken in g.and
$M=$ molecular weight of the solute.

The Beckmann apparatus of (Arther H. Thomas company, philadel phia, U.S.A) was used for determination of the molecular weight of samples (i.e.M.-LIX84 chclates). by the freezing point depression of the solent benzene. For this, benzene was previously dehydrated with an hydrous. CaO and then distilled. The apparatus was at firs calibrated by determining the molecular weight of pure naphthalene (several times) using the same solvent, and the percentage of error was found to be 2 approximately.

In operation, a weighed amount of benzene was introduced. into the imer tube and the side neck was stopped. It was then cooled near to its freezing point $\left(5.5^{\circ} \mathrm{C}\right)$ by ice-water mixture placed in the cooling bath. To this ncarly freezing solvent, a Beckmann thermometer was inserted so that the lower part of the thermometer read on the middle of the scale of the thermometer. This was adjusted by trial, each time by taking. out the thermometer. Transferring the mercury from the bottom to the top or vice versa by the application of heat or cooling by the freezing mixture.

Once the temperature on the Beckmann thermometer was fixed and the freezing point of the solvent was recorded, the thermometer was removed from the solvent tube and kept immersing in a beaker containing the freezing mixture. A weighed quantity of solute (M-LIX84) complex was introduced carcfully into the inner freezing tube and allowed to dissolve in benzene The freezing point of the solution was determined with continuous vigorous stirring (avoiding super cooling) by the associated stirrer using the pre-set Beckmann thermometer. The difference of the two readings gave the depression of the freczing point for the given amount of the solute-solvent system. The experiment was repeated for each solute several times taking different amounts of the complexes. The molecular weights were then calculated by using the above formula.


## RESULTS AND DISCUSSION

The transition metal complexes of Zr (IV), Ag (I) and Cd (II) with the ligand LIX84 have been synthesized as described previously. These complexes are characterized on the basis of conductance, magnetic measurements, infrared, molecular weight and electronic spectral studies and their other physical properties such as per cent of solubilities.

### 5.1 Molecular weight:

The molecular weight of the prepared complexes were determined by using cryoscopic method. Benzene was used as the solvent. The values obtained for the compounds have been used to determine the molar susceptibility and molar conductance values.These are shown in Table-4.

### 5.2 Molar Conductance:

The molar conductance data of the prepared transition metal-LIX 84 complexes along with that of the solvents are given in Table-1. These measured electrical conductances of $10^{-3} \mathrm{M}$ solutions indicate that the complexes 1, 2 and 3 do not behave as 1:1 electrolytes in DMSO or cyclohexane. The zero conductance values for all the prepared complexes 1, 2 and 3 show that they are non-electrolytes. However, it was not possible to measure the conductance values of the complexes in DMSO or cyclohexane as they were found insoluble in DMSO.

### 5.3 Magnetic Moments:

Molar susceptibilities of all the transition metal-LIX84 complexes were measured as stated as stated before and the data is shown in Table-4,
which also contains the calculated magnetic moments (spin only values) and the number of unpaired electron per metal atom. It is found that the complexes No. 2 and $\mathbf{3}$ are diamagnetic. The diamagnetism is supported by its measured small negative magnetic susceptibility values and also observed repulsion by the applied magnetic field. Complex 1 is found to be paramagnetic in nature having $\mu_{\text {eff }}$ value 5.30 B.M respectively. The complex 1 appears to be paramagnetic with four unpaired electrons but it may be a binuclear compound having Zr atoms with four unpaired electrons in their inner orbital configuration with high spin states respectively. Using the magnetic data, the stereochemistry of the complexes may be suggested as follows. The Zr (IV) ion $4 \mathrm{~d}^{10}$ electrons. But as the magnetic moment measurement of the complex 1 indicates the presence of four unpaired electrons for each Zr atom, it is suggested that this six coordinated paramagnetic Zr -LLX84 complex is octahedral with $4 \mathrm{~d}^{2} 5 \mathrm{~s} 5 \mathrm{p}^{3}$ hybridization as depicted below.

$\mathrm{d}^{2} \mathrm{sp}^{3}$ hybridization
The two coordinate silver complex 2 is also diamagnetic with $\mathrm{Ag}(\mathrm{I})$ ion having $d^{10}$ configuration and therefore its structure is probably linear with 5 s 5 p hybridization. Finally, the Cd (II) ion has $4 \mathrm{~d}^{10}$ configuration and hence the four-coordinated outerorbital diamagnetic cadmium complex 3
will be octahedral with $5 s 5 p^{3} 5 \mathrm{~d}^{2}$ hyridization. This hybridization may be represented by the following schematic diagram.


### 5.4 Per cent solumilinies:

The solubilities of the prepared transition metal complexes with LIX 84 were examined in different solvents ${ }^{56-57}$. Their solubility and percent solubility data are presented in table 2 and 3 respectively. The complexes are found to be insoluble in polar solvents, water, ethanol, acetone and sparingly soluble in other organic solvents. `

### 5.5 Infrared spectra:

Frequencies for LIX 84 are observed and M-LIX 84 Complexes are listed in the following Table- (5-6). The infrared spectrum of LIX84 shows Vibrational bands for-OH free ( $3379.1-3350 \mathrm{~cm}^{-1}$ ), $\mathrm{C}-\mathrm{H}$ streching (2968.6 $\mathrm{cm}^{-1}$ ) , $>\mathrm{C}=\mathrm{N}$....... streching $-\mathrm{CH}_{2}$-bending aromatic, $\mathrm{C}=\mathrm{C}$ streching (1461.9 $\mathrm{cm}^{-1}$ ), $\mathrm{N}-\mathrm{O}-$ bending band at ( $1373.2-13.65 \mathrm{~cm}^{-1}$ ). Medium absorption near ( $1022.2 \mathrm{~cm}^{-1}$ ) is caused by wagging of the group $-\mathrm{CH}_{3}$. for the interpretation of various absorption band found, the $\mathbb{R}$ spectrum of all the metal-LIX84 complexes have been compared with that of LIX84.

The presence of water molecule in coordination sphere ${ }^{58}$ is indicated by the very absorption band around $3583.5-3382.9 \mathrm{~cm}^{-1}$ in the IR spectrum of the complex 1.

Again the $\mathbb{R}$ spectrum of complexes 1 exhibits characteristic band at $3382.9 \mathrm{~cm}^{-1}$, (intra hydrozen) inside the coordination sphere. The $\mathbb{R}$ spectrum of complex 1 shows an indication band at $1000-1072.3 \mathrm{~cm}^{-1}$, which indicate of-OH bridging ${ }^{19}$ with metal to metal ions. However, the complexes have also bands in $1203.5-1257.5 \mathrm{~cm}^{-1}$ shifted towards lower frequencies. This is also an indication of oxygen atom of -OH group with the metal atom.

Moreover (M-O) band ${ }^{19}$ at $640.3-412.7 \mathrm{~cm}^{-1}$ is indicative of $\mathrm{M}-\mathrm{O}$ band formation in all the complexes. All complexes show $V(\mathrm{M}-\mathrm{N})$ band at $412.7-447.5 \mathrm{~cm}^{-1}$ which indicates the metal - nitrogen coordination band formation.

### 5.6 ELECTRONIC SPECTRA:

Information on probable geometries of the prepared matel-LIX84 Complexes may be obtained from their electronic spectra and that of LIX84. These spectra are shown in Fig. (5-8). The selected UV/VIS absorption band maximum are given in Table-(7-11). The absorption spectrum of 2-hydroxy5 -nonyl acetophenone oxime (i.e.LIX84) shows two bands at 251 nm and 350 nm which areas singed respectively ${ }^{51}$ to the $\mathrm{n} \rightarrow \pi^{*}$ transition of $\mathrm{C}-\mathrm{O}$ and $>\mathrm{N}-\mathrm{O}$ groups in cyclohaxane solvent. All the complexes have only one absorption maximum within $300-600 \mathrm{~nm}$ in their spectra of their solution in cyclohaxane. However, the absorption band maximum of the octahedral

Cadmium ( II ) Complex at about 300 nm ( $300-450 \mathrm{~nm}$ ) may be assigned to the $n \rightarrow \pi^{*}$ transition of $>\mathrm{N}-\mathrm{O}$ group.

The absorption band of octahedral binuclear Zirconium complex of LIX84 in the higher wavelengh i.e. at about 400 nm (300-500 num may be due to d-d transition assignable to $4 \mathrm{~A}_{2 g}(\mathrm{~F}) \leftarrow 4 \mathrm{~T}_{\mathrm{ig}}$. The complex 2 exhibic only the charge transfer bands ${ }^{51}$ because of the $\mathrm{d}^{10}$ configuration. In This spin-paired complex, charge transfer bands may obscure the ligand field bands and hence it does not show d-d transition ${ }^{51}$.

$\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$, Octahedral

$[\mathrm{Ag}(\mathrm{L})]$, Linear

$\left[\mathrm{Cd}(\mathrm{L})_{2} . \mathrm{HL}\right]$, Tetrahedral
Table-1

| No | Compound | Colour | Yield | Decompose Tem $\left.{ }^{\circ} \mathrm{C}\right)$ | Molar conductance <br> olm ${ }^{-1} \mathrm{~cm}^{2} \mathrm{~mol}^{-1}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ | Confee | 2.50 | 230 | 0 |
| 2 | $[\mathrm{Ag}(\mathrm{L})]$ | Black | 1.50 | 200 | 0 |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} . H L\right]$ | White | 3.69 | 280 | 0 |

Colour, yield, decomposition temperature and molar conductance of the complexes:
Table-2

| CompNo. | Complexes | DMSO | DMF | $\mathrm{C}_{2} \mathrm{H}_{2} \mathrm{Cl}_{2}$ | $\mathrm{C}_{6} \mathrm{H}_{6}$ | $\mathrm{OCl}_{4}$ | $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{OH}$ | $\mathrm{CHCH}_{3}$ | $\mathrm{C}_{6} \mathrm{H}_{12}$ | $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{CH}_{3} \mathrm{COCH}_{3}$ | $\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{CH}_{3}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | x |
| 2 | $[A g(L)]$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | $\times$ |
| 3 | $\left[\mathrm{Cd}(L)_{2} . H L\right]$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | x | $\checkmark$ | $\times$ | $\times$ | $\times$ |
| Ligand LIX 84 | $\mathrm{C}_{17} \mathrm{H}_{27} \mathrm{NO}_{2}$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | $\times$ | $\checkmark$ | $\times$ | $\times$ | $\times$ |

## Table 3

| Compound | DMSO | DMF | $\mathrm{C}_{3} \mathrm{H}_{5} \mathrm{CH}_{3}$ | $\mathrm{C}_{6} \mathrm{H}_{12}$ | $\mathrm{CHCl}_{3}$ | $\mathrm{CH}_{2} \mathrm{Cl}_{2}$ |
| :---: | :---: | :---: | :---: | :---: | :--- | :--- |
| $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ | - | - | 3.35 | 4.65 | 4.60 | 3.75 |
| $[\mathrm{Ag}(\mathrm{L})]$ | - | - | 3.8 | 3.65 | 3.00 | 2.85 |
| $\left[\mathrm{Cd}(\mathrm{L})_{2} \mathrm{HL}\right]$ | - | - | 6.00 | 6.50 | 4.55 | 3.50 |

-,means insoluble.
Table-4
Detailed magnetic data of the complexes $(T=303 \mathrm{~K})$ :

| $\begin{gathered} \text { Complexes } \\ \text { and Extractant } \end{gathered}$ | Length of the sample (l) in cm | Mass weight of the complex, in (m) $g$ | Susceptibil ity of the empty tube, $\mathrm{R}_{\mathrm{o}}$ | Susceptibility of the sample with tube, R | Mass <br> Suscep <br> tibility, <br> $x_{8} \times$ <br> $10^{-6}$ in <br> C.G.S <br> unit | Molecular <br> weight <br> M | $\begin{gathered} \text { Molar } \\ \text { Susceptib } \\ \text { ility, } \\ \chi_{M} \times 10^{-3} \\ \text { in } \\ \text { C.G.S } \\ \text { unit } \end{gathered}$ | Dia- magnetic correction, $10^{-6}$ in C.G.S unit | corr ${\underset{\chi}{M}}_{\text {corr }} \times 10^{-3}$ <br> in <br> C.G.S <br> unit | $\begin{gathered} \mu_{\mathrm{eff}} \\ \text { in } \\ \text { B.M } \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ | 3.2 | . 109 | -48 | +240 | 17.637 | 1277.44 | 22.53 | $-51238$ | 11.321 | $\begin{gathered} 5.30 \\ (\text { para }) \end{gathered}$ |
| [ $\mathrm{Ag}(\mathrm{L})$ ] | 2.9 | . 058 | -47 | -62 | $-1.564$ | 512 | -. 8002 | -193.63 | -. 9938 | $\begin{aligned} & -1.55 \\ & \text { (diam) } \end{aligned}$ |
| [Cd(L) 2 HL$]$ | 2.90 | . 044 | -50 | -64 | -1.924 | 941.24 | -1.8109 | -512.38 | -1.30 | $-1.78$ (diam) |
| $\mathrm{C}_{17} \mathrm{H}_{27} \mathrm{NO}_{2}$ |  |  |  |  |  |  |  |  |  |  |

Table - 7

## Electronic spectral data of the prepared complexes with assignment.

| No. | Compounds | Observed spectral band maxima in, nm | Significant spectral bands, $\mathrm{cm}{ }^{-1}$ |
| :---: | :---: | :---: | :---: |
|  | LLX\&4 | 251,350 | 21711,30275 |
| 1 | $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ | 400 | 34601 |
| 2 | $[\mathrm{Ag}(\mathrm{L})]$ | 290 | 25085 |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} \mathrm{HL}\right]$ | 300 | 25950 |

Table-8
Electronic spectral data of LIX 84



Ordinate



|  |  <br>  |
| :---: | :---: |
|  |  |
|  | - 人 <br>  |
|  |  |
|  |  <br>  |
|  |  |



Fig. 1 : Infrared spectrum of LD 84 in KBr cell.

Fig. 2 : Infrared spectrum of $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ in KBr cell.

Fig. 4 : Infrared spectrum of $\left[C d(L)_{2} \cdot H L\right]$ in KBr cell.

Fig. 5 : Electronic spectrum ofLIX 84 (in cychlohexane)

Fig. 6: Electronic spectrum of $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ (in cychlohexane)

Fig. 7 : Electronic spectrum of $[\mathrm{Ag}(\mathrm{L})]$ (in cychlohexane)

Fig. 8 : Electronic spectrum of $\left[C d(L)_{2} H L L\right]$ (in cyclibhexane)

## REFERENCES

1. F. A. Collon and G. Wilkinson," Advanced morganic Chemistry", 5th edn. Wiley and Sons Inc(1988).
2. "LIX84 extractant," A new reagent for metal cxtraction from anmoniacal solution, Technical bulletin, General mills
Ghemicals, Inc., Arizona, U.S.AA.
3. M. S. Islam, Ph. D. Thesis, Department ofChemistry, Rajshahi University(1089).
4. L. Pauling" The nature of the chemical bond", $3^{\text {rd }}$ edn., Cornell University Press, Itchaca, New york (1960).
5. M. Calvin and K. W. Wilson, J. Am. Chem. Soc., 62, 2003 (1945).
6. W. D. Johnston and H. Freiser, Anal. Chem. Acta. 11,201 (1945).
7. R. L. Gustafson and A. E. Martell, J. Am. Chem. 81, 525 (1959).
8. J. F. Steinbach Ph. D. Thesis, University of Pittsburgh (1954).
9. U. Mayer, Pure Appl. Chem., 41, 201 (1975).
10. U. Mayer, Monatsh Chem., 108, 1479 (1977).
11. U. Mayer, Monatsh Chem., 109, 421 (1.978).
12. U. Mayer, Monatsh Chem., 51, 1697 (1979).
13. Y. Fukda., A. Shimeera, M. Mukaida, D. Pujita and K. Sone J. Inorg. Nucl.Chem.; 36, 1265 (91974).
14. W. J. Geary., "Cuordination Chemistry", Rev.7,110 (1971).
15. George Ritsoulis, Can.j. Chem. 67, 1105 (1989).
16. A. Famshaw, "Introduction to Magneto Chemistry," Academi Press, London (1968).
17. W. L. Jolly, " Modern Inorganic Chemistry, " $2^{\text {nd }}$ Edn., Mc Graw-Hill, Inc (1991).
18. N. Elliott, J. Chem. Phys., 2,419 (1934).
19. Kazuo Nakamoto, " Infra-red and Raman Spectra of Inorganic and coordination Compounds, " 2nd Edn., John Wiley \& Sons, New York (1978).
20. J. Am. Chem. Soc., 79,1575 (1957).
21. A. D. Westland and M. T. H. Tarafder, Inorganic Chem., 21,3228 (1982).
22. M. T. H. Tarafder, Indian J. Chem., 26A, 878 (1987).
23. D. M. Addams, " Metal-ligand and Relative Vibrations, " Arnold, London 319 (1967).
24. D. M. Adams and J. B. Carrel, j. Chem. Soc., (A), 1299 (1968).
25. Indian J. Chem., 30A, 471-473 (1994).
26. J. Indian Chem. Soc., 65,710 (1988).
27. A. B. P. Lever "Inorganic Electronic Spectroscopy," Elsevier, New York 291 (1968)
28. C. R. Merigold, "Copper recovery from secondary sources using ammonia leaching, LIX84 solvent extraction and electroning," 2nd, International Symposium, Poland, 27-28th April (1977).
29. Law, H. Heny, Tiemey, Vidk Electro Chem. Process Group, Al and TBell Lab, Muray Hill, Nj, 07974 U.S.A PhinSurf. Finish 1987,74 (4). 747 (Eng).
30. M. S. Islam and M. A. Salam, M. Sc. Thesis 34-38 (1994).
31. Preliminary report about LIX84 published by Henkel Coorporation Minerals Industry Division. Tucson, Arizona 85745 U.S.A.
32. "Organic Iron standard for Atomic Absorption analysis," Henkel corporation. Minerals Industry Division, Tueson, Arizona 85745 U.S.A.
33. D. P. Graddon and E.C. Wlaton, Inorg. Chem., 21,49-57 (1961).
34. J. Member. Sci 1989,44(2-3), 273-87 (Eng).Sp. A.O. Hara, M.P. Boherer, (A.T and T Bell Lab. Murray Hill, NJ 07974 U.S.A.), Chem. abstracts October 2, 11,14, 118546 w (1989).
35. M. S. Islam M. L. Rahman and M. A. IsLam, J.Bangladesh Chem. Soc. 1 (2), 200-204 (1994).I
36. M. S. Islam, R. K. Das and M. N. Uddin, J.Bangladesh Chem. Soc., 10(2), 131-139 (1997).
37. M. S. Islam, M. N. Nahar, M. F. Islam, D. A. Begum, J. Bangladesh. Acad. Soc., 19(1), 33-(1995).
38. M. S. Islam and M. F. Islam, J. Acad. Soc., 14(1), 27-36(1990).
39. M S. Islam and M. F. Illam, J. Bangladesh Chem Soc, 6(1), 9-14(1993).
40. M. S. Islam and M. F. Islam, J. Bang. Acad Soc., 15(2),101-105 (19990).
41. M. S. Islam and M. F. Islam, J. Bang. Acad Soc., 15(2),101-105 (19990).
42. M. S. Islamand M. K. H. Talukder, J. Bangladesh Chem. Soc., 8(2), (1996).
43. F. Islam and R. K. Biswas, J. Bang. Acad. Sci., 15 (2), 61 (1981).
44. M. N. Nahar, M. F. Islam,D. A. Begrum and M. S. Islam, J. Bang, Acad. Sci.,
45. R. K. Das, M. Sc. Thesis, 38 (1995).
46. Malik, Tuli and Madan, "Selected Topics in Inorganic Chemistry", 5th edn., S. Chand and Comp. Ltd., New Delhi, (1991).
47. M. Singh, Synth. React. Inorg. Met. Org. Chem., 15,253 (1983).
48. B. M. Cate House, S. E. Livingstone and R. S. NyhoIm, J. Chem.

Soc., 4222 (1957).
49. W. W. Wendlandt and H. G. Hecht, " Refletance Spetroscopy, " Inter Science Plenum, New York., (1971).
50. J. R. Ferraro, "Low Frequency Vibration Spectra of Inorganic and Coordination Compounds," Plenum, New York (1971).
51. D. L. Pavia, G. M. Lampanam and G. S. Friz, "Introduction to Spectroscopy," Saunders College Publishing Co. (1979).
52. N. B. Colthup, L.H. Daly and S.E. Wibery," Introductionto Analytical Chemistry," John Wiley and Sons, New York, (1957).
53. C. N. R. Rao, " Chemical Application of Infrared Spectroscopy, " Second Printiong, Acad press Inc., U.S.A. (1967).
54. R.S. Whewell and C.Honson, " Ion Exchange and Solvent Extraction", Aseries of Advace. 8,1-92 (1981).
55. D. L. Pavia, G.M. Lampman and G.S. Kriz, Jr" Introduction to Spectroscopy", Sanders Golden Sundurst Series, U.S.A. (1979).
56. A. I. Vogel "Text book of Quantitative inorganic analysis", 4th edn., ELBS, Longman Green Ltd. (1978).
57. G. H. Morrison and H.Freiser, " Solvent Exrtaction in Analytical Chemistry, " John Wiley \& Sons, New York (1957).
58. N. B. Colthup, L. H. Daly and S. E. Wiberley", Introduction to Infrared and Raman Spectroscopy" Second Pinting Acad. Press., Inc., U.S.A. (1963).

PART-II


## ANTIBACTERIAL ACTIVITY TESTING

### 6.1 INTRODUCTION AND PRINCIPLE:

Any chemical or biological agent that either destroys or inhibits the growth of microorganisms is called antimicrobial agent.

The susceptibility of microorganism to antimicrobial agent can be determined in vitro by number of methods. The disc diffusion technique ${ }^{1.2}$ is widely acceptable for preliminary investigation of materials which are suspected to posses antimicrobial properties. Diffusion procedure, as normally used is essentially a qualitative test which allocates organism of the susceptible intermediate (moderately susceptible) or resistant categories.

In the disc diffusion technique, dried filter paper discs containing known amount of test materials are placed on agar plates seeded with test organisms. These are kept at low temperature $\left(4^{\circ} \mathrm{C}\right)$ for 4 hours.

Initially the dried discs absorb water from the surrounding test medium and the drug is dissolved. The drug migrates through the adjacent test medium by concentration gradient of the drug according to physical law that governs diffusion of molecules through an agar gel ${ }^{3}$. As a result, there is a gradual change of drug concentration in the agar surrounding each disc. Then the plates are incubated in an incubator at $37^{\circ} \mathrm{C}$ for 16 hours.

As the antibiotic diffusion progresses, microbial multiplication also proceeds. After an initial lag phase, a logarithmic growth phase is imitated, at that moment bacterial multiplication proceeds more rapidly than the drug can diffuse, and the bacterial cells which are not inhibited by the anti microbial agents will continue to multiply until a lawn of growth can be visualized. No growth will appear in the area where drug is present in inhibitory concentration.

Centrally, the more susceptible the test organism, the larger is the circular zone of inhibition. Antimicrobial activities of the test samples are expressed by measuring the zone of inhibition observed around the area. The diameter of the inhibition is usually measured to understand the extent of inhibition in different concentrations.

The size of the inhibitory zones dopends principally on the following factors.
a. Intrinsic antimicrobial sensitivity of the test sample.
b. Growth rate of the test microorganism.
c. Diffusion rate of the drug which is related to its water solubility.
d. Number of concentration of the freshly seeded test organisms.
e. Amount of the test sample on disc.
f. Thickness of the test medium in the petridishes.
g. Thickness of the filter paper disc.

### 6.2 APPARATUS AND REAGENTS :

a. Micropipette
b. Autoclave
c. Incubator
d. Refrigerator
e. Filter paper disc
f. Petridishes
g. Inoculating loop
h. Sterile cotton
i. Sterile forceps
j. Spirit Lamp
k. Laminar air flow unit

1. Nutrient agar

### 6.3 METHOD:

The test organisms are all human pathogenic. For this reason all steps of the work were done with high precaution and aseptic condition which are mentioned below. All steps of the work were carried out at Microbiology Laboratory, Pharmacy Department, Rajshahi University.

### 6.4 TEST ORGANISMS:

The following human pathogenic bacteria have been studied:

| Sl.No. | Gram Negative |
| :--- | :--- |
| 1 | Shigella dysenteriae |
| 2 | Escherichia coli |
| 3 | Pseudomonas aeruginosa |
|  | Gram Positive |
| 4 | Bacilhis subtilis |
| 5 | Sarcina lutea |
| 6 | Staphylococcus cureuts |

### 6.5 CULTURE MEDIA:

Nutrient agar medium was used as culture media. The formulation of Nutrient agar media (DIFCO) is as follows:
Nutrient Agar (mast Diagnostics)

| Formulation | Grams/liter |
| :--- | :--- |
| Peptone A | 6.0 |
| Yeast extract | 2.0 |
| Beef extract | 1.0 |
| Sodium Chloride | 5.0 |
| Agar A | 14.0 |
| Distilled water q.s. to 1000 ml |  | water, allowed to soak for 10 minutes, swirled to $\operatorname{mix}$ and then sterilized by

autoclaving for 15 minutes at $121^{\circ} \mathrm{C}$, and then the medium was cooled to 40 $45^{\circ} \mathrm{C}$ and mixed well, then poured into plates.

### 6.6 PREPARATION OF FRESH CULTURE:

The liquid culture is called broth culture. The culture media without agar powder per liter:

| Formulation | Grams/litre |
| :--- | :--- |
| Bacto-tryptone | 10.0 g |
| bacto-yeast extract | 5.0 g |
| NaCl | 10.0 g |

Adjusted pH to 7.5 with sodium hydroxide.
Tryptone, NaCl and yeast extract of calculated amount were taken in a conical flask and distilled water was added (volume should be less than 1 litre). The contents were put on a water bath to make a clear solution. The pH of the solution was then adjusted to 7.5 using NaOH or HCl as necessary. Distilled water was added sufficiently to make the final volume (1 litre). Again the total volume was heated on a water bath to obtain a clear solution. The conical flask was plugged with cotton and then autoclaved at 1 atm pressure for 15 minutes at $121^{\circ} \mathrm{C}$.

50 ml of both medium was transferred in a conical flask. The test microorganisms of pure culture were streaked on the mutrient both media with the help of sterile loop in an aseptic condition and incubated at $37^{\circ} \mathrm{C}$ for 24 hours. The both cultures thus obtained was considered as fresh culture. Fresh culture of this type was always used throughout the sensitivity testing.

### 6.7 PREPARATION OF PLATES:

Solid media in petridishes are often called 'plates'. The medium was poured into sterile petridishes in an aseptic condition on a level horizontal surface so as to give a uniform depth of approximately 4 mm . Then the medium had been allowed to cool at room temperature in order to solidify the medium.

### 6.8 PREPARATION OF DISCS :

A. Sample disc

1. Solutions of the compounds were prepared in respective solvents so that $20 \mu \mathrm{~L}$ contained 200 g of the compounds.
2. Filter paper discs were taken in petridish and sterilized by oven at $110^{\circ} \mathrm{C}$ for 1 hour.
3. 20 " L of the solutions were placed on the discs with the help of a micropipette, thus discs containing 200 g compounds were prepared.
4. These discs were then air dried.
B. Standard dics

Ready made Kanamycin K-30 discs containing 30 yg/disc of antibiotic Kanamycin were used as standard disc.

### 6.9 PLACEMENT OF THE DISCS AND INCUBATION :

The solidified agar plates were seeded with the $200 \mu \mathrm{~L}$ of fresh culture with the help of a micropipette and spread the microorganisms with the help of a sterile spreader in an aseptic condition.

The prepared discs of samples were placed gently on the freshly seeded solidified agar plates with a sterile forceps. Standard discs and control discs were also placed on the test to compare the effect of the test sample and to nullify the effect of solvent, respectively.

The plates were then kept in a refrigerator at $4^{\circ} \mathrm{C}$ for 4 hours in order that the materials had sufficient time to diffuse to a considered area of the plates. After this, the plates were incubated at $37^{\circ} \mathrm{C}$ for 16 hours.

### 6.10 CALCULATION OF THE ZONE OF INHIBITION:

After incubation, the diameter of the zone of inhibitions were observed and measured in mm by a transparent scale. Results obtained from these are listed in Tables (6.2-6.7).

### 6.11 RESULTS OF THE ANTIBACTERIAL ACTIVITY OF THE

 COMPLEXES 1, 2, 3 AND LIGALD AGAINST THE SIX PATHOGENIC BACTERIA (viz. Escherichia Coil, Shiella Dysenteries, Pseudomonas aeruginosa, Bacillus Subtilis. Sarcina lutea AND Staphylococcus aureus).Table 6.1: Complexes abbreviation for antibacterial activity.

| Complexes No | Complexes | Labelled |
| :---: | :---: | :---: |
| 1 | $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ | A |
| 2 | $[\mathrm{Ag}(\mathrm{L})]$ | B |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2}-\mathrm{HL}\right]$ | C |
| Ligand | 2-hybroxy-5-nonyl - <br> acetophenone oxime $=\mathrm{L}$ |  |

Table 6.2: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hybroxy-5-nonyl acetophenone, oxime against Escherichia coli.

| Complex <br> No | Complexes | Zone of inhibition of mycelia growth (in mm) |  |
| :---: | :---: | :---: | :---: |
|  |  | $200 \mu \mathrm{~g}$ disc | $30 \mu \mathrm{~g} /$ disc |
| 1 | $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ |  |  |
| 2 | [ $\operatorname{Ag}(L)$ ] | 24 |  |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} \cdot \mathrm{HL}\right]$ | 19 |  |
| Ligand | 2-hybroxy-5-nonylacetophenone, oxime $=\mathrm{L}$ | 9 |  |
|  | Control disc | Nil |  |
|  | Standard disc |  | 20 |

Where,
Kanamycin was used as standard compound labeled as K .
Control disc contains solvent only.

Table 6.3: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5-nonylacetophenone against Shigella dyscnteriae.

| Complex No. | Complexes | Zone of ingibition of mycelia growth (in mm) |  |
| :---: | :---: | :---: | :---: |
|  |  | $200 \mu \mathrm{~g} / \mathrm{disc}$ | $30 \mu \mathrm{~g} / \mathrm{disc}$ |
| 1 | $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ | 7 |  |
| 2 | $[\operatorname{Ag}(L)]$ | 23 |  |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} \cdot \mathrm{HL}\right]$ | 25 |  |
| Ligand | $\begin{aligned} & \text { 2-hybroxy-5-nonyl - } \\ & \text { acetophenone, oxime }=\mathrm{L} \end{aligned}$ | 8 |  |
|  | Control disc | Nil |  |
|  | Standard disc |  | 18 |

Table 6.4: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5-nonylacetophenone against $P$ seudomonas aeruginosa.

| Complex <br> No. | Complexes | Zone of inhibition of mycelia growth (in mm) |  |
| :---: | :---: | :---: | :---: |
|  |  | $200 \mu \mathrm{~g} / \mathrm{disc}$ | $30 \mu \mathrm{~g} / \mathrm{disc}$ |
| 1 | $\left[(\mathrm{ZrO})_{2}(\mathrm{~L})_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]$ |  |  |
| 2 | [ $\operatorname{Ag}(L)$ ] | 19 |  |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} . H L\right]$ | 22 |  |
| Ligand | 2-hybroxy-5-nonyl acetophenone, oxime $=\mathrm{L}$ | 8 |  |
|  | Control disc | Nil |  |
|  | Standard disc |  | 21 |

Table 6.5: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5-nonylacetophenone against. Bacillus subtilis.

| Complex <br> No. | Complexe | Zone of inhibition of mycelia <br> growth (in mm) |  |
| :--- | :---: | :---: | :---: |
|  |  | $200 \mu \mathrm{~g} / \mathrm{disc}$ | $30 \mu \mathrm{~g} / \mathrm{disc}$ |
| 1 | $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ | 7 |  |
| 2 | $[\mathrm{Ag}(\mathrm{L})]$ | 23 |  |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} \mathrm{HL}\right]$ | 17 |  |
| Ligand | 2-hybroxy-5-nonyl - <br> acetophenone, oxime <br> $=\mathrm{L}$ | 9 |  |
|  | Control disc | Nil |  |
|  | Standard disc | 20 |  |

Table 6.6: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5- nonylacetophenone against Sarcina lutea.

| Complex No. | Complexes | Zone of inhibition of mycelia growth (in mm) |  |
| :---: | :---: | :---: | :---: |
|  |  | $200 \mu \mathrm{~g} /$ disc | $30 \mu \mathrm{~g} / \mathrm{disc}$ |
| 1 | $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ |  |  |
| 2 | $[\mathrm{Ag}(\mathrm{L}) \mathrm{]}$ | 21 |  |
| 3 | [Cd( L$\left.)_{2} \cdot \mathrm{HL}\right]$ | 18 |  |
| Ligand | $\begin{aligned} & \text { 2-hybroxy-5-nonyl - } \\ & \text { acetophenone, oxime }=\mathrm{L} \end{aligned}$ | 9 |  |
|  | Control disc | Nil |  |
|  | Standard disc |  | 22 |

Table 6.7: Antibacterial activity of the complexes 1, 2, 3 and ligand 2-hydroxy-5-nonylacetophenone against Staphylococcus aureus.

| Complex No. | Complexes |  | Zone of inhibition of <br> mycelia growth (in mm) |  |
| :---: | :---: | :---: | :---: | :---: |
|  |  | $200 \mu \mathrm{~g} / \mathrm{disc}$ | $30 \mu \mathrm{~g} / \mathrm{disc}$ |  |
| 1 | $\left[(\mathrm{ZrO})_{2}\left(\mathrm{~L}_{4}\left(\mathrm{H}_{2} \mathrm{O}\right)_{2}\right]\right.$ | 8 |  |  |
| 2 | $[\mathrm{Ag}(\mathrm{L})]$ | 16 |  |  |
| 3 | $\left[\mathrm{Cd}(\mathrm{L})_{2} \cdot \mathrm{HL}\right]$ | 21 |  |  |
| Ligand | 2-hybroxy-5-nonyl - <br> acetophenone, oxime $=\mathrm{L}$ | 8 | 8 |  |
|  | Control disc | Nil |  |  |
|  | Standard disc | 20 |  |  |



Fig-6.1 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Escherichia coli.


Fig-6.2 Photographic representation of zone of inhibition of thecomplexes 1,2,3 and the standard compound kanamycin against Shigella dysenteriae.


Fig-6.3 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Pseudomonas aemiginosa.


Fig-6.4 Photographic representation of zone of inhibition of the complexes $1,2,3$ and the standard compound kanamycin against Bacillus subtilis.


Fig-6.5 Photographic representation of zone of inhibition of the complexes 1,2,3 and the standard compound kanamycin against Sarcina lutea.


Fig-6.6 Photographic representation of zone of inhibition of the complexes
1,2,3 and the standard compound kanamycin against Staphylococcus aureus.

### 6.12 DISCUSSION :

It has been observed that some drugs (Ligands) increase the activity when administered as metal complexes or their metal chelates ${ }^{4}$. The antibacterial activity of the metal compsexes $1,2,3$ and the ligand (2-hybroxy-5-nonyl acetophenone) are siudied against with six pathogenic bacteria viz.

1. Shigella dysenteriae
2. Escherchia coli
3. Pscudomonas aeraginosa.
4. Bacillus subtilis.
5. Sarcina lutea.
6. Staphylococcus aureus.

And the results are given in table (6.2-6.7). It is seen that the complex 2 and 3 showed the most activities above six pathogenic bacteria as shown in Table (6.2-6.7).

The Complex 2 showed the best activity against E. Coli and less activity against Staphylococcus aureus. The complexe 1 did not show activities against the six pathogenic bacteria.

All the results are compared with the standard compound, Kanamycin as seen in the Table (6.2-6.7). The ligand (oxime) did not show any activities against the above six pathogenic bacteria.

From the above discussion it is concluded that the complexe 2 and 3 showed good activities against the six pathogenic bacteria as compared to the stardard compound, Kanamycin. It is evident that the metal ion plays the key role to show good activities, because the ligand did not show any activity.

### 6.13 DETERMINATION OF MINIMUM INHIBITORY CONCENTRATIONS (MIC) OF COMPLEXES.

### 6.13.1 Introduction.

The lowest concentration of antimicrobial agent required to inhibit the growth of the organism in vitro is referred to as the minimum inhibitory concentration (MIC). There are two methods for determining this value:
i) Serial dilution technique or turbidimetric assay ${ }^{5,6}$.
ii) Paper disc technique or agar diffusion assay ${ }^{5}$.

Here, "serial dilution technique"5,6 was followed using nutrient broth medium. The MIC values of Complexes were determined against the following six test organisms:

| Gram Negative | Gram Positive |
| :--- | :--- |
| Shigella dysenteriae | Bacilhs subtilis |
| Escherichia coli | Sarcina lutea |
| Pseudomonas aeruginosa | Staphylococcus aureus |

### 6.13.2 Preparation of sample solution

2.048 mg of the test compound was taken in a vial. 2 mL of methanol was added the vial to dissolve the compound. Thus solutions with a concentration or i. $024 \mathrm{mg} / \mathrm{mL}$ were obtained.

### 6.13.3 Preparation of incolum

Over night cultures of the test bacteria grown at $37.5^{\circ} \mathrm{C}$ in nutrient broth medium was diluted in sterile nutrient broth medium in such a manner so that suspension contain about $3.5 \times 10^{6}$ cell/mL. This suspension was sued as the inoculum.

### 6.13.4 Procedure.

i) 12 test tubes were taken, nine of which were marked (1-9) and rest three were assigned as $C_{M}$ (nutrient broth medium), $\mathrm{C}_{\mathrm{s}}$ (nutrient broth medium +compound) Ci [nutrient broth medium+ inoculum (organism)].
ii) 1 ml of nutrient broth medium was poured to each of the 12 test tubes.
iii) These test tubes were cotton plugged and sterilized in a autoclave for 15 minutes at $121^{\circ} \mathrm{C}$ temperature and 1 atm . pressure.
iv) After cooling, 1 mL of the sample solution was added to the lst tube, mixed well and them 1 mL of this content was transferred to the second test tube.
v) The content of the 2 nd test tube was mixed well and again 1 mL of this mixture was transferred to the 3rd tube. This process of serial dilution was continued up to the 9th test tube.
vi) $10 \mu \mathrm{~L}$ of properly diluted inoculum was added to each of the nine test tubes and mixed well.
vii) To the control test tube, $\mathrm{C}_{8}, 1 \mathrm{~mL}$ of the sample solution was added; mixed well and 1 mL of this mixed content was discarded. This is to check the oleayyity of the medium in persence of diluted diluted solution of the compound.
viii) $10 \mu$ Lof the incolumns was added to the control test tube $C_{i}$, to observe the growth of the organism in the medium used.
ix) The control test tube $\mathrm{C}_{m}$, containing medium only was used to confirm the sterility of the medium.
x) All the test tubes were incubated at $37.5^{\circ} \mathrm{C}$ for MIC is the lowest drug concentration at which there in no growth.
6.14 RESULT OF THE MINIMUM INHBITORY CONCENTRATION OF THE COMPLEXE [Ag(L)] 2 AND [Cd(L) $)_{2}$ HL] 3 AGAINST THE SIX PATHOGENIC BACTERIA VIZ:

1. Escherchia coli.
2. Shigella dysenteriae.
3. Pscudomonas aeraginosa.
4. Bacillus subillls.
5. Sarcina lutea.
6. Staphylococcus aureus.
Table 5.8: Minimum inhibitory concentration of complexes [ $\mathrm{Ag}(\mathcal{L})] 2$ againest six pathogenic bacteria.

| Test tube No. | Nutrient both medium added (mL) | Diluted solution of complex ( $\mu \mathrm{g} / \mathrm{mL}$ ) | Inoculums added $(\mu L)$ | Bacterial Growth observation against |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | E. coli | S.dysenteriae | P.aeruginosa | B.subtilis | Sarcina lutea | S.aureus |
| 1 | 1 | 512 | 10 | -ve | -ve | -ve | -ve | -ve | -ve |
| 2 | 1 | 256 | 10 | -ve | -ve | -ve | -ve | -ve | +ve |
| 3 | 1 | 128 | 10 | -Ve | -ve | -ve |  |  |  |
| 4 | 1 | 64 | 10 | -ve | +ve | +ve | -ve | -ve | +ve |
| 5 | 1 | 32 | 10 | +ve | +ve | +ve | +ve | +ve | +ve |
| 6 | 1 | 16 | 10 | +ve | +ve | +ve | +ve | +ve | +ve |
| 7 | 1 | 8 | 10 | +ve | +ve | +ve | +ve | +ve | +ve |
| 8 | 1 | 4 | 10 | +ve | +ve | +ve | +ve | +ve | +ve |
| 9 | 1 | 2 | 10 | +ve | +ve | +ve | +ve | +ve | +ve |
| Cs | 1 | 512 | 10 | -ve | -ve | -ve | +ve | +ve | +ve |
| Ci | 1 | 0 | 10 | +ve | +ve | +ve | +ve | -ve | -ve |
| Cm | 1 | 0 | 10 | -ve | -ve | -ve | -ve | -ve | +ve |

Where, + ve $=$ Growth and $-v e=$ Not growth
Table 5.8: Minimum inhibitory concentration of complex [Cd(L) $\left.)_{2} H L\right] 3$ against six pathogenic bacteria.


### 6.15 DISCUSSION:

The minimum inhibitory comcentration (MIC) of the determined against six pathogenic bacteria viz.

1. Escherchia coli.
2. Shigella dysenteriae.
3. Pscudomonas aeraginosa.
4. Bacillus subtilis.
5. Sarcina lutea.
6. Stapliylococcus aureus by serial dilution technique. The results are shown in
Table. 6.10: MIC of the complexes $[\hat{A g}(\mathrm{~L})] 2$ and $\left[\mathrm{Cd}(\mathrm{L})_{2} \cdot \mathrm{HL}\right] 3$

| Test organisms | MIC $(\mu \mathrm{g} / \mathrm{ml})$ |  |
| :---: | :---: | :---: |
|  | Complex 2 | Complex 3 |
| Escherichia coli. | 64 | 64 |
| S. dysenteries | 128 | 128 |
| Pseudomonas | 128 | 128 |
| Bacillus subitisis | 128 | 128 |
| Staphylococcus caureus | 256 | 256 |
| Sarcina hutea | 256 | 256 |

## REFERENCES

1. A. W. Buer, W.M.M. Kirby, J.C. Sherris and M. Turck, Am. J. Clin. Pathol, 44,493 (1966).
2. S.S. Gnanamanickam and D.A. Smith, Selective toxicity of isoflavonoid phytoalenins to Gram positive bacteria, Phytoalenins to Gram positive bacteria, Phytoalenins to Gram posiitve bacteria, phytopathology, 70,894 (1980).
3. A. L. Barry, Principles and practice of Microbiology, Lea and Febgen, philadepbia, (1976).
4. William's, Chem. Rev, 72,209 (1972); A. Fenst and R.T. Harat, prog. Exp. Tumor, Res., 12,102 (1969).
5. Tyler, M.E.; Brady, L.R; Robbers, J.E. Pharmacognosy, Ninth edition, Lea and Febiger, Philadelphia, 1988, P-312-318.
6. Roland Reiners. Antibiotics, Chemotherapeutic agent and Development of Chemotherapy: Antibiotics An Introduction. Roche Scientific service, Switzerland 1982. 2-9 Detection of antibiotic activity P-21-25.

## APPENDIX

A) Hydrometallurgy:

The processes involved in the production of pure metals from ores(i.e., the natural minerals in metals or their compounds occur in the earth) are known as Metallurgy. It involves some chemical and mechanical steps. The process in which the metals present in the ore is made to go into aqueous solution due to some chemical changes and the metal is then separated by suitable means e.g. fractional crystallization, precipitation, ion-exchange or solvent extraction, is called Hydrometallurgy. Hydrometallurgy is concerned usually with the processing (large and small scale) of low grade metalliferous bodies; the primary purpose being the extraction of the metal in a suitable form in aqueous solution for further processing. At presen, hydrometallurgical operations have become of worldwide interest over the long known pyrometallurgical processes mainly for the following reasons: i) Very low grade ores, and the burden of required antipollution equipments have proven the pyrometallurgical processes uneconomical; ii) Environmental pollution has encouraged the treatment of metallurgical wastes by hydrometallurgical means; iii) Increased cost of raw materials and their diminishing supply encourage the reprocessing of scrap; hydrometallurgy has made an impact on the isolation of metals of metals of complex mixtures in solution; iv) Hydrometallurgy can handle the purification problem of the leach solution which is complecated by the presence of more than one metal in single solution.

The methods have been tried on a commercial scale are mainly (a) solvent extraction (liquid-liquid) and (b) ion - exchange of the two, the solvent extraction method has now become a method of major importance, because of high cost of ion - exchange matters, very low output or capacity $4 \mathrm{me}^{-1}$, fixed low rate (very low
process) and also very low selectivity of the ion exchange process as contrast to solvent extraction. Obviously, solvent extractant enjoys a favoured position among other exising powerful separating techniques because of its case, simplicity, selectivity, apeed and wide scope. It is applicable to both trace and macro levels of metal concentrations and gives high purity of the products. The cost of production is also low. Thus where fractional crystallization, precipitation and other usual techiques are tedious and gives high purity of the products. The cost of production is also low. Thus where fractional crystallization, precipitation and other usual techniques are tedious and costly, the solvent extraction process is very simple and cheap. The process (SX) is now well established in hydrometallurgy using more efficient and selective extractants developed, with fullscale plants with newly devised low-cost contactng devices. As a matter of face, a great variety of metals are now being treated hydrometallurgically, and extracted using the solvent extraction method.

During the last three decads solvent extraction finds increasing opplication in the extraction of the common metals (viz, coppir, Nickel, Cobalt and the like). A number of reports has of far been published. Most of these investions; but only a few of them are related to the kinetics of the processes. The chemical engineering principles underlying the techique of multistage counter-curent extraction are well understood and it is now possible to desing a full scale plant from simple distribution measurements. Moreover, the development of low cost new contacting devices has extended the applicability of the technique.

## B) SOLVENT EXTRACTION(SX)

In hydrometallurgy, the solvent extraction method is a chemical process which involves the mass transfer of complex species of a metal ion across the boundary between two practically immiscible phases on agitation of which one is an inert is an inert organic solvent containing the extracting reagent (i.e. the extractant), and the other, an aqueous feed material. The extractant reacts with the distribuent by processes such as solvation, chelation, ion - pairing ion - exchange etc., to extract the metal species from the aqueous phase. The extractant and the diluent together act as an efficient cxtracting solvent; the latter modifies the extratant properties of the farmer.

As a physical method of purification and separation, the extraction using a suitable solvent(organic) of renerally organic matters has long been familiar to especially the organic ${ }^{56}$ as well as analytical chemists. But there has been interest in solvent extraction in the field inorganic chemistry during the girst three decades of this century; although the extractability of several inorganic compoubds into organic solvents was early established. The extraction of uranyl nitrate from aqueous solution by ether ${ }^{57}$, for example, was discovered in 1842 .

Now a days, solvent extraction is not only used for the separation and production of rare earths but also finds increasing application in the extraction of commoner metals, especially from their low grade ores. Besides it has not extensive applications (laboratory and industrial) in the analytical field ${ }^{58}$ for the separation and purification of metals.

