# SYNTHESIS AND STUDIES OF SOME MIXED – LIGAND METAL COMPLEXES CONTAINING BENZOTRIAZOLE WITH SOME OTHER LIGANDS

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## Abstract

This paper presents the synthesis and studying of some mixed–ligands of Benzotriazole, Benzimidazole and thiocyanate ion. The reaction was carried out by using the appropriate molor ratios of Metal: ligand (1:1:2:2) (M: BT: BI: X) as required. The resulting products were found to solids which have been characterized by U.V-visible, I R spectra ,Molar conductivity, thermal stability, Determination of metal and magnetic susceptibility. From the obtained informations the general formula have been given for the prepared Complexes: [M (BT) (BI) X<sub>2</sub>].  $M = Mn^{2+}$ , Co<sup>2+</sup>, Ni<sup>2+</sup>, Cu<sup>2+</sup>, Cd<sup>2+</sup> and pb<sup>2+</sup>

## Introduction

Hetrocyclic nitrogens play in turn an important role in coordination chemistry <sup>(1)</sup>. The triazole derivatives have also of discrete or Supermolecules in particular, benzotriazole is largely employed as an efficient corrosion inhibitor for copper and its alloys<sup>(2)</sup> and the benzimidazole derivatives are very useful as fungicides and bacteriacides<sup>(3)</sup>. Although the reactivity and properties of transition metal complexes with imidazole and triazole derivatives have been studied a few cases involving the interaction of benzimidazole and benzotriazole ligands with iron (II, III) and ruthemium (II) (III) complexes<sup>(2)</sup>.

# Experimental

## Materials and Measurements

Metal salts (MnCl<sub>2</sub> 4H<sub>2</sub>O, CoCl<sub>2</sub> 6H<sub>2</sub>O, NiCl<sub>2</sub> 6H<sub>2</sub>O, CuCl<sub>2</sub>. 2H<sub>2</sub>O, CdCl<sub>2</sub>.H<sub>2</sub>O, Pb(NO<sub>3</sub>)<sub>2</sub>, NH<sub>4</sub> SCN and KBr) from Riedeldehaenage, ligands (benzotriazole and benzimidazole) from fluka, (ethanol and dimethyl sulfoxide) was obtained from BDH in high Purity.

The IR spectra in the region (4000 - 400)cm<sup>-1</sup> (KBRdisc) were recorded by using shimadzu. FTIR.8400S Fourier transform infrared spectrophotometer, the U.V.visible were carried out on shimadzu U.V.visible recorder spectrophotometer U.V-160.in DMSO solution (10<sup>-3</sup>M) and Philips PW 9526 digital conductivity meter was used to measure

Molar Conductivity, magnetic susceptibilities were measured by using Balance Magnetic susceptibility model MSB. MK1 at 25C° and the metal percentage in the complexes were determined following the gravimetric methods. Finally the melting Points were recorded by using stuart Melting Point Apparatus.

# General Procedure for the synthesis of the Complexes

To an aqueous solution of metal salts an ethanolic solution of benzotrizole (0.17-0.34 g) (1.42-2.85 m.mole) in 15 ml in ethanol was added follow by the addition of solution of benzimidazole (0.35-0.69 g) (2.96 -5.84 m.mole). The aqeous solution of NH<sub>4</sub>SCN (0.22-0.44 g) (2.89-5.78 m.mole) was added to the mixture with constant stirring, the products were precipitated immediately which were filtered off washed with (1:1) mixture of water: ethanol, recrystalized from ethanol ,and dried in an oven(60)C<sup>o</sup>.

## **Results and Discussion:-**

All the complexes prepared in this work were insoluble in water and methanol but soluble in dimethyl sulfoxide, the complexes are thermal stable as they decomposed above (300)  $^{\circ}$ C (except copper complex which decomposes at (265)  $^{\circ}$ C <sup>(4)</sup>.

The complexes are nonelctrolytes, the molar conductance are found to be in range

(5.6 - 8.4) ohm<sup>-1</sup> .cm<sup>2</sup> .mol<sup>-1</sup>(10<sup>-3</sup>)M in DMSO<sup>(5-7)</sup>.

The physical properties are listed on Table (1)

# **Magnetic Properties**

The magnetic moment ( $\mu$  eff) for the complexes of Mn<sup>+2</sup> (cl<sup>5</sup>) ,Ni<sup>+2</sup> (d<sup>8</sup>)were (4.920,2.602) B.M respectively , these values were in agreement with the values expected for spin only <sup>(8,9)</sup> while the values of  $\mu$  eff of Co<sup>+2</sup> (d<sup>7</sup>) complex was (4.561) B.M higher than the calculated value may be due to the orbital contribution <sup>(10,11)</sup>, the spin only ( $\mu$  eff) of Cu<sup>+2</sup> complex (d<sup>9)</sup> was found to be (1.670) B.M within the expected value for one .electron. Finally the complexes of Cd<sup>+2xv</sup>, Pb<sup>+2</sup> were diamagnetic as expected from their electron configuration .All data are included in Table (2).

# The Electronic Spectra

# $\lambda$ Electronic spectra of free ligands:

The electronic spectrum of benzotriazole in DMSO show strong bands at  $\lambda \max (273)$ n.m and  $\lambda \max (312)$ n.m(32051.282) cm<sup>-1</sup> due to the electronic transitions  $(\pi \rightarrow \pi^*)$  and  $(n \rightarrow \pi^*)$  respectively.

TheBenzimidazole spectrum exhibited absorption bands at  $\lambda \max (275)$  n.m (36363.636) <sup>cm-1</sup> caused by  $(\pi \rightarrow \pi^*)$ , and another bandsat  $\lambda \max (348)$  nm (28735.632) cm<sup>-1</sup> due to electronic transition of  $(n \rightarrow \pi^*)^{(12-14)}$ .

Finally the spectrum of Ammonium thiocyanate shows the electronic transition of  $(\pi \rightarrow \pi^*)$  and  $(n \rightarrow \pi^*)$  at  $\lambda \max (216)$  n.m and  $\lambda \max (351)$  n.m respectively <sup>(15)</sup>.

# The Electronic spectra of Complexes

# 1. $[Mn (BT)(BI)_2X_2]$

The spectrum of light brown  $(d^5)$ complex exhibited absorption band at  $\lambda \max (264) \operatorname{nm} (37878, 787) \operatorname{cm}^{-1}$  belong to  ${}^{6}A_{1}g \rightarrow {}^{4}T_{1}g_{(G)}$  another band appeared at the visible region at  $\lambda \max (634) \operatorname{n.m} (15772.87)$  cm<sup>-1</sup> assigned to  ${}^{6}A_{1}g \rightarrow {}^{4}T_{1}g_{(G)}$ . These rules are accepted for Mn<sup>+2</sup> Octahedral complexes<sup>(16)</sup>.

## 2. $[Co(BT)(BI)_2X_2]$

The spectrum of dark green complex (d<sup>7</sup>) exhibited three bands at  $\lambda \max (272)$  nm (36764.705) cm<sup>-1</sup> due to the electronic transition  ${}^{4}T_{1}g(F) \rightarrow {}^{4}A_{2}g(F)$  corresponding with charge transfer (C.T) Another bands in the visible region at  $\lambda \max (539)$  nm (18553.875) cm<sup>-1</sup> and  $\lambda \max (760)$  nm (13157.894) cm<sup>-1</sup> attributed to the  ${}^{4}T_{1}g(F) \rightarrow {}^{4}T_{2}g(F)$  respectively<sup>(16,17)</sup>.

# [Ni (BT) (BI) 2 X2]

The spectrum of the gray complex (d<sup>8</sup>) showed strong band at  $\lim (271)$  nm (36900.369) cm<sup>-1</sup>attributed to the transition <sup>3</sup>A<sub>2</sub>g (<sub>F</sub>) $\rightarrow$ <sup>3</sup>T<sub>1</sub>g (<sub>P</sub>) with charge transfer. Another band noticed at  $\lambda$  max (560) nm (17857.142) cm<sup>-1</sup>caused by <sup>3</sup>A<sub>2</sub>g(<sub>F</sub>) $\rightarrow$ <sup>3</sup>T<sub>1</sub>g (<sub>F</sub>) transition, the third band at  $\lim (649)$  nm (15408.32) cm<sup>-1</sup> due to<sup>3</sup>A<sub>2</sub>g (<sub>F</sub>) $\rightarrow$ <sup>3</sup>T<sub>2</sub>g (<sub>F</sub>) transition <sup>(18,19)</sup>.

Accordingly the ligand field splitting energy 10Dq was taken as equal to the electronic transition ( $v_1$ ) (15408.32) cm<sup>-1</sup> and the recah inter electronic repulsion parameter ( $\beta^-$ ) found to be (568.836) cm<sup>-1</sup>. The ratio of  $\beta = \frac{B'}{B^{\circ}}$  comes out to be (0.55) (B<sup>o</sup> is equal (1030) cm<sup>-1</sup>) the parameters are show the Ni (II) complex is octahedral <sup>(16,18,19)</sup>.

# [Cu (BT) (BI) 2 X2]

The spectrum of light green (d<sup>9</sup>) complex exhibited obsorption band at  $\lambda max$  (270) nm (37037.037) cm<sup>-1</sup>. Another bands in the visible region at  $\lambda$  max (507) nm (19723.865) cm<sup>-1</sup> assigned to <sup>2</sup>Eg (<sub>D</sub>) $\rightarrow$ <sup>2</sup>T<sub>2</sub>g (<sub>D</sub>) these transition are characteristic for octahedral complexes of Cu (II)<sup>(20,21)</sup>.

The complexes of  $Cu(II)^{(20,21)}$ .

The complexes of  $[Cd(BT) (BI)_2X_2]$ ,  $[Pb(BT) (BI)_2X_2]$  showed sbsorption bands at  $\lambda$  max (274) nm and  $\lambda$  max (284) nm  $\upsilon$ respectively attributed to charge transfer (ML) because the electronic configuration of  $Cd^{+2}$  and  $Pb^{+2}$  which confirm absence of any (d-d) transition <sup>(22)</sup>. But their spectrum suffered red shift with Hyper or Hypo chromic effect this absorption has been fully assigned in Table (3).

### **Infrared Spectra**

The spectrum of benzotriazole showed a weak band at (3247) cm<sup>-1</sup> which belongs to stretching vibration  $\nu$  (NH), another weak band at (3078) cm<sup>-1</sup> attributed to  $\nu$  (C-H) aromatic <sup>(23 -25)</sup>, finally the strong band belong to str.Vib  $\nu$  (N=N) was found at (1500) cm<sup>-1</sup>.

The spectrum of benzimidazole exhibited broad band at (3150) cm<sup>-1</sup> caused by str.vib  $\nu$  (NH), a weak band noticed at (3040) cm<sup>-1</sup> attributed to  $\nu$  (C-H) aromatic ,another weak sharp band appeared at (1640) cm<sup>-1</sup> to  $\nu$ (C= N) <sup>(26)</sup>. The medium band belong to str.vib of  $\nu$  (C=C) was found at (1605)cm<sup>-1</sup>.

The spectrum of thiocyanate ion showed a strong band at (2053) cm<sup>-1</sup> assigned to  $\nu$  (CN) and another band at (740) cm<sup>-1</sup> was attributed  $\nu$  (CS)<sup>(27)</sup>.

#### **Infrared Spectra of Complexes**

The infrared spectra of the prepared complexes exhibited the following bands:  $\nu$  (C= N) in the region (1620-1630) cm<sup>-1</sup> which was shifted to lower frequencies by (20-10) cm<sup>-1</sup> in comparison with benzimidazole spectrum indicates coordination with metal ions through nitrogen atom and the bands appeared in the region (1488-1496) cm<sup>-1</sup> belong to  $\nu$  (N=N) in benzotriazole moved to lower frequencies by (12-4) cm<sup>-1</sup> which indicated coordination with metal ions through the two nitrogen atoms.

The coordination of  $(SCN)^{-}$  ion through the nitrogen atom was comfired by the bands in the region  $(2067-2098)cm^{-1}$  shifted to higher frequencies by  $(14-45) cm^{-1} (27)$ .

#### **M-Ligand bonds**

The spectra of prepared complexes exhibited weak bands in the region (550-570) cm<sup>-1</sup> and (424-460) cm<sup>-1</sup> assigned to  $\nu$  (M–N) and bending viberation of s(M–NCS) respectively.

Table (4) gives the diagnostic absorption and their assignments.

#### The proposed geometrical structures:-

From all above studies the probable structure has been suggested:-

a. Octahedral geometry for the complexes of  $Mn^{+2}$ ,  $Co^{+2}$ ,  $Ni^{+2}$ ,  $Cu^{+2}$  and  $Cd^{+2}$ .

Distorted Octahedral geometry for the Pb<sup>+2</sup> complex <sup>(14, 20, 28).</sup>

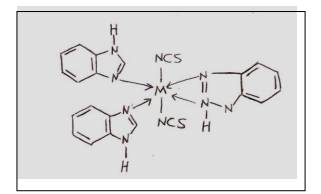
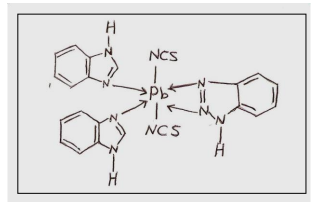


Fig. (1) :  $M = Mn^{+2}$ ,  $Co^{+2}$ ,  $Ni^{+2}$ ,  $Cu^{+2}$ ,  $Cd^{+2}$ 



**Fig.** (2)

 Table (1)

 The physical prosperities of the free ligands and their complexes.

Compound	Colour	M.P.(C°)	Dec (C°)	Elemental Analysis %M Calc.(found)	
BT	white	98		١	9
BI	Brown	172	240		7
NH <sub>4</sub> SCN	white	149	_	_	26
$[ \ Mn( \ BT) \ (BI)_2 \ X_2 ] \ (C_{22} \ H_{17} N_9 \\ S_2) \ Mn$	Light Brown	_	>300	10.43 (10.83)	8.4
$[\begin{array}{c} \text{Co(BT)} \ (\text{BI})_2 \ X_2 ] \ (\text{C}_{22} \ \text{H}_{17} \text{N}_9 \\ \text{S}_2) \ \text{Co} \end{array}$	Dark Green	Ι	>300	11.10 (10.78)	6.7
[ Ni( BT) (BI) <sub>2</sub> X <sub>2</sub> ] (C <sub>22</sub> H <sub>17</sub> N <sub>9</sub> S <sub>2</sub> ) Ni	Gray	_	>300	11.07 (10.15)	8.2
$[\begin{array}{c} Cu(\ BT)\ (BI)_2\ X_2]\ (C_{22}H_{17}N_9\\ S_2)\ Cu \end{array}$	Light green	185	265	11.87 (12.78)	5.6
$[ \ Cd( \ BT) \ (BI)_2 \ X_2 ] \ (C_{22} H_{17} N_9 \\ S_2) \ Cd$	white	—	>300	19.24 (18.46)	6.4
[ Pb ( BT) (BI)2 X2] (C22 H17N9 S2) Pb	white	—	>300	30.52 (30.99)	7.3

\* Calc % (found %)

Complexes	d <sup>n</sup>	Electron configuration	Term Symbole	Ground	$\mu$ eff		Orbital Contribution
				State	Found	Calc.	in Octahedral
[ Mn( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	d <sup>5</sup>	$t_2g^3eg^2$	<sup>6</sup> S	<sup>6</sup> A <sub>1</sub> g	4.920	5.916	No
[ Co( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	d <sup>7</sup>	$t_2g^5eg^2$	${}^{4}F$	${}^{4}T_{1}g$	4.561	3.872	Yes
[ Ni( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	d <sup>8</sup>	$t_2 g^6 eg^2$	<sup>3</sup> F	$^{3}A_{2}g$	2.602	2.828	NO
[ Cu( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	d <sup>9</sup>	$t_2 g^6 eg^3$	<sup>2</sup> D	<sup>2</sup> Eg	1.670	1.732	NO
[ Cd( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	d <sup>10</sup>	$t_2g^6eg^4$	$^{1}$ S		Zero	Zero	—
[ Pb ( BT) (BI) <sub>2</sub> X <sub>2</sub> ]			$^{1}$ S		Zero	Zero	—

Table (2)The magnetic susceptibilities for the complexes

Compound	Imax n.m	ABS	Wave number Cm <sup>-1</sup>	Emax	Transitions	Remarks
DT	273	2.447	36630.036	2447	$\pi \rightarrow \pi^*$	_
BT	312	0.012	32051.282	12	$n \rightarrow \pi^*$	_
	275	0.467	36363.636	467	$\pi \rightarrow \pi^*$	
BI	348	0.038	28735.632	38	$n \rightarrow \pi^*$	
•••	216	0.109	46296.296	109	$\pi \rightarrow \pi^*$	
X <sup>-</sup>	351	0.042	28490.028	42	$n \rightarrow \pi^*$	
	264	0.488	37878.787	488	$^{6}A_{1}g \rightarrow {}^{4}T_{1g}(P)$	—
[Mn (BT) (BI) <sub>2</sub> X <sub>2</sub> ]	634	0.010	15772.87	10	$^{6}A_{1}g \rightarrow {}^{4}T_{1g(G)}$	—
[Co(BT)(BI) <sub>2</sub> X <sub>2</sub> ]	272	2.466	36764.705	2466	${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}A_{2g(F)}$ $(V_{2})$	Corresponding with charge Transfer(C.T)
	539	0.050	18552.875	50	${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}T_{1g(P)}$ $(\nu_{3})$	New band
	760	0.043	13157.894	43	${}^{4}T_{1}g_{(F)} \rightarrow {}^{4T}T_{2g}(F) $ $(\nu_{1})$	New band
[ Ni( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	271	2.252	36900.369	2252	$^{3}A_{2}g_{(F)} \rightarrow ^{3}T_{1g}(P)$ $(\nu_{3})$	Corresponding with(C.T)
	560	0.038	17857.142	38	${}^{3}A_{2}g_{(F)} \rightarrow {}^{3}T_{1g(F)}$ $(\nu_{2})$	New band
	649	0.030	15408.32	30	${}^{3}A_{2}g_{(F)} \rightarrow {}^{3}T_{2g(F)}$ $(\nu_{1})$	New band
[ Cu( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	270	1.834	37037.037	1834	(C.T)	Blue shift
	507	0.016	19723.865	16	$^{2}\text{Eg}_{(D)} \rightarrow ~^{2}\text{T}_{2g(D)}$	-
[ Cd( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	274	1.325	36496.35	1325	$(C.T)$ $(M \to L)$	-
[ Pb ( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	284	2.500	35211.267	2500	$(C.T)$ $(M \rightarrow L)$	-

 Table (3)

 The electronic spectra of free ligands and their complexes

				i ð			•		
Compound	V (NH)	V (C-H) aromatic	V (NCS)	V (C=N)	V (N=N)	V (CS)	V (M-N)	δ (M - NCS)	
BT	3247.90 W.b	3078.18 v.w	_		1500 w	_	_		
BI	3150 b	3040 v.w	_	1640 w.sh	_	_			
NH <sub>4</sub> SCN	_	_	2053	_	_	_	_		
[Mn(BT) (BI) <sub>2</sub> X <sub>2</sub> ]	3333.06 v.w	3078.18 v.w	2075.26 v.s	1620.09 v.w	1488.94 w	833 v.s	555.46	432.03	
[Co(BT)(BI ) X <sub>2</sub> ]	3409.91 v.w	3109.04 v.w	2080 v.s	1620 w	1496.66 <sub>W.sh</sub>	748.33 v.s	563.18 w	424.31 w	
[ Ni( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	3394.48 W.b	3139.90 V.W	2090 V.S	1620.09 S	1496.66 W.sh	748.33 V.S	570 W	460 M	
[ Cu( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	3263.33 w.b	3147.61 sho	2098.41 v.s	1630 w	1490.37	748 S	570.89 w	440 w	
[ Cd( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	3340.48 v.w	3070.46 sho	2082.98 S	1620 m	1496 w.sh	750 v.s	555.46 w	450 m	
[ Pb ( BT) (BI) <sub>2</sub> X <sub>2</sub> ]	3456.20 v.w	3016.46 v.w	2067.55 v.s	1627.81 w	1490 v.w	745.61 S.b	550 sho	447.45 m.sh	

 Table (4)

 The characteristics infrared absorption free ligands and their complexes

\* S = strong, m = medium , W = weak , sho = shoulder , b = broad , v = very

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الخلاصة

تم تحضير و معقدات فلزية ذات لكاندات مختلفة من البنزوتر ايازول (BT) و البنز ايمدازول (BI) و ايون الثايوسيانات (X) مع الايونات الفلزية المنغنيز (II) و الرصاص (II) و بالرضبة المولية (1:2:2) شخصت المعقدات المحضرة على تقدير النسبة المئوية للفلز أطياف الأشعة فوق البنفسجية المرئية و اطياف الاشعة تحت الأشعة فوق البنفسجية المرئية و اطياف الاشعة تحت و قياسات الحساسية المعناطيسية و بناءً على ما تم استنتاجه فقد اقترحت الصيغة العامة للمعقدات المحضرة و كما يأتي :-Pb<sup>2+</sup>, Cd<sup>2+</sup> , Ni<sup>2+</sup> , Co<sup>2+</sup>, Mn<sup>2+</sup>=M