Advances in Physics Theories and Applications ISSN 2224-719X (Paper) ISSN 2225-0638 (Online) Vol.27, 2014



# Preparation, Characterization, and Antibacterial Properties of mixed ligand Complexes of L-leucine and Sulfamethoxazole with Mn(II),Co(II),Ni(II),Cu(II),Zn(II),Cd(II)and Hg(II) ions

Raheem Taher Mahdi , \*Taghreed H. Al- Noor , Ahmed .H. Ismail Chemistry Department, College of Science, Al-Mustansiriyah University, IRAQ \*Chemistry Department, Ibn -AI-Haithem College of Education, University of Baghdad, IRAQ \*Corres. author : <u>drtaghreed2@gmail.com</u>, <u>dr.raheem.T@gmail.com</u>

#### Abstract:

The research includes the synthesis and identification of the mixed ligands complexes of  $M^{+2}$  Ions in general composition ,[M(Leu)<sub>2</sub>(SMX)] Where *L-leucine*(C<sub>6</sub>H<sub>13</sub>NO<sub>2</sub>)symbolized (LeuH) as a primary ligand and Sulfamethoxazole (C<sub>10</sub>H<sub>11</sub>N<sub>3</sub>O<sub>3</sub>S)symbolized (SMX)) as a secondary ligand.

The ligands and the metal chlorides were brought in to reaction at room temperature in(v/v) ethanol /water as solvent containing NaOH. The reaction required the following [(metal: 2(Na<sup>+</sup> Leu<sup>-</sup>): (SMX)] molar ratios with M(II) ions, Were M(II) = Mn(II), Co(II), Ni(II), Cu(II), Zn(II), Cd(II) and Hg(II). The UV–Vis and magnetic moment data revealed an octahedral geometry around M(II), The conductivity data show a non-electrolytic nature of the complexes. The antimicrobial activities of ligands and their mixed ligand complexes were screened by disc diffusion method.

Keywords: Sulfamethoxazole(antibiotic), L-leucine, Mixed ligand, Metal complexes, Antimicrobial activity.

# **INTRODUCTION :**

L- leucine is one of the twenty major amino acids and is considered an essential amino acid [1]. (figure 1-Formula 1) It is branched chain amino acid and taken up by brain and muscle. In leucine metabolism, transamination gives -keto isocaproic acid, which is converted into corresponding CoA, this is similar to oxidative decarboxylation of alfaketoglutarate and pyruvate. The enzyme complex is very important in the body of living organism. A deficiency of the enzyme causes maple syrup urine disease. In this disease the urine gives odor of maple syrup or burnt sugar deterioration is rapid and results in mental retardation. [2].The amino acid L-leucine and various transition metals are important in the biological functions of humans, animals, and plants. [1-2].

 $(C_{10}H_{11}N_3O_3S)$  (IUPAC name = 4-amino-N-(5-methylisoxazol-3-yl)- Sulfamethoxazole 4-amino-N-(5-methylisoxazol-3-yl)-benzenesulfonamide (figure 1- Formula 2) is a sulfonamide bacteriostatic antibiotic. It is most often used as part of a synergistic combination with trimethoprim in a 5:1 ratio in co-trimoxazole, also It is commonly used to treat urinary tract infections. In addition it can be used as an alternative to amoxicillin-based antibiotics to treat sinusitis. It can also be used to treat toxoplasmosis.[4] It is also applicable for antiseptique, atitubercular and anti-inflammatory agent.





The investigation of metal sulfonamide compounds has received much attention due to the fact that sulfonamides were the first effective chemotherapeutic agents to be employed for the prevention and cure of bacterial infections in humans [4-5]. The sulphur containing ligands are well known for their anticarcinogenic, antibacterial,tuberculostatic, antifungal, insecticidal, and acaricidal activities [5]. Such sulfonamide derivatives widely used in clinical medicine as pharmacological agents with a wide variety of biological actions, were designed from the simple sulfanilamide lead molecule [6], also Schiff base known as anticancer and antiviral agents [7]., and its metal complexes have been widely studied because they have industrial, anti-fungal, anti-bacterial, anticancer herbicidal applications, anti-tubercular activities [8].N-Substituted sulfonamides are still among the most widely used antibacterial agents in the world, mainly because of their low cost, low toxicity, and excellent activity against bacterial diseases. [6], Literature survey shows that no studies on the synthesis and characterization of mixed ligand complexes of Leucine and Sulfamethoxazole(antibiotic) have been reported.

In this paper we present the synthesis and study of Mn(II),Co(II),Ni(II),Cu(II),Zn(II),

Cd(II) and Hg(II) complexes with amino acid (L-leucine) as a primary ligand and Sulfamethoxazole (antibiotic) as a secondary ligand.

- 2. Experimental
- 2.1. Materials and instruments

All chemicals were purchased from Merck / Aldrich. The reagents were used without further purification . Double distilled water was used.

b- Instruments: FT-I.R spectra were recorded as K Br discs using Fourier transform Infrared Spectrophotometer Shimadzu 24 FT-I.R 8400s. Electronic spectra of the prepared complexes were measured in the region (200- 1100) nm for  $10^{-3}$  M solutions in N, N-dimethylsulphoxide (DMSO) at 25 °C using shimadzu-U.V-160.A Ultra Violet Visible- Spectrophotometer with 1.000  $\pm$  0.001 cm matched quartz cell. While percentage of the metal in the complexes were determined by Atomic Absorption(A.A)Technique using Japan A.A-67G Shimadzu. Electrical conductivity measurements of the complexes were recorded at at room temperature for  $10^{-3}$  M solutions of the samples in (DMSO) using pw9527 Digital conductivity meter (Philips). Melting points were recorded by using Stuart melting point apparatus. "Magnetic susceptibility measurements were measured using Bruker magnet BM6 instrument at 298°K following the Farady's method. The proposed molecular structure of the complexes were drawing by using chem. office program, 3DX (2006).



# 2.2. Preparation of Complexes :

The complexes of the series  $[M (SMX)(Leu)_2]$ . were prepared by the following general method : (A) Sodium leucinate(Na<sup>+</sup> Leu<sup>-</sup>): The amino acid L-leucine [0.262 gm, 2 m mol] was dissolved in 10 ml H<sub>2</sub>O/ethanol (50%) mixture containing Na OH (2 m mol 0.8 g) in a flask and stirred at room temperature (20 °C)., the solution was deprotonated according to the Scheme (1).

# (B) General synthesis of the mixed ligand metal complexes[11]

A metal(II) chloride [(0.197g, MnCl<sub>2</sub>.4H<sub>2</sub>O, CoCl<sub>2</sub>.6H<sub>2</sub>O(0.237g,1mmol),NiCl<sub>2</sub>.6H<sub>2</sub>O (0.237g, 1mmol), CuCl<sub>2</sub>.2H<sub>2</sub>O(0.136g, 1mmol), ZnCl<sub>2</sub> (0.137g, 1mmol), CdCl<sub>2</sub> (0.183g, 1mmol),and HgCl<sub>2</sub> (0.271g,1mmol)]dissolved in in ethanol: water (1:1) 25ml respectively was added gradually with stirring to solution of Sodium leucinate (Na<sup>+</sup> Leu<sup>-</sup>).(0.253gm, 1mmole) of Sulfamethoxazole (SMX) was added to the mixture in each case by using stichiometric amount [(1:2:1) [(metal: 2(Na<sup>+</sup> Leu<sup>-</sup>): (SMX)] molar ratios, the above reaction mixture to raise the pH upto ~6.0 and the mixture was stirred for (20 -30mint)at room temperature. scheme (1) After one day a colored microcrystalline solid was obtained which was filtered. and washed with ethanol. The solid was recrystallized from a H<sub>2</sub>O/ethanol (50%) mixture. and dried in vacuum over anhydrous CaCl<sub>2</sub>. The yields range from 75 to 92 %. The decomposition temperatures range from: 216-340 °C.

# **3-Results and Discussion**

Resent study involves synthesis of seven new metal mixed ligand complexes with their Characterization using spectroscopic techniques and studying their Antibacterial Properties Application .A number of metallic complexes of L-leucine and Sulfamethoxazole with Mn(II),Co(II),Ni(II),Cu(II),Zn(II),Cd(II)and Hg(II) ions ,were synthesized from ethanol –aqueous medium 1:1 ratio ;General schem (1) of reaction as follow .The physical properties of the complexes are shown in (Table 1), All the complexes are colored, non-hygroscopic The complexes decomposed at high temperature on heating. These are insoluble in water or most of the organic solvents like methanol, benzene and carbon tetrachloride , DMF but soluble in DMSO.

The observed molar conductance ( $\Lambda M$ ) of 10<sup>-3</sup> solutions of the complexes in DMSO. lie in very low range (3-28)  $\Omega^{-1}$  cm<sup>2</sup>mol<sup>-1</sup> supporting their non-electrolytic behavior [9]. The atomic absorption measurements (Table-1-Calculated values in parentheses) for all complexes gave approximated values for theoretical values .The analysis data (Table-1) of metal complexes are consistent with their general formulation as 1:2:1, mixed ligand complexes of the type [M (Leu)<sub>2</sub> (SMX)]. Advances in Physics Theories and Applications ISSN 2224-719X (Paper) ISSN 2225-0638 (Online) Vol.27, 2014





Scheme (1): Preparation of [M(Leu)<sub>2</sub>(SMX)] complexes

Fourier-transform infrared spectra and m[of coordination:

The relevant vibration bands of the free ligands and the complexes are in the region  $400-4000 \text{ cm}^{-1}$  [8-15]. The most important infrared spectral bands that provide conclusive structural evidence for the coordination of the ligands to the central metal ions .

The assignment of some of the most characteristic FT-IR band of the complexes are shown in Tables (2-4) together with those of two ligands recorded for comparative purposes and facilitate the spectral analysis . In amino acids,  $\upsilon NH_3$ + appears in 3130-3030 cm<sup>-1</sup> region [9]. Table (2), displays the (FT-IR) spectrum for the (L-leucine )exhibited a band around  $\upsilon(3458)$  cm<sup>-1</sup> that corresponds to the stretching vibration of  $\upsilon(N-H) + \upsilon$  (O-H), while another strong absorption band at  $\upsilon$  (3055) cm<sup>-1</sup> is due to the  $\upsilon(N-H2)$ sym while the bands at (1581) cm<sup>-1</sup> and (1408) cm<sup>-1</sup> were assigned to the  $\upsilon(-COO)$ asy and  $\upsilon(-COO)$ sym respectively.  $\upsilon \Delta$  (-COO)asy-sym =171 cm<sup>-1</sup>. [8-10].

A general tendency in the relationship between v (COO<sup>-</sup>) (the difference between the wave numbers of the

asymmetric ( $\upsilon$  asym) and the symmetric ( $\upsilon$ sym) stretches of carboxylate group from the FT-IR spectra) and the types of coordination of the (COO<sup>-</sup>) group to metal ions by examining the structures.

In the experimental FT-IR spectra of leucinate the wave numbers of asymmetric and symmetric stretching vibrations of carboxylic anion change their position along the series of metals

M(II) = Mn(II), Co(II), Ni(II), Cu(II), Zn(II), Cd(II) and Hg(II) (see Table 4). Namely, the band of  $v_{asym}(COO^{-})$  is shifted toward higher wave numbers in spectra of mixed ligand, whereas the band of  $v_{asym}(COO^{-})$  is shifted towards lower wave numbers[11]. Thus, (L-leucine) molecule acts as bidentate monobasic ligands, coordinating through deprotonated carboxylic oxygen(-COO-) group, and its nitrogen of (-NH<sub>2</sub>) group [12, 13]. Table (3), displays the (FT-IR) spectrum for the (sulfamethoxazole).

In case of (SMX) molecule the  $\upsilon \delta$  (N–H) vibrations of –NH<sub>2</sub>(aromatic sec. amine) occur at

3468 and 3378 cm<sup>-1</sup> for free (SMX) due to vas (NH<sub>2</sub>) and vs (NH<sub>2</sub>),respectively. The hypochromic effect (decreasing in the intensity of v (NH) vibrations in case of mixed ligand complexes rather than (SMX) alone as well as the blue shifted in the wave numbers from 3299 cm<sup>-1</sup> [12, 13]

to range (  $3208 \text{ cm}^{-1}$ ) (mixed complex). Such these changes clearly indicate that the lone pair of electron of NH<sub>2</sub> and in sulfamethoxazole donor is participated in the complexation process with metals. acting as bidentate ligand. Where as sulfoxide (S=O) stretching occurs at , 1365 [v, vas(SO<sub>2</sub>], (1148and 1025)cm<sup>-1</sup>[s, vs(SO<sub>2</sub>)]. The band of C=C occurs at 1622-1469 cm<sup>-1</sup>, aromatic (C=C) at 1469 cm<sup>-1</sup> and alkenes (C=C) at 1622-1680 cm<sup>-1</sup>, cm<sup>-1</sup> [12]. and imine peak (C=N) 1622 cm<sup>-1</sup>. The v(C–N) of isoxazol-3-yl group occurs at 1309 cm<sup>-1</sup>, this band remains ~ (in the same region free ligand and in solid complexes. Thus, indicating its non involvement in coordination of the ligand with the metal ions .Finally, the bands at 1336 and 1162cm<sup>-1</sup> represent the asymmetric and symmetric stretching frequencies, respectively, of the sulfonyl group (S =O str) [12, 13]. In case of complexes, [M(Leu)<sub>2</sub>(SMX)], whereas sulfoxide stretching shifts to higher and lower frequencies respectively at ~106/55 cm<sup>-1</sup> due to coordination of the sulfonyl oxygen with metals in all the complexes. Hence in these complexes sulfamethoxazole, also acts as bidentate ligand coordinating through the oxygen of the sulphone group and nitrogen of the amine group. All the complexes are in agreement with octahedral geometry as proposed.[14,15]. In the IR spectra new bands have been observed in the region 600- 400cm<sup>-1</sup> which may be due to **v** (M-N) and **v** (M-O) bonds respectively. [14-17]

#### Magnetic Susceptibility:

Magnetic susceptibility was determined at room temperature using solid sample by Gouy method. Diamagnetic correction of metal- ligand system was calculated using the Pascal's constant. .The magnetic moments of the complexes shown in( Table -3) were calculated from the measured magnetic susceptibilities after employing diamagnetic corrections and revealed their diamagnetic nature as expected for Zn(II), Hg(II),Cd(II), with (3d<sup>10</sup>,4d<sup>10</sup>,5d<sup>10</sup>)configurations .The magnetic moment values o,[Mn(Leu)<sub>2</sub>(SMX)] lie in 6.182 BM indicating high spin octahedral stereochemistry of the complex. The magnetic moments of , [ Co (Leu)<sub>2</sub>(SMX)]complex lie in 4.635 BM which are in agreement with the generally accepted values for high spin octahedral Co(II) complexes. The magnetic moment values for the [Ni(Leu)<sub>2</sub>(SMX)]complex lie in 3,007 BM which reveals the octahedral stereochemistry of the ligands around Ni(II) ion. The observed magnetic moments of [Cu (Leu)<sub>2</sub>(SMX)] lie in 1.478BM showing one unpaired electron with paramagnetic nature and suggested a distorted octahedral geometry in teams of Jahn-Teller effect. [17- 18]

### **Electronic spectral :**

The (U.V- Vis) spectrum for the spectrum of Leucine show an absorption band at 305 nm (32786 cm<sup>-1</sup>) in (DMSO) solvent, this band is attributed to  $(\pi \rightarrow \pi^*)$ . The UV -Vis spectrum of the free ligand (SMX)in (DMSO) solvent appeared a high intense absorption bands at 275 nm (36363 cm<sup>-1</sup>) ( $\epsilon_{max}$ =1951 L. mol<sup>-1</sup>. cm<sup>-1</sup>), this band is attributed to ( $\pi \rightarrow \pi^*$ ).[19-20]

The parameters ligand Field Splitting Energy 10 Dq, Racah's Interelectronic Repulsion Parameters (B'), Nephelauxetic Ratio β and B' are calculated.[21]

The [Mn (SMX) (Leu)<sub>2</sub>] complex with d5 configuration exhibits absorption bands in the region 11135, 11402 & 19083 cm<sup>-1</sup>. These bands may be attributed to number of spin forbidden transitions involving 6A1g ground and several higher energy quartet states, which were in consistent with octahedral geometry around Mn (II) metal ions. 10 D q =11.135,  $\upsilon 1/\upsilon 2 = 0.97$ ,  $\upsilon 2/\upsilon 1 = 1.02$  from Tanabe-Sugano diagram for d<sup>5</sup> octahedral field, These values are found in accordance as reported in the literature for Mn(II) octahedral complexes.

The (U.V- Vis) [Co (SMX) (Leu)<sub>2</sub>] spectrum, exhibits four peaks , the first high intense peak at (257 nm)( 38910 cm<sup>-1</sup>)( $\varepsilon_{max}$  =1225 molar<sup>-1</sup>. cm<sup>-1</sup>), is due to the ligand field, and at 11135, 11402 and 19083 cm<sup>-1</sup>, there are assigned to 4T1g(F)  $\rightarrow$  4T2g(F) (v1) 4T1g(F)  $\rightarrow$  4A2g(F) (v2) and 4T1g(F)  $\rightarrow$  4T1g(p) (v3) transition respectively, which are characteristic of octahedral stereo geometry : .[ 22-24]. ,v1/v2 = 0.97, v2/v1=1.02 from Tanabe-Sugano diagram for d<sup>7</sup> octahedral field the value of 10Dq equal to 11.135 cm<sup>-1</sup>, B'=950 cm<sup>-1</sup>,  $\beta$  =0.997 cm<sup>-1</sup>. The Co(II) complex under present investigation process interelectronic repulsion parameter (B') 950 cm<sup>-1</sup>. The Racha parameter (B) is less than free ion value (973) suggesting a considerable orbital overlap and delocalization of electrons on the metal ion. The nephelauxetic ratio ( $\beta$ ) for the present Co(II) complex (0.997). This is less than one, suggesting partial covalence in the metal ligand bond. [23-25] suggest the octahedral geometry .

(U.V- Vis)  $[Cu(SMX) (Leu)_2]$  spectrum, exhibits two peaks, the first high peak at (265 nm)(37735 cm<sup>-1</sup>)( $\epsilon_{max} = 1905$  molar<sup>-1</sup>. cm<sup>-1</sup>) is due to the (CT), while the second weak broad peak at (610 nm)(16393 cm<sup>-1</sup>)( $\epsilon_{max} = 112$  molar<sup>-1</sup>. cm<sup>-1</sup>), which assigned to (2Eg $\rightarrow$ 2T2g), transition. Hence the Cu(II) mixed ligand complexes showed distorted octahedral geometry.

The (U.V- Vis) spectrum of [Ni (SMX) )(Leu)<sub>2</sub>], exhibits five peaks , the first middle peak at (251 nm)(37453cm<sup>-1</sup>)( $\varepsilon_{max}$  =1391 molar<sup>-1</sup>.cm<sup>-1</sup>) is due to the ligand field and the second high peak at (263 nm)( 38022 cm<sup>-1</sup>)( $\varepsilon_{max}$  =1.248 molar<sup>-1</sup>.cm<sup>-1</sup>) is due to (C.T), third, fourth, and fifth peaks assignment of the electronic spectral bands for, Ni(II) are 3A2g  $\rightarrow$ 3T2g(F)  $\upsilon$ 1=(891nm) 11223 cm<sup>-1</sup>)( $\varepsilon_{max}$  =72 molar<sup>-1</sup>. cm<sup>-1</sup>), 3A2g (F)  $\rightarrow$ 3T1g(F)  $\upsilon$ 2=(885 nm)( 11299 cm<sup>-1</sup>)( $\varepsilon_{max}$  =56 molar<sup>-1</sup>. cm<sup>-1</sup>), and at (873 nm)

(11454cm<sup>-1</sup>)( $\epsilon_{max} = 5 \text{ molar}^{-1} \text{ cm}^{-1}$ ) can be assigned to the  $,3A2g(F) \rightarrow 3T1g(p) (\upsilon 3), \upsilon 2/\upsilon 1 = 1.006, \upsilon 1/\upsilon 2 = 0.99, 10Dq=11.223, B'=966 \text{ cm}^{-1}, \beta=0.85$ , typical of Ni(II) ground state 3A2g(F)] octahedral complexes.

The  $\upsilon 2/\upsilon 1 = 1.006$  value are found in the usual range (1.056-1.082) reported for the majority of octahedral Ni(II) compounds. [21]

The electronic spectra of d<sup>10</sup>[Zn(II) ,C d(II)and Hg(II)]complexes do show the charge transfer , and the magnetic susceptibility shows that all complexes have diamagnetic moments., because d-d transitions are not possible hence electronic spectra did not give any fruitful information. in fact this result is a good agreement with previous work of octahedral geometry. [21-22]

#### **Antibacterial Activities studies:**

The zone inhibition of bacterial growth were measured in mm depending upon the diameter as shown in Table (6) Figure. (2)

Table 6 reveal that the synthesized compounds were potent as bacteriostatic agents.

The synthesized metal complexes were screened for their antimicrobial activity by well plate method in nutrient agar . The invitro antibacterial activity was carried against 4 hold cultures of pathogenic bacteria like gram (+)and gram (-) at  $37^{\circ}$  C. In order to ensure that solvent had no effect on bacteria, a control test was performed with DMSO and found inactive in culture medium. All the metal complexes showed good activity against gram (+)*Acineto* .Ni(II) complex have no effect on E-coli., *staphylococcus* and *Psedomonas*. *S-aureus* and gram (-) Except Ni(II) complex. Table (5).while the Mn(II) complex was active against two organisms *staphylococcus* and *Acineto*. The *staphylococcus* was affected vigorously by mixed ligand, Hg(II), Co(II), Cu (II) ,Zn(II)and Cd(II) respectively. *E-coli*. was moderately affected by the ligands, Zn(II),and Co(II), complexes.[23-24] .In metal complexes, on chelation the polarity of the metal ion with donor groups. Further, it increases the delocalization of  $\pi$ - electrons over the whole chelate ring. This increased lipophillicity enhances the penetration of the metal complexes into lipid membranes and block the metal binding lists in the enzymes [25] .

# **CONCLUSION:**

We have successfully synthesized the mixed ligand complex of

M (II)=Mn(II),Co(II),Ni(II),Cu(II),Zn(II),Cd(II)and Hg(II) containing O-N donor ligands. The complex was also characterized by molar conductance, magnetic susceptibility measurement and also by FT- IR, UV- visible spectroscopy. The UV–Vis and magnetic moment data revealed an octahedral geometry around M(II), (schem1 -3D).The conductivity data show a non-electrolytic nature of the complexes.

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	Table 1 : Analytical and some physical data of the complexes									
Ν	Complexes	Formula	Color	Yield	Decomposition	Λm	Metal%			
0.		weight		%	temperatures	$\Omega^{-1}$ .cm <sup>2</sup> .mole <sup>-1</sup>	(theory)			
		g/mol			°C					
1	[Mn(SMX)(Leu) <sub>2</sub> ]	569.55	Brown	86	261-264	28	(9.65)			
							9.98			
2	[Co(SMX)(Leu) <sub>2</sub> ]	573.54	Viol	75	254-258	19	(10.28)			
							11.08			
3	[Ni(SMX)(Leu) <sub>2</sub> ]	573.30	Green	78	240-247	18	(10.24)			
							8.63			
4	[Cu(SMX)(Leu) <sub>2</sub> ]	577.16	Blue	92	241-250	18	(10.99)			
							8.63			
5	[Zn(SMX)(Leu) <sub>2</sub> ]	580.02	White	80	216-220	17	(11.28)			
							9.98			
6	[Cd(SMX)(Leu) <sub>2</sub> ]	627.02	White	85	280-285	11	(17.93)			
							15.37			
7	[Hg(SMX)(Leu) <sub>2</sub> ]	715.20	White	90	291-296	3	(28.05)			

Table 2-FT-R spectral data of Leucine										
Leucine	υ (N-H)+ υ (O-H)	U (N-H <sub>2</sub> )sym	Δυ	υ (C-H) + ; CH <sub>3</sub>	υ (-COO)asy	υ (-COO)sym	υΔ (-COO) asy-sym			
C <sub>6</sub> H <sub>13</sub> NO <sub>2</sub>	3417m	3055s	347	2958,2874m	1516vs	1408vs	108			

Table 3-FT-R spectral data of         Sulfamethoxazole											
vas	υ	υ	υ	υδ def	υ	υSO <sub>2</sub>	υ	υ	υ	υ	υ
(N-H);	(C-H);	(C-H)	(C=C)	(N-H)	С–Н	asy	C–N	(C-O)	(SO <sub>2</sub> )	(S-N	(C-S)
-us NH <sub>2</sub>	aroma	+		Ring					sy		
& –NH	ticy	; CH <sub>3</sub>		breathing							
				bands							
3468vas	3378	2929 w,	1622	1597 vs	1504	1365 s	1309	1266	1157	987w	831
3300 <b>us</b>	3143 s	2858	vs		1469		S	ms	1143		vs
									1091		

	Table 4-FT-R spectral data of complexes												
No.	vas	υ (С-Н)	υ С–Н	υ	υδ def	υ С–Н	υ	υ	υ	υ	υ	M-N	М-О
	(N–H);	aromaticy	+	C=C	(N–H)	defor	$SO_2$	COO) <sub>a</sub>	-COO)	C–N	(SO <sub>2</sub> )		
	us NH <sub>2</sub>		; CH3		Ring	mation	asy	sy	sym		sy		
	& –NH				reathing								
					bands								
1	3419m	3392s	2956 vs,	1660	1581	1516 s	1384	1408	13611	1307	1172 m	515	497
Mn		, 3271vs	2872w	w	vs	1469 s			m	w	1049	522	
											m		459
											1025		
											mw		
2	3460s	3394vs	2958	1654	1621vs	-	1371	1467s	1333	1313	1179 m	640	574
Со		,3234	,2878w	vs						w		601	
3	3459vs	3298vs	2953vs	1635	1595vs	1469w	1365	1471s	1346s	1307	1120 m	609	522
Ni			,2929s	vs						w	1091 s	584	497
											1058m	m	
4	3387m	3246vs,	2958vs	1620	1566s	1494m	s	1496m	1375s	1307	1136vs	570	522
Cu	3317vs	3128s	,2929s	vs		1454m	1396	s		w	1111vs	557	w
											1053m		495
													w
5	3481w	3302vs	2958vs	1612	1560s	1469s	s	1452s	1344s	1307	1155m	642	561
Zn	3325s		,2928s	vs			1411			m	1116vs	586	497
			2870s								1078vs		
6	3350vs	3252vs	2956vs	1591	1560s	1471s	s	1452s	1342s	1305	1193m	651	555
Cd			,2920s	vs			1471			m	1122s	m	
			2874v								1078vs	599	
												m	
7	3470vs	3336	2926m	1674	1531vs	1502vs	1475	1431s	1359m	1307	1168m	684	530
Hg	3375vs	3242s		vs			s			m	1134	545	505
				1631							1078		



Table 5- Electronic Spectral data, magnetic momen, of the studied compounds									
Comp.	λnm	ABS	υ' ( cm <sup>-1</sup> )	Assignments	µeff (BM)	geometry			
C6H13NO <sub>2</sub> (leu)	305	1.951	32786	$\pi \rightarrow \pi^*$	-	-			
SMX	275		36363	$\pi \rightarrow \pi^*$	-	-			
[Mn(SMX) (Leu) <sub>2</sub> ]	246	1.362	42344	Lf	6.182	Octahedral			
	350	0.209	19083	$6A1g \rightarrow 4A1g$ (G) (v3)					
	881	0.128	11402	$6A1g \rightarrow 4T2g$ (G)(v2)					
	894	0.127	11135	$6A1g \rightarrow 4A1g$ (G))(v1)					
[Co (SMX) (Leu) <sub>2</sub> ]	257	1.225	38910	Lf	4.635	Octahedral			
	524	0.033	19083	$4T1g(F) \rightarrow 4T2g(F)(v3)$					
	877	0.013	11402	$4T1g(F) \rightarrow 4A2g(F)(v2)$					
	898	0.013	11135	$4T1g(F) \rightarrow 4T1g(p)(v1)$					
[Ni (SMX) )(Leu) <sub>2</sub> ]	251	0.544	38022	Lf	3.007	Octahedral			
	263	1.248	11454	СТ					
	873	0.054	11299	$3A2g(F) \rightarrow 3T1g(p)(v3)$					
	885	0.056	11223	$3A2g(F) \rightarrow 3T1g(F)(v2)$					
	891	0.072		$3A2g \rightarrow 3T2g(F)(v1)$					
[Cu (SMX) (Leu) <sub>2</sub> ]	265	1.905	37735	СТ	1.478	distorted			
	610	0.112	16393	$2Eg \rightarrow 2T2g$		Octahedral			
[Zn (SMX) )(Leu) <sub>2</sub> ]	268	0.355	37313	СТ	Dia	Octahedral			
[Cd (SMX) )(Leu) <sub>2</sub> ]	265	2.794	37735	СТ	Dia	Octahedral			
[Hg (SMX) )(Leu) <sub>2</sub> ]	262	1.419	38167	СТ	Dia	Octahedral			

Table 6- Represent the antimicrobial activity of         compounds									
Comp. Test Bacteria	E-coli.	staphylococcus	Psedomonas	Acineto					
SMX		40	27	45					
LeuH				23					
[Mn(SMX)(Leu)		21		21					
[Co((SMX) (Leu) <sub>2</sub> ]	10	26	16	17					
[Ni((SMX))(Leu) <sub>2</sub> ]				19					
[Cu((SMX) )(Leu) <sub>2</sub> ]		25	13	16					
[Zn((SMX))(Leu) <sub>2</sub> ]	8	20		17					
[Cd((SMX) )(Leu) <sub>2</sub> ]		18	18	21					
[Hg((SMX))(Leu) <sub>2</sub> ]	21	35	23	24					

