Synthesis and Theoretical Study of 4-(2-methyl-4oxoquinazoline-3(4H)-yl) Benzoic acid with Some Transition Metal Ions

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Abstract:

New complexes of the type $[ML_2(H_2O)_2]$, $[FeL_2(H_2O)Cl]$ and $[VOL_2]$ were M=Co(II),Ni(II) and Cu(II), L=4-(2-methyl-4-oxoquinazoline-3(4H)-yl) benzoic acid were synthesized and characterized by element analysis, magnetic susceptibility, molar conductance ,FT-IR and UV-visible. The studies indicate that the L acts as doubly monodentate bridge for metal ions and form mononuclear complexes. The complexes are found to be octahedral except V(IV) complex is square pyrimde shape . The structural geometries of compounds were also suggested in gas phase by theoretical treatments, using Hyper chem-6 program for the molecular mechanics and semi-empirical calculations, addition heat of formation(ΔH_f °) and binding energy (ΔE_b) for the free ligands and it's metal complexes were calculated by using PM3 method .PM3 was used to evaluate the vibration spectra of ligand and compare the theoretically calculated wave numbers with experimental values ,the theoretically obtained frequencies agreed calculation helped to assign unambiguously the most diagnostic bands.

Key word:Quinazoline, complexes of oxoquinazoline,Spectral studies

Introduction:

Heterocyclic compounds are organic compounds containing at least one atom of carbon ,and at least one element other than carbon .such as sulfur, oxygen or nitrogen within a ring structure[1,2].Heterocyclic that contains nitrogen atom is an important class of compounds, that displays a wide variety of biological activities ;therefore, these structures have received special attention in a considerable synthesis[3,4]. Much been paid to the attention has benzodiazepines, oxazepane, oxazepine, thiazepine and oxazine,for which abstracts have been reported in the libraries [5,6]. Among a wide variety of nitrogen heterocyclic that have been explored for developing pharmaceutically important molecules,

the quinazoline have played an important role in medicinal chemistry and subsequently have emerged as a pharmacophore[7].Recently ,nitraquazone, a quinazoline derivative has been found to possess potent phosphodiesterase inhibitory activity[7].

Material and Methods :

A-Materials, physical measurements and analysis :

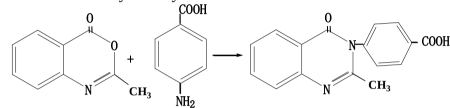
The chemicals used in this work BDH. were obtained from and Fluka. They were pure grade reagents . Metal contents of the complexes were determined using Shimadzu а A.A680G atomic absorption Spectrophotometer. FTIR spectra were using Shimadzo-8000 recorded spectrophotometer using CsI discs in

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the range (4000-200) cm⁻¹. Electronic spectra were recorded using Shimadza UV-Visible (160A) ultra violate spectrophotometer at wave length in the range (1100-200)nm. Electrical conductivity was measured using Coring conductometer and magnetic susceptibility was measured bv Johnson Matting Catalytic System Division and melting point was measured using Gallencamp M.F.B 600.01 . Elemental (C,H and N) analyses were carried out on a EUROEA-Elemental analyzer Italya.

B-Praperation of ligand:-

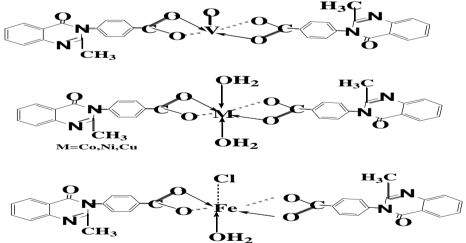
A mixture of 2-methyl -4-(H)-3,1benzoxazizin -4-one (1.79g,10mmol) ,para amino benzoic acid (10mmol) and pyridine (15 ml) was heated for 3 hours. Pyridine was evaporated at reduced pressure and the residue was first treated with dilute hydrochloric acid and then with aqueous (1 g) of sodium hydroxide .The product crystallized from heptanes to give 1.98 g (84%) of (L) [8].



C- Preparation of metal complexes

An ethanolic solution of ligand(L) (0.45, 0.02 mole) was added to a hot ethanolic and potassium hydroxide solution of metal chloride (0.01 mole). The reaction mixture was refluxed on water bath for 4h until a

precipitate appeared after cooling, the solution product was filtered off, washed with water and then little hot ethanol apparent dryness and dried in vacuum over anhydrous calcium chloride in a desiccators(yield 60-75%).



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Compound	Color	Melting	Yield%	C%	H%	N%	Metal%	Molar
_		point		Calcd	Calcd	Calcd	calculate	conductance
		_		(exp)	(Exp)	(Exp)	(found)	$Ohm^{-1} cm^2$
								mol^{-1}
L	White	190-192	86	68.5	4.28	10.00	-	-
				(67.9)	(3.98)	(11.02)		
$[Co(L)_2(H_2O)_2]$	Light-	182-184	74	4.90	4.97	0.61	9.02	15.15
	Green			(5.88)	(3.99)	(0.71)	(10.00)	
$[Ni(L)_{2}(H_2O)_2]$	Brown	178-180	75	4.91	4.30	0.61	9.00	17.38
×				(4.02)	(5.22)	(0.65)	(8.87)	
$[Cu(L)_2(H2O)_2]$	Light-	156-158	61	4.88	4.27	0.61	9.99	19.47
	Green			(5.74)	(5.02)	(0.55)	(10.69)	
$[Fe(L)_2(H_2O)Cl]$	Dark-	122-124	76	4.20	4.20	0.60	8.38	13.89
	Brown			(3.18)	(3.20)	(0.51)	(9.34)	
[VO(L) ₂]	Olive	132-134	67	5.13	3.21	0.64	8.17	16.36
	Olive			(6.09)	(4.09)	(0,59)	(9.16)	
	green							

 Table (1) shows the physical properties of prepared complexes

Hyper Chem.

Hyper Chem is a sophisticated molecular modeling environment that is known for its quality ,flexibility ,and ease of use.Uniting 3D visualization and animation with quantum chemical calculation ,molecular mechanics ,and dynamics,Hyper Chem puts more molecular modeling tools at your fingertips than any other windows program.[9].

Result and Discussion

All the complexes were sparingly soluble in common organic solvents but soluble in DMF,DMSO and acetonitrile.The analytical data indicates that the complexes are agree well with 1:2 metal to ligand stoichiometry shown in (Table1).The observed molar conductance values measured in DMF solution fall in the range (13.89-19.47) ohm⁻¹ cm² mol⁻¹ These observed values of the molar conductance are well within the expected range for non-electrolytes (Table1).

Magnetic Susceptibility:-

The magnetic moments obtained at room temperature for the complexes of Co(II),Ni(II) and Cu(II) are listed in (Table3).The cobalt complex shows magnetic moment of 4.86 B.M .the spin free octahedral complex of Co(II) is reported to exhibit magnetic moment range in the of 4.46-5.53B.M.[10]Hence observed magnetic moment for the Co(II)under study indicates that it has an octahedral configuration. The nickel complex shows magnetic moment of 2.95 B.M.The magnetic moment of octahedral Ni(II) complex is reported to exhibit magnetic moment in the range of 2.80-3.40B.M including spin orbital coupling[11]. Hence the observed magnetic moment for the Ni(II) complex suggests that it may have octahedral geometry. Copper complex exhibits magnetic moment 1.95 B.M.which is including the normal value [12] exhibit magnetic moment in the range of (1.84-2.20)B.M. including spin orbital coupling , suggests that it may have octahedral geometry.

Infrared spectroscopy:-

The FTIR spectrum for the ligand (L), shows a characteristic stretching absorption bands at 3186 cm^{-1} , 1693 cm^{-1} and 1362 cm^{-1} assigned to hydroxyl group. υ (COO)sym.. (COO)asym., υ respectively [13, 14]. The v COO stretching vibrations are important to predict the bonding mode of the ligand. The values of $[\Delta v = (COO)_{asym}$. υ (COO)_{sym.}] can be divided into 3

groups;(a)In compounds where Δ v cm⁻¹,the (COO)>350 carboxylate monodentate group binds in a fashion.However .other very weak intra- and intermolecular interactions cannot be excluded.(b)When Δυ (COO) <200 cm⁻¹, the carboxylate groups of these compounds can be considered to be bidentate.(c).In compounds where $\Delta \upsilon$ (COO) > 200 cm^{-1} and $< 350 cm^{-1}$ an intermediate between monodentate state and bidentate (anisobidentate) occurs. It has also been suggested that the Δv (COO) in the chelating mode is less than the Δ υ (COO) in a bridging mode[15,16].

The disappearance of hydrogen from hydroxyl group on complexation indicates that the complexation is through the oxygen atom.Streching of metal-oxygen bands of the complexes appeared in low frequency region [17] (467-421)cm⁻¹ .The IR data of the ligand and its complexes are shown in Table(2).In the present study of all prepared complexes of the ligand in the presence of weak band around 845-850 cm⁻¹ ,which may be assigned due to the presence of coordination water molecule in the complexes[18].

Table(2)The main diagnostic absorption bands of ligand and its complexes (cm⁻¹)

	U		0			
Compound	υOH	υ (COO) _{asy}	υ (COO) _{sym.}	υ M-O	υ M-Cl	Others
L	3186	1693	1362	-	-	-
$[\operatorname{Co}(\mathrm{L})_2(\mathrm{H}_2\mathrm{O})_2]$	-	1680	1345	467		. δ _{H2O} =845
						υ _{H2O} =3441
$[\mathrm{Ni}(\mathrm{L})_{2}(\mathrm{H}_{2}\mathrm{O})_{2}]$	-	1677	1340	432		δ_{H2O} =840
						υ _{(H2O} =3313
$[Cu(L)_2(H_2O)_2]$	-	1675	1375	421		$\delta_{H2O}=848$ $\upsilon_{(H2O)}=3387$
[Fe(L) ₂ (H ₂ O)Cl]	-	1710	1390	443	367	$\delta_{H2O} = 850$ $\upsilon_{(H2O)} = 3417$
[VO(L) ₂]	-	1671	1339	453		V=O = 972

Electronic Spectra:-

The electronic spectral data of Co(II),Ni(II),Cu(II),Fe(III) andV(IV) complexes were recorded in DMF as shown in Table(3). The Co(II) complex (light green), the electronic absorption bands appears at 9985 and 20089 cm⁻¹ ${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}A_{2}g$.due to and ${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}T_{1}g_{(P)}$ transition respectively in an octahedral environment[19].The electronic spectrum of Ni(II) complex shows two bands at 10810 and 25316 cm⁻¹ assignable to ${}^{3}A_{2}g \rightarrow {}^{3}T_{2}g_{(F)}$ and ${}^{3}A_{2}g \rightarrow {}^{3}T_{1}g_{(P)}$ transition respectively in octahedral environment[20].The light green colored of Cu(II) complex exhibits a broad band in the region

16395-12195 cm⁻¹ with maxima at 12210 cm⁻¹ in a distorted octahedral geometry [21]. The broadness of the band may be due to dynamic Jann-Teller distortion and is assigned to $^{2}\text{Eg}\rightarrow^{2}\text{T}_{2}\text{g}$ transitions. The electronic spectrum of Fe(III) complex shows three bands at 10857,15822 and 21740 cm^{-1} assignable to ${}^{6}A_{1}g \rightarrow {}^{4}T_{1}g(G), {}^{6}A_{1}g \rightarrow {}^{4}T_{2}g(G)$ and $^{6}A_{1}g \rightarrow ^{4}A_{1}g + ^{4}Eg(G)$ [22].The Vanadium complex colored of the electronic absorption bands appears at 13888 cm^{-1} and 18181 cm^{-1} , due to $^{2}B_{2}g \rightarrow ^{2}Eg$ and $^{2}B_{2}g$ $\rightarrow^2 B_1 g$ transition[23].

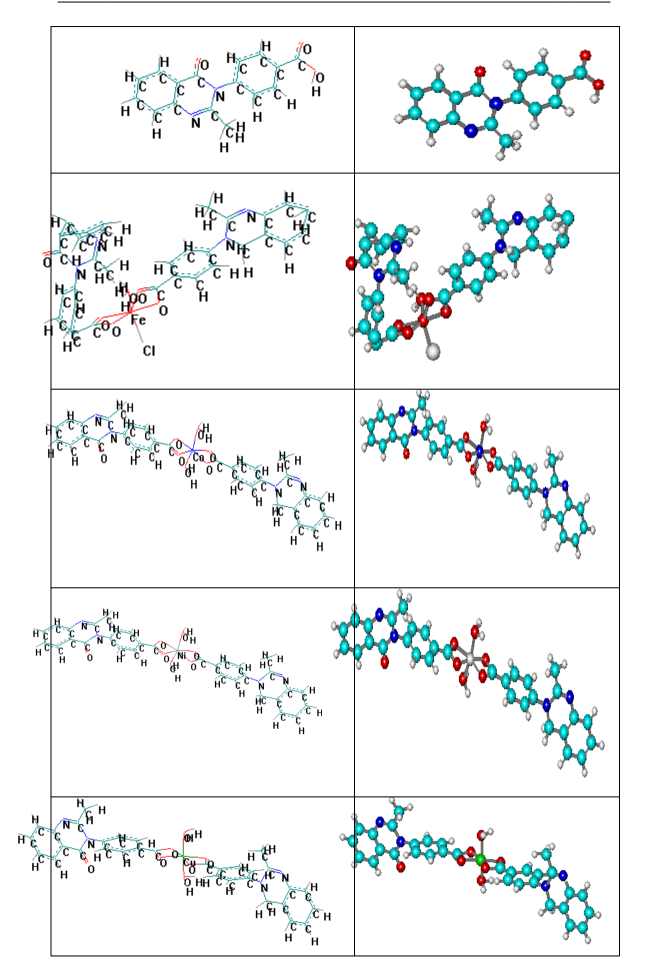
Suggested for the present prepared metal complexes of ligands								
Compound	Wave nu	(d-d) Transitions	В	Dq	10Dq	B	Structure	µeff
	mber cm- ¹	Transitions					Suggest	(BM)
L	325	-	-	-	-	-	-	-
	433							
$[Co(L)_2(H_2O)_2]$	9985	${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}A_{2}g,$	930	825	8250	0.958	octahedral	4.86
	15439 _(Cal.)	${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}A_{2}g_{(F)}$						
	20089	${}^{4}T_{1}g_{(F)} \rightarrow {}^{4}T_{1}g_{(P)}$						
$[Ni(L)_{2}(H_2O)_2]$	10810	$^{3}A_{2}g \rightarrow ^{3}T_{2}g_{(F)}$	834	932	9320	0.820	octahedral	2.95
	15151 _(Cal.)	${}^{3}A_{2}g \rightarrow {}^{3}A_{1}g_{(F)}$						
	25316	$^{3}A_{2}g \rightarrow ^{3}T_{1}g_{(P)}$						
$[Cu(L)_2(H_2O)_2]$	12210	$^{2}\text{Eg}\rightarrow^{2}\text{T}_{2}\text{g}$	-	-	-	-	octahedral	1.95
Fe(L) ₂ (H ₂ O)Cl]	10857	${}^{6}A_{1}g \rightarrow {}^{4}T_{1}g_{(G)}$					octahedral	
	15822	$^{\circ}A_1g \rightarrow {}^{4}T_2g_{(G)}$	1300	120	1200	0.53		5.01
	21740	$^{6}A_{1}g \rightarrow ^{4}A_{1}g + ^{4}Eg_{(G)}$						
[VO(L) ₂]	13888	$^{2}B_{2}g \rightarrow ^{2}Eg$					Square	1.53
	18181	$^{2}B_{2}g \rightarrow ^{2}B_{1}g$					Pyrimde	

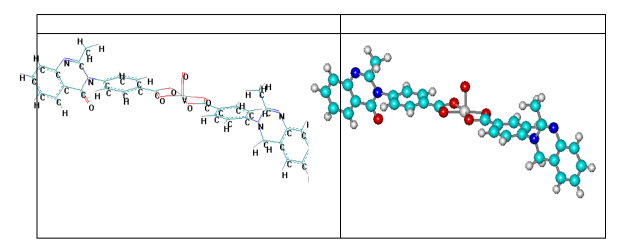
Table(2) Electronic spectra , magnetic moment(B.M) and The Structure Suggested for the present prepared metal complexes of ligands

HOMO and LUMO orbitals:-

(HOMO) to highest occupied molecular orbital and (LUMO) lowest unoccupied Molecular orbital .These are called orbitals the frontier orbitals, and determine the way the molecule interacts with other species. The HOMO is the orbital that could act as an electron donor, since it is the outermost(highest energy) orbital

containing electrons. The LUMO is the orbital that could act as the electron acceptor ,since it is the innermost(lowest energy)orbital that has room to accept electrons. In accordance with the above definitions, a single orbital may be both the LUMO and HOMO[24].





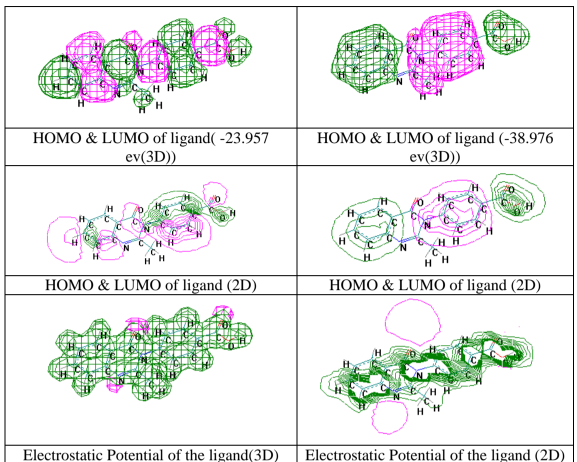
Fig(1) Conformational Structure of the ligand and their Complexes Table (4): Conformation energetic in $(KJ.Mol^{-1})$ for imines (L^{1}) and its metal

 Table (4): Conformation energetic in (KJ.Mol⁻¹) for imines (L¹) and its metal complexes

Conformation		PM3			
Comormation	$\Delta \mathbf{H}_{f}^{\circ}$	$\Delta \mathbf{E}_{b}$			
L	12.378933	-2198.8560667			
$[Co(L)_2(H_2O)_2]$	-65.0953082	-4280.6713082			
[Ni(L) ₂₍ H ₂ O) ₂]	- 8312.777312	-12695.2143127			
$[Cu(L)_2(H_2O)_2]$	-94.3439130	-4359.7960870			
Fe(L) ₂ (H ₂ O)Cl]	- 9238.608342	-13688.2983428			
[VO(L) ₂]	- 163.4977939	-244.507529544			

Table (5): Comparison of experimental and theoretical vibration Frequencies for the L^1 .

Comp.	υ (COO)asym.	υ (COO)sym.	υOH
L^1	(1693)* (1705)** (-0.70)***	(1369)* (1333)** (2.62)***	(3186)* (3100)** (2.00)***



Fig(2) HOMO & LUMO and Electrostatic Potential as 2D&3D Contours for the ligand

References:

- 1-Jrjun K.R. ,Nayan R.P.,Harish R.D. and Sunil S.N .2009. Novel synthesis and characterization of thiosemicarbazone compounds containing 4-acyl -2pyrazolin -5-ones.E-J. of chem...6 (3) 747-752,.
- 2-Vashi, R.T.V.Shelat.,C.D and Himanshu P.2010. Synthesis and Antifungal Activity of 6-Bromo-2[(4-(2,3Dichloro Phenyl))Piperazine -1yl)Methyl]-3-[8-Hydroxy Quinoline -5-yl]-3-Quinazoline-4-one Ligand and Transition its Metal Chelates.International J. Appl Bio. Pharma. Techn. 1 (3) 885-889.
- 3- Siddappa, K.Tukaram R., Mallikarjun ,M. and Reddy ,CV. 2008. Synthesis,Characterization and Antimicrobialu Studies of 3-[2-Hydroxy-quinolin-3-ylmethylene)amino-2-phenyl-3H-quinazolin-4-one

and its metal(II) Complexes"E-J. chem. 5 (1) 155-162,.

- 4-Varsha J., Pradeep M., and Sush K.2008. Synthesis and CNS depressant activity of some novel 3-[5-substituted 1,3,4thiadiazole-2-yl]-2-styryl quinazoline -4(3H)-ones"Eur. J. med. chem..43(1) 135-141.
- 5-Valerian D.and lleana D.2009. "Ruthenium Complexes Bearing N-Heterocyclic Carbene (NHC) Ligands"Pt. Metal Rev..49(4) 183-188.
- 6- Pandey R.N. and Nag A.K. 2009. spectral characterization of some oxometal complexes of 2-mercapto-3butyl –quinazoline-4-one.rasayan j.chem.,2 (4) 990-993.
- 7-Cathleen M.Crudden and Daryl P. Allen.2004. Stability and reactivity of N-heterocyclic d
- 8- Rabilloud G. and Sillion. B.1980.Synthesis of 4H-3,1-

- Benzoxazin -4-one and 4-(3H) Quinazolines from Anthranilic Acids and their Derivative by the Use of Triphenyl Phosphite and Pyridine. j.Hetr.Cycl.chem 17(29)1065-1068.
- 9- Mueller ,M. 2002. Fundamentals of Quantum Chemistry.,Kluwer Academic publishers, New York.
- 10- Nicholls, N. 1973. The chemistry of iron, cobalt and nickel .pengamou press, ltd Heading Ton Hill Hall, oxford, Eng. p. 1053, 1113.
- 11- Geary ,WJ. 1971. The use of conductivity measurements inorganic solvents for the characterization of coordination compounds.Coord. Chem. Rev. 7(1):81-122.
- 12-Sunmaz,M. and Sekerci,M. 2002. Synthesis and characterization of Cu(II),Co(II),Ni(II)and Zn(II)Schiff base complexes from 1-amino-5benzoyl -4-phenyl –H-pyrimidine -2one with salicylaldehyde.Polish J.Chem.76 (2) 907- 914.
- 13- Bakir, Sh.R.2007. Thesis,MSc ,College of Science for women ,University of Baghdad .Synthesis and Characterization of some new Schiff Bases and Their Complexes with Selection of Elements and Their Theoretical Study.
- 14 -.Bakir Sh. R., Kadhim N. J.and Alias M. F..2010. Synthesis, Structure Study and Theoretical Treatment of New Some Metal Complexes With 2hydroxy-4-nitro Phenyl 2-N(4-N,N dimethyl)Benzyliden. Al-Mustans. J.Sci.21 (2) 41-56.
- 15- Yousif E.and Rentschler E.2010 . Synthesis and characterization of some metal ions with [5-(4-cholrophenyl)-1,3,4-oxadiazol -2-yl]thio acetic acid. J. of Al-Nahr. univers. 13(2) 86-92.
- 16- El-Said A. I., A. Aly A. M., El-Meligy M. S. and Ibrahim M. A..2009. Mixed Ligand Zinc(II) and Cadmium(II) Complexes Containing Ceftriaxone or Cephradine Antibiotics

and Different Donors, The Journal of the Argentine Chemical Society Vol. 97 No 2,149-165.

- 17- Nakamoto, N. 2009. Infrared and Raman Spectra of Inorganic and Coordination Compounds. 6th Ed,Part2 John wiley and Sons,Inc.,New Jersy
- 18- Sliverstein, R.M. and Webser ,X.F. 2005. Spectrometric Identification of Organic Compounds . 7th Ed., Jon Wiley and Son's Inc, USA .
- 19- Soloman, E.I. and Lever, A.B.P.2006. Inorganic Electronic Structure and Spectroscopy.Applications and Case Studies.Vol.II,John Wiley and Sons.Inc, New York Chester ,Singapore,Toranto.
- 20-.Huheey J.H. 1994. Inorganic Chemistry Principle of Structure and Reactivity. Harper
- internation, Edition, Harper and Row papar And Row publishers, New York,.
- 21- Shriver D.F. and Atkins P.W. 2001. Inorganic Chemistry .3rd ed ., oxford university press.
- 22-Vashi,R.T.,and Shelat,C.D. 2010. Transition Metal Complexation Studies on Hetereo cyclic Ligands. Asian J.Chem.22(3) 1745-1750.
- 23- Vashi R.T. and Patel S.B.2009. Synthesis, Characterization and Novel Antifungal Activity of Quinazoline 4-One Derivatives Containing 8-Hydroxo Quinazoline Ligands and its Various Metal Complexes. E-J.Chem.6(S1)S445-S451.
- 24-, Bakir Sh.R., Kareem T.A.A., and, Alias M.F. 2010. Synthesis, structural study and antibacterial activity of some new complexes with 2-N(4-N,N-dimethyl benzyliden)-5-(P-iodo phenyl)-1,3,4-thiodiazole with Cr(III),Mn(II),and Cd(II)ions.,J. of Al-Nahr. Univers. .13(2):8-19,.

تحضير ودراسة H)-يل)حامض البنزوك مع بعض أيونات العناصر الانتقالية نظرية ل-4(2-مثيل-4-أوكساكونازولين-3-(4

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الخلاصة:

المعقدات الجديدة ذات الصيغة التالية [NL₂(H₂O)₂], [FeL₂(H₂O)₂], [FeL₂(H₂O)₂], [FeL₂(H₂O)₂], [IO₁Cl₂]) عندما وتشخيصها بأستخدام تحليل العناصر ،الحساسية المغناطيسية،التوصيلية المولارية ،طيف الاشعة تحت الحمراء وفوق البنفسجية ،تبين من هذه التحاليل بأن الليكاند يرتبط مع الايون الفلزي بشكل ثنائي الس لتكوين معقدات أحدية النواة. إن معقدات الكوبلت الثنائي،النيكا الثنائي،النحاس الثنائي والحديد الثلاثي تكون بشكل ثماني الس لتكوين معقدات أحدية النواة. إن معقدات الكوبلت الثنائي،النيكا الثنائي، النحاس الثنائي والحديد الثلاثي تكون بشكل ثماني الس لتكوين معقدات أحدية النواة. إن معقدات الكوبلت الثنائي،النيكا الثنائي،النحاس الثنائي والحديد الثلاثي تكون بشكل ثماني السطوح أما معقدات الفاديوم الرباعي التكافؤ فيكون بشكل هرم مربع القاعدة. اقترح الشكل الهندسي المعقدات المحضره في الطور الغازي ايضاً بأستخدام برنامج (Hyper Chem 6) بأستعمال الميكانيك الجزيئي والشبة والحضر، والشبة المحضره في الطور الغازي ايضاً بأستخدام برنامج (Byter Chem 6) بأستعمال الميكانيك الجزيئي والشبة والمعقدات الفناديوم الرباعي التكافؤ فيكون بشكل هرم مربع القاعدة. اقترح الشكل الهندسي المعقدات المحضره في الطور الغازي ايضاً بأستخدام برنامج (Hyper Chem 6) بأستعمال الميكانيك الجزيئي والشبة والمعتدات المحضره وذلك بأستخدام برنامج (PM3) التكافرين القياسية (My) وطاقة الترابط (AE) ليكاند والمعترات والمعترات أولي المور الغازي ايضاً بأستخدام برنامج (PM3) بأستعمال الميكانيك الجزيئي والشبة والمعقدات المحضره وذلك بأستخدام برنامج (PM3) ألمحسوبة وقد وجد ان هنالك تقارب بين القيام العملية والمين والمور الغاز وي المحسوب بالمساعدة لي تشخيص التوليسية (My) المعقدات المحضرة في حساب مارين القياسية (My) المعقدات المحضرة فيكاني والشبة والمورين القياسية (My) وطاقة الترايكا والمورين والمورين والمورين والموري والموري والموري والموري والتوريني والموري والمورية التربي والموري والموري والموري القيام والموري والموري والمور الغازي والموري والموريني والشبة والموري والمووري والمووي والموري والمووري والمووري والمووري والمووري والمووري