Synthesis,Structural and Biological studies of 4-[(1-phenyl-2,3-dimethyl-3-pyrozoline-5-one)azo]-N,N-dimethyl anline with some transition metal ions

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Abstract:

The present work includes the preparation and characterization of {Co(II), Ni(II), Pd(II), Fe(III), Ru(III), Rh(III), Os(III), Ir(III), Pt(IV) and VO(IV) complexes 4-[(1-phenyl-2,3-dimethyl-3-pyrozoline-5-one)azo]-N,Nof а new ligand dimethylanline (PAD). The product (PAD) was isolated, studies and characterized by measurements, i.e., (FT-IR), (UV) Spectroscopy and phsical elemental analysis(C.H.N). The prepared complexes were identified and their structural geometric were suggested in solid state by using flame atomic absorption, elemental analysis(C.H.N), (FT-IR) and (UV-Vis) Spectroscopy, as well as magnetic susceptibility and conductivity measurements . The study of the nature of the complexes formed in(ethanolic solution) following the mole ratio method, gave results which were compared successfully with those obtained from solid state studies. The apparent stability constant of the complexes have been studied with the were stable for more than (6) hours, as well as the molar time and their color absorptivities have been calculated. The antibacterial activity for the ligand (PAD) and their metal complexes were studied against two selected micro - organisms [(Klesbiella pneumonia) as gram negative] and[(Streptococcus faecalis) as gram positive]. Further more the antifungl activity against two micro-organism (Candida albicans and Aspergillus flavus) were studied for the ligand (PAD) and their metal complexes.

Key words: Azo compounds, Coordination compound of Azo , Synthesis and crystal structure of Azo compounds, Biological activity of Azo compounds .

Introduction: -

A series of N(1)-aryloxy (or thio) acetyl-3,5-dimethyl pyrazoles have been prepared in order to study their hypoglycemic and CNS activities[1,2].A slight reduction of blood sugar was observed when these compounds were screened on rats at an oral dose of 250mg/kg body weight as well as some of these compounds were found CNS stimulant, relatively nontoxic, induced writhing and piloerections in albino mice⁽¹⁾. Several N-substituted (3,5) or (2,3) –dimethyl

pyrazoles have recently been known to display significant hypoglycemic activity[3,5]. Some compounds having **N-acyl** and **N-aryloxyacetyl** [1] moieties have also evinced substantial hypoglycemic efficacy.

The present paper describes the preparation of a new azo compound derivative from (1-phenyl-2,3-dimethyl-4-amino-3-pyrozoline-5-one), in an attempt to introduce the azo (-N=N-) moiety in the structure of pyrozole ring to investigate the

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coordination behaviour of the new (PAD) ligand toward some transition metal ions {Co(II), Ni(II), Pd(II), Fe(III), Ru(III),Rh(III), Os(III), Ir(III), Pt(IV) and VO(IV)},which have been choosen to react with (PAD) and to compare the biological activity of (PAD) ligand and their metal complexes with the main ring structure.

Material and Methods: Physical measurements and analysis:-

Melting points were recorded on GallenKamp melting point apparatus and were uncorrected. FT-IR spectra were recorded using FT-IR.8300 Shimadzu in the range of (4000-200 cm⁻¹, samples were measured as (CsI disc). Electronic spectra were obtained UV-1650 using PC Shimadzu Spectrophotometer at room temperature. The measurements were recorded using a concentration of 10⁻ ^{3}M of the (PAD) ligand and their metal complexes in chloroform as a solvent.Microanalytical data for (C.H.N)were obtained using 1108CHN-O. The metal content was estimated Spectrophotometrically using Flame Atomic absorption Shimadzu A.A-670 Spectrophotometer. Conductivity measurements were obtained using Corning Conductivity Meter 220 with cell constant (1cm⁻¹), using DMF as a solvent and concentration of 10⁻³M at 25C°. Magnetic susceptibility measurements were obtained at 25C° on the solid state applying Faraday's method using Bruker BM6 instrument.

<u>A</u>-preparation of the AZO (PAD)ligand

(PAD) was prepared using the general procedure[6] .1-phenyl-2,3-dimethyl-4-amino-3-pyrozoline-5-

one(2.54gm,0.0142mol)was dissolved in 100ml of water and 10ml of concentrated hydrochloric acid and diazotized below 5C° with NaNO₂(1gm,0.0142mol) .The resulting diazoniumchloride solution was mixed N.N-dimethylanline with (1.72,0.0142mol) dissolved in alcoholicNaOH solution (3gm,100ml) below 0C°. The mixture was left in the refrigerator over night ,the solid product was filtered off, washed with cold water 250ml and crystallized from ethanol and dried over CaCl₂.

<u>**B-Preparation of the metal complexes**</u> (1-10)

The complexes metal were prepared by mixing the ethanolic the solutions of (PAD) ligand (0.01 mol) with the appropriate metal ion salts [CoCl₂.6H₂O, NiCl₂.6H₂O, PdCl₂(PhCN)₂, FeCl₃.9H₂O, RuCl₃. H₂O, RhCl₃. H₂O, OsCl₃. H₂O, IrCl₃. H₂O. H₂PtCl₆.6H₂O and $VOSO_4.5H_2O$], (0.01mol) .The pH of the reaction mixture was adjusted to~8 and it was refluxed for (2-3hr.) on a metal water bath.The chelates precipitated on refluxing the reaction mixture were filtered, washed thoroughly with hot water then ethanol several times, till the filtrate was colourless .The colored precipitates were finally washed with pet.ether (60- 80°) and then dried under vacuum .

<u>C--Study of complex formation in</u> solution

Complexes of (PAD) with metal ions were studied in solution using ethanol as a solvent, in order to determined [M : (PAD)]ratio in the complex following Molar ratio method [7]. A series of solutions were prepared having a constant concentration [10-3M] of the metal ion and (PAD).The [M : (PAD)]ratio was determined from the relationship between the absorption of the absorbed light and the mole ratio of [M : (PAD). <u>D-</u> Stability constant of azo complexes

The stability constant (**K**)[7] of the (1:1) or (1:2) [Metal: (PAD)] complex were evaluated as fallows: Two sets of solutions were prepared, the first set of solutions were formulated to contain stoichiometric amount (1ml) of (10⁻ ³M) ligand to (1 ml) of (10^{-3}M) of metal ion by placing in to a three series of 10ml volumetric flasks. The solutions of the coloured complexes were diluted to the mark with ethanol(As). The second set were formulated to contain five fold excess (5 ml) of $(10^{-3}M)$ ligand, by placing in to a three series of (10ml) volumetric flasks followed by addition of (1ml) of $(10^{-3}M)$ of metal ion solution, the volumes were then completed to the mark with ethanol(**Am**). The absorbance (As and Am) of the solutions, were measured at λ max of maximum absorption . The stability constant (**K**) and the molar absorptivity (ε_{max}) have been calculated.

<u>*E-*</u> Biological activity for (PAD) and its complexes

The biological activity of the prepared (PAD) and their respective complexes were studied against selected types of bacteria which include[(*Klesbiella pneumonia*) as gram negative] and[(Streptococcus faecalis) as gram positive], were cultivated in Nutrient agar medium, as well as DMSO was used as a solvent and as a control, the concentrations of the compounds in this solvent were 10^{-1} ³M, by using **Disc Sensitivity Test** [8,9], this method involves the exposure of the zone of inhibition toward the diffusion of micro-organism on agar plate. The plates were

incubated for 24hr. at 37 °C, the zone of inhibition of bacterial growth around the disc was observed.

In order to complete this study, the new (PAD) ligand and their metal complexes were tested for their *in vitro* growth inhibitory activity against a pathogenic fungi, i.e., (Candida albicans and Aspergillus flavus) on Potato dextrose agar medium and incubated at 30 C° for 72 hr., DMSO was used as a solvent and as a control, for both techniques, the concentration of the compounds in this solvent were 10⁻³M. The inhibition of fugal growth, expressed in percentage terms, were determined on the growth in test plates compared to the respective control plates, as given by the Vincent equation [8-10], Table (5).

Results and Discussion:-

(A)-Elemental Analysis : -

The analytical and physical properties of the (PAD) and its metal complexes are summarised in Table (1), indicate [1:1] [Metal:Ligand] stoichiometry in case [1- 4 and 10] complexes except [5-9] complexes where the molar ratio are [1:2] .The new (PAD) ligand was soluble in common organic solvents such as (ethanol,aceton and methanol) whereas the [1-10] of the new azo coloured crystallin solids complexes were soluble in (CH₂Cl₂,CHCl₃,DMF and **DMSO**)They are thermally stable and unaffected by atmospheric oxygen and moisture. The elemental analyses data are in agreement with the proposed stoichiometry.

		-		Elemental analyses					
Comp. No	Color	Melting	Yield %		Found (Calc.) %	Suggested Formula for isolated precipitate		
110.	Color	Point C°		С	Η	Ν	М	isolateu precipitate	
(PAD)	Orange	187-189	95	68.34 (68.05)	5.93 (6.27)	20.07 (20.89)	-	$C_{19}H_{21}N_5O$	
[1]	Greenish-Blue	208	84	46.95 (47.30)	4.58 (4.35)	15.26 (14.52)	12.33 (12.02)	[Co (PAD)Cl ₂]. H ₂ O	
[2]	Reddish brown	211	72	44.71 (45.53)	3.44 (4.19)	13.82 (13.98)	10.77 (11.72)	[Ni (PAD)Cl ₂] 2 H ₂ O	
[3]	Brown	215	86	43.61 (44.49)	4.23 (4.09)	12.24 (13.66)	19.82 (20.76)	[Pd (PAD)Cl ₂]	
[4]	Dark brown	219	70	25.04 (25.21)	3.60 (2.32)	7.11 (7.74)	6.49 (6.17)	[Fe (PAD)Cl ₂ (H ₂ O) ₂] .Cl.2H ₂ O	
[5]	Dull grey	233	78	52.88 (51.96)	4.13 (4.78)	16.27 (15.95)	10.69 (11.51)	[Ru (PAD) 2Cl2] Cl	
[6]	Yellowish-red	230	80	50.07 (50.81)	3.91 (4.68)	15.41 (15.60)	11.58 (11.46)	[Rh (PAD) 2Cl2] Cl. H2O	
[7]	Blackish brown	238	72	43.77 (44.67)	4.02 (4.11)	12.89 (13.71)	18.13 (18.63)	[Os (PAD) 2 Cl2] Cl. 3H2O	
[8]	Orangish- red	232	78	43.21 (42.99)	3.17 (3.95)	3.84 (3.95)	18.01 (18.12)	[Ir(PAD) 2 Cl2] Cl. 2 EtOH	
[9]	Dark red	238	81	46.19 (45.27)	4.24 (4.17)	14.46 (13.90)	19.31 (19.37)	[Pt (PAD) 2 Cl2] Cl2	
[10]	Dark green	220	83	45.68 (45.78)	3.36 (4.21)	14.29 (14.05)	11.12 (10.23)	[VO (PAD) SO ₄]	

Table (1)Physical data for (PAD) and it's metal complexes

<u>(B)-</u>Infra red spectra : -

The FT-IR.spectrum of the free (PAD) ligand was compared with the spectra of metal complexes and data is shown in Table (2).A strong bands is observed in the free (PAD) ligand at (1568cm^{-1}) and (1775cm^{-1}) , which are assigned to **vN=N** and vC=O respectively [11,12].Coordination of the (PAD) ligand to the metal through the azo nitrogen atom and oxygen atom of the carbonyl group are expected to reduce the electron density in the azo and carbonyl links and lower the vN=N and vC=O absorptions frequency.In the spectra of all the new azo complexes ,the bands due to vN=N and vC=O were shifted to lower frequencies (1550-1556) cm^{-1} and

(1763-1768) cm⁻¹ respectively as is shown in table (2). On the basis of this evidence, it is concluded that (PAD) acts as bidentate ligand in all the complexes . In the far-infrared region the azo complexes exhibited bands around (513 - 520) cm⁻¹, (460 - 508) cm⁻¹ and (370 - 405) cm⁻¹, which are assignable to v(M - O), v(M - N) and v(M - Cl) modes respectively[13]. A strong band was observed at (978) cm⁻¹ due to $(\mathbf{v} \mathbf{V} = \mathbf{O})$ stretching mode in [10] complex[13], furthermore a band related to (SO_4) sulfate anion in the [10] complex was observed at (1504) cm⁻¹, which indicated а bidentate behavior[13,14].

Table (2)Characteristic Stretching Vibrational Frequencies (cm⁻¹) locatedin the FT-IR of (PAD) and their metal complexes

Comp.No.	vN=N	vC=O	vM-O	vM-N	vM-Cl	Others
(PAD)	1568(s)	1775(s)	-	-	-	-
[1]	1553(m)	1763(m)	514(mw)	483(mw)	387(w)	3453(vO-H of water)
[2]	1553(m)	1766(m)	516(mw)	488(mw)	401(w)	3456(vO-H of water)
[3]	1556(m)	1763(m)	513(mw)	504(mw)	405(w)	-
[4]	1550(m)	1765(m)	520(mw)	508(mw)	398(w)	1615 of coordinated H ₂ O 3456(vO-H of water)
[5]	1552(m)	1763(m)	518(mw)	495(mw)	400(w)	-
[6]	1550(m)	1766(m)	516(mw)	508(mw)	403(w)	3458(vO-H of water)
[7]	1554(m)	1766(m)	519(mw)	506(mw)	405(w)	3458(vO-H of water)
[8]	1552(m)	1764(m)	516(mw)	508(mw)	400(w)	3511(vO-H of water)
[9]	1555(m)	1768(m)	520(mw)	504(mw)	405(w)	-
[10]	1550(m)	1766(m)	518(mw)	500(mw)	-	1504 of (SO ₄)anion coordination

Where:-(s)=strong, (m)=medium, (mw)=medium week, (w)=week

(C)- Electroinc spectra , magnetic susceptibility and conductivity measurements :-The electronic spectrum of the(**PAD**) ligand {4-[(1-phenyl-2,3-dimethyl-3pyrozoline-5-one)azo]-N,N-dimethyl anline} shows an intense band at (40816) cm^{-1} with a weak shoulder at (36680) cm⁻¹ followed by a strong intense band at (31125) cm⁻¹ and (29411) cm⁻¹. The first three bands be due $(\pi \rightarrow$ **π**) may to transitions, while the fourth band is due to $(\mathbf{n} \rightarrow \pi^*)$ transitions which arises due to greater delocalisation of the (C=C, C=O and N=N) π - electron resulting in a bathochromic shift[12,15].

The electronic spectra of the **[1-10]** metal complexes solutions in chloroform were recorded in the range (200 - 1100) nm and the data of these complexes are reported in Table(3).

[1]:-The greenish-blue cobalt(II) complex gave a magnetic moment value of (4.53)B.M, which indicates a type complex.Electronic high-spin spectrum in chloroform solvent exhibited a splitted band in the range of (17452-14184)cm⁻¹ [16,17], Fig. (1) .These bands can be assigned to the transition ${}^{4}A_{2} \rightarrow {}^{4}T_{1 (p)} (v_{3})$. A broad band was observed at 3244cm⁻¹ in infrared spectrum can be assigned to the ${}^{4}A_{2} \rightarrow {}^{4}T_{2}$ (f) (v₁), while the transition of (v_2) expected in the range (5000 - 6000) cm⁻¹ can not be measured[18]. The various ligand field parameters (10Dq, \mathbf{B}^{-} and \mathbf{v}_{2}) have been calculated by refered to Tanaba-Sugano diagram for (\mathbf{d}') configuration[19-21], to be (3244, 720.8 and 5453) respectively, as well as the calculation of the spin-orbit coupling constant (λ) was calculated .The resulting value ($\lambda = -198.4$) show the present complex to be distorted tetrahedral[20,22]. The nephelauxetic factor (β) was calculated and found to be (0.64) indicating high degree of

covalence in bonding of ligand donor atoms with cobalt (II) ion[21,22] .The molar conductance showed that the complex was nonelectrolyte, table(3).

[2]:-The nickel (II) complex is diamagnetic suggesting square planar geometry.The electronic spectrum of Ni(II) complex shows two main absorption bands, at (18520)cm⁻¹ and (25000)cm⁻¹, which are assigned to¹A_{1g} \rightarrow ¹A_{2g} and ¹A_{1g} \rightarrow ¹B_{1g} transition respectively ,in a square planar disposition[17,19,23-25]. Conductivity measurement in (DMF) showed that the complex was non conducting behavior, table (3).

[3]:-The prepared brown Pd (II) complex showed a strong charge transfer band at (29630) cm⁻¹, as well as two ligand field bands at (22580) cm⁻¹ and (26334) cm⁻¹ which may be assigned to :- ${}^{1}A_{1g} \rightarrow {}^{1}B_{1g}$ and ${}^{1}A_{1g} \rightarrow$ ${}^{1}E_{g}$, respectively ,in a square planar environment[19,22,26]. The magnetic moment value was (0.82)B.M. Conductivity measurement in (DMF) showed that the complex was non ionic, table (3).

[4]:-The Fe(III) complex displays three main absorption bands which are assigned to :- ${}^{6}A_{1g} \rightarrow {}^{4}T_{1g}$, $\rightarrow {}^{4}T_{2g}$, (PAD) \rightarrow Fe ⁶A_{1g} (C.T) , transition[16,19,23] respectively. The field parameters ligand calculated[19,21,24] for this complex are as follows : $(10Dq = 6634cm^{-1}), (B^{-1})$ $= 620 \text{ cm}^{-1}$) and ($\beta = 0.45$). The magnetic moment is (5.83)B.M. with electrons five unpaired and an octahedral configuration [23,24]. Conductivity measurement showed that the complex was to be ionic ,table(3).

[5]:- The spectrum of this complex exhibited four absorption bands at (15898,19708,27568 and 35144) cm⁻¹. These bands correspond to those of octahedral Ru (III) complexes[19,21,26-28], and were assigned to :- ${}^{2}T_{2g} \rightarrow {}^{4}T_{1g}$, ${}^{2}T_{2g} \rightarrow {}^{4}T_{2g}$, ${}^{2}T_{2g} \rightarrow {}^{2}E_{g}$ and ${}^{2}T_{2g} \rightarrow {}^{2}A_{1g}$ transitions[16,19] respectively. The value of (**10Dq** and μ_{eff}) came out (27568 and 1.69) respectively, revealing inner-orbital low-spin distorted octahedral stereochemistry of the ligand around Ru(III) ion[29]. Coductivity measurement showed that complex was ionic , table (3).

[6]:-Electronic spectrum of the yellowish - red [6] complex in chloroform solution, showed four absorption bands, the first band as a weak shoulder at (18203) cm⁻¹,a second band as a broad one at (20812) cm⁻¹ and the last two absorption bands as a shoulder at (23122) and (25068) cm⁻¹, which might be assigned to the cm⁻¹, which might be assigned to the transitions :- ${}^{1}A_{1g} \rightarrow {}^{3}T_{1g}$, ${}^{1}A_{1g} \rightarrow {}^{3}T_{2g}$, ${}^{1}A_{1g} \rightarrow {}^{1}T_{1g}$ and ${}^{1}A_{1g} \rightarrow {}^{1}T_{2g}$ respectively , which came in accordance with the published data for Rh(III) octahedral complexes [22,26,28]. The band at 23122 cm⁻¹can be taken as (10Dq) value[18,19,22]. Magnetic moment of solid complex ,table(3) ,showed a higer orbital contribution.Conductivity

measurement in DMF showd that the complex was ionic, table (3).

[7]:-The spectral behavior of this complex ,table(3), is identical with that of [4] complex, which agree with octahedral geometry around Os(III) complex [19,24,30].The magnetic moment that was measured at room temperature was found to be (1.98)B.M.this value refer to high spin complex ,which is the only case in octahedral environment [24,30]. The conductance measurement indicate that the complex to be ionic , table (3) .

[8]:-The electronic spectrum of the prepared orange-red Ir(III) complex in chloroform solution , showed three absorption bands at (17625,22280 and 25461) cm⁻¹, which might be assigned to the transitions :- ${}^{1}A_{1g} \rightarrow {}^{3}T_{1g}$,

 ${}^{1}A_{1g} \rightarrow {}^{1}T_{1}g$ and ${}^{1}A_{1g} \rightarrow {}^{1}T_{2g}$ respectively indicate ,these an octahedral geometry[19,31] .The different ligand parameters (10Dq, B and β) have been calculated using **Tanaba-Sugano** diagram of (**d**⁶) system and found to be (24160, 270 respectively[19-21].The and **0.41**) observed high magnetic moment value(1.83B.M.) of the present Ir(III) $(\mathbf{d}^{\mathbf{0}})$ configuration , which agree with octahedral geometry around Ir(III), this result indicate a higher orbital contribution around metal ion[19,31]. Conductivity measurement in DMF showed that the complex was to be ionic, table (3).

[9]:-The spectral and magnetic moment behaviors of this complex is identical with that of [6] and [8] complexes ,which agree with octahedral geometry around Pt(IV) complex[18,19,21] , table(3).Conductivity measurements in DMF(182.18µs.cm⁻¹) showed electrolytic nature of 1:2.

[10]:-The electronic spectrum of vanadium (IV) complex, showed three prominent (d-d) absorption bands, the first one as a strong broad band at (11130) cm⁻¹, and the second and third bands as a weak shoulders at (19053 and 24112) cm^{-1} , these bands can be attributed to the following transitions [14,16,19,22]. The magnetic moment (1.98B.M) is higher than spin value of the vanadium metal only, this result indicates a higher orbital contribution [19,22] .These transitions and magnetic moment value came in accordance with the published data for square pyramidal VO(IV) complexes [14,19,22]. Conductivity measurement in **DMF** showed that the complex was non-ionic, Fig. (2), table(3).



Fig. (1): - UV-Vis.- Spectrum of [1] complex



Fig. (2): -UV-Vis.- Spectrum of [10] complex

Table (3) Electronic Spectra (CHCl ₃), Conductance (in DMF), and
Magnetic moment (B.M) for Azo metal complexes

Comp No.	Bands cm ⁻¹	Assignment	$\begin{array}{c} \mathbf{Molar}\\ \mathbf{cond.}\\ \boldsymbol{\mu}\\ \mathbf{S.cm}^{-1}\\ \mathbf{.mol}^{-1} \end{array}$	$\mu_{ ext{eff.}}$ B.M	Suggested Structure
[1]	3244 5453 (cal) 15733(av.)		13.02	4.53	Tetrahedral
[2]	18520 25000	$^{1}A_{1g} \longrightarrow ^{1}A_{2g}$ $^{1}A_{1g} \longrightarrow ^{1}B_{1g}$	18.32	0.36	Square planer
[3]	22580 26334 29630	$^{1}A_{1}g \longrightarrow ^{1}B_{1}g$ $^{1}A_{1}g \longrightarrow ^{1}Eg$ $^{PAD} \longrightarrow ^{Pd}(C.T)$	11.68	0.14	Square planer
[4]	15084 17811 29344	$\overset{^{6}}{\overset{^{6}}{_{A_{1}g}}} \overset{\overset{\overset{}{\longrightarrow}}{\longrightarrow}} \overset{^{4}}{\overset{^{4}}{_{T_{2}g}}} \overset{^{4}}{\overset{^{4}}{_{F_{e}(C.T)}}} \overset{^{6}}{\overset{^{6}}{}} \overset{^{6}}{\overset{^{6}}{\overset{^{6}}{}} \overset{^{6}}{\overset{^{6}}{}} \overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}}}{$	68.58	5.83	Octahedral
[5]	15898 19708 27568 31144	$ \begin{array}{c} {}^{2}T_{2}g \longrightarrow {}^{4}T_{1}g \\ {}^{2}T_{2}g \longrightarrow {}^{4}T_{2}g \\ {}^{2}T_{2}g \longrightarrow {}^{2}Eg \\ {}^{2}T_{2}g \longrightarrow {}^{2}A_{1}g \end{array} $	70.15	1.69	Octahedral
[6]	18203 20812 23122 25068	$ \overset{^{1}}{\overset{^{1}}A_{1}g} \xrightarrow{\overset{3}{\longrightarrow}} \overset{^{3}}{\overset{^{3}}T_{1}g} \\ \overset{^{1}}{\overset{^{1}}A_{1}g} \xrightarrow{\overset{3}{\longrightarrow}} \overset{^{3}}{\overset{^{1}}T_{1}g} \\ \overset{^{1}}{\overset{^{1}}A_{1}g} \xrightarrow{\overset{1}{\longrightarrow}} \overset{^{1}}{\overset{^{1}}T_{2}g} $	70.15	1.69	Octahedral
[7]	16811 18238 31480	$ \overset{^{6}}{\overset{^{6}}{\underset{(PAD)}{\overset{^{6}}{\longrightarrow}}}} \overset{^{4}}{\overset{^{4}}{\underset{(Pad)}{\overset{^{6}}{\longrightarrow}}}} \overset{^{6}}{\overset{^{4}}{\underset{(Pad)}{\overset{^{6}}{\longrightarrow}}}} \overset{^{6}}{\overset{^{6}}{\underset{(Pad)}{\overset{^{6}}{\longrightarrow}}}} \overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\longrightarrow}}}}} \overset{^{6}}{\overset{^{6}}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}{\overset{^{6}}}{\overset{^{6}$	75.39	1.98	Octahedral
[8]	17625 22280 25461	$ \overset{^{1}}{\overset{^{1}}{A_{1}g}} \xrightarrow{\overset{^{3}}{\longrightarrow}} \overset{^{3}}{\overset{^{1}}{T_{1}g}} \overset{^{1}}{\overset{^{1}}{A_{1}g}} \xrightarrow{\overset{^{1}}{\longrightarrow}} \overset{^{1}}{\overset{^{1}}{T_{2}g}} $	70.73	1.83	Octahedral
[9]	18998 24023 30158	$\stackrel{^{1}}{\overset{^{1}}{\underset{(PAD)}{\overset{1}{\longrightarrow}}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\longrightarrow}}} \stackrel{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{\overset{^{3}}{\underset{Pt(C.T)}{$	182.18	2.04	Octahedral
[10]	11130 19053 24112	$2B_{2}g \longrightarrow 2Eg 2B_{2}g \longrightarrow 2B_{1}g 2B_{2}g \longrightarrow 2A_{1}g$	10.25	1.98	Square pyramideal

<u>Molar ratio and Stability constant</u> <u>for Azo metal complexes</u> :-

The molar ratio method was follow to determine the [M:(PAD)]

ratio . The results of complex in ethanol as a solvent, table (4), suggest that the metal to ligand ratio was [1:1] for [1-4 and 10] complexes, while [1:2]

for [5-9] complexes, which were comparable to those obtained from solid state study ,table(1).

Chelate stability constant (K) $\{eq.(1) \text{ and } (2)\}$, as well as molar absorptivity $(\epsilon_{max}) \{eq.(3)\}$ for all complexes were obtained spectrophotometrically by using the following equations :-

K=($1-\alpha / \alpha^2 C$) Where[1:1]of [M:(PAD)]...(1)

K=(
$$1-\alpha/4 \alpha^{3}$$
C) Where[1:2]of [M:(PAD)]...(2)

Am = (ε_{max} bC)...(3)

The degree of dissociation (α) for

the complex is obtained from the relation ship (4).

 $\alpha = (Am-As/Am)....(4)$

Where (As) and (Am) are the absorption of the partially and fully formed complex respectively at optimum concentration and (λ max) of maximum absorption, Fig. (3), table (4).

The results in Table(4) ,show that mole ratio of (1:2) for [5-9] complexes yielded rather high, this probably due

to the presence of a bulk aromatic groups and a higher oxidation state of the metal ions. As well as the developed color for all complexes become stable after one hour up to six hours.



Fig. (3): - Mole ratio plot of (PAD [) complexes (5 and 9)

metur complexes at room temperature										
Complex	[M:(PAD)	A_s	Am 🤉	K L. mol ⁻¹	Emax L.mol ⁻¹ cm ⁻¹	α	λmax nm			
[1]	1:1	0.278	0.308	9.59 *10 ⁵	3081	0.097	571			
[2]	1:1	0.373	0.428	$6.32 * 10^{10}$	4281	0.128	660			
[3]	1:1	0.398	0.453	$6.00 * 10^5$	4532	0.121	403			
[4]	1:1	0.356	0.409	$2.42 * 10^5$	4093	0.130	486			
[5]	1:2	0.495	0.551	$2.18 * 10^{10}$	5511	0.101	408			
[6]	1:2	0.294	0.336	$4.48 * 10^{10}$	3363	0.125	432			
[7]	1:2	0.355	0.403	$1.62 * 10^{10}$	3090	0.119	443			
[8]	1:2	0.259	0.309	5.03 *10 ⁹	3106	0.161	418			
[9]	1:2	0.374	0.418	1.93 *10 ¹⁰	6682	4180	515			
[10]	1:1	0.324	0.378	4.25 *10 ⁵	5720	3785	756			

 Table (4) Molar ratio, Stability constant and Molar absorptivities of Azo

 metal complexes at room temperature

Structure of [PAD] and it's Complexes [1-10]:- the structure of the above mentioned **[PAD]** and **[1-10]** their complexes can be illustrated as follow :-

According to the results obtained ,



Antimicrobial activities:-

The azo (PAD) ligand and [1-10]their metal chelates were screend in vitro in order to evaluate their antibacterial action against[(*Klesbiella pneumonia*) as gram negative] and[(*Streptococcus faecalis*) as gram **positive] species** as wll as antifungl activity against two micro-organism *(Candida albicans* and *Aspergillus flavus)*, table (5) at two different concentration. The results from these studies ,showed the followings:-

1) All the **azo metal complexes** are more toxic as compared with their **[PAD]** parent ligand against the same micro-organsim and under the identical experimental conditions, table (5).

2) The increase in the antibacterial and antifungal activites of metal chelates may be due to the effect of the metal ion on the normal cell process. The also increases toxicity with the increasing concentration of test solution containing the new azo complexes .These activities may be explained by Tweedy,s Chelation Theory [34,35].

3). The[1,2 and 10] complexes showed a moderate activities , while [3-9] complexes exhibited а great enhancement of activity against types of bacteria, Table (6). This difference in synergistic effect between the metal ion and the ligand molecule may be attributed to many factors, of these in that [3-9] complexes are considered to be a hard metal ions making their complexes to be less lipophilic, which relatively retard their permeation through the lipid part of the cell membrane. On the other hand the [1] and 2] are considered to be soft metal ions, which render their complexes to be more lipophilic and will facilitate the penetration through the cell wall and affect the invironment of cells.

 Table (5) Antibacterial and antifungal activities data for the Azo metal complexes and their parent (PAD) ligand

	Klesbiella pneumonia		Streptococcus faecali		Aspergillus flavus		Candida albicans	
Compo.	100	200	100	200	100	200	100	200
	ppm	ppm	ppm	ppm	ppm	ppm	ppm	ppm
Control (DMSO)	-		-			-		-
(PAD)	6	10	-	6	35	27	40	30
[1]	6	12	8	10	30	18	20	15
[2]	8	12	10	16	31	20	25	20
[3]	10	16	8	10	27	18	23	14
[4]	6	14	10	16	24	12	20	11
[5]	6	18	6	10	22	16	27	15
[6]	8	10	6	12	30	18	22	16
[7]	6	8	8	18	25	12	31	23
[8]	8	12	10	16	22	16	18	13
[9]	10	16	6	20	20	18	20	11
[10]	6	8	8	10	30	23	21	18
When	30-40: (+++) , 20-30 : (++++) , 10-20: (+++++)							

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تحضير ودراسة كلاً من تراكيب و الفعالية البايولوجية لـ4-[(1-فنيل-2و3-ثنائي المثيل-3- بايروزولين-5-أون)أزو]-N،N-ثنائي مثيل- أنيلين مع بعض أيونات العناصر الانتقالية

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الخلاصة: -

تم في هذا البحث تحضير و تشخيص معقدات الليكاند الجديد :-