SYNTHESIS AND CHARACTERIZATION OF THE HEAVY METALS; AU (III) ,Pd (II),Pt(IV) Rh(III) COMPLEXES OF S – PROPYNYL 2 –THIOBENZIMIDAZOLE

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ABSTRACT:

Complexes of Au(III) ,Pd (II) , Pt (IV) and Rh(III) with S – propynyle -2-thiobenzimidazole (BENZA) have been prepared and characterized by IR and UV-Visible spectral methods in addition to magnetic and conductivity measurements and micro – elemental analysis (CHN). The probable structures of the new complexes have been suggested.

INTRODUCTION:

The aromatic benzimidazole (BNZ) nucleus form ,an important class of biologically active compounds with a variety of pharmacological actions. Benzimidazole and especially 3 –thio derivatives do belong mainly for their activity to the systemically active class of antibacterial agent (1) .Some derivatives ,however show a broad spectrum of activity to be as fungicides ,acaricides and nematocides as well (2).

compounds BNZ and their derivatives were used also in rubber industry to improve its efficiency and increase its durability (3,4). It was found that when it used with the α -olefine polymer it inhibited the degradation of the later ,when it exposed to the UV light (5). Therefore it is regarded as a stabilizer for the polymer. Its heavy salts with silver and lead were used in printing field ,and as stabilizers in photographic films(6,7). They were used also in agriculture ,where they increase the growth activity activity (8).

BNZ compounds containing acetylenic moieties show awide range of biological activities of which are antimicrobial ,germicidal and disinfectant (9-11) .Moreover the interaction of the unsaturated compounds especially allylic have been

Studie with some transition metals (12). Impotents of metal complexes have been also established and some of them were found possess biological to activity.Some drugs have increased activity when administrated as metal complexes ,this attributed to the synergetic effect between the metal and the ligand (10,13).

EXPERMINTAL:

A: Chemicals and Appartus

All chemicals used were of high purity available from Fluka Company. Melting points were recorded uncorrected on Gallen Visual melting apparatus. I.R.spectra point obtained using Pye-Unicam spectrophotometer in the range 4000 -200 cm-1 ,U. V.-Vis. spectra were obtained using Pye-unicam 8700 series UV/Vis, spectrophotometer .Microanlytical data (CHN) were obtained using Perkin - Elmer 240 B elemental analyzer ,Magnetic susceptibility measurements were done Bruker using magnet BM₆ .Conducativity measurements were obtained utilizing PW9527 **Digital** conductivity Philips apparatus.

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B:Methods of Preparation

1. Preparation of BNZA

The BNZA was prepared according to the literature methods (14), starting from 2- hydroxy aniline .Crystals were obtained with a melting point 52oC.

2. Preparation of the BNZA

Metal Complexes

a.[$Au_2(BNZA)Cl_4)$]($AuCl_4$)₂ (I)

To a hot stirred solution of **BNZA** (2 mmole) in ethanol (50ml),(1.0mmole)of chaloroauric acid dissolved in 30ml ethanol was added .the mixture was refluxed for one hour .then cooled to room temperature .Brown precipitate was filtred and washed two to three times with ethanol and dried under identified vacuum was .It characterized by elemental analysis ,I.R.,UV -Vis .,magnetic susceptibility and conductivity measurements .The results are shown in table(1)

$b.[Pd(BNZA)Cl_2].C_2H_5OH(II)$

0.208 mmole of dichloro bis(benzonitrile)palladium (II)(dissolved in(10ml) ethanol) ,was 0.45mmole added to of (dissolved in 20ml of the same solvent) .The mixture was stirred for 30min .at room temperature and kept for one day at room temperature, an orange -brown precipitate was formed ,which was separated by filtration and washed several times with little amount of ethanol ,then dried under vacuum .The formation complex was studied by the same way as for complex(1)above.

c. [Pt (BNZA)₂ Cl₂]Cl₂.H₂O(III)

The platinum (IV) complex was prepared by the reaction of the potassium hexachloroplatinate (IV) with BENZA in (1:2) mole ratio ,using (1:1)acetone /water mixture as a solvent ,at room temperature. The product was washed with water and little acetone after filtration and drying under vacuum ,it was identified by the same techniques described as above.

$d.[Rh(BNZA)_2] Cl_3. 2H_2O (IV)$

The reaction of Rhodium trichloride hydrate (0.2635g,1) mmole with BNZA (0.406 g,2mmole) in a hot ethanol —water mixture yielded an orange precipitate ,moderately solube in ethanol ,+water and sparingly soluble in benzene .It was characterized by the same techniques used for complexes (mentioned before) .

RESULTS AND DISSCUSSION

Stable complexes were isolated in all cases .Based on the micro analytical data ,the general formula of the complexes can be suggested as [M(BNZA)X Cl y] Clz .X,where M= Pt and Pd ions $x=1,2,\ y=1,2$, z=0,2 and X=H₂O or C₂H $_5$ OH respictabively .When M = Au or Rh ions ; $x=1,2,\ y=0,4$, z=0,3 and $X=2AuCl_4$, $2H_2O$ respectively .

BNZA is a potential ligand which may coordinate through either the acetylenic or thioamide moieties or both .BNZA has the ability to form complexes organometallic coordination through the acetylenic group with a number of metal ions (12) ,therefore it is expected that I.R. measurements are highly informative with respect to the complexation behavior with various metal ions .The characteristic frequencies of the free ligand and its metal complexes were readily assigned based on comparison with literature references (11,15). The vibrational frequency corresponding to the $v c \equiv c$ was clearly characterized and is of basic importance. The spectra of all complexes showed similar behavior where a red shift by $\sim 570 \text{ cm} - 1 \text{ took}$ place ,which indicates a strong pi bonding between metal the acetylenic group (16,17). In addition the stretching frequency of the terminal C --H bond was noticed unchanged in the of all complexes spectra (2), which further indicates a pi -bonding (15)

The spectrum of Au (III)complex shows the appearance of the band belonding to the $v c \equiv c$, with a change in the shape of the band of 1540 cm - 1 ,where it appeared at the same position of another band found at this position , so a strong π – coordination can be suggested between gold atom the acetylenic bond. It can be also concluded that the sulfur atom coordinates with the gold ion because of the shift of the vc - sto lower frequency by 15 cm - 1, and the shift of the thioamide band (II) to higher frequency in addition to the appearance of vAu - S band at 410 cm - 1. The N atom also coordinates with gold due to the appearance of a band at 510 cm - 1.. The presence of two bands at 290 and 356 cm - 1 indicate the coordination of chloride with gold as seen in table (2).

When chloroauic acid was treated with BNZA, a brown product ,insoluble was formed dichloromethane and acetone **DMSO** moderately soluble in .According to the results of elemental analysis ,i.r., UV- Vis ., and conductivity measurements the following formula may be assigned.

This formula reflects the following facts about the complex behavior (a) its appreciable conductance ,(b) square geometry about gold ,(c)noticeably low percentage of C ,H and N and (d)the behavior of gold (III) as soft acid .The tendency of Au(III) to N and O donors in addition to acetylenic group as donors ligands leads to the conclusion that (a) not all the chlorides of AuCl4— are displaced by the ligand in most cases ,(b) the coordination ligand ,even when behaves as bidentate like 2,2— bipyridyl

(18) ,are readily replaced by chloride or bromide ,e.g.

$$\begin{aligned} \left[AuCl_2(Phen.) \right]^+ &+ 2Cl & \qquad \left[AuCl \right]_2^+ &+ phen. \\ \left[AuCl_3(Py.) \right]^+ &+ Cl^- & \qquad \left[AuCl \right]_4^- &+ py. \end{aligned}$$

Many Au(III) complexes with N O and S donor chelate ligands were synthetic (19).Gold (III)forms complexes with soft ligand ,e.g. S ,Se,I and CN, which are more stable than those with O – or N — donor ligands and both O and S donor sites the S atom is always preferred by Au (III) center, also ocatahedral geometry could be obtained for soft ligands e.g the tetragonally distorted octahedral trans [Aul2 (diars)2]+ I- (20).In the present case Au (III) coordinates to N,O,S and acetylenic group of BNZA and it seems that an equilibrium exists for the complexation reaction ,this is an evident by the presence of AuCl4- anion in the present structure ,also only two chlorideced was displaced by BNZA.

In the case of Pd(II)complex a shift of the stretching vibration of the triple bond took place by 390 cm— 1 toward lower region, which refers to the coordination of the π -bond with the Pd (II)ion .This is more indicated by the presence of terminal CH bond stretching at 3300 cm-1 .A new strong band appeared at 340 cm— 1 which indicates the formation of Pd - Cl bond.In addition another weak band was noticed at 235 cm—1 .Both are characterized as belonging to assymmetric and symmetric frequencies of two chloride atoms coordinating in a cis – position (15) .No changes in the positions of thioamude bands were noticed .This may show that the ring atoms are not participating in coordination with the atom .In addition a new band at 415 cm— 1 was detected which belongs to Pd- bond .Therefore the following structural formula can be suggested:

$$R = O$$
 $R = O$
 R

The spectrum of Pt(IV)complex show

the v c \equiv c to be shifted by 380 cm— 1 toward lower frequency ,which indicates the coordination of the metal through π – bond with the ligand .This is more indicated by the presence of the acetylenic v CH of 3300 cm— 1 .Also the position of the C-S band absorption was shifted by 20 cm— 1 and the appearance of Pt –S absorption band at 440 cm— 1 .This indicates the coordination of Pt(IV)with the sulfer atom of BNZA .

This bonding behavior of Pt(IV)and S is expected, due to the softness of both atoms, the strong band appeared at 330 cm—1 which refers to the coordination of two chlorides at trans-position (15), Table (2) .According to this, the following structural formula can be suugested:

$$\begin{bmatrix} R & C & C \\ S & C & N \\ C & C & C \\ C & C & S \\ C & C & S \\ C & C & C \end{bmatrix} + 2$$

$$R = \begin{bmatrix} O & N \\ O & N \\ O & N \\ O & N \\ O & O \end{bmatrix}$$

The thioamide band (II)was shifted to higher freguency by 35 cm— 1 in the spectrum of Rh (III) complex ,which indicates the bonding of the N –atom of the SCN moiety with the metal ion .Also Rh –N bond stretching appeared at 503 cm— 1 ,and another one at 420 cm— 1 which belongs to Rh –S bond.

The disappearance of the band belonging to $v c \equiv c$ and the appearance of a new one at 1680 cm— 1 indicates the coordination of Rh with BNZA through Pi—system ,which is further indicated by presence of v CH band at 3300 cm— 1 ,Table (2) ,therefore, the following structure can be suggested:

$$\begin{bmatrix} N & CH \\ S & Rh \end{bmatrix} \qquad R = \begin{bmatrix} O & N \\ O & S - CH_2 - I \end{bmatrix}$$

The broad bands at ~ 3400 -3350 cm— 1 and at 1610 cm— 1 in the case of Pt (IV)

and Rh (III) complexes indicate the presence of water in the crystal lattice. UV -vis Spectroscopy ,conductivity and magnetic susceptibility measurements Little information is available about the electronic spectra of the square planar gold complexes, and a reference to PdCl 4-2 ion is made because the later is a good example of the absorption behavior of the d8 square planar environment .On comparism of the spectra of this ion and that of the new gold complex which was obtained from it,s ethanolic solution, a number of absorption bands have been assigned, which came in good accordance between the two spectra, and as follows (21):

$$^{1}A_{1}g \longrightarrow ^{3}B_{1}g$$
 17999 cm⁻¹
 $^{1}A_{1}g \longrightarrow ^{1}B_{1}g$ 24908 cm⁻¹
 $^{1}A_{1}g \longrightarrow ^{1}Eg$ 31760 cm⁻¹
L \longrightarrow Au(C.T) 35000 cm⁻¹

Therefore square planar geometry can be suggested for the Au (III)complex .Also the value of magnetic moment for the prepared complex was diamagnetic and the data of electrical conductivity measurements in DMSO solvent (95) µs . cm—1 agree with the suggested formula to be ionic .

The spectrum of the palladium complex shows absorption bands at 22988 cm—1 belonging to the ${}^{1}A_{1}g \longrightarrow {}^{1}B_{1}g$ transition, which is equal to the 10 Dq and an other band at 27777 cm—1which corresponds to the ${}^{1}A_{1}g \longrightarrow {}^{1}E_{1}g$ transition .Therefore square planar geometry can be assigned for the structure (22) .This geometry is common for Pd (II) complexes ,due to the effect of the large size of the ion on it, s ligand field strength ,which is greater than the pairing energy (7,23). The value of magnetic moment for the prepared complex was (0.75) B.M, which indicates the diamagnetic nature of the complex. The electric conduced measurement in DMSO agreed with the suggested formula to be non - conducting Table,

(3).

The electronic spectrume of the platinum (IV) shows two bands ,the first weak one appeared as a shoulder at 21739 cm—1 ,the second band appeared at 27322 cm—1 with higher intensity .These two bands can be assigned to the following transition in ocatahedral environment (21):

$${}^{1}A_{1}g (t_{2}g^{6}) \longrightarrow {}^{3}T_{1}g (t_{2}g^{5}eg^{1})$$
 ${}^{1}A_{1}g (t_{2}g^{6}) \longrightarrow {}^{1}T_{1}g (t_{2}g^{5}eg^{1})$

This coordination type is common for pt (IV) complexes (21,22),study of conductivity behavior in DMSO shows the complex to be ionic (105)µ.cm—1.The magnetic measurements data is 1.02 B.M shows which the complex to be diamagnetic.

The UV-Vis. spectrum of Rh(III)complex showed a main strong band at 34.200 cm—1 and two shoulders at 30.200 and 25.300 cm-1 .The tetrahedral geometry was excluded because no bands appeared below 20.000 cm—1 (21), therefore spinpaired square planar configuration may be assigned to the complex in a monomeric state as suggested by the ready solubility in benzene geometry was converted to distorted octahedral due to the cooredination of

two DMSO molecules as the complexes was dissolved in DMSO. Conductivity measurement in DMSO show the complex to be conducting, therefore the chloride is not considered to be coordinated with the metal and is located outside the coordination sphere.

The distination between ligand field and charge transfer bands cannot be made sharply ,due to mixing of the ligand orbital with the metal orbitals .Therefore, the ligand field bands cannot be easily identified in the spectra of spin -paired square planar complexes (22,23). Neverthless and by referring to the available data for low spin ions,i.e.Co+ 3,Rh+3 and Ir+3, published in the literature (22),the band at 25.300 cm—1 can be assigned to the transition $a_1 g \longrightarrow b_1 g$ which can be taken as the 10Dq value .The measured effective magnetic moment was (0.89)B.M.of behavior .The electric diamagnetic conductivity measurements show the complex to be conducting (130)µs. cm—1, Table (3).

Table (1): Physical data for the metal complexes.

Compounds	color	Reaction Solvent	Recrystal. Solvent	Melting Point ^o C	%yield	Microanalytical data Found(calc.)		
						C%	Н%	N%
[Au ₂ (BNZA)Cl ₄)].2AuCl ₄	Brown					10.80	0.430	1.08
I	DIOWII	EtOH	EtOH	119_D	35	(11.23)	(0.655)	(1.31)
[Pd(BNZA)Cl ₂)].C ₂ H ₅ OH II	Light brown	EtOH	EtOH	70	40	33.03 (34.99)	3.010 (3.159)	3.520 (3.402)
[Pt(BNZA) ₂ Cl ₂)]Cl ₂ . H ₂ O III	Yellow	(CH ₃) ₂ CO/H ₂ O	(CH ₃)CO	110 _D	46	33.10 (32.78)	2.10 (2.18)	3.950 (3.825)
[Rh(BNZA) ₂]Cl ₃ .2H ₂ O IV	Orange	EtOH/H ₂ O	EtOH	87	32	46.10 (46.46)	3.40 (2.71)	5.37 (5.42)

Table (2):selected I.R bands of S-propynyl -2 -thiobenzimidazole and it ,s metal complexes

Compounds	Thioamide(II) $\mathbf{v}_{c_{=N+C=S}}$	Thioamide(IV) Vc - s	V _{C ≅C}	V _{C ≅CIII}	S ≡ _{CIII}	V _{M-N}	V _{M-C}	V _{M-S}	V _{M-O}	$\mathbf{V}_{ ext{M-CI}}$
BNZA	1275 w	770 s	2100m	3300w	620w	-	-	-	-	-
I	1295 m	760 s	1540 s	3300 w	620 w	510	420w	470w	-	295 356
II	1275m	770m	1710s	3300w	620w	-	415	-	-	340 325
III	1275m	740m	1720s	3300	620	-	415	440	-	330
IV	1305m	760m	1680s	3300	620	503	415	420	-	1

Table (3): Electronic specteral data of BNZA complexes and their probable assignments, and magnetic moments.

Compands	Bands ,cm ⁻¹	Assignment	μ _{eff.} (B.M.)		
I	17999	$^{1}A_{1}g$ $^{3}B_{1}g$	0.0		
	24908	$^{1}A_{1}g$ $^{1}B_{1}g$	0.0		
	31760	¹ A ₁ g ¹ Eg			
	35000	L A u(C.T)			
II	22988	$^{1}A_{1}g$ $^{1}B_{1}g$	0.75		
	27777	¹ A ₁ g ¹ Eg	0.75		
II	21739	$^{1}A_{1}g$ $^{3}T_{1}g$	1.02		
	27322	¹A ₁g → ¹T ₁g	1.02		
IV	34.350		0.00		
	30.200 &25300		0.89		

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تحضير وتشخيص بعض معقدات العناصر الثقيلة الذهب (III) البلاديوم (IV) البلاتين (IV) والراديوم (III) مع الليكاند ثايوبروبانيل -2 حثايوبنزاميدازول

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الخلاصة:

تم في هذا البحث تحضير معقدات الذهب (III) والبلاديوم (II) والبلاتين (IV) والرديوم (III) مع الليكاند ثايوبروبانيل -2 -ثايوبنزاميدازول وتشخيصها بطرائق التحليل الدقيق للعناصر والطرق الطيفية ، الاشعة تحت الحمراء والاشعة فوق البنفسجية - المرئية بالاضافة الى القياسات الحساسية المغناطيسية والتوصيلية الكهربائية .وتم اقتراح التراكيب المحتملة للمعقدات المحضرة الجديدة.