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Characterization and biological effectiveness of synthesized complexes of Palladium (II) from imine compounds



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Abstract

This article involved synthesize a novel Palladium (II) complexes as follows: The imine compounds were prepared by reacting of aldehydes with nitrogen compounds: 4-methylbenzene-1,2-diamine, naphthalene-1,8-diamine, and 4-chloro-5-methyl benzene -1,2-di amine and sublimated into ethanol. The reaction was continued by the thin layer chromatography (TLC), where the imine compounds were determined by spectrophotometry of FT-IR UV-Visible, ¹H-NMR. The palladium complexes were synthesized by the reaction of the prepared imine compounds (after they were dissolved) using absolute ethanol and palladium salt (PdCl₂), which was dissolved by using absolute ethanol with 4 drops of 11.6 N of HCl acid, then the mixture was raised for 3 hours. When the amine compounds reacted with the Palladium ion (II), this interaction leads to formation of palladium complexes, the reaction was continued by the thin layer chromatography (TLC), the complexes were spectrophotometrically characterized by measurement the ultraviolet-visible, infrared rays, mass spectrometry and molar conductivity. The geometric shape of the complexes had been proven, which the palladium complexes have a square planer shape. The biological activity of some synthesized complexes was determined using two different types of bacteria (Gram-Positive and Gram-Negative), namely *Staphylococcus Aureus* and *Escherichia coli*. The results show that some concentrations have an intense inhibitory effect on the target bacteria. Keywords: Imine compounds, complex of Palladium (II);

1. Introduction

The organometallic compounds are important synthesized compounds in organic chemistry [1-3]. The nitrogen atom of ligands (**Imine compounds**) have the ability to form complex [4, 5]. The Condensation reaction is considered one of the most important and common methods of preparing amine compounds, which it occurs by the direct condensation between primary aromatic amines and aldehydes or ketones, the reaction is done by using glacial acetic acid as a catalyst, leading to displacement of the water molecule to produce imine compound [6-10] below in scheme 1:

The Imine compounds are organic compounds that contain the active group (-HC=N-), Imine produced by the interactions between ketones or aldehydes with primary amines, which considered important organic compounds in the synthesis process of cyclic

compounds and organometallic compounds, Imine compounds are effective materials in the medical field [11]. For the first time, imine compounds were prepared by Hugo Schiff [12]. Imine compounds are used as ligand compounds, where it's more soluble with metallic salts [13]. Imine compounds are one of the compounds that can be used to make a Palladium (II) complex [14].

$$Ar-NH_2 + Ar'$$

Absolut Ethanol/H⁺

Reflux for 3hrs

 Ar
 Ar
 Ar
 Ar
 Ar

Scheme 1. General reaction of synthesized imine compounds

One of the important palladium complexes was prepared from the reaction of 2-(4-ethyl) phenyl amino acetyl-N-phenyl hydrazine carbothioamide

compounds with the palladium (II) salt, the diagnosis was made by spectroscopic methods to suggest the geometry of the prepared complex, where the shape was a square planer with dsp² hybridization, there were effects of the synthesized complex on the growth process of *Bacillus Escherichia* (Gram-Positive) and *Escherichia coli* (Gram-negative), this complex was shown to have anti-activity for these types of bacteria, the following figure shows the formula for the synthesized complex (palladium (II) complex) [15]:

Scheme 2. Synthesis of palladium (II) complex

2. Experimental

2.1. Synthesis of imine compounds

0.004 mol $(0.5 \, \text{gm})$ of 3,4-diaminotoluene was dissolved in $(15 \, \text{mL})$ of EtOH and mixed with $0.008 \, \text{mol}$ $(1.15 \, \text{gm})$ of para-chloro benzaldehyde, which was also dissolved in $(12 \, \text{mL})$ of EtOH. The solution was cooled after the refluxing process $(3 \, \text{hours})$ and filtered. The precipitant was recrystallized from EtOH to obtain the imine (S_2) , at the same process the other imine compounds were synthesized [16].

2.2. Synthesis the complexes of palladium (II)

0.001mol (0.4gm) from S_2 was dissolved in (20 mL) EtOH and mixed with 0.0005mol (0.09gm) of PdCl₂ which was also dissolved in (15 mL) of EtOH by assisting few drops of HCl (11.6 N). The solution was cooled after the refluxing process (3 hours) and filtered. The precipitant was recrystallized from EtOH to obtain the (P_2) complex, at the same process,

all the palladium (II) complexes were synthesized [4].

2.3. Biological Evaluation

Antibacterial activity of the palladium complexes was done by using Mueller Hinton Agar against *Staphylococcuus Aureus* and *Escherichia coli*. The holes have a diameter of 6 mm. The biological activity was dependent on calculated of the inhibition zone.

3. DISCUSSION THE RESULTS

3.1. Imine compounds

Imines were synthesized from the reaction between aldehyde compounds and diamine compounds, the reaction showed in the following equations (scheme 3):

The mechanism of imine compound synthesis includes the addition of the proton from glacial acid to the carbonyl group (C=O) of aromatic aldehyde, which leads to the formation of the intermediate compound (carbonium ion). The second step includes the attack of the nitrogen atom of the amino group (-NH₂) in the aromatic amines as a nucleophilic attack on the carbon of the carbonium ion to form the intermediate compound (N-Subtituted Hemiaminal). In the third step, the proton is withdrawn from the nitrogen atom by the water molecule to form a carbinolamine. The fourth step involves adding a proton from the acid to the hydroxyl group. The fifth step involves the displacement of a water molecule from the compound. The final step is the displacement of a proton of nitrogen to yield the stable imine compound, as shown in Scheme 4:

Table $\ 1$. Properties of synthesized imine compounds

Comp. Code	Molecular Formula	m.p.°C	Yield %	Colour	M. wt	Time of Reaction	$\mathbf{R}_{\mathbf{f}}$
S ₂	C21H16N4O4	64-66	64	Light yellow	122.17	2 hrs.	0.8
S ₃	C21H14Cl2F2N2	100-102	52	Light nutty	122.17	2 hrs.	0.4
S ₄	C23H16N4	98-100	71	Yellow	122.17	1 hr.	0.8
S 5	C24H16Cl2N2	130-132	50	Light nutty	158.2	3 hr.	0.5
S_6	C24H16N4O4	240-242	55	Orange	158.2	3 hrs.	0.7
S9	C21H15Cl3N2	118-120	65	Light nutty	154.6	1 hrs.	0.8
S ₁₀	C19H13ClN4O4	235-238	75	Light nutty	154.6	30 min	0.6
S_{11}	$C_{21}H_{15}Br_2ClN_2$	178-180	50	Light nutty	154.06	3 hrs.	0.8
S ₁₂	C24H16Br2N2	118-120	82	Green yellow	158.2	3 hrs.	0.7
S ₁₃	C25H27ClN4	194-196	51	Dark nutty	154.06	2 hrs.	0.7

Table 2. Structure and nomenclature of synthesized imine compounds

Table 2 . Structure and nomenclature of synthesized imine compounds							
Code	Structure nomenclature	Code	Structure nomenclature				
S ₂	CH3 CH NO ₂ NO ₂ NO ₂ (1E,1'E)-N,V'-(4-methyl-1,2-phenylene)bis(1-(4-nitrophenyl)methanimine)	S9	CH CH (1E,1'E)-N,N'-(4-chloro-5-methyl-1,2-phenylene)bis(1-(4-chlorophenyl)methanimine)				
S ₃	H ₃ C CH Cl F F N,N'-(4-methyl-1,2-phenylene)bis(1-(2-chloro-4-fluorophenyl)methanimine)	S ₁₀	CI HC CH NO ₂ NO ₂ NO ₂ (1E,1'E)-N,N'-(4-chloro-5-methyl-1,2-phenylene)bis(1-(4-nitrophenyl)methanimine)				
S4	H ₃ C H ₃ C C ₅ N 4,4'-((1E,1'E)-((4-methyl-1,2-phenylene) bis(azaneylylidene)) bis(methaneylylidene))dibenzonitrile	Sii	CI CH ₃ HC CH Br Br (1E,1'E)-N,N'-(4-chloro-5-methyl-1,2-phenylene)bis(1-(4-bromophenyl)methanimine)				
S ₅	(1E,1'E)-N,N'-(naphthalene-1,8-diyl)bis(1-(4-chlorophenyl)methanimine)	S ₁₂	HC CH Br Br (1E,1'E)-N,N'-(naphthalene-1,8-diyl)bis(1-(4-bromophenyl)methanimine)				
S ₆	HC CH NO ₂ NO ₂ (1E,1'E)-N,N'-(naphthalene-1,8-diyl)bis(1- (4-nitrophenyl)methanimine)	S13	H ₃ C N _C H ₃ H ₃ C N _C H ₃ 4.4'-((1E,1'E)-((4-ch)toro-5-methyl-1,2-phenylene)bis(azaneylyildene)bis(methaneylyildene))bis (XA'-dimethyl-nilm)				

3.1.1. Ultra violet - Visible of imines

The compounds were dissolved in DMSO and characterized by Uv-Visible spectra, the transitions of S_2 showed the transition at (238 nm and 296nm) of the type $\pi \rightarrow \pi^*$ caused by C=C bonds of aromatic structure, transition at (480 nm) of the type $n \rightarrow \pi^*$ for C=N [17]. Table 3 showed all the types of transitions and wavelengths for all synthesized imines. See figures 1-4 for some imines compounds.

3.1.2. FT-IR of imine compounds

The FT-IR spectrum of the compound S_2 showed the absorption absorption band at $(1600 \, \text{cm}^{-1})$ for C=N imine, absorption band at $(1440 \, \text{cm}^{-1})$ for C=C aromatic, stretch absorption band at $(3070 \, \text{cm}^{-1})$ refers to aromatic C-H, absorption band $(3109 \, \text{cm}^{-1})$ refers to C-H of imine, absorption band at $(2856 \, \text{and} \, 2922 \, \text{cm}^{-1})$ refers to symmetric and asymmetric respectively of aliphatic C-H [18, 19]. Table 4 showed the other stretch absorption bands of all synthesized imine compounds. See the figures 5-8 of some imines compounds.

Table 3. T	Γhe Ultra violet -	Visible of synthesized imines	
2	3.6 / ^	TD •4•	

Comp.	Max/nmλ	Transition
S ₂	238	π-π* of Aromatic C=C
~2	296	π-π*
	480	n-π*of C=N
S ₃	238	π-π* of Aromatic C=C
-	296	π-π*
	438	n-π*of C=N
S ₄	238	π-π* of Aromatic C=C
	295	π-π*
	475	n-π*of C=N
S ₅	237	π-π* of Aromatic C=C
	295	π-π*
	460	n-π*of C=N
S 6	238	π-π* of Aromatic C=C
	295	π-π*
	435	n-π*of C=N
S_9	235	π-π* of Aromatic C=C
	297	π-π*
	480	n-π*of C=N
S ₁₀	240	π-π* of Aromatic C=C
	295	π-π*
	560	n-π*of C=N
S_{11}	235	π-π* of Aromatic C=C
	295	π-π*
	435	n-π*of C=N
S ₁₂	235	π-π* of Aromatic C=C
	295	π-π*
	380	n-π*of C=N
S ₁₃	235	π-π* of Aromatic C=C
	295	π-π*
	465	n-π*of C=N

3.1.3. ¹H-NMR spectra of imine compounds

The ^1H - NMR of imine (S₂) showed the chemical shift (δ ppm): singlet signal at (δ = 2.45) refers to the methyl group (-CH₃), singlet at (δ = 10.14) refers to the proton of imine group and a multiplet at (δ =7.96-8.42) refers to the protons of aromatic system. Table 5 shows the ^1H -NMR of the synthesized imines [20, 21]. See the figures 9-12 of some imines compounds

3.2. Complexes of Palladium (II)

The structure, formula, and other properties of

synthesized Palladium recorded in the table (6), Palladium complexes were characterized with molar conductivity, atomic absorption and (Uv-Visible, FT-IR, Lc-mass) spectroscopies, The good yield of the synthesized Palladium complexes was for P₉ 90%, the highest melting points of synthesized Palladium complexes were for compounds P₁₁ and P₁₃ (>300 °C). Table 7 showed the time of reaction and Rf (the ratio between the distance of the complexes and the mixtures of solvents in the thin chromatography) of all synthesized Palladium (II) complexes.

3.2.1. Ultra violet - Visible of complexes of the Palladium (II)

Palladium complexes were dissolved in solvent (DMSO) and showed the transitions: P_2 showed the transition at (230 nm and 295nm) of the type $\pi \rightarrow \pi^*$ caused by C=C bonds of aromatic structure, transition at (438nm) of the type $n \rightarrow \pi^*$ for C=N, transition at (880 nm) of the $^1A_1g \rightarrow ^1A_2g$ transition, all transition proved the shape of P_2 was Square planer [17]. Table 8 showed all the transitions for all synthesized Palladium complexes. See the figures 13-16 of some Palladium (II) complexes

3.2.2. FT - IR spectra complexes of Palladium (II)

The FT-IR of the synthesized complexes of Palladium (II) appeared the absorption: Complex (P₂) showed the band at (1606 cm⁻¹) for C=N imine, band at (1460cm⁻¹) for C=C aromatic, band at (3070 cm⁻¹) refers to aromatic C-H, band at (3120cm⁻¹) refers to C-H of imine, band at (2800 and 2924 cm⁻¹) refers to symmetric and an asymmetric respectively of aliphatic C-H, the band at (599 cm⁻¹) refers to presence of Palladium – Nitrogen band (M-N) [18]. Table 9 showed the other absorption bands of all synthesized Palladium (II) complexes. See the figures 17-20 of some Palladium (II) complexes.

Table 4. The	FT-IR	of synthesized	imine compounds

Comp	oC N	υC=C	»C-N	υ=C-H Aliphatic		υ=С-Н	υ=С-Н	Other
Comp.	υC-N	Ar	υC=N	Symmetric	Asymmetric	Ar	Imine	Groups
S_2	1109	1440	1600	2856	2922	3070	3109	NO ₂ : 1344, 1517
S ₃	1118	1456	1602	2862	2922	3072	3111	Cl: 1041, F: 1259
S ₄	1118	1440	1610	2922	2974	3072	3090	CN: 2227
S ₅	1124	1419	1598			3034	3068	Cl: 1087
S_6	1165	1512	1600			3068	3078	NO ₂ : 1348,1512
S9	1093	1446	1600	2972	2924	3010	3120	Cl: 1008
S ₁₀	1105	1595	1610	2922	2080	3109	3109	NO ₂ : (1344, 1514)/Cl:999
S ₁₁	1109	1446	1627	2922	2890	3070	3140	Br: 686, Cl: 1006
S ₁₂	1165	1421	1597			3041	3070	Br: 644
S ₁₃	1195	1442	1612	2889	2804	3030	3075	C1: 999

Table 5. The ¹H-NMR spectra of some synthesized imine compounds

imine co	mpounds			
Comp.	Group	No. of	Chemical	Type of
No.		proton	shift	Single
		1	(ppm)δ	
	-С <u>Н</u> 3	3	2.45	singlet
	Aromatic	11	7.96-8.42	multiplet
G		11	7.90-0.42	munipiei
S_2	protons		10.11	
	2(-	2	10.14	singlet
	C <u>H</u> =N-)			
	-С <u>Н3</u>	3	2.53	singlet
	Aromatic	9	6.73-7.97	multiplet
S_3	protons			_
	2(-	2	10.20	singlet
	C <u>H</u> =N-)	_	10.20	Singier
	-CH ₃	3	2.53	singlet
	Aromatic	11	7.07-8.62	
		11	7.07-8.02	multiplet
S4	protons			
	2(-	2	10.10	singlet
	C <u>H</u> =N-)			
	Aromatic	14	6.48-8.06	multiplet
	protons			•
S ₅	2(-	2	10.85	singlet
	C <u>H</u> =N-)	_	10.05	Singice
		1.4	(50, 0.20	
	Aromatic	14	6.50-8.29	multiplet
S_6	protons			
20	2(-	2	10.20	singlet
	C <u>H</u> =N-)			
	-C <u>H</u> 3	3	1.08	singlet
	Aromatic	11	6.96-8.18	multiplet
S_9	protons			•
,	2(-	2	10.00	singlet
	C <u>H</u> =N-)	_	10.00	Singici
		2	2.46	ain alat
	-C <u>H</u> 3	3	2.46	singlet
_	Aromatic	11	6.70-8.87	multiplet
S_{10}	protons			
	2(-	2	10.20	singlet
	C <u>H</u> =N-)			
	-C <u>H</u> 3	3	2.61	singlet
	Aromatic	11	6.90-8.11	multiplet
S ₁₁	protons			
	2(-	2	10.00	singlet
	`		10.00	singlet
	<u>CH</u> =N-)	4.	< 40 = 00	7,0 7
	Aromatic	14	6.48-7.88	multiplet
S ₁₂	protons			
512	2(-	2	10.00	singlet
	C <u>H</u> =N-)			
	-CH ₃	3	2.60	singlet
	4 (-	12	3.07	singlet
	NCH ₃)	14	5.07	Singici
C		10	6 62 7 00	multiplet
S ₁₃	Aromatic	10	6.63-7.98	multiplet
	protons			
	2(-	2	9.67	singlet
	C <u>H</u> =N-)			

3.2.3. Lc - mass of the complexes of Palladium (II) [4]

The Lc - mass spectra showed the fragments of the complexes and proved the synthesizing process and the square planer shape, the tables 10-19 showed the fragments of the complexes. See the figures 21-24 of some Palladium (II) complexes and scheme 5-14.

3.2.4. Molar conductance measurements of the complexes of Palladium (II)

The measured of molar conductivity of the complexes uses to know the ionic formulas of complexes, the complexes concentration are 1×10^{-3} molarity, the measured completed by dissolving the samples in DMF at the room temperature. If the value of conductance is more than 70 (Ohm⁻¹. cm⁻¹. mol⁻¹ x 10^{-6}), this proves that the negative charge of chloride is outside of the coordination [22]. See table 20.

3.2.5. Atomic absorption of the Palladium Complexes [23]

Standard solutions of metal chlorides were used for titration. A specific weight of the solid compounds was digested with a mixture (5 mL) of concentrated nitric acid and perchloric acid, this issue was repeated several times until completely dissolve of all organic matter. The reaction was evaporated to drying. After cooling, the remaining salt was dissolved in deionized water where the proportion of the presence of palladium in the synthesized complexes was measured and compared with the theoretically calculated ratio, as shown in the following table 21.

3.2.6. Biological activity of some Palladium complexes

Antibacterial activity of Palladium complexes was done by using Mueller Hinton Agar against *Staphylococcuus Aureus* and *Escherichia Coli*. The Palladium complexes dissolved in DMSO Solvent, the holes diameter was 6 mm, the concentrations of Palladium complexes are 50% and 100% [24], the inhibition zone recorded in table 22. See pictures 1 and 2.

Table 6. Properties of synthesized complexes of Palladium (II)

Comp.	Structure	Formula	m.p. °C	Yield%	Color
P ₂	O ₂ N CH NO ₂ Pd ⁻² CH NO ₂ CH NO ₂	C42H32N8O8Pd	280-284	58	Gray
P ₃	F CH3 CH CI	C ₄₂ H ₂₈ Cl ₄ F ₄ N ₄ Pd	196-200	52	Gray
P ₄	NC CH3 CN HC N CH3 CH3 CN NC CH3 CN	C46H32N8Pd	>300	60	Gray
P ₅	CH CH CH	C48H32Cl4N4Pd	230-234	55	Brawn
P ₆	O ₂ N C ₁ C ₁ C ₁ NO ₂	C48H32Cl2N8O8Pd	260-263	30	Gray
P9	CI CHS CI	C42H30Cl6N4Pd	230-234	90	Gray
P ₁₀	O ₂ N CH3 NO ₂ Pd ⁻² CH3 NO ₂ O ₃ N CH3 NO ₂	C42H30Cl2N8O8Pd	220-223	68	Green yellow
P ₁₁	C CH3 WF	C42H30Br4Cl2N4Pd	>300	40	Light nutty
P ₁₂	Br CH Br	C42H32Br4N4Pd	206-209	97	Brown
P ₁₃	H ₂ C H ₃ C H ₃ C H ₃ C H ₃ C CH	C50H54Cl2N8Pd	300>	31	Gray

Table	Table 7. Rf of synthesized Palladium complexes									
Co	Wt.	Wt.	Wt. of	M.wt of	Tim	R				
mp.	of	of	Compl	complex	e of	f				
Co	ligan	Sal	ex(g)	(g/mol)	Reac					
de	d(g)	t(g)			tion					
\mathbf{P}_2	0.4	0.0	0.26	883	2	0				
		9			hrs.					
						7				
\mathbf{P}_3	0.25	0.0	0.16	912	2	0				
		6			hrs.					
						7				
P4	0.35	0.0	0.26	803	3	0				
		4			hrs.					
						7				
P ₅	0.1	0.0	0.17	913	3	0				
		2			hrs.					
						8				
P ₆	0.25	0.0	0.09	1026	3	0				
		5			hrs.					
						5				
P9	0.25	0.0	0.18	909	3	0				
		5			hrs.					
						8				
P_{10}	0.4	0.0	0.34	952	2	0				
		9			hrs.					
						8				
P ₁₁	0.1	0.0	0.04	1087	3	0				
		2			hrs.					
						4				
P ₁₂	0.25	0.0	0.3	1090	2	0				
		5			hrs.					
						8				
P_{13}	0.25	0.0	0.1	944	2	0				
		6			hrs.					
						5				

 $\label{thm:condition} Table~8.~Transitions~and~Wavelengths~and~the~suggested~Structure~of~synthesized~Palladium~(II)~complexes~$

Comp. No	Max/nmλ nm	Transition	Suggested Structure
	230 and 290	π-π* of Aromatic C=C	Square
P ₂	435	$n \rightarrow \pi^*$ of C=N - C.T	planer
	880	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	
P ₃	232 and 295	π-π* of Aromatic C=C	Square
	420	$n\rightarrow\pi^*$ of C=N - C.T	planer
	883	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	•
P ₅	230 and 296	π-π* of Aromatic C=C	Square
	470	$n\rightarrow\pi^*$ of C=N - C.T	planer
	886	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	F
P ₆	230 and 295	π-π* of Aromatic C=C	Square
	435	$n\rightarrow\pi^*$ of C=N - C.T	planer
	887	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	•
P ₉	235 and 295	π-π* of Aromatic C=C	Square
	465	$n\rightarrow\pi^*$ of C=N - C.T	planer
	886	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	•
P ₁₀	235 and	π-π* of Aromatic C=C	Square

	295		planer
	440	$n{\rightarrow}\pi^*$ of C=N - C.T]
	880	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	
P ₁₁	234 and 296	π-π* of Aromatic C=C	Square
	420	$n{\rightarrow}\pi^*$ of C=N - C.T	planer
	886	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	
P ₁₂	232 and 294	π-π* of Aromatic C=C	Square
	480	$n\rightarrow\pi^*$ of C=N - C.T	planer
	890	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	1
P ₁₃	234 and 294	π-π* of Aromatic C=C	Square
	480	$n\rightarrow\pi^*$ of C=N - C.T	planer
	886	$^{1}A_{1}g \rightarrow ^{1}A_{2}g$	

C.T=Charge transfer

Table 9. FT - IR spectra of the synthesized complexes of Palladium (II) $\,$

	υC-	υC=	υC=	υ=С-Н	Aliphatic	υ=С	υ=C-	vM	Other
Comp	N	C	N	Symmetri	Asymmetr	-H	H	-N	Groups
		Ar		c	ic	Ar	Imin		
							e		
P_2	111	1460	1606	2800	2924	3070	3120	599	NO2: 1346,
	2								1525
P_3	113	1454	1602	2892	2926	3078	3221	597	F:1215, Cl: 900
	2								
P ₄	113	1523	1633	2856	2924	3060	3217	480	CN: 2231
	0								
P ₅		1595	1637			3051	3180	586	Cl:964
P ₆	110	1417	1589	2856	2924	3050	3080	505	NO ₂ :
	7								1519,1344
P9	109	1456	1616			3064	3200	503	Cl: 1012
-	3								
P ₁₀	111	1446	1608	2860	2922	3075	3095	592	NO2:(1523,144
	2								2)/ Cl:1016
P ₁₁	107	1454	1633	2890	2926	3060	3072	590	Br: 719, Cl:
	0								1010
P_{12}	107	1587	1633			3040	3065	493	Br: 823
	8								
P ₁₃	112	1454	1612	2858	2920	3015	3040	476	Cl:941
10	2								

Table 10. Lc -Mass fragments of P2

Fragments	m / z
$\mathbf{M}^{+} = \mathbf{C}_{42}\mathbf{H}_{32}\mathbf{N}_{8}\mathbf{O}_{8}\mathbf{P}\mathbf{d}^{2+}$	883
$C_{15}H_{12}N_2Pd^{2,2+}$	327
$C_{11}H_{10}N_4Pd^{6.2+}$	306
$C_2H_2N_2Pd^{4\cdot 2+}$	158
C ₈ H ₇ N ₂ ³ •	131

Table 11. Lc -Mass fragments of P₃

Fragments	m / z
$M^{+}=C_{42}H_{28}Cl_{4}N_{4}Pd^{2+}$	913
C ₁₅ H11ClFN ₂ Pd ^{,2+}	376
C ₁₄ H ₁₀ ClFN·	245
$C_2H_2N_2^{4.2+}$	158
$C_8H_7N_2^{3.}$	131
$C_7H_6N^{3.}$	105

Table 12. Lc -Mass fragments of P₄

Tuble 12. Le Musbingments of 14				
Fragments	m / z			
$M^+ = C_{46}H_{32}N_8Pd^{+2}$	803			
C ₃₅ H ₂₄ N ₅ Pd ^{5. 2+}	628			
C9H8N4Pd ^{8. 2+}	277			
$C_{14}H_{11}N^{2}$	193			
$C_2H_2N_2Pd^{4.+2}$	163			

Table	13.	Lc	-Mass	fragments	of	P
Lanc	IJ.	L	-1V1a55	H azments	VI.	

Fragments	m / z
$M^+ = C_{48}H_{32}Cl_4N_4Pd^{2+}$	913
C ₁₈ H ₁₂ ClN ₂ Pd· ²⁺	393
C ₁₀ H ₆ N ₂ Pd ^{4. 2+}	259
$C_{10}H_6N_2^{4.}$	150
C ₇ H ₅ ClN·	136
C ₆ H ₄ Cl·	110

Table 14. LC-Mass fragments of P₆

Fragments	m / z
$M^+ = C_{48}H_{32}N_8O_8Pd^{2+}$	955
C ₃₀ H ₂₀ N ₄ Pd ^{4. 2+}	550
$C_{10}H_8N_5O_2Pd^{7.2+}$	333
C ₁₁ H ₇ N ₂ Pd ^{3, 2+}	275
C8H6N3O2 ^{3.}	173
C ₇ H ₅ N ^{2.}	103

Table 15. Lc -Mass fragments of P9

Fragments	m / z
$M^{+}=C_{42}H_{30}Cl_{6}N_{4}Pd^{2+}$	910
C ₂₃ H ₁₅ Cl ₃ N ₄ Pd ^{4. 2+}	553
C ₁₄ H ₁₀ ClNPd ^{2. 2+}	335
C ₈ H ₆ ClN ₂ Pd ^{3, 2+}	273
C9H7 ClN2 ² ·	173
C7H5ClN2 ^{4.}	153

Table 16. Lc -Mass fragments of P₁₀

Table 10. Le -Mass Hagments of 110			
Fragments	m / z		
$M^{+}=C_{42}H_{30}Cl_{2}N_{8}O_{8}Pd^{2+}$	952		
C ₂₁ H ₁₅ N ₅ O ₄ Pd ^{4. 2+}	502		
C ₁₈ H ₁₄ Cl ₂ N ₄ Pd ^{4, 2+}	467		
$C_{14}H_{10}N_3O_2^{3}$.	250		
$C_{14}H_{11}N^{2}$.	191		
C ₈ H ₆ N ₃ ³ .	173		
C ₇ H ₅ N ² .	100		

Table 17. Lc -mass fragments of P₁₁

Tuble 17. Le mass magments of 111			
Fragments	m / z		
$M^{+}=C_{42}H_{30}Br_{4}Cl_{2}N_{4}Pd^{2+}$	1088		
C ₂₁ H ₁₃ BrClN ₃ Pd ^{4. 2+}	532		
$C_{15}H_{11}N^{3}$.	221		
$C_8H_6BrN_2^{3.}$	207		
C ₇ H ₅ ^{3.}	93		

Table 18. Lc -mass fragments of P_{12}

Tuble 10: De muss it ugments of 112			
Fragments	m / z		
$M^{+}=C_{48}H_{32}Br_{4}N_{4}Pd^{2+}$	1091		
C ₂₅ H ₁₇ Br ₂ N ₃ Pd ^{2. 2+}	622		
C ₂₀ H ₁₄ BrN ₄ Pd ^{5. 2+}	498		
$C_{10}H_6N_2Pd^{4.2+}$	262		
$C_{11}H_7N^2$.	158		
C7H5N ^{2.}	102		

Table 19. Lc -mass fragments of P₁₃

Fragments	m / z
$M^{+}=C_{50}H_{54}Cl_{2}N_{8}Pd^{2+}$	944
C ₂₅ H ₂₇ ClN ₄ Pd ²⁺	519
$C_{21}H_{16}N_3^{3.}$	310
$C_{14}H_{10}N^{3}$.	194
C ₇ H ₅ ClN ₂ ⁴ ·	157
$C_7H_5N^{4.}$	105

Table 20. Molar conductance measurements of the complexes of Palladium (II)

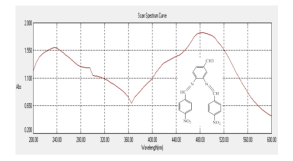
N o.	Comple xes	Concentra tion Molarity	Tempera ture °C	Molar Condacti vity Am (Ohm ⁻¹ .Cm ² .mo
P ₂	$[Pd(S_2)_2 \\]^{+2}$	1x10 ⁻³	28	1 ⁻¹)*10 ⁻⁶
P ₃	[Pd(S ₃) ₂	1x10 ⁻³	28	77
P ₄	[Pd(S ₄) ₂	1x10 ⁻³	28	71
P ₅	$[Pd(S_5)_2]^{+2}$	1x10 ⁻³	28	75
P ₆	S ₆) ₂] [Pd(+2	1x10 ⁻³	28	80
P ₉	[Pd(S ₉) ₂] +2	1x10 ⁻³	28	77
P ₁	[Pd(S ₁₀)	1x10 ⁻³	28	80
P ₁	[Pd(S ₁₁)	1x10 ⁻³	28	84
P ₁	[Pd(S ₁₂)	1x10 ⁻³	28	75
P ₁	[Pd(S ₁₃) 2] +2	1x10 ⁻³	28	79

Table 21. Ratio of Theoretical and Practically of Palladium (II)

No.	Practically	Theoretical
\mathbf{P}_2	10.17	11.15
\mathbf{P}_3	11.24	10.82
P_4	10.69	12.17
P ₅	12.05	11.66
P ₆	12.02	11.14
P 9	11.98	11.70
P ₁₀	11.78	11.18
P ₁₁	8.17	9.19
P ₁₂	8.13	9.16
P ₁₃	11.28	11.27

 $\label{eq:complexes} \textbf{Table 22. Inhibition diameter (mm) of some Palladium complexes}$

No.	Staphylococcuus Aureus			erichia oli	
	Concentration				
	50%	100%	50%	100%	
P ₂	10mm	10mm	18mm	12mm	
P 3	11mm	15mm	14mm	10mm	
P 4	16mm	15mm	16mm	17mm	
P ₉	12mm	12mm	16mm	0	
P ₁₀	12mm	12mm	15mm	9mm	



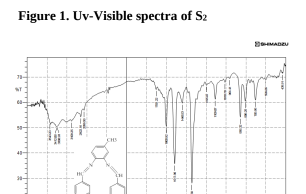


Figure 11. FT-IR spectra of S₂

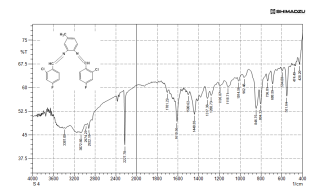


Figure 12. FT-IR spectra of S₃

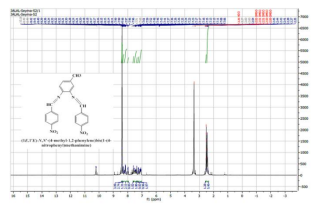


Figure 21. $^{1}\text{H-NMR}$ spectra of S_{2}

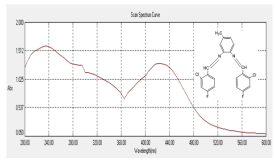


Figure 2. Uv-Visible spectra S₃

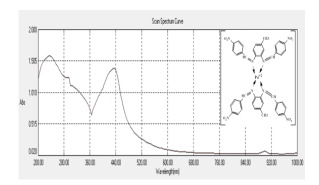


Figure 31. Uv-Visible spectra of P₂

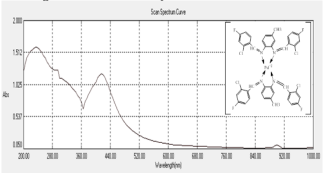


Figure 32. Uv-Visible spectra of P₃

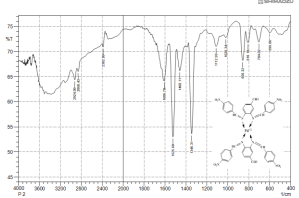


Figure 41. FT-IR spectra of P₂

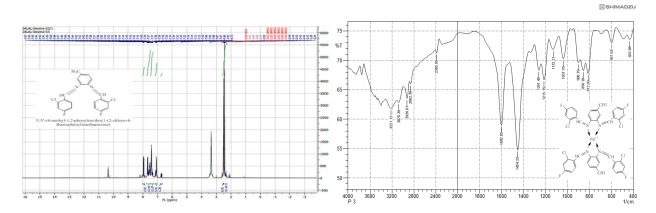


Figure 42. FT-IR spectra of P₃

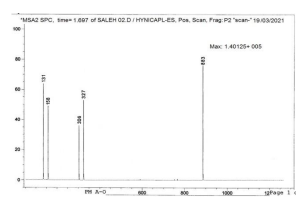


Figure 22. ¹H-NMR spectra of S₃

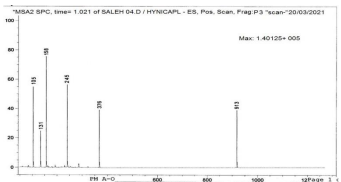


Figure 51. LC-Mass spectra of P2

Scheme 3. The general equation for imine compounds synthesis

Figure 52. LC-Mass spectra of P₃

Scheme 4. Mechanism of imine compounds synthesizing

4. Conclusions

In this study, the intermediate compounds (imine compounds) in synthesizing the palladium complexes was confirmed by the disappearance of the infrared bands and the proton spectrum signals of the amine group (-NH2) in the primary aromatic amines, the disappearance of the carbonyl (C=O) bands in the aromatic aldehydes of the primary materials and the appears of new bands and signals of the imine group (-C=N-). The result of the thin layer chromatography (TLC) and the spectrophotometrically characterized by measuring ultraviolet-visible and infrared rays, mass spectrometry and molar conductivity measurement proved the geometric shape of palladium (II) complexes exhibits a square planer. antibacterial activity of Palladium complexes was done against Staphylococcuus Aureus and Escherichia Coli, where our obtained results proved that the best activity of Palladium (II) complexes was for P₄ (16 mm, 50%) against *Staphylococcuus Aureus* and was for P₂ (18 mm, 50%) against Escherichia Coli.

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