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Synthesis, characterization, thermal studies, and DFT calculations on Pd(II) complexes containing N-methylbenzylamine

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Abstract This work describes the synthesis, characterization, and the thermal behavior investigation of four palladium(II) complexes with general formulae $[PdX_2(mba)_2]$, in which mba = N-methylbenzylamine and $X = OAc^{-}(1)$, Cl⁻(2), Br⁻(3) or I⁻(4). The complexes were characterized by elemental analysis, infrared vibrational spectroscopy, and ¹H nuclear magnetic resonance. The stoichiometry of the complexes was established by means of elemental analysis and thermogravimetry (TG). TG/DTA curves showed that the thermodecomposition of the four complexes occurred in 3-4 steps, leading to metallic palladium as final residue. The palladium content found in all curves was in agreement with the mass percentages calculated for the complexes. The following thermal stability sequence was found: 3 > 2 > 4 > 1. The geometry optimization of 1, 2, 3, and 4, calculated using the DFT/B3LYP method, yielded a slightly distorted square planar environment around the Pd(II) ion made by two anionic groups and two nitrogen atoms from the mba ligand (N1 and N2), in a transrelationship.

Keywords Palladium(II) \cdot *N*-Methylbenzylamine \cdot DFT \cdot TG and DTA

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Introduction

Cyclopalladated complexes represent an important branch in modern organometallic chemistry since they play important roles not only in organic synthesis [1], photochemistry [2], and homogeneous catalysis [3], but also in the design of liquid crystal materials [4] and new biologically active species [5]. Five-membered palladacycles bearing N-donor organic ligands such as *N*,*N*-dimethylbenzylamine (dmba) and *N*-benzylideneaniline (bzan) as well as halide and pseudohalides as coligands have been one of our major research interests [6–8]. This class of compounds exhibits promising antimycobacterial and antitumoral activities [9–11], interesting supramolecular assemblies in solid state [12], and a plethora of reactivity such as 1,3-dipolar cycloaddition of multiply bonded molecules [13, 14] and insertion reactions of acetylenes into Pd–C bond [15].

During our attempts to prepare cyclometallated complexes from the reaction between N-methylbenzylamine (mba) and palladium(II) acetate (Fuchita's synthesis [16]), we have obtained analytical and spectroscopic evidences of the formation of coordination compounds of the type $[PdX_2(mba)_2]$, in which $X = OAc^-(1)$, $Cl^-(2)$, $Br^-(3)$, $I^-(4)$. In pursuing our interest in the coordination chemistry [17–20], biological activity [21, 22], and thermal behavior [23–27] of transition metal compounds containing N-based ligands, we present herein the synthesis, characterization, DFT calculations, and thermal studies on compounds 1–4 (Scheme 1) by means of thermogravimetry (TG) and differential thermal analysis (DTA).

Experimental

General comments

All reagents were obtained from commercial suppliers and used without further purification.



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Scheme 1 Structural representation of the compounds $[PdX_2(mba)_2](X = OAc^-(1), Cl^-(2), Br^-(3), I^-(4))$

Synthesis of the complexes

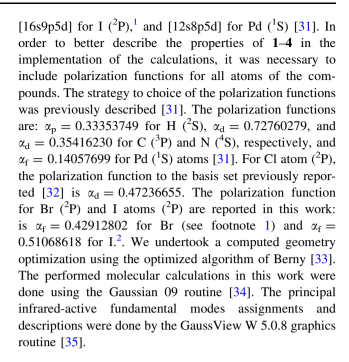
Complex 1 was prepared as follows: 4.91 mmol (595 mg) of *N*-methylbenzylamine were added to a suspension of [Pd(OAc)₂] (4.44 mmol, 995.6 mg) in 50 mL of benzene. The mixture was stirred at 60 °C for 1 day. The resulting yellow suspension was filtered off. The solid material was washed with benzene and dried under vacuum. Yield: 77%. Compounds 2–4 were readily obtained by metathetical reactions of [Pd(OAc)₂(mba)₂] with the appropriate amount of KX salts in a mixture of methanol:acetone (2:1). The resulting suspensions were filtered off and the solids were washed and dried under vacuum. Yield 62–87%.

Instrumentation

Elemental analyses of carbon, nitrogen, and hydrogen were performed on a microanalyser elemental analyser CHN, model 1112 FLASH EA. Infrared spectra were recorded in KBr pellets on a Nicolet model SX-FT-Impact 400 spectrophotometer in the 4000–400 cm⁻¹ spectral range. ¹H NMR spectra were recorded in CDCl₃ solutions at room temperature on a Varian INOVA 500 spectrometer. Melting points were determined on a MAPFQ apparatus. Thermal analyses (TG) and differential thermal analyses (DTA) were carried out using a TA Instruments model SDQ 600, under flow of dry synthetic air (50 mL min⁻¹), temperature up to 900 °C and at heating rate of 20 °C min⁻¹ in α -alumina sample holders. The reference substance was pure α-alumina in DTA measurements. X-ray powder diffraction patterns were measured on a Siemens D-5000 X-ray diffractometer using Cu K_α radiation ($\lambda = 1.541 \text{ Å}$) and setting of 34 kV and 20 mA. The peaks were identified using ICDD bases [28].

Computational strategy

In this work, the employed quantum chemical approach to determining the molecular structures was Becke three-parameter hybrid theory [29] using the Lee-Yang-Par (LYP) correlation functional [30] and the basis sets used for calculations were: [4s] for H (²S) [31], [5s4p] for C (³P), N (⁴S), and O(³P) [31], [11s7p] for Cl (²P) [32],



Results and discussion

The elemental analysis and thermogravimetric results together with IR spectroscopy and ¹H NMR data confirmed the proposed formulae for the complexes 1–4. Table 1 represents the colors, melting points and the results of elemental analysis.

Infrared spectra

Table 2 represents some selected absorptions of the IR spectra of the complexes along with their assignments. The overall pattern of the IR spectra of 1–4 resembles very closely to that of the free ligand. The coordination of



Table 1 Results of elemental analyses and melting points of the compounds 1-4

Complex	Color	M.p./°C	Found (calcd.)/%			
			С	Н	N	
$C_{20}H_{28}N_2O_4Pd$ (1)	Yellow	129	51.9 (51.5)	5.9 (6.1)	6.3 (6.0)	
$C_{16}H_{22}N_2Cl_2Pd$ (2)	Yellow	170	45.5 (45.8)	5.5 (5.3)	6.6 (6.7)	
$C_{16}H_{22}N_2Br_2Pd$ (3)	Yellow	154	37.9 (37.8)	4.7 (4.4)	5.5 (5.7)	
$C_{16}H_{22}N_2I_2Pd$ (4)	Orange	155	32.2 (31.9)	3.6 (3.7)	4.7 (4.7)	

N-methylbenzylamine by the nitrogen atom in 1-4 was evidenced by the appearance of the typical absorptions of the -NHCH₃ moiety at ~ 3200 (vNH), 2920 (v_{as} CH₃), and 2862 cm⁻¹ (v_sCH₃). The shift of the absorption band attributed to the $\delta NH + \delta CH_3$ mode to lower frequency (ca. 1052 cm⁻¹) when compared with that one of the free ligand (1103 cm⁻¹) is also indicative of coordination. In addition, two out-of-plane C-H bending vibrational modes (γCH) were observed as very intense bands at 750 and 700 cm⁻¹ which strongly supports the presence of monosubstituted phenyl rings. The $v_{as}COO^-$ and $v_sCOO^$ vibrational modes for coordinated acetate groups in 1 appeared as two broadened and intense absorptions over the spectral range of 1650–1350 cm⁻¹, respectively. The vPd-X and vPd-N band frequencies are expected to occur below 350 cm⁻¹ [36]. However, the vPd-X and vPd-N absorptions could not be detected since the spectrophotometer used in this work operates in the 4000–400 cm⁻¹ range.

¹H NMR spectra

The observed chemical shifts in the ¹H NMR spectra of *N*-methylbenzylamine and the complexes are shown in Table 3.

The ¹H NMR spectra of **1–4** showed that the *N*-methylbenzylamine is coordinated to the metallic center by the nitrogen atom due to the downfield shift of NH signals from 1.35 ppm (mba ligand) to 2.42–3.48 ppm. In addition, the methylene group attached to NH in the complexes **1–4** appeared as multiplets whereas in the ¹H NMR spectrum of mba a singlet was noticed at 3.69 ppm. The observed loss of magnetic equivalence of methylene groups the ¹H NMR spectra of the complexes clearly indicates the coordination of mba.

The analytical and IR results obtained for compounds 1–4 suggest a square planar environment around the Pd atom whose coordination sites are occupied by two nitrogen atoms from mba ligands, and two anionic X⁻ groups (X⁻ = Cl, Br, I, OAc). The *trans*-configuration was attributed to these complexes on basis of known X-ray structures of similar compounds of general formulae *trans*-[PdX₂L₂] {L = N-based ligands; X⁻ = Cl [37], Br [38], I [39], OAc [40]}. As no single crystal for X-ray diffraction studies could be obtained, the structures of the Pd(II) compounds 1–4 were optimized using DFT theory (B3LYP method) and are shown in Fig. 1. A selection of calculated bond lengths and angles is shown in Table 4.

The calculated structures for 1-4 show a satisfactory agreement with the available crystal structure data of similar compounds of the type $[PdX_2L_2]$ (L = N-based ligands) [37-40], mainly in terms of bond lengths. The calculations of vibrational frequencies were employed to determine whether optimized geometries constitute

Table 2 Selected experimental (exp) and calculated (calc) IR frequencies (cm⁻¹) together with percentual error (p.e.) for complexes 1-4

Assignment	Wavenumber/cm ⁻¹											
	1			2		3			4			
	exp	calc	p.e.	exp	calc	p.e.	exp	calc	p.e.	exp	calc	p.e.
vNH	3178	3100	-2.5	3221	3391	+5.3	3198	3382	+5.7	3178	3375	+6.2
νCH_{ring}	3049	3181	+4.3	3032	3172	+4.6	3026	3172	+4.8	3049	3171	+4.0
vCH ₃	2920	3097	+6.0	2921	3106	+6.3	2914	3028	+3.9	2920	3032	+3.8
$v_{\rm as} {\rm COO}$	1630	1613	-1.0	_	_	_	_	_	_	_	_	_
$v_{\rm ring}$	a	1532	_	1495	1487	-0.5	1491	1488	-0.6	1491	1488	-0.2
$\delta \mathrm{CH}_2$	a	1379	_	1423	1377	-3.2	1421	1377	-3.1	1454	1385	-4.7
$\delta \text{CH}_3 + v_s \text{COO}$	1315	1350	+2.7	_	_	_	_	_	_	_	_	_
ν CN + γ CH	908	891	-1.9	910	881	-3.2	905	880	-2.8	905	876	-3.2
γCH_{ring}	748	760	+1.6	746	767	+2.8	748	765	+2.3	748	768	+2.7
γCH_{ring}	a	716	_	698	716	+2.6	702	716	+2.0	696	723	+3.9
δ OCO	688	681	-1.0	_	_	_	_	_	_	_	_	_

a Obscured



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Table 3 ¹H NMR data (ppm) for compounds **1–4** at 298 K, in CDCl₃-d given as δ (ppm), multiplicity [integration]

Scheme		¹ H NMR data						
		NH	-CH ₂ -	N-CH ₃	-C ₆ H ₅			
нн	mba	a	3.69 s, [2H]	2.40 s, [3H]	7.20–7.27 m, [5H]			
H H H₃Ç ¶	1	a	3.14-3.09 m, [1H]	2.12 s, [3H]	7.42–7.25 m, [5H]			
			3.90-3.85 m, [1H]					
Pd H	2	3.38 s, [1H]	3.47-3.51 m, [1H]	2.43 s, [3H]	7.36–7.43 m, [5H]			
	J.		4.21–4.25 m, [1H]					
•	3	3.39 al, [1H]	3.50-3.58 m, [1H]	2.47-2,42 m, [3H]	7.36–7.42 m, [5H]			
			4.21-4.28 m, [1H]					
	4	3.40 al, [1H]	3.44-3.49 m, [1H]	2.45 s, [3H]	7.40-7.48 m, [5H]			
			4.01–4.07 m, [1H]					

s singlet, m multiplet

Fig. 1 Calculated structures of trans-[PdX₂(mba)₂] (X = OAc⁻ (1), Cl⁻ (2), Br⁻ (3), I⁻ (4))

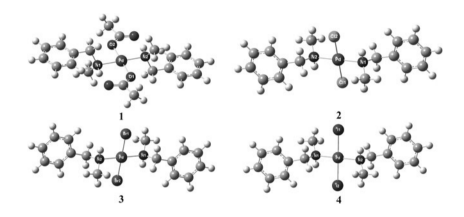


Table 4 Selected calculated bond lengths (\mathring{A}), and bond angles ($^{\circ}$) for 1, 2, 3, and 4

$[Pd(OAc)_2(mba)_2] (1)$		[PdCl ₂ (mba) ₂] ([PdCl2(mba)2] (2)		[PdBr2(mba)2] (3)		[PdI ₂ (mba) ₂] (4)	
Bond lengths/Å								
Pd-N1	2.11213	Pd-N1	2.10607	Pd-N1	2.11459	Pd-N1	2.11469	
Pd-N2	2.11261	Pd-N2	2.10603	Pd-N2	2.11448	Pd-N2	2.11470	
Pd-O1	2.07235	Pd-Cl1	2.38488	Pd-Br1	2.50963	Pd-I1	2.59335	
Pd-O2	2.07217	Pd-C12	2.38499	Pd-Br2	2.50974	Pd-I2	2.59329	
Bond angles/°								
N1-Pd-O1	84.932	N2-Pd-C12	93.718	N2-Pd-Br2	93.740	N2-Pd-I2	94.253	
O1-Pd-N2	95.020	Cl2-Pd-N1	86.273	Br2-Pd-N1	86.256	I2-Pd-N1	85.758	
N2-Pd-O2	84.947	Cl1-Pd-N1	93.725	Br1-Pd-N1	93.745	I1-Pd-N1	94.231	
O2-Pd-N1	95.100	Cl1-Pd-N2	86.284	Br1-Pd-N2	86.260	I1-Pd-N2	85.757	
O1-Pd-O2	179.962	Cl1-Pd-Cl2	179.970	Br1-Pd-Br2	179.945	I1-Pd-I2	179.989	
N1-Pd-N2	179.940	N1-Pd-N2	179.982	N1-Pd-N2	179.977	N1-Pd-N2	179.975	

minimum or saddle points. The most important theoretical frequencies values calculated for the compounds are also depicted in Table 2. The calculated frequencies are in good agreement with the experimental values, with a percentual error less than 6%, supporting the suggested *trans*-[PdX₂(mba)₂] structure.

Thermogravimetric analysis

TG and DTA curves obtained for the compounds 1–4 are shown in Fig. 2 and Table 5 lists the results of the thermal studies of these compounds together with the assignments of each decomposition stage based on mass calculation.



a Not observed

Fig. 2 TG and DTA curves of the complexes $[PdX_2(mba)_2]$ $(X = OAc^-(1), Cl^-(2), Br^-(3), I^-(4))$

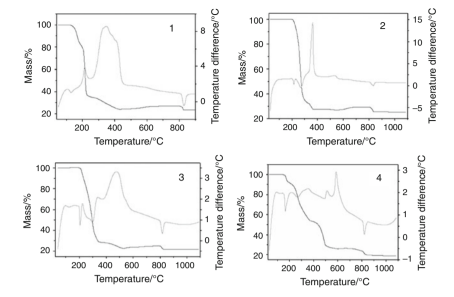


Table 5 Thermal analysis data for compounds $[PdX_2(mba)_2]$ $(X = OAc^-(1), Cl^-(2), Br^-(3), I^-(4))$

Complex	Step	$\Delta T/^{\circ}\mathrm{C}$	$\Delta m/\%$	DTA peak/°C		
				Endo	Exo	
1	1	111–421	-75.71	125.3	213; 251–468	
	2	421-813	+2.42	_	_	
	3	813-829	-3.21	828	_	
	Residue		23.50	_	_	
2	1	189-365	-72.53	214; 269	362	
	2	365-799	+1.86	_	532	
	3	799-842	-3.93	833	_	
	Residue		25.40	_	_	
3	1	197-355	-71.29	204; 224–337	_	
	2	355-529	-5.95	_	476	
	3	529-789	+2.30	_	_	
	4	789-832	-3.07	817	_	
	Residue		21.99	_	_	
4	1	134-368	-45.89	164; 265	352	
	2	368-612	-28.15	_	511; 585	
	3	612-809	-2.73	_	_	
	4	809-875	-3.70	822	_	
	Residue		19.53	_	_	

Therefore, the groups indicated at the right column of Table 2 do not correspond necessarily to the gaseous final products of decomposition. The X-ray powder diffractograms of the final products, obtained after the decomposition of PdO, showed the characteristic peaks of Pd (ASTM 05-0681).

Complex 1 and 2 present similar decomposition patterns. The first step of decomposition corresponds to the release of organic and inorganic ligands together with

uptake of O_2 , leading to the partial conversion of Pd to PdO. In complex 1 this step occurs in the range 111–421 °C and is accompanied by one endothermic peak at 125 °C in the DTA curve and two exothermic signals at 213 °C and 251–468 °C. In the second step there is an increase in mass of 2.42% which is attributed to the uptake of O_2 and conversion of the remaining Pd to PdO. This process is completed at 813 °C. The third and final step corresponds to the decomposition of PdO to Pd, which finishes at 829 °C and is accompanied by an endothermic peak at 828 °C in the DTA curve. This behavior was already observed for other palladium complexes [21].

For the complex **2**, the first step of decomposition occurs in the range 189–365 °C, accompanied by two endothermic peaks at 214 and 269 °C and one exothermic signal at 362 °C. This mass variation is ascribed, by mass calculations, to the release of organic and inorganic ligands and partial conversion of Pd to PdO. The second step $(365–799 \,^{\circ}\text{C})$ is attributed to an increase in mass of 1.86%, related to the uptake of O_2 and conversion of the remaining Pd to PdO. This process is related to the exothermic peak at 532 °C in the DTA curve. The third step is completed at 842 °C and corresponds to the decomposition of PdO to Pd which is associated with the endothermic peak at 828 °C.

The decomposition of complex 3 occurs in four steps. The release of organic and inorganic ligands together with uptake of O_2 occurs in the two initial steps: 197–355 and 355–529 °C. These events are accomplished by endothermic signals at 204 and 224–337 °C and by an exothermic peak at 476 °C in the DTA curves. The final residue of the second step consists of a mixture of Pd and PdO. The third step (529–789 °C) corresponds to the uptake of O_2 and conversion of the remaining Pd to PdO. The last mass loss is characterized by the decomposition of PdO to Pd and is



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completed at 832 °C (endothermic peak in the DTA curve at 817 °C).

Complex 4 displays a different thermal decomposition pattern. TG curve of compound 4 shows two consecutive mass loss in the range 134–368 °C assigned, by mass calculation, to the loss of one organic ligand unit (mba) and one iodide. This process is related to the endothermic peaks at 164 and 265 °C and one exothermic peak at 352 °C. The following steps are overlapped: a mass loss at 368–612 °C, followed by a slight mass increase over the range 612–669 °C and a final mass decrease which is completed at 809 °C. Due to the complexity of the decomposition, the assignments of these steps could not be done by mass calculations. A further increase of temperature to 879 °C results in the last mass loss which is ascribed to the reduction of PdO to Pd. The palladium content found in the decomposition is in agreement with the mass percentage calculated for the complex.

Taking into account the initial decomposition temperatures, the thermal stability of the complexes $[PdX_2(mba)_2]$ varies in the sequence: X = 3 > 2 > 4 > 1. The thermodecomposition of the complex $[Pd(OAc)_2(mba)_2]$ (1) initiates at the lowest temperature among all the complexes studied. The low thermal stability of complex 1 was already expected in agreement with our previous results [8]. On the other hand, bromo- and chloro-derivatives exhibit the highest thermal stability.

Conclusions

Synthesis, characterization, and thermal behavior of four $[PdX_2(mba)_2]$ type complexes in which mba = N-methylbenzylamine and $X = OAC^{-}(1)$, $Cl^{-}(2)$, $Br^{-}(3)$, and $I^{-}(3)$ (4), were investigated in this work. IR and ¹H NMR spectra indicated that in the four complexes the organic ligand is coordinated to the metal by the nitrogen atom in a neutral monodentate mode. Taking into account the initial decomposition temperatures, the following thermal stability sequence could be established: 3 > 2 > 4 > 1. The thermogravimetric data showed that all decompositions initiate with the release of organic and inorganic ligands, followed by the uptake of O2, leading to the formation of a mixture of Pd and PdO. Afterwards, all PdO is decomposed to Pd⁰, which is the final residue of the thermal decomposition of 1-4. The computational strategy used in this work represents a good alternative for calculations on vibrational frequencies of metal-based compounds.

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