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Synthesis, X-ray structure, Hirshfeld analysis, and DFT studies of a new Pd(II) complex with an anionic *s*-triazine *NNO* donor ligand

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Abstract

A new Pd(II) complex, [Pd(Triaz)Cl], with the hydrazono-*s*-triazine ligand, 2,4-di-*tert*butyl-6-((2-(4-morpholino-6-(phenylamino)-1,3,5-triazin-2-yl)hydrazono)methyl)phenol, was synthesized by the reaction of PdCl₂ with the organic ligand (1:1) in acetone under isothermal conditions. The molecular structure of the [Pd(Triaz)Cl] complex was determined using FTIR and ¹H NMR spectroscopic techniques, and single-crystal X-ray diffraction. Moreover, using Hirshfeld surface analysis, the percentages of the intermolecular interactions were determined. The obtained values were 60.6%, 11.6%, 8.1%, 3.6%, and 5.0% for the H····H, C···H, O···H, N···H, and Cl···H interactions, respectively. Among them, the O···H, C···H and C···N interactions are considered extremely important. Natural bond orbital calculations have been used to calculate the amount of electron transfer from the ligand to the metal ion and to evaluate the Pd-N, Pd-O, and Pd-Cl coordination bonding interactions.

Keywords: S-triazine; Hydrazone; Pd(II) complex; Hirshfeld surface analysis; NBO

Introduction

s-Triazine is a heterocyclic motif for multiple synthetic and natural products [1]. Being an important scaffold, *s*-triazine derivatives have been extensively studied in both pharmaceutical and coordination chemistry [2,3]. Some of these derivatives, e.g., 2-amino-4-morphlino-*s*-triazine, and hexamethylmelamine, exhibited a wide range of pharmaceutical features such as antitumor activity [4], antioxidant [5], antiviral [6], herbicidal, antimicrobial, and antibacterial [7] properties. Furthermore, the syntheses of various mono-, di-, and trisubstituted *s*-triazines derivatives from cyanuric chloride have been reported-with remarkable pharmaceutical activities [8-13] In particular, hydrazine-based *s*-triazines derivatives have shown potency activities as anticancer agents for A431 (epidermoid carcinoma) and A549 (lung carcinoma) compared to Lapatinib, HKI-272, and AST-1306 as standard references [14]. In addition, they have been used as inhibitors for wtEGFR, EGFR, and mTOR receptors [15,16].

Recently, our research group has reported several complexes bearing *NNN*-tridentate ligands derived from *s*-triazine with divalent metal ions, such as Co(II) [17], Ni(II) [18], Cd(II) [19a] and Mn(II) [19b]. Furthermore, *s*-triazine ligands have been used for recovering Pd(II) from strongly acidic solutions [20].

In this work, we studied the structural aspects of a new Pd(II) complex, [**Pd(Triaz)Cl**], which contains the hydrazono-*s*-triazine ligand (HTriaz) 2,4-di-*tert*-butyl-6-((2-(4-morpholino-6-(phenylamino)-1,3,5-triazin-2-yl)hydrazono)methyl)phenol (**Scheme 1**). Both molecular and supramolecular structures were investigated using X-ray crystallography, FTIR ¹H NMR spectra, Hirshfeld surface analysis, and DFT calculations. The anti-cancer activity was also explored against two cancer cell lines including breast cancer (MCF-7) and human prostate cancer (PC3).



Scheme 1. Synthesis of the [Pd(Triaz)Cl] complex.

Experimental

General

FTIR spectra were measured on a PerkinElmer Spectrum 100 FT-IR spectrophotometer (PerkinElmer Inc., Waltham, MA, USA). ¹H NMR spectra of **HTriaz** and [**Pd**(**Triaz**)**Cl**] were recorded on DMSO- d_6 using a JEOL 400 MHz spectrometer (JEOL Ltd., Tokyo, Japan) at room temperature. Mass spectra were recorded on JMS-600 H JEOL spectrometer (JEOL Ltd., Tokyo, Japan).

Synthesis of the ligand (*E*)-2,4-di-*tert*-butyl-6-((2-(4-morpholino-6-(phenylamino)-1,3,5-triazin-2-yl)hydrazono)methyl)phenol (HTriaz)

The ligand (HTriaz) was prepared as previously reported [21].

Synthesis of [Pd(Triaz)Cl] complex

Palladium(II) chloride (21.1 mg, 0.119 mmol) was added to 20 mL of an acetone solution of (**HTriaz**) (60.0 mg, 0.119 mmol). The reaction mixture was stirred at 50 °C for 4 days. Afterwards, the mixture was filtered and maintained at room temperature to allow for the slow evaporation of the solvent, yielding the final product [**Pd**(**Triaz**)**Cl**] as reddish-brown block crystals.

¹H NMR (400 MHz, DMSO- d_6 , ppm): δ 10.84 (s, 1H, NH), 8.22 (s, 1H, NH), 7.97 (s, 1H, CH=N), 7.60 (d, 2H, $J_{\text{HH}} = 8.1$ Hz, phenyl), 7.39 (t, 2H, $J_{\text{HH}} = 7.6$ Hz, phenyl), 7.30 (s, 1H, C₆H₂), 7.22 (s, 1H, C₆H₂), 7.13 (t, 1H, $J_{\text{HH}} = 7.6$ Hz, phenyl), 3.74 (d, 4H, $J_{\text{HH}} = 4.5$ Hz,

morpholine), 3.68 (d, 4H, $J_{\rm HH} = 5.0$ Hz, morpholine), 1.37 (s, 9H, *tert*-butyl), 1.26 (s, 9H, *tert*-butyl). IR (KBr) cm⁻¹: 3435 (NH), 3263, 3102, 2956, 2856, and 1626 (C=N), and 1579 (C=C). LC-MS (ESI⁺-MS): 627.43 [M]⁺. Anal. calcd. for C₂₈H₃₆ClN₇O₂Pd: C, 52.18; H, 5.63; N, 15.21. Found: C, 52.15; H, 5.59; N, 15.25.

X-Ray structural determinations

The crystal of **[Pd(Triaz)CI]** was immersed in cryo-oil, mounted in a loop, and measured at 170 K. XRD data were collected on a Bruker Kappa Apex II diffractometer with MoK*a* radiation. For cell refinement and data reduction, the *Denzo-Scalepack* [22] software package was used. A numerical absorption correction (*SADABS* [23]) was applied to the data before structure solution. The structure of **[Pd(Triaz)CI]** was solved by intrinsic phasing using the SHELXT software [24]. Structural refinement was then performed using the SHELXL-2017 package [24] and the *SHELXLE* [25] graphical user interface. A disordered acetone molecule of as solvent of crystallization in **[Pd(Triaz)CI]** was disordered over two sites sharing locations of C28 and C28B carbon atoms. The occupancy ratio of the disordered components was 0.51/0.49. The NH hydrogen atoms were located from the difference Fourier map and isotropically refined. The rest of the hydrogen atoms were geometrically positioned and constrained to ride on their parent atoms with C-H = 0.95–0.99 Å and U_{iso} = 1.2–1.5_{eq} (parent atom). Topology analyses were performed using the Crystal Explorer 17.5 program [26] to determine the contribution percentages of the different intermolecular interactions in the crystal structure of **[Pd(Triaz)CI]**.

Computational details

Density functional theory (DFT) single point calculations were performed using the Gaussian 09 software package [27] with the MPW1PW91 and Wb97XD methods [28] combined with the cc-PVTZ and cc-PVTZ-PP [29] basis sets for nonmetal atoms and Pd, respectively. The multiplicity of the system was set to be 1 because the well known fact that tetra-coordinated

Pd(II) complexes are square planar and diamagnetic. The natural charge populations and interaction energies between the donor atoms and the central metal ion, Pd(II), were computed using NBO 3.1 [30] program.

Results and discussion

Synthesis and characterization of [Pd(Triaz)Cl]

The Pd(II) complex, **[Pd(Triaz)Cl]**, was synthesized as shown in **Scheme 1** and characterized by FTIR, and ¹H NMR spectroscopy, ESI⁺-MS, elemental analysis, and single crystal X-ray diffraction.

The FTIR spectrum of the Pd(II) complex [**Pd**(**Triaz**)**Cl**] exhibits signals characteristic for the functional groups as follow: NH (3435), C=N (1628), and C=C (1579 cm⁻¹). In the ¹H NMR spectrum, the five phenyl protons were observed as doublet, triplet, and triplet at δ 7.60, 7.39, and 7.13 ppm with coupling constants (*J*_{HH}) of 8.1, 7.6, and 7.6 Hz (integrations 2:2:1), respectively. Furthermore, the two protons of the tetra substituted phenyl ring (C₆H₂) were observed as two singlets at δ 7.30 and 7.22 ppm. The eight morpholine protons were then detected as doublets at δ 3.74 and 3.68 ppm with coupling constants (*J*_{HH}) of 4.5 and 5.0 Hz, respectively. The signals of the two *tert*-butyl groups were detected as singlets at δ 1.37 and 1.26 ppm. The two N*H* protons, which appeared at δ 11.30 and 9.34 ppm in the (**HTriaz**) ligand, were slightly shifted in the Pd(II) complex to δ 10.84 and 8.22 ppm. The C*H*=N proton was observed at δ 8.28 ppm in (**HTriaz**); however, it was detected at δ 7.97 ppm in the Pd(II) complex (**Figs. 1–3**).



Fig. 1. ¹H NMR spectrum of the (**HTriaz**) ligand in DMSO- d_6 .



Fig. 2. ¹H NMR spectrum of the [Pd(Triaz)Cl] complex in DMSO-*d*₆.



2.0 11.5 11.0 10.5 10.0 9.5 9.0 8.5 8.0 7.5 7.0 6.5 6.0 5.5 5.0 4.5 4.0 3.5 3.0 2.5 2.0 1.5 1.0 0.5 (f1 (ppm)

Fig. 3. ¹H NMR spectra of (HTriaz) and [Pd(Triaz)Cl] in DMSO-*d*₆.

Crystal structure

Fig. 4 shows the X-ray structure of the [**Pd**(**Triaz**)**Cl**] complex that crystallized in the monoclinic crystal system and C2/c space group with one molecular unit in the asymmetric unit and Z=8 (**Table 1**). The deprotonated *s*-triazine ligand (**Triaz**[¬]) acted as a mononegative *NNO*-donor tridentate chelate coordinating with Pd(II) ion *via* one nitrogen from the *s*-triazine core, one nitrogen from the hydrazone moiety, and the phenolic oxygen atom. The distances between the Pd and donor atoms are 2.054(3), 1.948(3), and 1.964(2) Å for the Pd1-N1, Pd1-N7, and Pd1-O1 bonds, respectively. The coordination sphere of this complex is completed by one chloride ion located *trans* to the N7 atom with a Pd1-Cl1 distance of 2.3270(11) Å. The bite angles of the organic ligand are 80.78°(12) and 91.81°(11) for N7-Pd1-N1 and N7-Pd1-O1, respectively. **Table 2** provides a list of the most important distances

and angles. The angle between the *cis* bonds are in the range of 80.78(12) to $102.14(9)^{\circ}$ for the N7-Pd1-N1 and N1-Pd1-Cl1, respectively; and the angles of the *trans* O1-Pd1-N1 and N7-Pd1-Cl1 bonds are 172.46(11) and $177.08(9)^{\circ}$, respectively. The coordination geometry of the Pd(II) ion is considered to be distorted square planar.

CCDC	1989545
Empirical formula	$C_{31}H_{42}ClN_7O_3Pd$
Fw	702.56
Temp (K)	170(2)
λ (Å)	0.71073
Crystal system	Monoclinic
Space group	C2/c
<i>a</i> (Å)	34.4695(9)
<i>b</i> (Å)	10.1613(3)
<i>c</i> (Å)	22.4498(6)
β (deg)	122.8340(10)
$V(\text{\AA}^3)$	6607.0(3)
Z	8
$\rho_{\rm calc}~({\rm Mg/m}^3)$	1.413
μ (MoK α) (mm ⁻¹)	0.685
No. reflns.	45096
Unique reflns.	7566
GOOF (F ²)	1.086
R _{int}	0.0749
$R1^{a} (I \ge 2\sigma)$	0.0528
wR2 ^b ($I \ge 2\sigma$)	0.0880

Table 1. Crystallographic data of [Pd(Triaz)Cl].

^{*a*} $R1 = \Sigma |\overline{|F_{o}| - |F_{c}||/\Sigma|F_{o}|}$. ^{*b*} wR2 = $[\Sigma[w(F_{o}^{2} - F_{c}^{2})^{2}]/\Sigma[w(F_{o}^{2})^{2}]]^{1/2}$.



Fig. 4. X-ray structure of **[Pd(Triaz)Cl]**. Thermal ellipsoids were drawn at the 50% probability level.

Atoms	Distance	Atoms	Angle (°)
Pd1-N7	1.948(3)	N7-Pd1-O1	91.81(11)
Pd1-O1	1.964(2)	N7-Pd1-N1	80.78(12)
Pd1-N1	2.054(3)	O1-Pd1-N1	172.46(11)
Pd1-Cl1	2.3270(11)	N7-Pd1-Cl1	177.08(9)
		O1-Pd1-Cl1	85.27(8)
		N1-Pd1-Cl1	102.14(9)

Table 2. Selected bond lengths [Å] and angles [°] for [Pd(Triaz)Cl].

Analysis of molecular packing

Hirshfeld d_{norm} surfaces and fingerprint plots were used to quantify the contributions of the different intermolecular interactions (**Fig. 5**). Because of the presence of a disordered solvent molecule, the Hirshfeld analysis results are presented for the two disordered components. **Fig. 6** shows a summary of the most important contacts and their contribution percentages. The two components yielded similar results; hence, the discussion was focused only on part A for simplicity. The Hirshfeld d_{norm} maps of the Pd(II) complex demonstrated that its units

are mainly packed by the nonpolar H····H interactions, which correspond to 60.6% from the whole fingerprint area. The second most abundant atom-atom interactions were the C····H contacts, accounting for 11.6%. The polar O····H, N····H, and Cl····H interactions comprised 8.1%, 3.6% and 5.0%, respectively. In Fig. 6, most of the intermolecular interactions have contact distances longer than the sum of the van der Waals radii of the two elements of each contact, except for the O····H, C···H, and C···N interactions, which appeared as red spots in the d_{norm} map (**Figs. 7** and **8**). As shown in **Fig. 7**, the O···H contacts is represented as an intense red circle in its d_{norm} map, indicating the hydrogen bond between the acetone oxygen atom and the proton from the N-H group of the organic ligand. According to Hirshfeld analysis, the corresponding O···H interaction distance is 1.839 Å (O3···H6); however, the refined model indicates the distance is 2.06(4) Å. The shortest C···H contacts, which belong to the C-H··· π interactions, were C4···H30B and C16···H9B (2.608 and 2.696 Å). Furthermore, the shortest C···N contact was the C14···N3 (3.184 Å), indicating the existence of weak π – π stacking interactions.







Fig. 5. Hirshfeld surfaces of the [Pd(Triaz)Cl] complex.



Fig. 6. Contribution percentages of the intermolecular interactions in the crystal structure of **[Pd(Triaz)Cl]**.



Fig. 7 2D fingerprint plot (left) and d_{norm} map (right) of the O···H contacts of [Pd(Triaz)Cl].



Fig. 8. Decomposed d_{norm} (-0.06 to 3.72) and FP plots of the C···H (orange) and C···N (black) contacts of [Pd(Triaz)Cl].

DFT studies

Natural population analysis

The divalent Pd ion was coordinated with two negatively charged ligand groups (Cl⁻ and Triaz⁻). Their charges as isolated ions were +2, -1, and -1, respectively. Because of the interaction between the Pd(II) ion (Lewis acid) and the ligand groups (Lewis bases), part of the negative charge was transferred from ligand to the Pd(II) (**Table 3**). The chloride ion transferred 0.506–0.520 e⁻ to Pd(II); however, the anionic organic ligand transferred almost all of its negative charge to the metal center (0.963–0.974 e⁻). Therefore, Triaz⁻ has a net charge very close to zero (-0.026 to -0.037 e⁻) and the charge of Pd(II) decreased to 0.505–0.530 e⁻.

Table 3. Natural charges at the Pd(II) atom, and the coordinated Cl⁻ and Triaz⁻.

Atom	MPW1PW91	WB97XD	
Pd	0.5053	0.5303	
CΓ	-0.4797	-0.4937	
Triaz ⁻	-0.0256	-0.0367	

Using natural bond orbital (NBO) calculations, the strength of the interactions between the Pd(II) center and ligand donor atoms was estimated (**Table 4**). The interaction energy of the Pd-N(hydrazone) was significantly higher than that for the Pd-N(*s*-triazine). There is only one anti-bonding natural orbital (LP*(6)) from the Pd(II) that contributes in the interaction with the N1 lone pair filled nonbonding natural orbital. The LP*(6) NBO has mainly a *s*-orbital character with some contribution from *p*-orbitals and very little contribution from *d*-orbitals. This can be inferred from the nearly spherical shape of the isodensity surface shown in **Fig. 9**. Similarly, the Pd-N(hydrazone) bonds are attributed to the interaction of the filled lone pair NBO from the corresponding nitrogen atom with the empty LP*(5)Pd anti-bonding NBO. The orbital character compared to the latter. The LP*(6) NBO contributed to all of the Pd-N, Pd-O, and Pd-Cl interactions. The Pd-O coordination interaction was

attributed to the overlap between the LP(3)O filled NBO and LP*(5)Pd empty anti-bonding NBO. The net interaction energy of the Pd-O bond was 193.41 and 168.99 kcal/mol, using WB97XD and MPW1PW91 methods, respectively. The Pd-Cl coordination was attributed to the mixed interactions between the filled NBOs of the Cl⁻ ligand and the anti-bonding NBOs of the Pd(II) central metal ion (**Table 4**). **Fig. 9** shows the anti-bonding NBOs of Pd(II), and their corresponding occupancies and energies are listed in **Table 5**. The filled donor NBOs of the isolated ligand have almost 2.0 e⁻, which are significantly lowered in the complex because of the interactions with the Pd(II) anti-bonding NBOs. The latter have almost zero occupancy in the free (isolated) Pd(II) and increased up to 0.4662 e⁻ in LP*(5)Pd because of the electron donation from the ligand donor atoms to the metal ion.

Table 4. The donor (NBO_i)-acceptor (NBO_j) interaction energies of Pd-N, Pd-O, and Pd-Cl coordination interactions.

NBO _i	NBOj	WB97XD	MPW1PW91
LP(1)O1	LP*(5)Pd1	13.75	15.38
LP(1)O1	LP*(7)Pd1	20.41	20.00
LP(2)O1	LP*(8)Pd1	6.06	4.47
LP(3)O1	LP*(5)Pd1	116.79	97.98
LP(3)O1	LP*(6)Pd1	8.53	7.23
LP(3)O1	LP*(7)Pd1	27.87	23.93
O1→Pd1		193.41	168.99
LP(1)N7	LP*(5)Pd1	118.81	101.16
LP(1)N7	LP*(6)Pd1	11.98	10.74
LP(1)N7	LP*(7)Pd1	27.60	24.05
N7→Pd1		158.39	135.95
LP(1)N1	LP*(6)Pd1	104.23	90.88
N1→Pd1		104.23	90.88
LP(1)Cl1	LP*(5)Pd1	5.24	5.64
LP(1)Cl1	LP*(6)Pd1	4.16	4.78
LP(1)Cl1	LP*(7)Pd1	5.18	4.30
LP(2)Cl1	LP*(5)Pd1	9.63	9.38
LP(2)Cl1	LP*(6)Pd1	9.83	10.04
LP(3)Cl1	LP*(5)Pd1	1.50	1.21
LP(3)Cl1	LP*(6)Pd1	1.22	0.91
LP(3)Cl1	LP*(8)Pd1	7.17	5.68
Cl1→Pd1		43.93	41.94

Table 5.	Occupancy	and energy of	of the differe	nt NBOs inc	luded in the P	d-O, Pd-Cl,	and Pd-N
interactio	ons.						

	MPW1PW91	WB97XD		
NBO	Occupancy	Energy	Occupancy	Energy
LP(1)O1	1.9428	-0.5755	1.9435	-0.6305
LP(2)O1	1.7117	-0.2829	1.7186	-0.3335
LP(3)O1	1.6309	-0.4595	1.6390	-0.5106
LP(1)N1	1.6532	-0.4398	1.6604	-0.4862
LP(1)N7	1.6303	-0.4641	1.6368	-0.5104
LP(1)Cl1	1.9739	-0.4264	1.9751	-0.4851
LP(2)Cl1	1.9689	-0.7465	1.9703	-0.7951
LP(3)Cl1	1.9663	-0.2911	1.9681	-0.3683
LP*(5)Pd1	0.4662	0.0215	0.4607	0.0584
LP*(6)Pd1	0.1617	0.4265	0.1640	0.4942
LP*(7)Pd1	0.1059	0.2893	0.1075	0.3447
LP*(8)Pd1	0.0644	0.1112	0.0637	0.1653



Fig. 9. Anti-bonding NBOs of Pd(II) contributing in the interactions with the NBOs of the donor ligand groups.

Antiproliferative activity

The anti-cancer activity of the synthesized metal complex [Pd(Triaz)Cl], was further explored against two cancer cell lines (MCF-7 and PC3) followed by MTT assay. Initially, the cytotoxicity of starting ligand (HTriaz) was previously examined and reported which possess very week activity against the MCF-7 breast cancer and HCT-116 colon cancer [21]. The results shown for the antiproliferative activity of the [Pd(Triaz)Cl], 18.23% inhibition against PC3 cell line at 50μ g/mL. On the other hands, exhibited 7.1% inhibition when used 30μ g/mL against the MCF-7 breast cancer. It can be concluded that both ligand and its Pd-(II) complex have week activity against tumor.

Conclusions

A Pd(II) complex, [Pd(Triaz)CI], with a *NNO*-donor chelate derived from hydrazono-*s*triazine ligand was synthesized and its molecular structure investigated using different spectroscopic techniques and single-crystal X-ray diffraction combined with Hirshfeld topology analysis. The latter was performed to describe the supramolecular structure of the complex. The neutral square planar [Pd(Triaz)CI] complex comprised one mono-negative tridentate ligand, Triaz⁻, and one chloride ion, Cl⁻, in the inner sphere. The O…H, C…H, and C…N interactions are the most important ones, controlling the molecular packing of the complex. DFT calculations with the aid of NBO analysis were used to describe the Pd-N, Pd-O and Pd-Cl coordination interactions. Due to the week anti-cancer activity for the metal complex and the ligand further investigation and screen for different cancer lines well be carried out in the future.

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Highlights

- The new Pd(II) complex [Pd(Triaz)Cl] bearing hydrazono-*s*-triazine ligand explored.
- The molecular structure assigned by X-ray single crystal technique.
- Molecular insights were also investigated.

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author statement

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Conflicts of Interest

"The authors declare no conflict of interest."

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