Thermal behavior of volatile palladium(II) complexes with tetradentate Schiff bases containing propylene-diimine bridge

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Abstract

Palladium(II) complexes with tetradentate Schiff bases, Pd(acacpda)(I) and Pd(acacdmpda)(II) (acacpda^{2–}—*N*,*N*⁻(propylene)-bis(acetylacetoneiminato-), acacdmpda^{2–}—*N*,*N*^{'-}(2,2-dimethylpropylene)-bis(acetylacetoneiminato-)), have been synthesized, purified and characterized by elemental analysis, IR and NMR spectroscopy. Complex I was obtained for the first time, and its solid-state structure was investigated by single-crystal X-ray diffraction. It has a discrete molecular structure with the Pd²⁺ ion being in a distorted square-planar coordination environment formed by N and O atoms of the acacpda^{2–} ligand. The thermal properties of complexes I and II in condensed phase were investigated by thermogravimetry. Compound II transfers into the gas phase almost completely, while compound I evaporates with a partial decomposition. The temperature dependences of saturated vapor pressure were measured by the flow method in the temperature intervals (423–457) and (422–478) K for I and II, respectively. Both complexes were shown to exhibit an excellent stability upon sublimation. The thermodynamic parameters of sublimation processes were calculated to be: $\Delta_{sub}H_{440} = (133 \pm 4)$ kJ mol⁻¹, $\Delta_{sub}S_{440}^{0} = (210 \pm 9)$ J (mol K)⁻¹ for I, $\Delta_{sub}H_{450} = (135 \pm 1)$ kJ mol⁻¹, $\Delta_{sub}S_{450}^{0} = (218 \pm 3)$ J (mol K)⁻¹ for II. It has been found that the complex II with a more bulky Schiff base ligand possesses slightly higher vapor pressure values as compared to I, which is probably due to the lower contribution of weak intermolecular interactions occurring in the crystal structure of II.

Keywords Palladium(II) complexes · Schiff bases · MOCVD precursors · Crystal structure · Vapor pressure

Introduction

Nowadays, bimetallic films and nanoparticles based on palladium are widely used in different fields of application, for example, as materials for magnetooptical recording devices [1, 2], biosensors for methotrexate detection [3], water purification and decolorization systems [4, 5], hydrogen purification membranes [6–8], hydrogen sensors

[9, 10] and in catalysis of a number of chemical reactions such as oxidation, reduction, hydrogenation, cross-coupling and others [10-15].

The perspective techniques to obtain such bimetallic structures are metal organic chemical vapor deposition (MOCVD) and related methods such as atomic layer deposition (ALD), etc. The main advantages of these methods are complete preside control over the properties (composition and microstructure/size) of the produced films and nanoparticles by means of variation a number of deposition parameters as well as the possibility of formation of target structures on the complicated shape objects and the absence of side components (for example, protecting agents in the case of the colloidal synthesis methods). The investigation of the formation of Pd-based bimetallic systems by chemical vapor-phase deposition techniques has been developing ever more actively since 1990s: various structures have been obtained with a number of metals including silver [16, 17], platinum [18–21],



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ruthenium [21], gold [22], copper [23] and nickel [24–26]. However, it should be noted that in order to realize all the advantages of the MOCVD and related methods, the volatile precursors of both components apart from the standard requirements for thermal properties (sufficient volatility, thermal stability and others [27]) should possess their experimental compatibility. Thus, in order to ensure further progress in the area discussed, it is necessary to expand a number of thermochemically characterized volatile palladium complexes to select the optimal combinations of the precursors for effective deposition of each bimetallic system.

To date, palladium complexes with the following ligands have been proposed as volatile MOCVD/ALD precursors: β -diketonates, β -iminoketonates, β -enaminodiones, dialkyldithiocarbamates, dialkyldithiophosphates, cyclopentadienyls, allyls, alkenols, etc. [28-36]. This work is devoted to a new class of palladium MOCVD precursors-Pd(II) complexes with tetradentate Schiff bases (Fig. 1). Preliminary research has shown that the replacement of the ethylene diimine bridge in the ligand with propylene one increases the volatility of the formed complexes [37]. So, the objects of the present investigation are palladium complexes with propylene-diimine bridge containing Schiff bases: Pd(acacpda) I and Pd(acacdmpda) II (Fig. 1, $acacpda^{2}$ —N,N'-(propylene)-bis(acetylaceacacdmpda^{2–}—N,N'-(2,2-dimethylpropytoneiminato-), lene)-bis(acetylacetoneiminato-)). The paper presents the synthesis, detailed characterization and thermochemical investigation of the complexes including the crystal structure of compound I obtained for the first time.



Fig. 1 General structure formula of palladium(II) complexes with tetradentate Schiff bases studied

Experimental

Materials and methods

The following reagents were used in the presented work: palladium chloride (Krastsvetmet, palladium content 59.845%), acetylacetone (Acros Organics, purity > 99%), propylenediamine (Merck, purity > 99%), 2,2-dimethyl-propylenediamine (Aldrich, purity > 99%). Acetonitrile (Kriokhrom, purity > 99%), hexane ("Reaktiv", purity > 98%), toluene ("Reaktiv", purity > 98%) were used as a solvents. All of the reactants were used as supplied.

The elemental analysis was carried out on a CHNSanalyzer vario MICRO cube with the accuracy 0.5%. IR spectra were recorded on a Scimitar FTS 2000 spectrometer in the 400–4000 cm⁻¹ range; the samples were prepared by pressing the pellets with KBr. NMR spectra were recorded on an Avance 500 spectrometer at 500 MHz for ¹H NMR and 125 MHz for ¹³C NMR. The primary spectral data are presented in Supplementary Materials. The melting points (*m.p.*) were determined by the visual method using the Kofler bench.

Synthesis of Schiff bases and palladium complexes

*H*₂*acacpda* was synthesized according to the method [38] through the interaction of 2.15 mL (21.0 mmol) acetylacetone and 0.90 mL (10.5 mmol) propylenediamine by means of stirring under reflux in the absence of a solvent. The product was purified by recrystallization from hexane, yield 80%. Anal. Calcd. for $C_{13}H_{22}N_2O_2$ (mass. %): C 65.5, H 9.3, N 11.8. Found C 65.2, H 9.2, N 11.7. IR spectrum (cm⁻¹): 726, 751, 1026, 1088, 1138, 1282, 1353, 1440, 1514, 1568, 1607, 2875, 2953, 3547.

 $H_{2}acacdmpda$ was synthesized and purified similarly to H₂acacpda from 1.95 mL (18.8 mmol) of Hacac and 1.20 mL (9.41 mmol) of 2,2-dimethylpropylenediamine. Yield 80%. Anal. Calcd. for C₁₅H₂₆N₂O₂: C 67.6, H 9.8, N 10.5. Found: C 67.7, H 9.7, N 10.7. IR spectrum (cm⁻¹): 751, 807, 975, 1023, 1104, 1197, 1290, 1359, 1442, 1521, 1572, 1611, 2880, 2970, 3388.

Pd(acacpda)(I) was obtained according to modified procedure [39]. Firstly, $Pd(CH_3CN)_2Cl_2$ complex was prepared by dissolving of 0.518 g (2.92 mmol) $PdCl_2$ in 70 mL of acetonitrile at slight heating and then the 20 mL of acetonitrile solution of H₂acacpda (0.696 g, 2.92 mmol) was added to reaction mixture which was further kept for 30 min. Then, the NaOH aqueous solution (0.234 g, 5.85 mmol) was added. The product was separated as a precipitate after 30 min of stirring at room temperature by adding of 500 mL of water. An additional amount of the product was obtained by toluene extraction of the filtrate. The complex was purified by vacuum sublimation (443 K, 6.67 Pa). Yield 62% (0.615 g). *m. p.* 468 K. Anal. Calcd. for PdC₁₃H₂₀N₂O₂(**I**): C 45.6, H 5.9, N 8.2. Found: C 45.4, H 6.0, N 8.1. IR spectrum (cm⁻¹): 423, 460, 758, 952, 1015, 1119, 1252, 1274, 1359, 1405, 1466, 1512, 1577, 2913, 2938. ¹H NMR (CDCl₃, 7.26 ppm): 1.9564 (s, 6H, CH₃C=N), 1.9767 (s, 6H, CH₃C=O), 2.0532 (m, 2H, (CH₂)CH₂(CH₂)), 3.0389 (t, 4H, (CH₂)CH₂(CH₂)), 4.8548 (s, 2H, CH). ¹³C NMR (CDCl₃, 77.2 ppm): 23.0504 (<u>CH₃C=N</u>), 24.5162 (<u>CH₃C=O</u>), 32.4997 (CH₂)<u>CH₂(CH₂), 51.2816 (<u>C</u>H), 99.4074 (<u>CH₂-N</u>), 164.3894 (<u>C</u>=N), 176.7554 (C=O).</u>

Pd(acacdmpda)(II) was synthesized in a similar way to I from 0.333 g (1.88 mmol) of PdCl₂, 0.500 g (1.88 mmol) of H₂acacdmpda, and 0.180 g (4.50 mmol) of NaOH. The complex was purified by vacuum sublimation ((433-443) K, 6.67 Pa). Yield 88% (0.543 g). m. p. 487 K. Anal. Calcd. for PdC₁₅H₂₄N₂O₂(**II**): C 48.4, H 6.5, N 7.6. Found: C 48.0, H 6.5, N 7.6. IR spectrum (cm⁻¹): 449, 469, 756, 952, 1013, 1112, 1232, 1273, 1332, 1365, 1398, 1456, 1511, 1571, 2916, 2956. ¹H NMR (CDCl₃, 7.26 ppm): 1.1141 (s, 6H, C(CH₃)₂), 1.9522 (s, 6H, CH₃C=N), 1.9899 (s, 6H, CH₃C=O), 3.0442 (s, 4H, CH₂C-N), 4.8332 (s, 2H, CH). ¹³C NMR (CDCl₃, 77.2 ppm): 22.4577 (CH₃C=N), 24.7277 (CH_3)₂C(CH_2)₂, 25.9650 (CH_3 C=O), 36.5668 (CH₃)₂C(CH₂)₂, 61.1032 (CH), 99.6479 (CH₂–N), 163.3507 (C=N), 176.6955 (C=O).

Single-crystal X-ray diffraction

A yellow needle crystal of I was selected under a microscope and then mounted to the tip of the thin glass fiber with epoxy resin. X-ray intensity data were collected on a Bruker-Nonius X8 Apex CCD diffractometer at 296(2) K graphite monochromatized MoKa radiation using $(\lambda = 0.71,073 \text{ Å})$. The standard technique was used (combined φ and ω scans of narrow frames). Data reduction and multi-scan absorption were carried out using the SADABS [40]. The structure was solved by direct methods and refined by full-matrix least-squares on F^2 using the SHELXTL software package [40]. All non-hydrogen atoms were refined anisotropically. Hydrogen atoms of acacpda²⁻ ligand were placed in geometrically calculated positions and refined as riding on their parent carbon atoms with $U_{iso}(H) = 1.5U_{eq}(C)$ for methyl hydrogen atoms and $U_{\rm iso}({\rm H}) = 1.2 U_{\rm eq}({\rm C})$ for methylene and β -hydrogen atoms. The molecular graphics were performed using DIAMOND program [41]. Crystallographic data and selected refinement details are given in Table 1. The main bond lengths and angles are presented in Table 2. The atomic positional and thermal parameters, full lists of bond lengths and angles were deposited at the Cambridge Crystallographic Data Centre under the reference number CCDC 1576778 and are available free of charge via www.ccdc.cam.ac.uk/data_request/cif.

Thermal investigation

The thermogravitmetric investigation (TG/DTA) was performed on a Netzsch TG 209 F1 Iris thermal analyzer in an open Al₂O₃ crucible in a helium flow (30.0 mL min⁻¹). The heating rate was 10 K min⁻¹; sample mass was (10 \pm 1) mg.

The temperature dependencies of saturated vapor pressure of the complexes were measured by flow (transpiration) method in the atmosphere of dry inert gas carrier (helium) using the equipment presented in Fig. 2. The principles of the method and the experimental procedure used are described in [42, 43] and [44], respectively. During the experiment, the substance in source 1 is transferred to the gas phase by heating and carried by the carrier gas flow through the porous plugs 5 to the receiver 2 where it condenses on the cold walls. The amounts of substance in source and receiver were determined gravimetrically (uncertainty is 5×10^{-4} g). The sublimation is performed by a calibrated heater 6 with a thermocouple sensor 7. The temperature was measured by thermal control device 8 with uncertainty 0.5 K, the carrier gas flow was measured by flow mass (11, 12), and helium flow rate was measured by device 13 with uncertainty 2%. The total relative standard uncertainty of this method was not above 5%. The measurements were taken in the guasi-equilibrium conditions, which were proved by the independence of the vapor pressure on the helium flow rate. The partial pressure of saturated vapor over complexes was calculated according to Eq. 1:

$$P = P_{\text{total}} \frac{n}{n+N} \tag{1}$$

 P_{total} total pressure in a system; *n* number of moles of carried compound; *N* number of moles of helium. The calculations were based on the assumption that the substance vaporizes in the monomolecular form. The experimental results were processed statistically using the objective function recommended in [45]. Data obtained are presented in Table 1S (Supplementary Materials).

Results and discussion

Synthesis and characterization

Palladium complexes I and II were synthesized by the interaction of $Pd(CH_3CN)_2Cl_2$ prepared in situ by dissolving of $PdCl_2$ in CH_3CN under reflux with

 Table 1
 Crystallographic data,

 data collection and refinement
 parameters for I

Empirical formula	$C_{13}H_{20}N_2O_2Pd$
Formula weight	342.71
Temperature/K	296(2)
Crystal size/mm ³	$0.37 \times 0.05 \times 0.05$
Crystal system	Orthorhombic
Space group	$P2_{1}2_{1}2_{1}$
Ζ	4
a/Å	6.39800(10)
b/Å	12.9005(4)
c/Å	16.9119(6)
V/Å ³	1395.86(7)
$D_{\rm calcd}/{\rm g}~{\rm cm}^{-3}$	1.631
μ (Mo K α)/mm ⁻¹	1.326
<i>F</i> (000)	696
θ -range for data collection/°	1.99–27.74
Ranges of h, k, l	$-7 \le h \le 5; -10 \le k \le 16; -22 \le l \le 22$
Reflections collected	7235
Independent reflections	$3164 \ (R_{\rm int} = 0.0211)$
Observed reflections $[I > 2\sigma(I)]$	3015
Parameters refined	168
$R [F^2 > 2\sigma(F^2)]$	$R_1 = 0.0425; \ wR_2 = 0.0981$
$R(F^2)$ (all data)	$R_1 = 0.0444, wR_2 = 0.0999$
Goodness of fit on F^2	1.055

Table 2 Selected bond lengths(Å), bond angles (°) anddihedral angles (°) for I

Bond lengths		
Pd(1)–O(1)		1.994(3)
Pd(1)-O(2)		1.996(3)
Pd(1)-N(1)		2.023(3)
Pd(1)–N(2)		2.023(3)
Bond angles		
O(1)-Pd(1)-O(2)		79.73(13)
O(1)-Pd(1)-N(1)		91.98(14)
O(1)-Pd(1)-N(2)		171.59(14)
O(2)-Pd(1)-N(1)		171.52(14)
O(2)-Pd(1)-N(2)		92.13(13)
N(1)-Pd(1)-N(2)		96.08(14)
Atoms in plane		Angle
Dihedral angles		
O(1)Pd(1)N(1)	O(1)C(1)C(3)C(4)N(1)	17.12(12)
O(2)Pd(1)N(2)	O(2)C(12)C(11)C(9)N(2)	20.14(13)
N(1)Pd(1)N(2)	N(1)C(6)C(8)N(2)	17.91(16)
N(1)C(6)C(8)N(2)	C(6)C(7)C(8)	63.0(3)

corresponding Schiff base in acetonitrile and an aqueous solution of alkali (Eq. 2). The products were precipitated by adding a large amount of water. The complexes I and II

are yellow crystalline powders stable on storage in air at reduced temperature for at least 6 months.



Fig. 2 Experimental setup for tensimetric measurements by flow method: 1—source, 2—receiver, 3—analyzed substance in source, 4—condensed substance in receiver, 5—porous plugs, 6—heater, 7—thermocouple sensors, 8—thermal control device, 9—carrier gas cylinder, 10—gas drying unit, 11, 12—gas consumption regulator (flow mass), 13—carrier gas flow rate measuring device

$$PdCl_{2} \xrightarrow[CH_{3}CN]{} Pd(CH_{3}CN)_{2}Cl_{2} \xrightarrow[CH_{3}]{} PdL;$$
(2)

HL-Hacacpda, Hacacdmpda

The comparison of IR spectra of the complexes with IR spectra of corresponded Schiff bases is presented in Table 3. The presence of stretching vibrational bands v(Pd-N), v(Pd-O) in the IR spectra of the complexes as well as the absence of bands v(N-H), v(O-H) indicates the complexation. Shifts of absorption bands v(C=O), v(C=N) toward lower frequencies also confirm the formation of the complexes.

The expected signal positions in the ¹H NMR and ¹³C NMR spectra, as well as the integrated intensities of the ¹H NMR spectra signals of the complexes, also confirm their structure in solution (Fig. 1). The proton signals of the CH₃C=N, CH₃C=O groups are located closely, with a larger shift of CH₃C=O proton group toward the stronger field. The signals of the protons of other aliphatic fragments are easily recognizable and typical. Comparison of the ¹H NMR spectra of complexes I and II with the spectra



Fig. 3 Thermal ellipsoid representation of I with the atomic numbering scheme (ellipsoids are drawn at the 50% probability level)

of corresponding Schiff bases [45] shows the absence of acidic proton signal at about 11 ppm, indicating the complexation.

According to single-crystal X-ray diffraction analysis, the complex I has a molecular structure consisting of discrete Pd(acacpda) molecules (Fig. 3), which can be best compared with the structure of the related complex Pd(acacdmpda), II [37]. All atoms of the complex molecule are crystallographycally independent and occupy general positions. Within the molecule, the Pd^{2+} ion has a distorted square-planar coordination environment formed by imino N and carbonyl O atoms of the tetradentate acacpda²⁻ ligand. The bond lengths and angles around the palladium center are, as expected, close to the corresponding values found in II (Table 2). In particular, the average values for Pd-O and Pd-N bond lengths are 1.995(1) and 2.023(1) Å, respectively, while the chelate N-Pd-O angles are 91.98(14) and 92.13(13)°. The N-Pd-N and O-Pd-O angles are 96.08(14) and 79.73(13)°, respectively. Due to the tetradentate coordination of the $acacpda^{2-}$ ligand, the Pd(acacpda) molecule contains two conjugated six-membered metallocycles (PdOCCCN) and one six-membered metallocycle (PdNCCCN). Both (PdOCCCN) metallocycles are rather nonplanar, with the angles of folding along the O(1)N(1) and O(2)N(2) lines

Table 3 The assignment of absorption bands (cm^{-1}) in IR spectra of the complexes I and II and corresponded Schiff bases

Compound v(Pd–N), v(Pd–O)		$\delta(\text{CCC})$	v(C=C)	v(C=O), v(C=N)	v(С–H)	v(N–H), v(O–H)	
H ₂ acacpda	_	726, 751, 1026	1282, 1353, 1440	1514, 1568, 1607	2875, 2953	3547	
Pd(acacpda)(I)	423, 460	758, 952, 1015	1252, 1274, 1405	1466, 1512, 1577	2913, 2938	_	
H ₂ acacdmpda	-	751, 807, 975, 1023	1290, 1359, 1442	1521, 1572, 1611	2880, 2970	3493	
Pd(acacdmpda)(II)	449, 469	756, 962, 1013	1232, 1273, 1398	1456, 1511, 1571	2916, 2956	-	

Fig. 4 Packing diagram of I viewed along the a axis. H atoms are omitted for clarity. Hydrogen bonds C-H···O are shown as dashed lines



being 17.1 and 20.1°, respectively. The (PdNCCCN) metallocycle contains no conjugated bonds and deviates considerably from planarity; its geometry is characterized by the angles of folding along the C(6)C(8) and N(1)N(2) lines, which are 63.0° and 17.9° , respectively. Therefore, the (PdNCCCN) metallocycle can be best described as having a distorted boat conformation with the palladium atom at the front and the opposing carbon atom at the back.

At a supramolecular level, both complexes **I** and **II** exhibit C–H···O hydrogen bonds, leading to a chain-like arrangement of molecules in the crystals (Fig. 4). Despite the weakness of these interactions, their relative strength can be compared based on the C···O distances. In **I**, the central (α) hydrogen atom of the β -diketone unit participates in the formation of hydrogen bond, showing the C···O distance of 3.32 Å, while in **II**, the methylene hydrogen

atoms make weaker hydrogen bonds with the C...O distances of 3.48 and 3.51 Å. Moreover, the hydrogen-bonded chains of Pd(acacpda) and Pd(acacdmpda) molecules are organized in different ways and, as one can expect, their packing motifs and densities are rather different. In particular, the chains of Pd(acacpda) molecules in I have a zigzag shape and are characterized by the following Pd---Pd distances: 7.89 Å between the adjacent Pd(acacpda) molecules within the chain and 6.40 Å between the Pd(acacpda) molecules from the adjacent chains. In II, the chains of Pd(acacdmpda) molecules have a more linear shape with the corresponding Pd…Pd distances of 6.18 and 8.05 Å [37]. Most likely, the reason for the differences in hydrogen-bonding patterns in I and II lies in the higher bulkiness of acacdmpda²⁻ ligand as compared to that of acacpda²⁻ ligand. The presence of two methyl groups in



Fig. 5 Mass loss curves of complexes Pd(acacpda)(I) and Pd(acacdmpda)(II)



Fig. 6 Temperature dependences of the saturated vapor pressure of complexes Pd(acacpda)(I) and Pd(acacdmpda)(II)

propylene-diimine bridge of acacdmpda^{2–} ligand results in a less dense packing of the Pd(acacdmpda) molecules in the crystal of II.

Thermal properties

Thermal properties of the complexes in condensed phase were investigated by TG/DTA. The endoeffects on the

DTA curves at 467 K and 485 K (onset temperatures) for I and II, respectively, correspond to the melting process, that have been confirmed visually on the Kofler's table. The mass loss curves are presented in Fig. 5. Under the conditions of the TG experiments, the complexes evaporate in the temperature range (498–598) K. Compound II transfers into gas phase almost quantitatively (mass loss 98.0%), while compound I evaporates with a partial decomposition (mass loss 87.3%). Temperatures of 50% mass loss ($T_{50\%}$) are close (576 and 573 K for I and II, respectively); consequently, the complexes appear to be comparable in volatility.

In quasi-equilibrium conditions of tensimetric experiments, in the temperature intervals (423-457) K and (422–478) K for I and II, respectively, the difference in volatility of the complexes was more accurately revealed. The mass of substance in receiver was equal to the mass of evaporated one in the all investigated temperature range, indicating the stability of the complexes during sublimation processes. The results obtained are presented in Fig. 6 and Table 1S (Supplementary Materials). The thermodynamic parameters of sublimation and logarithmic temperature dependences of the saturated vapor pressure are listed in Table 4. It should be noted that the differences in experimental pressure values from the calculated ones (Table 1S) are not exceed standard uncertainties in temperature and pressure measurements, while the random character of these deviations indicates the absence of a systematic error in the measurements.

The data obtained allow us to conclude that the insertion of methyl substituents in the diimine bridge slightly increases the volatility of the complexes: vapor pressure values of complexes I and II at 433 K are 0.82 and 1.35 Pa, respectively. It should be noted that the same regularities were observed for analogous copper(II) complexes by means of TG method [46]. The slight difference in volatility observed for the complexes under study could be associated with the weak intermolecular interactions revealed in their crystal packing. In fact, the structure of less volatile complex I is characterized by stronger C– H…O hydrogen bonds described above. Moreover, the packing of complex I is more dense as it could be also concluded from the higher calculated density of this compound (1.631 g cm⁻¹) in comparison with that of complex

Table 4The thermodynamicparameters of sublimation andtemperature dependences ofsaturated vapor pressure ofcomplexes I and II

Compound	$\Delta T/K$	Ν	$\ln(p/p_0) = A - B/T$		$\Delta_{\rm sub}H_{\rm T*}/{\rm kJ}~{\rm mol}^{-1}$	$\Delta_{sub}S^0_{T^*}/J \ (mol \ K^{-1})$
			A	В		
Pd(acacpda)(I)	423–457	11	25.34	15,994	133 ± 4	210 ± 9
Pd(acacdmpda)(II)	422–478	13	26.23	16,238	135 ± 1	218 ± 3

 ΔT temperature interval; T^* middle temperature of the interval investigated; N number of experimental points; $p_0 = 1$ atm = 760 Torr = 101,325 Pa

II (1.526 g cm⁻¹), whereas the molecular weight of complex I is lower, and the number of molecules in the unit cell is the same. The loosening of the crystalline package due to methyl substituents introduced into the diimine bridge of the Schiff base ligand could also be one of the factors that lead to an increase in compound volatility.

At the same time, palladium chelates I and II are characterized by the lower volatility in comparison with palladium complexes with bidentate Schiff bases (β -iminoketonates). For example, the saturated vapor pressure of the simplest representative of palladium β -iminoketonates with alkyl substituent at the nitrogen atom, Pd(Mei-acac)₂ (Mei-acac⁻—4-methylamino-3-penten-2-onato-), is almost 30 times higher (34.85 Pa at 433 K) [32]. The same regularity seems to be typical for complexes of 3d-transition metals with coordination polyhedrons close to the squareplanar namely Cu(II), Co(II), Ni(II) [46-51]. However, prior to the present work, the quantitative tensimetric data were available only for nickel complexes (saturated vapor pressure of Ni(acacen) at 433 K is 0.83 Pa, Ni(i-acac)2-32.1 Pa (acacen^{2–}-N,N'-(ethylene)-bis(acetylacetoneiminato-), *i*-acac⁻-4-amino-3-penten-2-onato-) [50, 51], whereas for the other complexes it was shown only qualitatively (by TG experiments).

Nevertheless, it is worth noting that the vapor pressure values and stability of complexes I and II during sublimation make it possible to use these precursors in the typical MOCVD reactors. In this aspect, the tensimetric data obtained are useful to select the temperature parameters of the evaporator (precursor vapor source) in MOCVD experiments to control the partial pressure of palladium precursor during deposition of bimetallic nanostructures.

Conclusions

Volatile palladium(II) complexes Pd(acacpda)(I) and Pd(acacdmpda)(II) with tetradentate Schiff bases containing propylene-diimine bridge were synthesized and investigated as potential MOCVD precursors. The compounds obtained were fully characterized by elemental analysis, IR and NMR spectroscopy, and solid-state structure of complex I obtained for the first time was investigated by single-crystal X-ray diffraction. The influence of the ligand structure on the complexes thermal properties (volatility and stability) were studied using thermogravimetric analysis and tensimetry (flow method). Temperature dependences of saturated vapor pressure have been measured in the temperature intervals (423–457) K and (422–478) K for I and II, respectively, and thermodynamic parameters of sublimation were calculated.

It has been shown that the introducing methyl groups to the central atom of diimine bridge in Schiff base ligand (complex II vs. complex I) does not lead to significant changes in the palladium coordination polyhedron being a distorted square-planar formed by imino N and carbonyl O atoms. However, the modification of the ligand under consideration results in noticeable changes in the crystal packing, which is reflected in the thermal properties of the complexes. In particular, complex II is characterized by slightly higher volatility than I (the difference in vapor pressure values is about 0.2 order of magnitude). This observation could be explained in terms of extenuation of weak hydrogen intermolecular interactions in the crystal structure of II with a more bulky Schiff base ligand and its lower packing density.

Although the evaporation of complex I is accompanied by a partial decomposition, the sublimation processes for both complexes have been shown to proceed purely. Consequently, the thermal stability of the complexes I and II as well as the vapor pressure values are sufficient for the use in MOCVD processes.

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