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A redox-active ligand combines a PCP pincer site with a bidentate N–N donor in opposition†

Derek W. Leong,‡ Yanwu Shao,‡ Nattamai Bhuvanesh and Oleg V. Ozerov *

A binucleating ligand (**2**) combining a monoanionic PCP pincer cleft with a monoanionic N–N cleft has been prepared on the basis of bis(imidazolyl)methane (**1**). Installation of divalent Pd or Pt into the PCP cleft proceeded smoothly, with the formation of square planar (PCP)MCl complexes (M = Pd or Pt). The “C” in the PCP is the central carbon of the β -diketimate-like N–N cleft. The N–N cleft in these complexes was obtained in three levels of protonation. In **4-Pd** and **4-Pt**, the N–N cleft carries one proton and is neutral; in **3-Pd** or **3-Pt**, it is additionally protonated by HCl, and in **6-Pd** or **6-Pt**, the proton has been removed by the $(\text{Me}_3\text{Si})_2\text{NLi}$ base and replaced by Li. The chloride in **4-Pd/Pt** was replaced with an iodide to make **6-Pd/Pt** in a metathesis with Me_3SiI . Compounds **4-Pd/Pt** and **6-Pd/Pt** exhibited tautomerism reminiscent of acetylacetone and β -diketimines, possessing either an NH or a CH bond. Compounds **6-Pd/Pt** possess a lithiated N–N cleft and serve as convenient transmetallation agents. Their reactions with $\text{BF}_3(\text{OEt}_2)$, $\frac{1}{2}\text{ZnCl}_2$, or TaCl_5 produced new compounds **7-Pd/Pt**, **8-Pd/Pt**, and **9-Pd/Pt** that carry a BF_2 , $\frac{1}{2}\text{Zn}$, or TaCl_4 units in the N–N cleft. Compounds **8-Pd/Pt** are trimetallic with a central Zn coordinated by two N–N clefts each of which carries a (PCP)MCl on the opposite side. Structures of **3-Pd**, **7-Pt**, **8-Pd**, and **9-Pd** in the solid state were established by X-ray diffractometry. They demonstrate the remarkable planarity of the extended conjugated organic π -system into which the N–N and the PCP binding sites are incorporated. Electrochemical studies on **7–9** established that each of these extended π -systems can be oxidized twice quasi-reversibly. Compounds **9-Pd/Pt** could be additionally reduced quasi-reversibly by a single electron, which was ascribed to the reduction of the Ta^{V} center to Ta^{IV} . They also differed from **7** and **8** in possessing apparent lower-energy LMCT bands responsible for the blue or purple colors.

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10th anniversary statement

I (Oleg Ozerov) was among the first crop of associate editors for ICF, serving during 2013–2022. I recall that at one of our first Board meetings taking place in Beijing, there was a spirited discussion regarding the place and the future for ICF in the chemistry publishing world. We optimistically resolved to make ICF the best inorganic chemistry journal because any other outlook seemed lacking. Now, ten years later, that resolve seems well justified and I am glad I got to play a modest role in the journal's rise. Being an ICF editor allowed me to expand my scientific horizons, and to learn to face the challenges of this different facet of academic life. But it also allowed me to make a number of new friends, and to experience new places in China and Asia in general. For that I am grateful and wish ICF a continued path of success.

Introduction

Pincer ligands have become ubiquitous in the studies of organometallic structure and reactivity.^{1–3} Pincer-supported

transition metal complexes have been studied as catalysts for a large variety of applications,^{1–3} including alkane dehydrogenation,^{4,5} transfer (de)hydrogenation of compounds with C=N/C–N and C=O/C–O bonds,^{6–8} C–C coupling,^{9,10} C–H borylation,^{11,12} and nucleophilic addition to ketones and aldehydes.¹³ The tridentate pincer cleft provides a robust environment for the transition metal, in some cases enabling catalytic performance at a temperature of 200 °C or higher,¹⁴ which is unusual for organometallic transformations. The most obvious way to influence the transition metal center in a pincer environment is through the modification of the pincer donor sites. However, substantial attention has been paid to

Department of Chemistry, Texas A&M University, 3255 TAMU, College Station, TX 77842, USA. E-mail: ozerov@chem.tamu.edu

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‡These authors contributed equally.



the more remote influences, including the use of redox non-innocent ligands^{15–20} and the provision of distal binding sites for the coordination of additional metals.^{21,22} The so-called Janus bis-pincer systems (A–D, Fig. 1) can be included within this context, with one pincer-metal unit providing a remote influence on the other.^{23–30} The Janus bis-pincers provide two opposing binding cleft sites of the same kind, naturally leading to symmetric bimetallic complexes. There also exist examples of non-Janus bis(pincer) ligands with a shared atom or unit linking the two pincer domains.^{31–34} However, it is synthetically difficult to prepare heterobimetallic complexes with symmetric bis(pincer) ligands. Considering ways to introduce two very different metals into the same molecule, we became interested in preparing a “dissymmetric Janus” type of ligand, with one phosphine-anchored pincer site suitable for a late transition metal, and another N–N bidentate site amenable to binding a broad variety of metals and metalloids. This arrangement provides a way to influence the structure and reactivity of the pincer ligated transition metal center through variation of the other element in the N–N cleft, but without changing the immediate coordination environment about the transition metal. We envisaged an N–N site similar to β -diketiminates (also known as “nacnac” ligands)³⁵ that is integrated into a redox-active pincer framework. In considering this design, we were influenced by the binucleating “nindigo” ligands (E,

Fig. 1) pioneered by the Hicks group,³⁶ and also explored by Caulton and Mindiola,³⁷ and by Kaim and Lahiri with co-workers.³⁸ The “nindigo” ligands are essentially a redox non-innocent Janus bis-“nacnac” system. Herein we report on the chemistry of a binucleating ligand fusing a PCP pincer with a “nacnac”-like N–N site. A similar binucleating ligand combining an NCN pincer with a bidentate N–N site (F, Fig. 1) was recently explored by Ong, Frenking, Zhao, and coworkers.³⁹ In this work, we report our exploration of how the nature of the element in the N–N cleft influences the coordination environment about the transition metal in the pincer cleft, along with the electrochemical and spectroscopic properties of the molecule as a whole. This synthetic solution for the remote modulation of the properties of a transition metal pincer complex may provide new way for exploring and tuning catalytic activity in the future.

Results and discussion

Synthesis of the Janus ligand

Kamble *et al.* recently reported a convenient synthesis of benzimidazoles by a thermal or microwave reaction of acid amides with diaminobenzenes in aqueous hydrochloric acid.⁴⁰ Substituting hydrochloric acid with 85% H₃PO₄ allowed us to carry out an analogous synthesis of bis(3-benzimidazolyl) methane (**1**) at a higher temperature without a need for a microwave reactor. We were able to isolate **1** in near-quantitative yield. Treatment of **1** with 2.4 equiv. of ClPrP₂¹ and 2.4 equiv. of Et₃N in THF overnight at ambient temperature resulted in the formation of the proto-ligand **2**, (Scheme 1) which was isolated in excellent yield upon workup. Compound **2** was characterized by ¹H, ¹³C, and ³¹P NMR spectroscopy, with a singlet resonance at 68.2 ppm in the ³¹P{¹H} NMR spectrum. The protons of the methylene linkage give rise to a singlet at δ 5.21 ppm in the ¹H NMR spectrum and a triplet ($J_{C-P} = 17$ Hz) at δ 31.8 ppm in the ¹³C{¹H} spectrum. The PCP cleft of compound **2** can be seen as a member of a class of proto-pincers with a central CH₂ moiety that connects (hetero) arene rings bearing the two side phosphine arms.^{41–47} On the other hand, the N–N cleft of **2** is related to a number of mono-anionic N–N bidentate ligands.⁴⁸

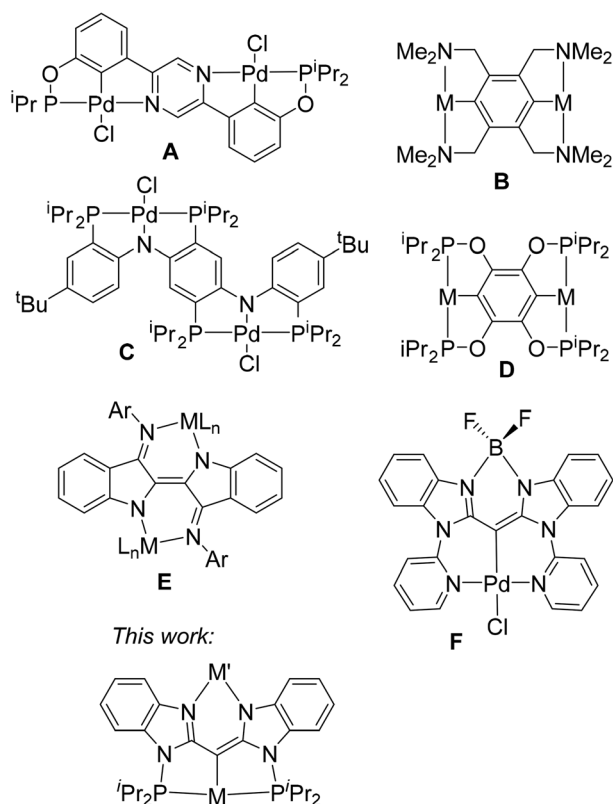
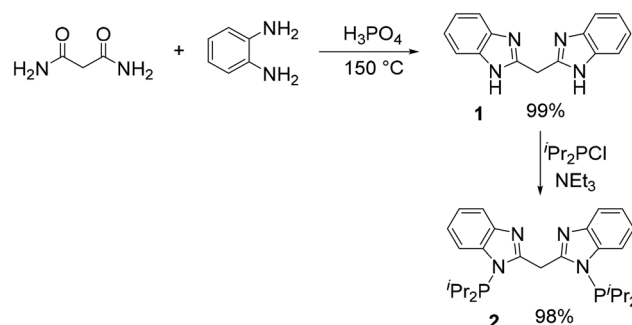


Fig. 1 Literature examples of Janus binucleating pincer (A–D) and “nacnac” (E) ligands, Ong’s compound F, and the target of investigation in this work.



Scheme 1 Synthesis of binucleating ligand **2**.



Synthesis and spectroscopic characterization of bi- and trimetallic compounds

The metalation of the PCP pincer can be carried out with or without the addition of external base, such as Et₃N. In the absence of added Et₃N, compounds **3-Pd** and **3-Pt** were isolated from reactions with (COD)MCl₂ (M = Pd, Pt). The synthesis of **3-Pd** required a rather high temperature (180 °C in *o*-C₆H₄Cl₂) to ensure high conversion, but the Pt reaction proceeded well at 120 °C. The use of Et₃N during the metalation with (COD)PdCl₂ resulted in the formation of **4-Pd** after 12 h at 120 °C in toluene; however, in the case of Pt, use of NEt₃ resulted in decomposition to multiple unidentified products. Compound **4-Pt** was accessed by deprotonation of isolated **3-Pt** with NEt₃ (Scheme 2). Treatment of **4-Pd** with Me₃SiI in toluene resulted in halogen exchange, allowing for the isolation of **5-Pd** in 82% yield after workup.

Complexes **4-Pd/4-Pt/5-Pd** exist as an equilibrium mixture of two tautomers, in which the proton is located either on the nitrogens of the N–N cleft (**4a-Pd/4a-Pt/5a-Pd**), or on the central carbon (**4b-Pd/4b-Pt/5b-Pd**). The different locations of H are evident from the ¹H and ¹³C NMR spectra. For example, in **4a-Pd**, this N–H proton resonates at δ 10.23 ppm (C₆D₆ solvent) as a broad singlet, while the C–H of **4b-Pd** resonates at δ 5.74 ppm as a sharper singlet. The Pd-bound carbons of the two isomers also have distinct chemical shifts (δ 68.0 ppm, **4a-Pd**; δ 48.9 ppm, **4b-Pd**, CDCl₃ solvent). In **4b-Pt**, the C–H resonance (δ 5.51 in CDCl₃) displays telltale ¹⁹⁵Pt satellites (²J_{H–Pt} = 155 Hz). The two tautomers also possess ³¹P NMR chemical shifts that are different by ca. 15–20 ppm in each pair, with the more upfield resonance belonging to the C–H tautomers (**4b-Pd/4b-Pt/5b-Pd**). The observed equilibrium ratios are solvent-dependent, with chloroform favoring the C–H tautomer to a greater degree than benzene or toluene. Compounds **4-Pt** and **5-Pd** displayed a slightly greater predilection towards the NH tautomer (**4a** or **5a**) than did **4-Pd**, in both arene solvents and CDCl₃. It should be noted that we saw no evidence for tautomerism in compounds **2** or **3**.

Deprotonation of **4a/b-Pd** or **4a/b-Pt** with (Me₃Si)₂NLi led to the formation of the Li–Pd/Pt compounds **6-Pd/6-Pt**. Both **6-Pd/Pt** were isolated in high yields as bis-THF adducts, with presumed THF binding to Li. The N–N cleft of the Janus ligand in **3** can be viewed as accepting both protons of the CH₂ group in **2** upon its metalation (and one of the chlorides from PdCl₂ or PtCl₂). Compounds **4** and **6** can be viewed as products of successive dehydrochlorination and then deprotonation of **3**. In other words, **3**, **4**, and **6** represent three different protonation states with 2, 1, or 0 protons from the original CH₂ group retained. The presence of chloride in the isolated **6-Pd** and **6-Pt** was confirmed by observing formation of one equiv. of Me₃SiCl when each was treated with two equiv. of Me₃SiI.

The Li compounds **6** served as convenient precursors for the synthesis of bi- and trimetallic compounds **7**, **8**, and **9** in decent yields *via* reactions with BF₃·OEt₂, ZnCl₂ (0.5 equiv.), or TaCl₅, respectively. Compounds **7–9** show the maximum possible symmetry (C_{2v} for **7** and **9**, and for each pincer fragment

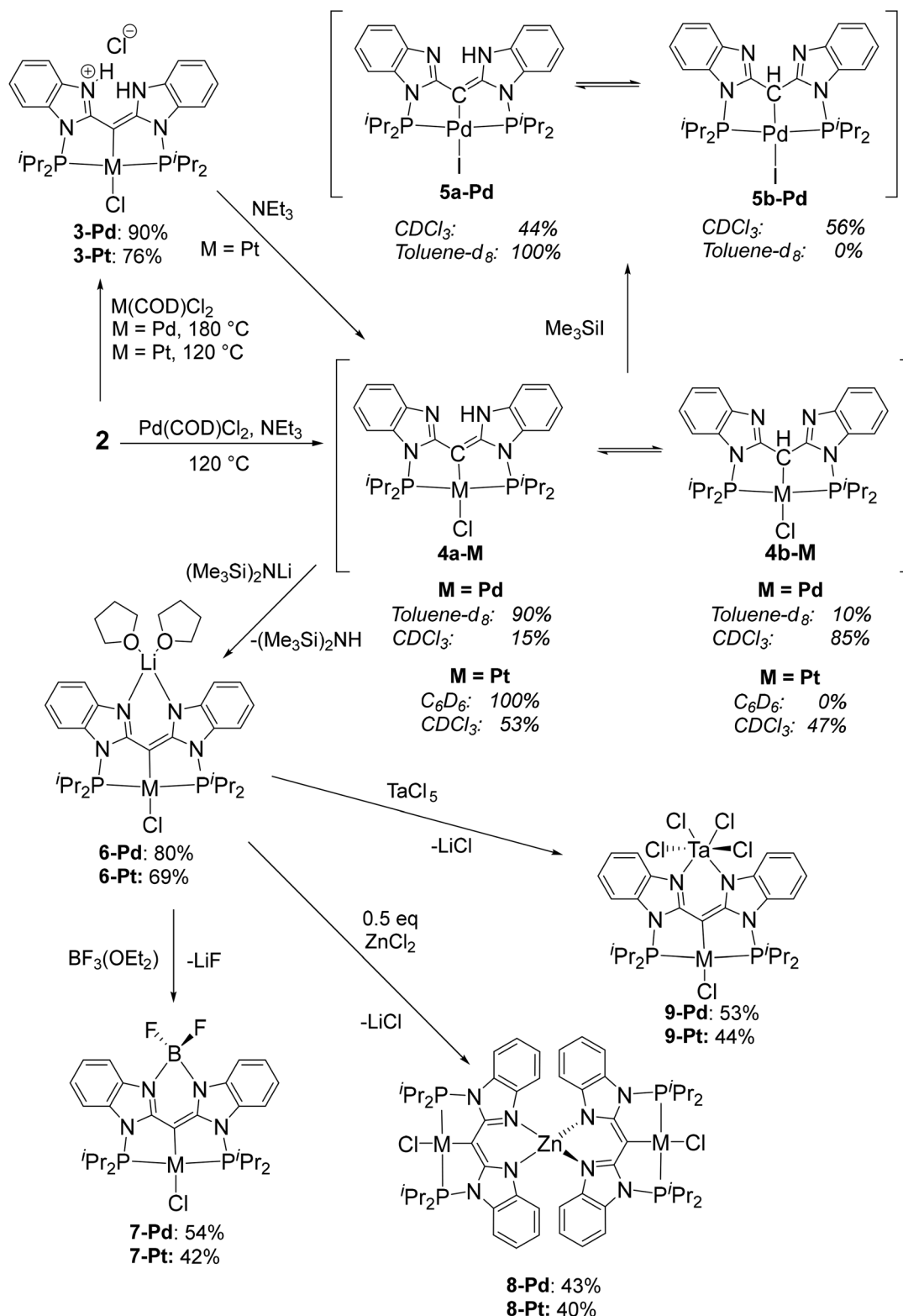
in **8** locally; D_{2d} for **8** as a whole) in their NMR spectra at ambient temperature. The ¹H and ¹³C NMR resonances for the PCP/NN ligand backbone in these compounds are quite similar (Tables S1 and S2†). Interestingly, the ³¹P NMR chemical shifts and the ³¹P–¹⁹⁵Pt coupling constant vary considerably in the **3–9** series, Table 1. These differences may be related to the perturbation of the P–Pd–P/P–Pt–P angles and the Pd/Pt–P distances as a function of the size of the moiety in the N–N cleft, as well as the different electronic effects of those moieties. The ¹¹B NMR chemical shifts in **7-Pd** and **7-Pt** of ca. 3 ppm, the ¹¹B–¹⁹F coupling constants of 29–30 Hz, and the ¹⁹F NMR chemical shift of ca. –138 ppm are similar to those observed in BODIPY and the related compounds with a BF₂ unit in an amido/enamine cleft.^{39,49–51}

Compounds **3–8** are yellow, orange, or red, while **9-Pd** is purple, and **9-Pt** is blue. The properties of **4** and of **7–9** were further studied by UV-Vis spectroscopy. All of these compounds possess strong absorption features in the 300–450 nm range. We surmise that they correspond to the π–π* transitions in the extended organic π-system of the ligand. The UV-Vis spectra of **4** were collected in toluene where **4a-Pd** and **4a-Pt** are the dominant species in solution that contain an extended organic π-system that is not present in **4b-Pd** and **4b-Pt**. It appears that a switch from Pd to Pt incurs a bathochromic shift for analogous compounds by about 15–20 nm. For the same metal in the PCP cleft, the order of increasing absorption energy for these features is **8** < **4** < **7** < **9**, which can be correlated with the increasing electron-withdrawing power in the series of $\frac{1}{2}\text{Zn}^{2+} < \text{H}^+ < \text{BF}_2^+ < \text{TaCl}_4^+$. In addition to these π–π* transition features, distinct lower energy broad bands were also observed for **9-Pd** (λ = 581 nm) and **9-Pt** (λ = 602 nm). These are presumably responsible for the blue or purple colors of these compounds. We tentatively ascribe LMCT character to them, for the transition from the filled π-orbital of the ligand to an empty d-orbital at the d⁰ Ta center.

Structural characterization

The structures of **3-Pd**, **7-Pt**, **8-Pd**, and **9-Pd** were determined by single-crystal X-ray diffractometry (Fig. 2 and Table 2). In all compounds, the geometry about Pd or Pt is approximately square-planar, and that plane is also approximately the plane of the flat, conjugated organic framework. The H₂Cl unit in **3-Pd** and the boron atom in **7-Pt** are also positioned in that plane. The structure of **3-Pd** in the N–N cleft is that of two N–H bonds, with these protons hydrogen-bonded to the chloride. The coordination geometry about boron in **7-Pt** closely mimics BODIPY and other related compounds where a monoanionic, bidentate N–N ligand is bound to the BF₂ unit.^{49,51} The Zn atom in **8-Pd** is approximately in the plane with one “half” of the ligand framework, but is displaced by ca. 0.65 Å from the other half (shown in Fig. 2C). It is likely a consequence of the need to minimize the considerable crowding brought upon by the two metallaligands binding to the same Zn atom. This also manifests itself in the twisting distortion of one the conjugated “halves”, in that the two “halves” are not strictly perpendicular to each other, and ultimately in the dissymmetry of the





Scheme 2 Synthesis of mono-metallic complexes at the PCP-pincer cleft with solvent-dependent tautomer ratios and the synthesis of bi- and tri-metallic complexes.

distorted tetrahedral environment about Zn. The coordination environment about Ta in **9-Pd** is quite close to octahedral, but Ta is displaced from the ligand plane by *ca.* 0.62 Å. We believe

this can be partly attributed to that the hypothetical positioning of Ta “in-plane” would result in the steric clash between the pair of “in-plane” chlorides and the hydrogens *ortho* to the



Table 1 Selected NMR data for palladium and platinum compounds

	$\delta^{13}\text{C}$ for C-M, ppm	$\delta^{31}\text{P}$, ppm	$^1J_{\text{Pt-P}}$, Hz
3-Pd^a	70.2	110.7	—
3-Pt^a	59.4	102.6	3077
4a-Pd	75.2	108.6	—
4b-Pd^a	45.9	93.1	—
4a-Pt	64.0	98.9	3182
4b-Pt^a	— ^d	90.3	3048
5a-Pd^a	68.0	114.6	—
5b-Pd^b	— ^d	97.8	—
6-Pd	76.6	99.1	—
6-Pt	64.9	92.9	3214
7-Pd	64.7	124.4	—
7-Pt	55.3	115.3	3268
8-Pd	74.3	106.2	—
8-Pt	62.4	99.0	3182
9-Pd	— ^d	114.4	—
9-Pt^c	— ^d	106.8	3074

Spectra were collected in C_6D_6 unless otherwise noted. ^aSpectra collected in CDCl_3 . ^bSpectra collected in C_7D_8 . ^cSpectra collected in CD_2Cl_2 . ^dThe Pd/Pt-C resonances were not observed owing to the poor solubility of **9-Pd/Pt** and the low achievable concentration of **4b-Pt** and **5b-Pd**.

nitrogens. In a way, this demonstrates that although the N-N cleft is not “bulky” in the usual three-dimensional sense, it can be thought of as “2D-bulky” in its own plane. Thus, there is little steric congestion with a tetrahedral moiety bound in

Table 2 Selected XRD-derived bond distances in compounds **3-Pd**, **7-Pt**, **8-Pd**, and **9-Pd** (M = Pd or Pt, M' = B, Zn, or Ta) that were studied in solid state by X-ray crystallography

	3-Pd	7-Pt^a	7-Pt^a	8-Pd^b	8-Pd^b	9-Pd
M-P	2.280(2)	2.2897(12)	2.2974(13)	2.297(3)	2.296(3)	2.2862(10)
M-P	2.271(2)	2.2962(12)	2.3098(13)	2.272(3)	2.299(3)	2.2917(10)
M-C	2.028(2)	1.971(4)	1.964(5)	2.008(10)	2.018(10)	1.991(4)
N-M'	N/A	1.561(6)	1.562(7)	1.996(10)	1.963(9)	2.107(3)
N-M'	N/A	1.572(6)	1.569(7)	1.984(8)	2.008(8)	2.104(3)

^aData for both independent molecules of **7-Pt** in the unit cell are given. ^bData for the two crystallographically different Pd-bound ligand frameworks in **8-Pd** are given.

the N-N cleft, but an octahedral moiety must necessarily contend with this “2D-bulkiness”.

Nonetheless, the series of prepared complexes demonstrate well the ability of the ligand to accommodate a variety of sizes in the N-N cleft of the ligand. The C-C-C bond angle that is highlighted in Fig. 2B serves as a good metric to showcase the flexibility. The angle varies by almost 10° across the series of four structures with H_2Cl in **3-Pd** giving rise to the largest angle, followed by the similar values engendered by Zn in **8-Pd** and TaCl_4 in **9-Pd**, and finally the BF_2 moiety in **7-Pt** giving the smallest C-C-C bond angle. The flexing of the N-N cleft portion of the ligand can be seen to affect the geometry on the

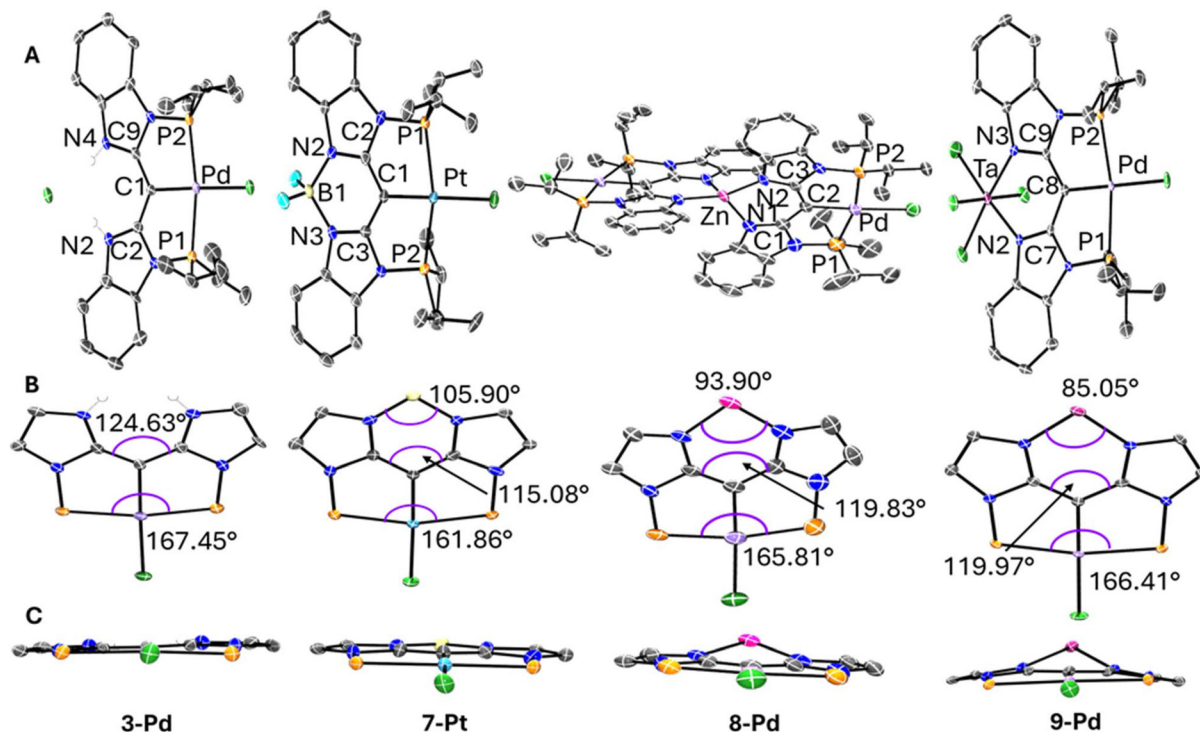


Fig. 2 (A) POV-Ray renditions of ORTEP plots (50% probability ellipsoids) of **3-Pd**, **7-Pt**, **8-Pd**, and **9-Pd** showing selected atom labeling. Hydrogen atoms, solvent molecules, disorder in isopropyl arms are omitted for clarity. An additional molecule of **7-Pt** in the asymmetric unit is also omitted for clarity in **7-Pt** structure. (B) Highlights of selected approximate bond angles for **3-Pd**, **7-Pt**, **8-Pd**, and **9-Pd**. Arenes, isopropyl arms, hydrogen atoms, and disorder are omitted for clarity. (C) Alternate view of each structure in (B) down the M-Cl axis (M = Pd or Pt) to highlight the planarity or lack thereof.



opposing side of the ligand, in the PCP portion, as illustrated by the changes in the P–M–P angle (Fig. 2B). When the N–N cleft of the ligand has to contract when binding the smaller boron atom in **7-Pt**, this causes the phosphines used in the PCP cleft to be pulled back which contracts the P–M–P angle. The binding of the larger Zn (**8-Pd**) or Ta (**9-Pd**), or especially the more open N–N cleft in **3-Pd**, corresponds to larger P–M–P angles. The changes in the N–N cleft can clearly influence the PCP side without changes in the groups immediately connected to the PCP-bound transition metal. However, the spread of the P–M–P angles is only *ca.* 5°, and they are all well within the common range^{52,53} for the [5,5-PXP]⁵⁴ pincer complexes. The modest degree of PCP perturbation illustrates that the NN/PCP ligand system is flexible enough to adapt to the significant variation in the N–N cleft without making the binding of the transition metal on the PCP side impossible.

Electrochemistry

The electrochemical properties of compounds **4**, and **7–9** were studied by means of cyclic voltammetry (CV) experiments (Table 3). A similar picture was observed within each pair of Pd *vs.* Pt analogs, with two quasi-reversible oxidations per ligand. The Pt compounds generally appeared easier to oxidize by *ca.* 0.1 V. Previously studied oxidations of conventional (POCOP)MCl or (PCP)MCl complexes were found to be irreversible.^{55,56} These considerations suggest that the oxi-

dation events in **7–9** are primarily ligand- and not Pd- or Pt-based. DFT calculations performed on the closely related **F** (Fig. 1) indicated that its HOMO is primarily ligand-based.³⁹ We previously studied the reversible oxidation of Pd and Pt complexes supported by diarylamido-based pincer ligands, which was also ligand-based, and where the Pt analogs were also easier to oxidize by several hundredths of a volt.^{57,58} The Pd–Pt difference likely reflects the greater π -interaction of the filled 5d-orbitals of Pt with the π -system of the ligand *vs.* the 4d-orbitals of Pd, driving the energy of the ligand HOMO slightly higher. This notion is also consistent with the slightly lower energy of the LMCT transition in **9-Pt** compared to **9-Pd** (*vide supra*). Compounds **4-Pd** and **4-Pt** appeared to give rise to two quasi-reversible oxidations; however, the recorded CVs had additional unidentified features (see ESI†). They may be owing to the presence of both the **4a** and **4b** tautomers, or to the facile loss of H⁺ after oxidation. With the BF₂ unit in the N–N cleft, the CV waves were well-defined and quasi-reversible for both **7-Pd** and **7-Pt**. The trimetallic species **8-Pd** and **8-Pt** each presented three quasi-reversible waves, one of which was a two-electron wave. We assign the two one-electron waves closer to 0 V as corresponding to the **8⁺/8** and **8²⁺/8⁺** events, with the third wave corresponding to the **8⁴⁺/8²⁺** event. Ostensibly, initial one-electron oxidation of one of the PCP/NN ligands influences the subsequent oxidation of the other PCP/NN ligand. It is interesting that the second oxidation events are not likewise differentiated. It is possible that the second oxidation affects the electron density farther away from the N–N cleft.

In the case of the Ta complexes **9-Pd** and **9-Pt**, while the first oxidation is well-behaved, the second oxidation appeared only partially reversible. It is possible that the higher potential needed for the second oxidation of **9** leads to a less stable dicationic species than in the case of **7** (or the tetracationic in the case of **8**). In addition, both **9-Pd** and **9-Pt** also displayed a quasi-reversible wave corresponding to a **9/9[–]** event at –0.87 and –0.88 V. We tentatively ascribe it to the reduction of the Ta^V center to Ta^{IV}. These values for **9** fall in between the reduction potentials for the [TaCl₆][–]/[TaCl₆]^{2–} (–1.11 V) and the [TaCl₅(NCMe)]⁰/[TaCl₅(NCMe)][–] (–0.65 V) couples, as reported by Bursten *et al.*⁵⁹

Two oxidations per ligand in **7–9** can be understood by using simple resonance structures, a few of which are shown in Fig. 3 for **7-Pd**. The lone pair localized on the central carbon in resonance structure **II** is delocalized, primarily in the “nacnac” fashion (**I** and **III**), but also beyond, into the six-

Table 3 Reduction potentials (V) of the events observed by cyclic voltammetry for compounds **7–9**

Complex N	N ²⁺ /N ⁺	N ⁺ /N ⁰	N ⁰ /N [–]
7-Pd	1.18	0.25	—
7-Pt	1.03	0.11	—
8-Pd	0.92 ^a	0.02 ^b	—
		–0.22 ^b	—
8-Pt	0.82 ^a	–0.09 ^b	—
		–0.32 ^b	—
9-Pd^c	1.18	0.57	–0.87
9-Pt^c	1.07	0.47	–0.88

Cyclic voltammograms were recorded in CH₂Cl₂ at 1 mM of analyte and referenced to the Fc⁺/Fc redox couple. Unless otherwise noted CVs were run with 0.1 M of [Bu₄N][PF₆] as the supporting electrolyte. ^aCorresponds to a two-electron event N⁴⁺/N²⁺. ^bTwo events corresponding to N²⁺/N⁺ and N⁺/N⁰ because of the presence of two ligands in the molecules. ^cCVs run with 0.1 M of [Bu₄N][BARF₂₄] as the supporting electrolyte.

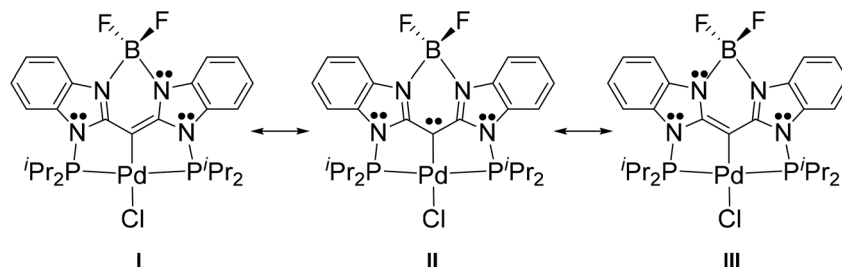


Fig. 3 Main resonance structures for **7-Pd**.



membered aromatic rings (not shown). The two successive oxidations can be thought to convert this central lone pair first into a radical, and then into an empty orbital site, with extensive delocalization. The importance of the resonance structure similar to **II** in compound **F** (Fig. 1) led Ong, Frenking, Zhao, and coworkers to point out the similarity of the central carbon-metal interaction in **F** to the transition metal complexes of “carbodicarbenes”.³⁹ That similarity also exists for 7–9. The importance of the resonance structure **II** is also related to the PCP pincer complexes featuring a central diaryl-carbene, explored by the Piers^{43–45,60} and Iluc⁴⁶ groups.

Conclusion

In summary, a new binucleating ligand system has been presented. It combines a monoanionic PCP pincer on one side with a monoanionic N–N bidentate cleft on the opposite side. The PCP pincer cleft was accommodated with square-planar, d⁸ Pd or Pt centers, while the N–N cleft has proven adaptable to binding such divergent set of elements as boron, zinc, or tantalum. Two N–N clefts can bind a single Zn center, resulting in trimetallic (ZnPd₂ or ZnPt₂) complexes. Structural studies showed that the ligand stays nearly flat in the coordination plane of Pd or Pt, indicating a high degree of conjugation throughout. When bound to metals (or metalloids) *via* both the N–N and the PCP clefts, the ligand system can undergo two quasi-reversible oxidations, ascribed primarily to the oxidation of the extended organic π -system.

Data availability

Details of experimental procedures, graphical NMR spectra, and details of X-ray structural determination, CV experiments, and graphic UV-Vis spectra have been included as ESI.† CCDC 2351193–2351196† contain crystallographic data for **3-Pd**, **7-Pt**, **8-Pd**, and **9-Pd**.

Conflicts of interest

There are no conflicts to declare.

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