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An Isolable Mononuclear Palladium(I) Amido Complex

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Characterized by XRD, EPR, XAS

been characterized by X-ray crystallography, electron paramagnetic resonance, and multiedge Pd X-ray absorption spectroscopy. Theoretical study revealed that, while the three-electron-two-center π -interaction between Pd and N in the Pd(I) complex imposes severe Pauli repulsion in its Pd-N bond, pronounced attractive interligand dispersion force aids its stabilization. In accord with its electronic features, reactions of homolytic Pd-N bond cleavage and deprotonation of primary amines are observed on the Pd(I) amido complex.

INTRODUCTION

Palladium is a frequently used metal in the modern molecular chemists' catalyst arsenal and can be found in a variety of useful organic reactions, such as Wacker oxidations, crosscouplings, C-H bond functionalization, and hydrogenation reactions.¹ Underlying the broad utility of Pd are the multiple oxidation states accessible, which suit them well to the formation and cleavage of covalent bonds. Pd compounds most commonly occur as diamagnetic, closed-shell Pd(0), Pd(II), and Pd(IV) complexes. Paramagnetic, open-shell Pd compounds are far less prevalent, but are now receiving increased attention.²⁻⁵ Paramagnetic Pd(I) species, for example, are implicated as intermediates in some Pd-catalyzed reactions, but sparse data are available concerning their electronic structures and reactivity. In Pd-catalyzed C-C bond-forming reactions, Pd(I) halide species L_nPd^IX (L = phosphine; X = Br, I) are proposed to form via single-electron transfer reactions of photoexcited Pd(0) phosphine complexes with organic halides, $^{6-8}$ as well as in the reactions of Pd(0) phosphine complexes with fluoroalkyl halides (Figure 1A).⁹ In the reactions of Pd(0) species with dioxygen, which is a key step in Pd-catalyzed aerobic oxidation reactions, Pd(I) superoxide species $L_n Pd^{I}(\eta^1 - OO^{\bullet})$ are proposed as intermediates en route to Pd(II) peroxides $L_n Pd^{II}(\eta^2 - O_2)$ (Figure 1A).¹⁰ The formation of the Pd(I) intermediates has also been envisaged in thermolysis or photolysis of Pd(II) compounds. For example, photolysis-induced decomposition of (PNP)Pd-(II) alkyl (PNP = phosphine-amido-phosphine) complexes was proposed to proceed through (PNP)Pd(I) intermediates that dimerized to yield diamagnetic, dinuclear Pd(I) complexes (Figure 1A).¹¹

(BINAP)PdCl₂ with LiNHAr^{Trip}. This Pd(I) amido species has

A common feature of the aforementioned reactive Pd(I)intermediates is the presence of π -donors—ligands whose coordinating atoms bear nonbonding lone pairs, e.g., halides, superoxide, and amido in the aforementioned cases-within Pd's inner coordination sphere. Such mononuclear Pd(I) species reported to date are exceedingly reactive and have only been observed using spectroscopic methods in γ -ray-irradiated $Pd(II)Cl_2$ and $Pd(II)(acac)_2$ and thermalized Pd(II)-exchanged zeolites,¹²⁻¹⁴ where their stability is conferred by immobilization within the lattice framework. Certain Pd(0)and Pd(II) complexes are reported to undergo chemical and electrochemical redox reactions to produce mononuclear paramagnetic Pd(I) complexes.^{15–19} These include three isolable complexes, $[(PBu_3)_2Pd]X$ (X = PF₆, $(CB_{11}Cl_{11}H)^{18,19}$ and $[(PBu_{3}^{t})_{2}Pd(NCMe)](CB_{11}Cl_{11}H)$. These mononuclear Pd(I) complexes, however, are not supported by π -donating ligands. To our knowledge, isolable Pd(I) complexes bearing π -donating ligand have remained elusive. The challenge of accessing mononuclear Pd(I)complexes bearing π -donating ligands likely originates from severe Pauli repulsion: conventional wisdom dictates that π interactions between the nd orbitals of low-valent platinumgroup metals (Ru, Rh, Pd, Os, Ir, Pt) and the lone pairs of π donors (Figure 1B) should enforce severe Pauli repulsion,^{20,21} leading to the destabilization of the metal-ligand multiple

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Figure 1. Research background on Pd(I) species bearing π -donating ligands. (A) Examples of reactive Pd(I) intermediates containing π -donating ligands. (B) Pauli repulsion in metal-ligand π -interaction. (C) Cartoon illustrating how London dispersion forces stabilize an M-X bond.



Figure 2. Synthetic route to the Pd(I) amido complex and its molecular structure. (A) Reaction giving Pd(I) amido complex [(BINAP)Pd(NHAr^{Trip})] (1) and the byproducts 2–4. (B) Molecular structure of 1, showing one of the two crystallographically independent molecules in the unit cell with 30% thermal ellipsoids and a partial atom-numbering scheme. Hydrogen atoms, except the one on N1, are omitted for clarity. Selected distances (Å) and angles (deg): Pd1–N1 2.063(2), Pd1–P1 2.3314(7), Pd1–P2 2.2883(7), N1–C45 1.377(3), P2–Pd1–P1 92.20(2), N1–Pd1–P1 108.06(6), N1–Pd1–P2 154.04(6), C45–N1–Pd1 135.29(18). (C) Proposed route for the formation of 1 and the byproducts 2–4.

bond. Pauli repulsion has been used to explain the high reactivity of low-valent platinum-group metal amido, alkoxide, imido, oxo, and nitride species.^{22–24} So far, low-valent platinum-group metal complexes containing π -donor ligands are restricted to the nd^6-nd^8 metal ions.^{25–27} Analogous isolable complexes containing a nd^9 -platinum-group metal center are unknown.

Herein we report the first nd^9 platinum-group metal terminal amido complex, namely, the Pd(I) complex [(BINAP)Pd-(NHAr^{Trip})] (BINAP = 2,2'-bis(diphenylphosphino)-1,1'binaphthalene, Ar^{Trip} = 2,6-bis(2',4',6'-triisopropylphenyl)- phenyl). The Pd(I) amido complex is isolated from the reaction of (BINAP)PdCl₂ with LiNHAr^{Trip} and has been characterized by single-crystal X-ray diffraction, electron paramagnetic resonance spectroscopy, and multiedge Pd X-ray absorption spectroscopy. Electronic structure calculations indicate that the Pd(I) amido complex is S = 1/2 with a three-electron two-center π -interaction between the Pd and N atoms and that the Pd–N bond indeed presents severe destabilizing Pauli repulsion. However, dispersion forces²⁸ (Figure 1C) between BINAP and the bulky amido ligand aids the stabilization of the complex. Further reactivity studies evaluate

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Figure 3. Spectroscopic data of Pd complexes. (A–C) EPR spectra of 1 (black line) and simulations (red line) at 9 GHz CW, 34 GHz electron spin echo field sweep (derivative), and 94 GHz CW EPR. The minor features at the right in the 94 GHz EPR spectrum come from the Mn^{2+} impurities from Critoseal in the W-band tube. (D) Overlay of the smoothed second derivatives of Pd K-edge XAS spectra of $Pd^{0}(IPr)_{2}$ (5, blue), 1 (red), and (BINAP)Pd^{II}(OAc)₂ (6, black). (E) Experimental (red) and TDDFT-calculated (black) Pd L₃-edge XAS spectra of 1. The B3LYP functional and SARC-ZORA basis set were used for Pd, with ZORA-def2-TZVP(-f) used for all other atoms. Contributions of individual orbitals were obtained from Löwdin population analysis of the QROs. Orbitals are plotted at an isovalue of 0.03 au with hydrogen atoms removed for clarity.

the lability of the Pd–N bond. When subjected to thermolysis, photoirradiation, or coordination of exogenous ligands, the Pd(I) amido complex undergoes homolytic Pd–N bond cleavage to release the aminyl radical $[NHAr^{Trip}]^{\bullet}$. In addition, the Pd(I) amido complex can react with primary alkyl and aryl amines to release NH_2Ar^{Trip} . These observations showcase the unique reactivity of low-valent, odd-electron platinum-groupmetal amido complexes.

RESULTS AND DISCUSSION

Synthesis and Molecular Structure of [(BINAP)Pd-(**NHAr**^{Trip})]. The Pd(I) amido complex [(BINAP)Pd-(NHAr^{Trip})] (1) was obtained via the reaction of (BINAP)-PdCl₂ with two equivalents of LiNHAr^{Trip}. Mixing the two reagents in toluene at -78 °C produced a blue solution, whose ³¹P NMR spectrum features two signals with an AB-splitting pattern, suggesting the formation of a Pd(II) monoamido intermediate, presumably (BINAP)Pd(NHAr^{Trip})Cl (A). When warmed to room temperature, the solution turned green. Recrystallization led to the isolation of the Pd(I) amido complex 1 as a green crystalline solid in 51% yield (Figure 2A). The composition of 1 is supported by elemental analysis (C, H, N) and cold electrospray mass spectroscopy (Figure S6), and its structure has been confirmed by single-crystal X-ray diffraction (Figure 2B). The infrared spectrum of 1 features an N-H stretch at 3295 cm⁻¹. The deuterated isotopologue

[(BINAP)Pd(NDAr^{Trip})] (1-d) shows the N-D stretch at 2538 cm^{-1} (Figure S5). In addition to 1, three amine byproducts, the amino-imine 2, the aniline featuring an alkene side arm 3, and the free aniline 4 (Figure 2A), are observed. These amines are likely formed from the decomposition reactions of the aminyl radical [NHAr^{Trip}]. Consequently, 1 is proposed to form via a homolytic Pd-N bond cleavage reaction of a Pd(II) bisamido intermediate, (BINAP)Pd- $(NHAr^{Trip})_{2}$, that can be produced by the interaction of **A** with LiNHAr^{Trip} (Figure 2C). Single crystals of 1 were obtained by cooling a diethyl ether solution at -30 °C for 2 days. An X-ray diffraction study revealed that there are two crystallographically independent molecules in the unit cell, with similar structure metrics. As the representative, Figure 2B depicts the structure of one of the molecules. The three-coordinate Pd(I) complex has a P-Pd-P angle of $92.20(2)^{\circ}$ and two unequal P-Pd-N angles of $108.06(6)^{\circ}$ and $154.04(6)^{\circ}$. While the sum of the three angles (354.3°) indicates an idealized planar geometry for the Pd center, the presence of two unequal P-Pd-N angles differentiates 1 from trigonal-planar threecoordinate Pd(0) complexes $(Pr_2^iPCH_2CH_2CH_2PPr_2^i)Pd$ - $(PCy_3)^{29}$ and $Pd(PPh_3)_3^{30}$ and T-shaped three-coordinate Pd(II) complexes $(PBu_3^t)Pd(C_6H_4-4-OCH_3)(N(C_6H_3-3,5 (CF_3)_2)_2^{31}$ and $(IPr)Pd(3-methylnorborn-2-yl)(NHC_6H_4-4 CH_3$) (IPr = 1,3-bis(2',6'-diisopropylphenyl)imidazole-2-ylidene).³² The Pd–N bond distance in 1 is 2.063(2) Å, which is slightly longer than those of the three-coordinate Pd(II) amido complexes (sIPr)Pd(3-methylnorborn-2-yl)(NHC₆H₄-4- OCH_3) (sIPr = 1,3-bis(2',6'-diisopropylphenyl)imidazolin-2ylidene) (2.037(3) Å) and (IPr)Pd(3-methylnorborn-2-yl)- $(NHC_6H_4-4-CH_3)$ (2.011(3) Å).³² Counter to the knowledge that the *trans* influence of the amido ligand would elongate the transoid Pd-P bond, the 2.2883(7) Å Pd1-P2 bond is shorter than the 2.3314(7) Å Pd1-P1 bond. The shortening of the Pd-P bond distance might be caused by the joint effect of Pdto-phosphine π -backdonation (vide infra) and the attractive dispersion forces between the phenyl groups on the P atom and the proximal 2,4,6-triisopropylphenyl group on the amido ligand (vide infra). In addition, the bond distances within the amido ligand of 1 itself are typical of anilido ligands in metal complexes^{31,32} and are different from those of the parent phenylaminyl radical [NHPh]^{•33} and the arylaminyl complexes (Table S1),³⁴⁻³⁷ which hints at the monoanionic nature of $[NHAr^{Trip}]^{1-}$ in 1. Notably, the core structure of 1 is reminiscent of Hillhouse's nickel(I) amido complexes [(dtbpe)Ni(NHDipp)] (dtbpe = 1,2-bis(di-tertbutylphosphino)ethane) and [(dtbpe)Ni(NHDmp)] (Dmp = 2,6-dimesitylphenyl).^{38,3}

Spectroscopic and Electronic Structure Features of [(BINAP)Pd(NHAr^{Trip})]. Complex 1 features an $S = \frac{1}{2}$ ground spin-state as indicated by the measured magnetic moment of 2.1(1) $\mu_{\rm B}$. Continuous wave (CW) and field-swept echodetected (FSE) electron paramagnetic resonance (EPR) spectra were obtained for 1 (Figures 3A-C and S74). The X-band (9 GHz) CW-EPR spectrum obtained at room temperature shows a four-line pattern centered at $g_{iso} \approx$ 2.052 with $A_{iso}(^{31}\text{P1}) = 202 \text{ G} (567 \text{ MHz}) \text{ and } A_{iso}(^{31}\text{P2}) = 80$ G (225 MHz) (Figure S74). The anisotropy of the g-values and the ³¹P A-values were resolved at low temperatures by obtaining spectra at the X-, Q- (35 GHz), and W-band (94 GHz). Figure 3A–C shows the frozen-solution multifrequency EPR spectra obtained for 1 along with simulations using globally fit spin-Hamiltonian parameters. The g-values obtained are $g_1 = 2.0085$, $g_2 = 2.0350$, and $g_3 = 2.1115$. The small g-value is consistent with the density functional theory (DFT)-calculated electronic structure, as discussed below (Table 2), but is unusual among the few known Pd(I)

Table 1. Experimental Pd K-Edge and L_{2,3}-Edge Energies

	K-edge (eV)	L ₃ pre-edge max (eV)	L ₂ pre-edge max (eV)	total L ₃ + L ₂ mainline area
5	24 347	nd	nd	nd
1	24 351	3174.5	3331.7	1.66(0.03)
6	24 353	3176.4	3333.6	4.02(0.16)

 Table 2. Experimental and DFT-Calculated EPR Parameters

 for 1

	experimental	calculated ^a
g tensor	[2.0085, 2.0350, 2.1115]	[2.0216, 2.0408, 2.1065]
$A(^{31}\text{P1})/\text{MHz}$	[205, 210, 260]	[237, 242, 298]
$A(^{31}\text{P2})/\text{MHz}$	[535, 535, 631]	[535, 537, 650]
$A(^{105}\text{Pd})/\text{MHz}$	NR	[105, 31.1, 80.8]
$A(^{14}N)/MHz$	NR	[2.6, 3.3, 24.7]

"Calculations used crystallographic coordinates with the B3LYP hybrid density functional, D3 correction for dispersion, the SARC-ZORA basis set on Pd, and the ZORA-def2-TZVP(-f) basis set on all other atoms. complexes. For example, the spectra of the Pd(I) species generated by γ -ray irradiation on Pd(II) compounds have their g_{\perp} -values in the range 2.0 to 2.1 and g_{\parallel} -values in the range 2.1 to 2.6,⁴⁰ the spectrum of $[(PBu_3^t)_2Pd][CB_{11}Cl_{11}H]$ has the gvalues of $g_{\perp} = 2.338$, $g_{\parallel} = 1.971$,¹⁹ and that of $[(PBu^{t}_{3})_{2}Pd$ - $(NCMe)][CB_{11}Cl_{11}H]$ shows $g_{\perp} = 2.088$, $g_{\mu} = 2.314$.¹⁹ For the reference, EPR of Pd(III) generally displays axial spectra with g_{\parallel} and g_{\perp} values of 2.01–2.05 and 2.28–2.31, respectively, although isotropic and rhombic EPR spectra have also been reported.⁴¹ The ³¹P hyperfine constant values (HFCs) determined by globally fitting the frozen-solution EPR spectra of 1 for ³¹P1 are $A_1 = 535$, $A_2 = 535$, and $A_3 = 631$ MHz. The values for ${}^{31}P2$ are $A_1 = 205$, $A_2 = 210$, and $A_3 = 260$ MHz. Deduction of atomic spin density from the isotropic (A_{iso}) and dipolar (T) contributions to the HFCs⁴² yields a spin density of 13.0% on P1, of which 4.3% resides in the 3s orbital and 8.7% in a 3p orbital, and a spin density on P2 is 6.4%, where 1.7% resides in the 3s orbital and 4.7% in the 3p orbital. Thus, total spin density on two P nuclei is ca. 20% based on the EPR experiments. $A(^{105}Pd)$ and $A(^{14}N)$ contributions to the spectra could not be resolved.

To probe the electronics of the Pd center in 1, Pd K- and Ledge XAS data for 1, the Pd(0) complex $[Pd^{0}(IPr)_{2}]$ (5), and the Pd(II) complex $[(BINAP)Pd^{II}(OAc)_2]$ (6) were collected. As shown in Figures 3 D and S75 and Table 1, the rising-edge energies obtained from the K-edge ($1s \rightarrow valence/continuum$) for the series of compounds shift toward higher energies with increasing formal oxidation state, although it has been noted that factors such as coordination environment and ligand identity can shift rising-edge energies to degrees similar to shifts caused by oxidation state changes.⁴³ Pre-edge features arising from the metal $1s \rightarrow nd$ transitions are also conventionally used to assess physical oxidation states. Such excitations are not expected for the Pd(0) complexes because of their d¹⁰ configuration, but will occur in the case of formally Pd(I) and Pd(II) complexes. Such features are not resolved in the experimental Pd K-edge spectra likely owing to the weak intensity of these quadrupole-allowed but dipole-forbidden transitions as well as the core-hole lifetime broadening endemic to second- and third-row metal K-edge XAS.

To complement the K-edge data, the Pd L_{2.3}-edge (2p \rightarrow valence/continuum) XAS data were obtained, where dipoleallowed Pd $2p \rightarrow 4d$ transitions gives rise to intense features that probe the energetics and covalency of the Pd ligand field.⁴⁴ Experimental Pd L_{2,3}-edge spectra are shown in Figures 3 E and S75-S77 for 1 and 6. L₃ and L₂ mainlines in these spectra can be assigned to excitations from the Pd 2p orbitals into singly or unoccupied Pd 4d orbitals. Satellite features at higher energies relative to the mainline result from transitions into higher lying levels of primarily ligand character (vide *infra*). The L_3 and L_2 mainlines for 1 are shifted by ca. 2 eV to lower energy relative to 6 (Table 1). Areas of $L_{2,3}$ -edge mainlines can be used to quantify metal *n*d character in valence acceptor orbitals, providing a direct probe of covalency and physical oxidation state.⁴³ This approach has been applied to Pd L_{2.3}-edges, where an increase in the L_{2.3}-edge area of PdAl₃ and PdCl₂ compared to Pd metal was attributed to d-count depletion.⁴⁵ In the present case, the total L_{2,3}-edge mainline area of 1 is ca. half that of 6, consistent with Pd-localized reduction.

Hybrid DFT calculations were carried out to interrogate the electronic structure of **1**. All calculations used the dispersion-



Figure 4. DFT calculations of the Pd(I) amido complex. (A) Selected molecular orbitals of 1; hydrogen atoms on ligands were omitted. (B) EDA-NOCV results (in kcal/mol). Fragments of (BINAP)Pd and NHAr^{Trip} are given in the table in singlet (S) or doublet (D) electronic states. (C) Short interligand H…H contacts with distances of less than 2.4 Å (purple lines).

corrected B3LYP-D3 hybrid density functional^{46,47} with the segmented all-electron relativistically contracted (SARC)-ZORA basis on Pd48 and ZORA-def2-TZVP(-f) basis set on all other atoms. All calculations used crystallographic coordinates. Veracity of the electronic structure calculations was judged by comparison of calculated spectroscopic parameters to experiment. Spin-Hamiltonian parameters calculated for 1 agree well with experiment (Table 2). Specifically, the g-values as well as the magnitudes and asymmetry of the ³¹P HFCs are effectively reproduced by experiment. Calculated Mulliken spin densities comprise 9.5% 3p and 3.9% 3s on P1 and 5.7% 3p and 2.7% 3s on P2, in splendid agreement with analysis of the experimentally obtained HFCs (vide supra). The calculation yielded low ¹⁴N and ¹⁰⁵Pd HFC values, consistent with the fact that these features are unresolved in the EPR data.

The single-point DFT solutions were used as starting points for time-dependent DFT (TDDFT) calculations of the X-ray absorption spectroscopy (XAS) data obtained for 1 and 6. TDDFT has been used previously to calculate L2.3-edge XAS spectra for second-row transition metals and was shown to adequately reproduce the structure of experimental L3-edges without including the contributions from spin-orbit coupling that yield the L3-L2 splitting.49 TDDFT reproduced the overall envelope of L3 features in the experimental spectra (Figures 3E and S80). The satellite features to higher energy of the L₃ mainlines comprise transitions into ligand-localized molecular orbitals with minimal Pd 4d character. The mainline transition in 1 primarily comprises the Pd 2p \rightarrow SOMO excitation, while for (BINAP)Pd^{II}(OAc)₂ the mainline consists of the Pd 2p LUMO excitation as well as transitions to two additional low-lying MOs with ca. 10% contributions from Pd 4d orbitals. Calculated total Pd 4d character in all acceptor

orbital holes against summed L₃ and L₂ mainline areas gives a correlation with $R^2 = 0.97$ (Figure S82). Formally Pd^I complex 1 exhibits 30% lower overall d vacancy than the Pd^{II} complex, providing further support for metal-centered reduction relative to formally Pd^{II}-containing **6**.

The experimentally validated DFT calculations indicate that 1 bears a three-electron two-center Pd-N π -bond, and its electronic configuration can be described as $(\pi_{d_{xy}+p_N})^2 (d_{z^2}, d_{xzy}, d_{z^2}, d_{x^{-y^2}})^8 (\pi^*_{d_{xy}, p_N})^1$. Löwdin orbital populations show that the SOMO is highly delocalized, although there is a substantial (40%) Pd 4d contribution, consistent with the observed anisotropy in g-values obtained for 1 (Figure 4A). Major contributors to the SOMO come from the Pdonors (10.3% 3p) and the amide (10.4% N 2p). The sum of the spin density on the NHAr^{Trip} fragment, 0.22, precludes description of 1 as an aminyl radical complex. Relevant comparisons include the aminyl radical species [(bipy)Rh(N- $(\text{Trop})_2)$ [OTf] (0.56, bipy = bipyridine, Trop = 5*H*-dibenzo[*a*,*d*]cyclohepten-5-yl),²⁶ [(Ph₂B(CH₂PBu^t₂)₂)Cu(N- $(tolyl-p)_2$ (0.69),³⁶ [(Me₃NN)Cu(NPh₂)] (0.58, Me₃NN = 2,4-bis(2,4,6-trimethylphenylimido)pentyl),³⁷ [(PNP)NiCl]-[OTf] (0.69),³⁵ and [(PNP)Re(CO)₃][OTf] (0.83),³⁴ which feature higher spin density on their N-ligands. On the other hand, the spin-density distribution on 1 is closer to that of Hillhouse's Ni(I) amido complex [(dtbpe)Ni(NHDmp)],³⁹ where the unpaired spin locates essentially on the nickel center (0.76 on nickel and 0.07 on nitrogen, as shown in Figure S89).

To gain further insight into the nature of the Pd–N bond in 1, we did bonding analysis by using the state-of-the-art energy decomposition analysis (EDA)⁵⁰ with the natural orbitals for the chemical valency (NOCV) approach (see SI for details).^{51–53} The most important numerical results are provided in Figure 4B, while more detailed information can



Figure 5. Reactivity of 1. (A) Thermolysis-, light irradiation-, and ligand-coordination-induced decomposition reactions of 1, as well as its reactions with primary amines and a phenol. (B) Hammett plot with *para*-substituted 2,6-dimethylaniline (k_X/k_H = relative rate constants for *para-X*-substituted 2,6-dimethylanilines versus 2,6-dimethylaniline; σ_p = Hammett substituent constant, ρ = reaction constant). (C) Kinetic isotope effect k_H/k_D for the reactions of 1 with 2,6-dimethylaniline.

be found in Table S3 and Figures S84-S88. The interacting neutral fragments of (BINAP)Pd⁰ and NHAr^{Trip} have comparable orbital energy (i.e., $\Delta E_{\rm orb}$ = -72.0 kcal/mol) with that of the ionic fragments of [(BINAP)Pd^I]⁺ and- $[NHAr^{Trip}]^-$ (i.e., $\Delta E_{orb} = -73.7$ kcal/mol), implying the two resonant structures exist for 1, but the neutral fragments are slightly favored because of the smaller orbital energy (see Figure S88a). This also agrees very well with the spectroscopic data and spin density analysis as shown in Figure S88b. It revealed that there is significant Pauli repulsion (i.e., $\Delta E_{\text{Pauli}} =$ 123.1 kcal/mol) within the Pd-N interaction by using the neutral interacting fragments. Nonetheless, the favorable attractive interactions, including the electrostatic energy $(\Delta E_{\text{elstat}} = -92.3 \text{ kcal/mol})$ and orbital interaction energy $(\Delta E_{\rm orb} = -72.0 \text{ kcal/mol})$, can compensate and even exceed the Pauli repulsion ΔE_{Pauli} by 41.2 kcal/mol. It should be emphasized that the dispersion energy ($\Delta E_{disp} = -31.3$ kcal/ mol) contributing 16% of total attractive forces leads to a summed bond dissociation energy (D_e) of 50.9 kcal/mol, which helps to stabilize the Pd(I) amido species at ambient temperature. The dispersion forces can be discerned from the short interligand H…H contacts between BINAP and NHAr^{Trip} in the optimized structure of 1 (Figure 4C). The ionic interacting fragments have similar Pauli repulsion and orbital energies but with a significant strong electrostatic energy $(\Delta E_{\text{elstat}} = -163.5 \text{ kcal/mol})$, which is understandable between the cation [(BINAP)Pd^I]⁺ and anion [NHAr^{Trip}]⁻ fragments and thus leads to a larger intrinsic energy $\Delta E_{\rm int}$ of -139.0 kcal/ mol and bond dissociation energy (D_e) of 121.3 kcal/mol. It should be mentioned that, in addition to the thermodynamic factors, the stabilization of 1 apparently benefits from steric protection of the Pd-N bond by the bulky substituents.

Reactivity of [(BINAP)Pd(NHAr^{Trip})]. Complex 1 is the first isolable d^9 platinum-group-metal complex bearing a π -

donating ligand. Being different from the aforementioned fleeting Pd(I) species, it is stable at ambient temperature, which facilitates studies of its reactivity. When **1** is subjected to thermolysis, light irradiation, or coordination with exogenous ligands, it undergoes Pd–N bond homolytic cleavage reactions to yield anilines and Pd(0) species (Figure 5A). In these reactions, the arylaminyl radical dimerization product **2**, the alkane-chain dehydrogenation product **3**, and the aniline NH₂Ar^{Trip} (**4**) are observed, implicating the radical intermediate [NHAr^{Trip}][•] in these reactions (Figure S19). In addition, two Pd(0) complexes, [(BINAP)Pd(2,6-(CH₃)₂-C₆H₃NC)₂] (7) and [(BINAP)Pd(η^2 -3,5-(CF₃)₂-C₆H₃CHCH₂)] (**8**), have been isolated in the corresponding reaction and structurally authenticated by X-ray diffraction studies (Figure S29 and S34).

The decomposition reaction of 1 under white light irradiation (6 W LED) at room temperature shows zeroorder kinetics with a kinetic constant $k_{irri} = 1.6 \times 10^{-8} \text{ mol} \cdot \text{L}^{-1} \cdot$ s^{-1} (Figure S23), whereas the thermal decomposition reaction at 348 K is a first-order reaction with a rate constant $k_{\text{therm}} =$ $1.1 \times 10^{-4} \text{ s}^{-1}$ (Figure S27). The thermal- and lightirradiation-induced decomposition reactions of 1 might result from amido-to-palladium(I) electron transfer to form $(BINAP)Pd^{0}(NHAr^{Trip})^{\bullet}$. The absorption spectrum of 1 in toluene shows an intense feature at 808 nm (7300 mol⁻¹·L· cm^{-1}). TDDFT calculations suggest that the band principally arises due to what is best described as a $\pi \to \pi^*$ transition with ligand-to-metal charge transfer (LMCT) character: the principal donor orbital is the Pd–N π -bonding SOMO–1, whose largest contributor is the N 2p, while the acceptor orbital is the Pd–N π^* SOMO, which is dominated by Pd 4d character (Figure S79). Either transition fully populates the Pd-N antibonding MO and thus promotes bond cleavage. Kinetics study on the reactions of 1 with an excess amount of

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c04965.

Experimental procedures, characterization, data, computational details, and Cartesian coordinates (PDF)

Accession Codes

CCDC 2059309–2059313 and 2070924 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/ cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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3,5-bis(trifluoromethyl)styrene in benzene at 30 $^{\circ}$ C revealed pseudo-first-order kinetics on the concentrations of both 1 and the alkene (Figures S41–S47). This suggests that the ligand-coordination-induced-decomposition reaction might proceed via an associative mechanism or involve ligand exchange equilibria.

When 1 is treated with phenols and amines, e.g., 2,6-di(tertbutyl)-4-methylphenol, 2,6-di(isopropyl)aniline, 2,6-dimethylaniline, *n*-butyl amine, and *n*-octyl amine, the free aniline NH_2Ar^{Trip} (4) was formed in high yields. The reaction with the phenol allows the isolation of the Pd(0) complex (2,6-di-tertbutyl-4-methylenecyclohexa-2.5-dienone)palladium(0) (9) in 67% yield (Figure 5A), whose structure has been confirmed by X-ray diffraction (Figures S48 and S49). The reactions with amines produced blue-purple mixtures, and the attempts to isolate palladium-containing species from the mixture were unsuccessful. These reactions likely start with a proton-transfer step between the basic amido ligand in 1 and the H-X (X = O, N) bonds in phenols and amines, rather than a hydrogenatom-abstraction mechanism as (i) a linear Hammett plot with a positive reaction constant $\rho = +0.77$ was obtained for the reactions of 1 with anilines bearing different para-substituents (Figure 5B), (ii) no obvious kinetic isotope effect was observed in the reactions 1 with 2,6-Me₂C₆H₃NH₂ and 2,6- $Me_2C_6H_3ND_2$ (Figure 5C), and (iii) no reaction took place when 1 was treated with substrates bearing weak C-H bonds, e.g., xanthene (73.3 kcal/mol in DMSO) and cyclohexa-1,4diene (67 kcal/mol in the gas phase), whose C-H bond dissociation energies are much lower than those of the O-H bond of 2,6-di(tert-butyl)-4-methylphenol (78 kcal/mol in DMSO) and the N-H bonds of amines (\geq 89 kcal/mol in DMSO).⁵⁴ Thus, the formation of the Pd(0) complex 9 can be explained by the sequential steps of proton transfer followed by electron transfer and hydrogen atom abstraction reactions (Figure S55). Notably, as anilido anions are generally less basic than aliphatic amido anions,⁵⁴ the capability of 1 to react with aliphatic amines to form NH_2Ar^{Trip} demonstrates the unique reactivity of the d⁹ Pd amido species. The elucidation of the mechanism of the reaction of 1 with aliphatic amines needs further study.

CONCLUSION

In summary, the first isolable, room-temperature-stable Pd(I)complex bearing a π -donating ligand [(BINAP)Pd-(NHAr^{Trip})], which also represents the first platinum-groupmetal amido complex with an nd^9 electronic configuration, has been synthesized from the salt elimination reaction of (BINAP)PdCl₂ with LiNHAr^{Trip}. The Pd(I) amido complex features a long Pd–N bond and inequivalent N–Pd–P angles. EPR and XAS spectroscopies indicate that the Pd-N bond in the Pd(I) amido complex is highly covalent. Theoretical studies accord with the spectroscopic data and further revealed that pronounced dispersion forces between the BINAP and the bulky amido ligand NHAr^{Trip} help the stabilization of this complex. Reactivity studies revealed that the Pd-N bond in [(BINAP)Pd(NHAr^{Trip})] can undergo homolytic cleavage reaction to release the aminyl radical [NHAr^{Trip}]• when the complex is subjected to heat, light irradiation, or ligand coordination. The Pd(I) complex can also react with primary amines to yield NH_2Ar^{Trip} . The reactivity profile of the Pd(I) amido complex points out the potential synthetic utility of lowvalent platinum-group-metal species bearing π -donating ligands for the design of new reactions.

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Notes

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