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Reversible Modulation of the Electronic and Spatial Environment around Ni(0) Centers Bearing Multifunctional Carbene Ligands with Triarylaluminum

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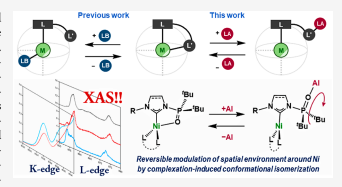
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ABSTRACT: Designing and modulating the electronic and spatial environments surrounding metal centers is a crucial issue in a wide range of chemistry fields that use organometallic compounds. Herein, we demonstrate a Lewis-acid-mediated reversible expansion, contraction, and transformation of the spatial environment surrounding nickel(0) centers that bear *N*-phosphine oxide-substituted *N*-heterocyclic carbenes (henceforth referred to as (S)PoxIms). Reaction between tetrahedral ($syn-\kappa$ -C,O-(S)PoxIm)-Ni(CO)₂ and Al(C₆F₅)₃ smoothly afforded heterobimetallic Ni/Al species such as trigonal-planar { κ -C-Ni(CO)₂}(μ -anti-(S)PoxIm)-{ κ -O-Al(C₆F₅)₃} via a complexation-induced rotation of the *N*-phosphine oxide moieties, while the addition of 4-dimethylamino-pyridine resulted in the quantitative regeneration of the former Ni



complexes. The corresponding interconversion also occurred between (SPoxIm)Ni(η^2 : η^2 -diphenyldivinylsilane) and { κ -C-Ni(η^2 : η^2 -diene)}(μ -anti-SPoxIm){ κ -O-Al(C₆F₅)₃} via the coordination and dissociation of Al(C₆F₅)₃. The shape and size of the space around the Ni(0) center was drastically changed through this Lewis-acid-mediated interconversion. Moreover, the multinuclear NMR, IR, and XAS analyses of the aforementioned carbonyl complexes clarified the details of the changes in the electronic states on the Ni centers; i.e., the electron delocalization was effectively enhanced among the Ni atom and CO ligands in the heterobimetallic Ni/Al species. The results presented in this work thus provide a strategy for reversibly modulating both the electronic and spatial environment of organometallic complexes, in addition to the well-accepted Lewis-base-mediated ligand-substitution methods.

■ INTRODUCTION

Designing the spatial environment around a metal center is a critical issue in terms of the electronic and steric properties and hence the reactivity of organometallic complexes. Chemists have thus focused on the design of the structural, electronic, and dynamic properties of supported ligands, as demonstrated in the field of, e.g., homogeneous catalysis, 1-6 supramolecular chemistry, ^{7,8} and materials science. ⁹ One well-known strategy for reversibly modulating the electronic and spatial environment around metal centers is based on a ligand-substitution process on metal centers bearing multifunctional ligands including a hemilabile coordination moiety (L') in the presence of an external Lewis base (LB) (Figure 1A, left). 10-14 However, these processes often yield an equilibrium mixture that is difficult to separate. In this context, we have reported the reversible, recyclable, and pressure-responsive room-temperature-chemisorption of carbon monoxide (CO) on a zerovalent nickel complex that bears multifunctional multipurpose carbene ligands, 14 namely N-phosphine oxidesubstituted imidazolinylidenes (SPoxIms) and the corresponding imidazolylidenes (PoxIms) (Figure 1B, left). 15,16 (S)PoxIm ligands in tetrahedral (S)PoxImNi(CO)_n complexes selectively adopt either a κ -C- (n=3) or κ -C,O- (n=2) coordination mode, which can be interconverted through the addition/exclusion of CO.

A strategy based on the use of Lewis acids (LAs) is another potential option for reversibly modulating the spatial environment around metal centers that bear multifunctional ligands (Figure 1A, right). The LA-mediated process can cause a substantial change in both the electronic state and spatial environment of the metal center due to the change in the number of ligands. This feature clearly distinguishes LA-mediated processes from LB-mediated ones as the latter commonly proceeds without changing the number of ligands.

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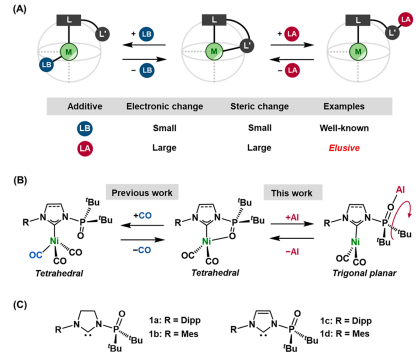


Figure 1. (A) A simplified scheme of the reversible modulation of the spatial environment around a metal center in an organometallic complex mediated by Lewis bases (LBs, left) or Lewis acids (LAs, right). (B) Reversible transformation of Ni/(S)PoxIm complexes using Lewis-basic CO (our previous work, left) and Lewis-acidic aluminum species (this work, right). (C) The (S)PoxIms used in this work (Dipp = 2,6diisopropylphenyl, Mes = 2,4,6-trimethylphenyl).

However, examples of such LA-mediated reactions have been severely limited because LAs often trigger the irreversible decomposition of the organometallic complex via the abstraction of ligands to form an LA-LB adduct. A judicious combination of multifunctional ligands and LAs is thus essential for constructing an LA-mediated system that can reversibly regulate the electronic and spatial environments around the metal centers. It should be noted that Fan et al. have reported an example that relies predominantly on an electrostatic interactions^{21,22} involving the reversible complexation of Na+ and a crown-ether moiety included in a Rh complex that bears Aza-CrownPhos; the reactivity of the Rh complex can be regulated through the Na+-mediated interconversion.²³ The authors evaluated the change in the spatial environment around the Rh center using multinuclear NMR spectroscopy and ESI mass spectrometry, even though a quantitative evaluation of how much the field expanded and diminished via the reaction was not discussed.

Herein, we present a novel strategy to reversibly transform the spatial environment around a nickel center by combining (S)PoxIms 1a-d (Figure 1C) as multifunctional ligands and tris(pentafluorophenyl)aluminum as an LA, where the geometry of the Ni center is interconverted between tetrahedral and trigonal planar (Figure 1B, right). We quantitatively evaluated the variation of the spatial environment, i.e., the volume and shape of the space, and the electronic state around the nickel center based on single-crystal X-ray diffraction (SC-XRD) and X-ray absorption spectroscopy (XAS) analyses.

■ RESULTS AND DISCUSSION

Previously, we have reported that (S)PoxIms include syn- and anti-conformers with respect to the N-P bonds, which can interconvert easily at room temperature via rotation of the Nphosphinoyl groups ($\Delta E^{\ddagger} \sim 12 \text{ kcal mol}^{-1}$ for previously reported PoxIms), even though the anti-conformers are thermodynamically favorable (Figure 2). 15,16 Importantly, the

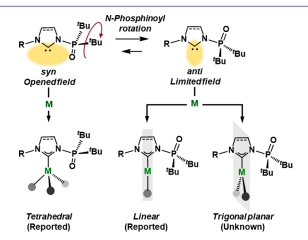


Figure 2. Interconversion between syn- and anti-conformers of (S)PoxIms, as well as complexation between (S)PoxIms and transition metals that prefer different coordination geometries.

volume and shape of the space can be drastically varied via rotation of the N-phosphinoyl groups, and a relatively limited space is present around the carbene carbon atoms in the anti-(S)PoxIms. Due to this limited spatial environment, only group-11 metals such as Cu²⁴ and Au²⁵ have been confirmed to form complexes that bear anti-(S)PoxIms in a κ -C fashion, given that these metals tend to adopt two-coordinated linear complexation geometries (Figure 2). We have also reported the synthesis of heterobimetallic complexes of the type (κ -C-

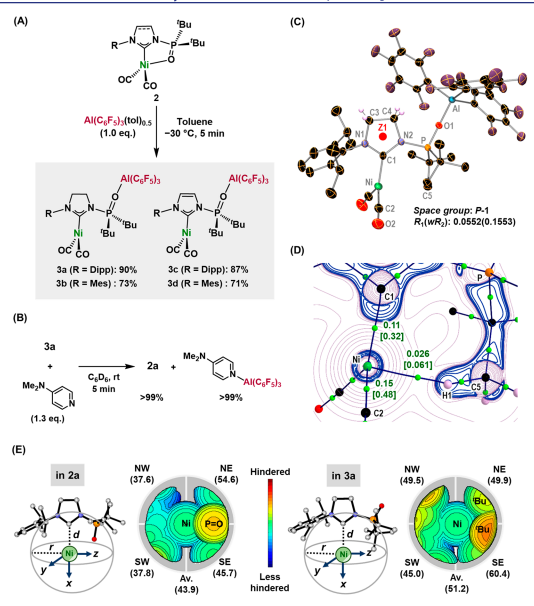


Figure 3. (A) Synthesis of the heterobimetallic Ni/Al complexes 3 via the reaction between (syn - κ - C,O - 1)Ni(CO)₂ (2) and an equimolar amount of Al(C₆F₅)₃(tol)_{0.5}. Yield of isolated products is shown. (B) Reaction of 3a with DMAP. (C) Molecular structure of 3a with a thermal ellipsoid at 30% probability; H atoms (except those bound to C3 and C4) are omitted for clarity. Selected bond lengths (Å) and angles (deg): Ni–C1 1.945(3), Ni–C2 1.759(3), Ni–C5 3.240(4), C2–O2 1.140(4), P–O1 1.524(2), O1–Al 1.799(2), Z1–C1–Ni 169.9, C1–N2–P–O1 159.7(2). (D) The quantum theory of AIM bond paths (black lines) and bond-critical points (green dots) are overlaid with the contour plots of $\nabla^2 \rho$ through the plane defined by the C1, Ni, and C5 atoms. Selected atoms and values of the electron density ($e r_{Bohr}^{-3}$) are shown. The values of $\nabla^2 \rho$ ($e r_{Bohr}^{-5}$) are shown in square brackets. (E) Topographic steric maps and W_{bur} values of 2a/3a calculated using the program SambVca (r = 3.5 Å; d = 2.0 Å; bond radii scaled by 1.17; mesh spacing 0.05; H atoms are omitted) based on the structural parameters obtained from the SC-XRD analysis.

Cu)(μ -anti-(S)PoxIm)(κ -O-Al) through complexation between anti-(κ -O-(S)PoxIm)Al(C₆F₅)₃ and CuO^tBu followed by an irreversible intramolecular Cu-O^tBu/Al-C₆F₅ transmetalation triggered by the rotation of the N-phosphinoyl moiety. However, reversible spatial modifications around the metal center between isolable species remain challenging.

In this work, we envisioned an $Al(C_6F_5)_3$ -mediated reversible interconversion between complexes bearing *syn*-and *anti-*(S)PoxIms based on the use of Ni(0), i.e., an interconversion between tetrahedral (κ - $C_7O_7(S)$ PoxIm)Ni-(CO)₂ and trigonal-planar { κ -C-Ni(CO)₂}(μ -*anti-*(S)PoxIm)-{ κ -O-Al(C_6F_5)₃} (Figure 1B, right). The preparation of trigonal-planar Ni(0) dicarbonyl complexes that bear bulky

N-heterocyclic carbenes (NHCs) such as (I^tBu)Ni(CO)₂ and (IAd)Ni(CO)₂^{26,27} (I^tBu = 1,3-di-*tert*-butylimidazol-2-ylidene; IAd = 1,3-diadamantylimidazol-2-ylidene) has already been reported, which inspired us to explore the preparation of Ni(0) complexes bearing bulky *anti*-(S)PoxIm ligands (Figure 2).

We started our study with the reaction between $(syn-\kappa-C,O-1a)$ Ni(CO)₂ (2a) and an equimolar amount of Al- $(C_6F_5)_3(tol)_{0.5}$, which resulted in the selective formation of heterobimetallic Ni/Al complex 3a in 90% yield (Figure 3A). We also confirmed that the Al($C_6F_5)_3$ -mediated rotation of *N*-phosphinoyl moieties in 2b-d afforded the corresponding heterobimetallic Ni/Al complexes (3b-d) in good-to-high yield. The differences in steric bulkiness between the Dipp and

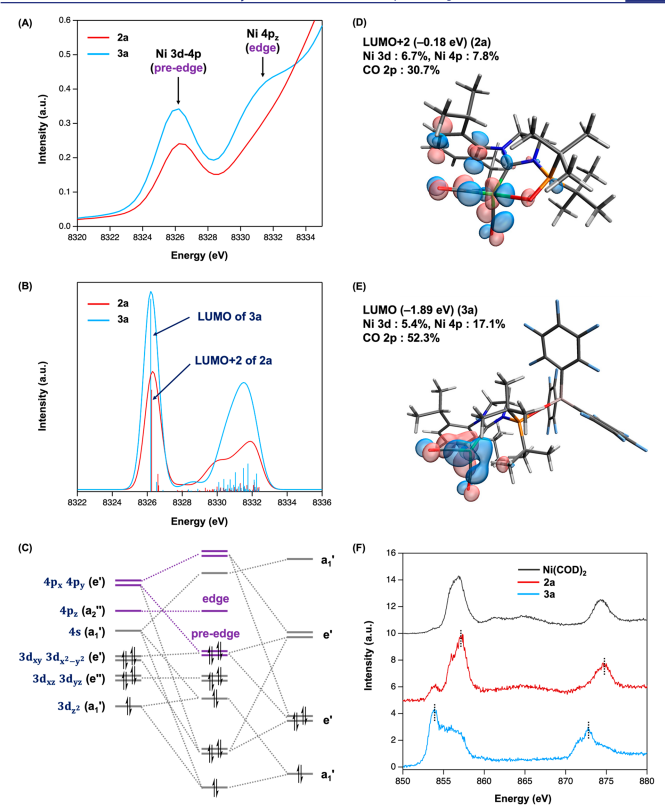


Figure 4. (A) Ni K-edge XANES spectra of 2a/3a in toluene. (B) The simulated Ni K-edge XAS spectra of 2a/3a using the TDDFT method at the ZORA-B3LYP/CP(PPP) (for Ni) and ZORA-def2-TZVP(-f) (for C, H, O, N, F, P, and Al) level. (C) Qualitative MO diagram of the trigonal-planar complex. The purple bars denote electric dipole allowed, and the gray bars forbidden transition orbitals in the Ni K-edge XAS. (D) Kohn—Sham LUMO+2 for 2a. (E) Kohn—Sham LUMO for 3a. (F) Ni L_{2,3}-edge XAS spectra of Ni(cod)₂, 2a, and 3a in the PFY method.

Mes groups, as well as the structural flexibility variations between saturated SPoxIms and unsaturated PoxIms, had no significant effect on the progress of the reaction. The molecular structures of 3a-d were fully characterized using NMR and

ATR-infrared (IR) absorption spectroscopy as well as SC-XRD analysis (vide infra). Subsequently, we treated 3a with slight excess amounts of 4-dimethylaminopyridine (DMAP) and confirmed the quantitative regeneration of 2a with the

concomitant formation of the adduct DMAP-Al(C_6F_5)₃ (Figure 3B). These results demonstrated that the interconversion between 2a and 3a can be successfully mediated by the addition and removal of Al(C_6F_5)₃.

Next, we compared the electronic and geometric parameters of 2a and 3a. In the ³¹P NMR spectrum of 3a, the resonance corresponding to the N-phosphinoyl moiety is observed at $\delta_{\rm p}$ 82.0 in toluene- d_8 , which represents a significant downfield shift compared to that of 2a ($\delta_{\rm p}$ 68.0 in THF- $d_{\rm s}$), ¹⁴ while it is almost identical to that of $(\kappa$ -O-1a)Al(C₆F₅)₃ (δ _P 79.8 in CD₂Cl₂).²⁴ These results indicate that the 1a moiety in 3a should predominantly adopt the anti-conformation. The geometric parameters of 3a obtained from the SC-XRD analyses are shown in Figure 3C. As previously reported, the Ni center in 2a, which bears the κ - C_1O -syn-1a ligand (C-N-P-O torsion angle: 8.5(1)°), adopts a tetrahedral geometry. 14 In stark contrast, 3a features a trigonal-planar Ni center (sum of bond angles around Ni: 358.9°) bearing *anti-***1a** (C1-N2-P-O1 torsion angle: $159.7(2)^{\circ}$), as shown in Figure 3C. The Ni(CO)₂ unit is located slightly out of ideal alignment with the carbene lone pair (Z1-C1-Ni angle: 169.9°; Z1 = centroid of the imidazolinylidene ring), most likely due to the high steric demand of the ^tBu groups.

Interestingly, a noncovalent interaction between the Ni and H1 atoms was confirmed in the gas-phase-optimized structure of 3a (level: PBE0/def2-TZVPD//M06L/def2-SVPD for O, F, and Ni; def2-SVP for all other atoms) via a topological analysis of the electron density, which was calculated using the quantum theory of atoms in molecule (AIM) method (Figure 3D).^{29,30} It should also be noted here that the geometric parameters of the solid state structure of 3a confirmed by SC-XRD analysis were closely reproduced in the theoretically calculated gas-phase-optimized structure; e.g., the interatomic distances between the Ni and C5 atoms were 3.18 Å in the gasphase and 3.240(4) Å in the crystalline state. In fact, the Ni··· H1 distance $(2.13 \text{ Å})^{31}$ and the relatively high electron density $(\rho = 0.026 \ e \ r_{\rm Bohr}^{-3})$ as well as the positive value of the Laplacian of the electron density ($\nabla^2 \rho = +0.061 \ e \ r_{Bohr}^{-5}$) at the bond-critical point (BCP) between these atoms suggests the participation of an agostic Ni···H interaction.^{29,30}

Dissociation of the N-phosphinoyl moiety from tetrahedral 2a to afford trigonal planar 3a also caused obvious changes in the wavenumbers of the stretching vibration of the CO ligands. In fact, the signal corresponding to the stretching vibration of the CO ligands in 3a was observed at 2019 cm⁻¹, i.e., at a higher wavenumber than that of 2a (1967 cm⁻¹). The AIM analysis clarifies the increase in the delocalization indices $\delta(N_i)$ C_{CO}) and $\delta(Ni, O_{CO})$ in 3a compared to those in 2a, which represent a number of electron pairs delocalized between the Ni and carbonyl carbon atoms (C_{CO}) and the Ni and carbonyl oxygen atoms (O_{CO}), respectively (Figures S7-S8).³⁰ The mean values of $\delta(Ni, C_{CO})$ and $\delta(Ni, O_{CO})$ increase from 0.51 (2a) to 1.20 (3a) and from 0.063 (2a) to 0.23 (3a), respectively, indicating that electrons become effectively delocalized through the Ni, C_{CO}, and O_{CO} atoms in 3a. On the other hand, the values of $\delta(C_{CO}, O_{CO})$ are nearly identical between 2a (1.70–1.71) and 3a (1.60–1.62). Thus, we attribute the aforementioned blue-shift to the increased polarization of the π CO bonding orbitals due to an increased cationic nature on the Ni in 3a (AIM atomic net charge = 0.50 e) compared to 2a (AIM atomic net charge = -0.1 e).³²

Based on the aforementioned geometric parameters of 2a and 3a in the crystalline state, the impact of the *N*-phosphinoyl

rotation on the spatial environment surrounding the Ni center was quantitatively evaluated on the basis of percent buried volume (%V_{bur}) (Figure 3E).^{2,33} Topographic steric maps that visualize the % $V_{\rm bur}$ values in the northwestern (NW), northeastern (NE), southwestern (SW), and southeastern (SE) quadrants, as well as their average (Av), are also depicted; these maps were produced using the program SambVca. 2.1.² The Av(% V_{bur}) values in 2a and 3a were calculated to be 43.9 and 51.2, respectively, clearly demonstrating that the space around the Ni center is significantly altered via rotation of the N-phosphinoyl moiety. Importantly, drastic expansion/contraction of the space can be precisely and reversibly triggered by the addition/removal of $Al(C_6F_5)_3$. In addition, this rotational transformation alters the shape of the space around the Ni center, as is clearly confirmed by the comparison of both the $\%V_{\rm bur}({\rm NE})$ and $\%V_{\rm bur}({\rm SE})$ values of ${\bf 2a}$ and ${\bf 3a}$.

We then carried out solution-phase XAS measurements to determine whether the results obtained from the SC-XRD analysis could be extended to the solvated states of 2a and 3a. A magnified view of the pre-edge and edge region of the Ni Kedge X-ray absorption near edge structure (XANES) is shown in Figure 4A. In the case of 2a, a pre-edge peak corresponding to the electric dipole transition from the Ni 1s orbital to the Ni 4p(-3d) orbitals was observed at 8326 eV and no characteristic peak was detected at the absorption edge, which is typical for nickel complexes with tetrahedral geometry. 14 Meanwhile, a remarkable peak appeared in the edge region (8331 eV) for 3a, along with a greater pre-edge peak intensity compared to 2a. We attribute the characteristic edge peak to the Ni 1s \rightarrow 4p. transition in 3a, given the presence of the nonbonding $4p_z$ orbital (a2" symmetry) (Figure 4C). Considering that such nonbonding 4pz orbitals are commonly found in trigonalplanar 16-electron complexes, the results of the solution-phase XAS experiments clearly rationalize the adoption of a trigonalplanar geometry by 3a, even in toluene, as observed in the crystal structure.

To gain further insight, the Ni K-edge XAS spectra of 2a and 3a were simulated using the time-dependent DFT (TDDFT) calculation at the B3LYP level with the zeroth-order regular approximation $(ZORA)^{34-36}$ to take the relativistic effect into consideration.³⁷ The CP(PPP) basis set was used for Ni, while the ZORA-def2-TZVP(-f) basis set was used for all of the other elements. The intensities of the pre-edge and edge peaks of the simulated XAS spectra show good agreement with those of the experimental spectra (Figure 4A and 4B). To identify the predominant contribution to the pre-edge peaks, we then analyzed the molecular orbitals (MOs) of 2a and 3a. In the case of 2a, the transition from the Ni 1s orbital to LUMO+2 significantly contributes to the pre-edge peak, where LUMO+2 mainly consists of the Ni 3d (6.7%), Ni 4p (7.8%), and CO 2p (30.7%) orbitals (Figure 4D). On the other hand, in the case of 3a, the increased intensity of the corresponding pre-edge peak can be rationalized predominantly by the transition from the Ni 1s orbital to the LUMO. The LUMO of 3a features increased contribution from the Ni 4p orbitals (17.1%) and CO ligands (52.3%) compared to their contributions to the LUMO+2 in 2a (Figure 4E), while the Ni 3d orbitals exhibit a nearly identical degree of contribution to the respective MOs in 2a and 3a. We therefore attributed the increase in the preedge peak intensity of 3a induced by the $Al(C_6F_5)_3$ -mediated geometric change to the increased efficiency of the Ni 3d-4p mixing and contribution of CO ligands (e' symmetry) in the trigonal planar geometry.

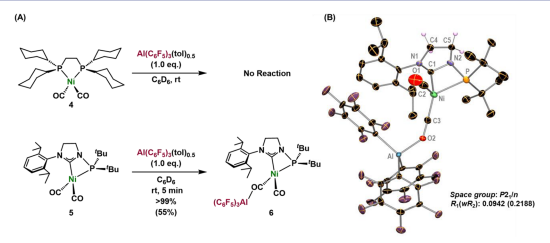


Figure 5. (A) Reactions between complexes 4 and 5 and $Al(C_6F_5)_3(tol)_{0.5}$. The yield was determined by NMR analysis. The yield of isolated products is shown in parentheses. (B) Molecular structure of 6 with thermal ellipsoids at 30% probability; H atoms (except those bound to C4 and C5) are omitted for clarity. Selected bond lengths (Å) and angles (deg): Ni–C1 1.991(6), Ni–P 2.314(2), Ni–C2 1.814(8), Ni–C3 1.691(5), C2–O1 1.12(1), C3–O2 1.197(6), O2–Al 1.861(4), and C3–O2–Al 139.3(4).

We further explored the electronic structures of the Ni 3d orbitals in the solid state for 2a and 3a based on Ni L_{2,3}-edge XAS experiments using the partial fluorescence yield (PFY) method. In the cases of 2a and Ni(cod)2, the absorption maxima on the Ni L₃-edge appear at 856.0 and 856.2 eV, respectively, with a shoulder peak in their lower-energy regions (Figure 4F). These Ni L_{2,3}-edge XAS spectra represent spectroscopic features characteristic of tetrahedral nickel complexes with a d10 electron configuration (formally Ni(0)). In contrast, for 3a, the peak maximum shifts toward the lower-energy region to 852.9 eV, clearly indicating that the 3d orbitals are stabilized through the formation of the 16electron complex with a vacant 4p, orbital. In fact, the energy levels of the unoccupied frontier MOs of 3a are substantially lower than those of 2a, showing good agreement with the experimental Ni L_{2,3}-edge XAS data (Table S4). Furthermore, a broad signal is observed in the higher energy region at 854-858 eV. Based on the results of the Cu L_{2,3}-edge XAS experiments previously reported by Lancaster et al., we attribute this signal to the transitions from the Ni 2p_{3/2} orbital to the ligand and Ni 3d orbitals, given the occupied d10 electron configuration in 3a.³⁷

To clarify the key role of (S)PoxIms, we subsequently explored reactions between $Al(C_6F_5)_3$ and Ni(0) dicarbonyl complexes 4 and 5 (Figure 5A). Complex 4, which carries the bis(dicyclohexylphosphino)ethane ligand, did not react with $Al(C_6F_5)_3$ under the applied conditions. On the other hand, 6 was quantitatively formed without complexation between the phosphine moiety and $Al(C_6F_5)_3$ when N-phosphanylsubstituted imidazolidine-2-ylidene was used as a hemilabile ligand instead of 1a. The molecular structure of 6 was unambiguously determined by a SC-XRD analysis and is shown in Figure 5B. Even though our targeted spatial modulation was not confirmed, these results are noteworthy to demonstrate that CO ligands are sufficiently Lewis-basic to form classical Lewis-acid/base adducts with Lewis-acidic triarylaluminum species.³⁸⁻⁴⁰ The aforementioned results suggest that to realize the Al(C₆F₅)₃-mediated reversible modulation of the spatial environment surrounding Ni(0) it is crucial to use a hemilabile ligand that includes both hard and soft (or mild) Lewis-basic sites and that is responsive to external stimuli in order to induce a preprogrammable

structural isomerization. (S)PoxIms with their distinctive Lewis-basic sites, i.e., the harder phosphinoyl oxygen moiety and the softer carbene moiety, as well as the complexation-induced rotation of the *N*-phosphinoyl moiety may thus successfully serve as pioneering examples for this purpose.

Finally, we confirmed that the present strategy can be expanded to $(syn-\kappa-C,O-1a)$ Ni complex 7a, which contains an $\eta^2:\eta^2$ -diphenyldivinylsilane ligand instead of CO (Figure 6A). Treatment of 7a with $Al(C_6F_5)_3(tol)_{0.5}$ resulted in the quantitative formation of the heterobimetallic Ni/Al complex 8a via the complexation-induced rotation of the N-phosphinoyl moiety. Moreover, the addition of DMAP again caused the dissociation of $Al(C_6F_5)_3$ to regenerate 7a.

The molecular structures of compounds 7a and 8a were confirmed by SC-XRD analyses (Figure 6B). Although the C1-Ni lengths remained almost unchanged upon rotation of the N-phosphinoyl moiety, the sterically demanding ^tBu groups cause a significant deviation of the Ni(η^2 : η^2 diphenyldivinylsilane) unit from the plane that is defined by the C1, N1, C6, C7, and N2 atoms; i.e., the deviated distances of Ni atoms from the plane are 0.02 Å in 7a and 0.41 Å in 8a. Both 7a and 8a include an unsymmetrical coordination environment around the Ni center in their crystalline states, and the two olefin moieties are thus inequivalent. Nevertheless, the bond lengths of C2=C3 and C4=C5 are nearly identical, even when compared between 7a and 8a, i.e., C2=C3: 1.393(2) Å in 7a and 1.380(8) Å in 8a; C4=C5: 1.404(2) Å in 7a and 1.402(7) Å in 8a, implying a negligible variation of the electron density in these olefin moieties upon the aforementioned interconversion. However, an obvious elongation occurred in the bonds between the Ni and internal carbon atoms (C2 and C4), which should be caused by the structural constraint to adopt the distorted trigonal-planar geometry. The Ni-O1 bond (2.481(1) Å) is longer than the corresponding one in 2a (2.227(1) Å), ¹⁴ whereas the participation of the bonding interaction between the Ni and O1 atoms in 7a was confirmed by the AIM analysis (Figure S12).

We also found that ¹H NMR resonances of the coordinated olefin moieties, which are magnetically equivalent in solution at room temperature, upfield shifted upon the coordination change around the Ni center from four-coordinated 7a to distorted trigonal-planar 8a (Figure 6C). In the ³¹P NMR

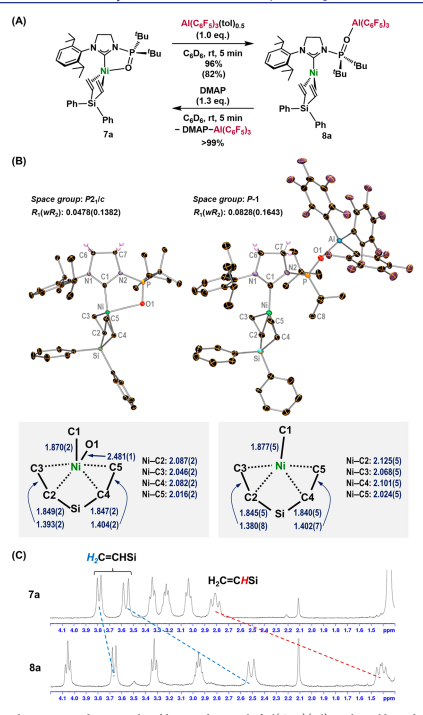


Figure 6. (A) Interconversion between 7a and 8a upon the addition and removal of $Al(C_6F_5)(tol)_{0.5}$. The yield was determined by NMR analyses. The yield of isolated products is shown in parentheses. (B) Molecular structure of 7a (left) and 8a (right) with thermal ellipsoids at 30% probability; H atoms (except those bound to C6 and C7) are omitted for clarity. Selected bond lengths (Å) are also shown. (C) The ¹H NMR spectra of 7a and 8a in C_6D_6 .

spectrum, a significant downfield shift from δ_P 68.3 (7a) to 83.3 (8a) corroborates the formation of the P=O-Al bond. In their entirety, the results of the NMR and SC-XRD analyses suggest that the $Al(C_6F_5)_3$ -mediated rotation of the N-phosphinoyl group can modulate not only the spatial environment around the Ni center but also the chemical environment surrounding the coordinated 1,4-diene unit through the interconversion between 7a and 8a.

CONCLUSIONS

In summary, we have reported a strategy that allows reversible modulation of the electronic state and spatial environment around metal centers that bear multifunctional ligands based on the use of Lewis acids. To this end, a series of N-phosphine oxide-substituted N-heterocyclic carbenes, referred to as (S)PoxIms, were employed, as (S)PoxIms can afford heterobimetallic species in κ -C-M¹ and κ -O-M² fashions without the formation of quenched NHC—Lewis acid adducts. The reaction between tetrahedral (syn- κ -C,O-(S)PoxIm)Ni-

 $(CO)_2$ and $Al(C_6F_5)_3$ smoothly afforded heterobimetallic Ni/ Al species such as trigonal-planar $\{\kappa\text{-C-Ni(CO)}_2\}(\mu\text{-anti-$ (S)PoxIm) $\{\kappa$ -O-Al(C₆F₅)₃ $\}$ through complexation-induced rotation of the N-phosphine oxide moiety, while the addition of DMAP quantitatively triggered the formation of the former complex. We also confirmed that the interconversion between $(syn-\kappa-C,O-SPoxIm)Ni(\eta^2:\eta^2-1,4-diene)$ and the corresponding heterobimetallic Ni/Al species effectively proceeded upon addition and removal of Al(C₆F₅)₃, which successfully demonstrates that the present strategy can be applied for modulating the spatial environment around not only metal centers but also the coordinated substrates. We experimentally and theoretically confirmed that the shape and size of the space around the Ni(0) center drastically expanded/contracted using this Lewis-acid-mediated procedure. Furthermore, a detailed discussion based on multinuclear NMR, IR absorption, and Xray absorption spectroscopy shed light on the changes in the electronic states of the Ni centers. Thus, this work manifests a conceptually novel and effective approach to design and modulate the electronic and spatial environment surrounding metal centers in organometallic compounds using a combination of multifunctional ligands and Lewis acids.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.3c06267.

> Full details pertaining to the experimental methods, identification of the compounds, and DFT calculations. (PDF)

Accession Codes

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Notes

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REFERENCES

- (1) Wu, K.; Doyle, A. G. Parameterization of phosphine ligands demonstrates enhancement of nickel catalysis via remote steric effects. Nat. Chem. 2017, 9, 779-784.
- (2) Falivene, L.; Cao, Z.; Petta, A.; Serra, L.; Poater, A.; Oliva, R.; Scarano, V.; Cavallo, L. Towards the online computer-aided design of catalytic pockets. Nat. Chem. 2019, 11, 872-879.
- (3) Durand, D. J.; Fey, N. Computational Ligand Descriptors for Catalyst Design. Chem. Rev. 2019, 119, 6561-6594.
- (4) Zhang, Z.; Shao, Y.; Tang, J.; Jiang, J.; Wang, L.; Li, S. Supramolecular asymmetric catalysis mediated by crown ethers and related recognition systems. Green Synth. Catal. 2021, 2, 156-164.

- (5) Trouvé, J.; Gramage-Doria, R. Beyond hydrogen bonding: recent trends of outer sphere interactions in transition metal catalysis. *Chem. Soc. Rev.* **2021**, *50*, 3565–3584.
- (6) Blanco, V.; Leigh, D. A.; Marcos, V. Artificial switchable catalysts. Chem. Soc. Rev. 2015, 44, 5341-5370.
- (7) Miller, A. J. M. Controlling ligand binding for tunable and switchable catalysis: cation-modulated hemilability in pincer-crown ether ligands. *Dalton Trans.* **2017**, *46*, 11987–12000.
- (8) Yoo, C.; Dodge, H. M.; Miller, A. J. M. Cation-controlled catalysis with crown ether-containing transition metal complexes. *Chem. Commun.* **2019**, *55*, 5047–5059.
- (9) Peris, E. Smart N-Heterocyclic Carbene Ligands in Catalysis. Chem. Rev. 2018, 118, 9988–10031.
- (10) Braunstein, P.; Naud, F. Hemilability of Hybrid Ligands and the Coordination Chemistry of Oxazoline-Based Systems. *Angew. Chem., Int. Ed.* **2001**, *40*, 680–699.
- (11) Fliedel, C.; Schnee, G.; Braunstein, P. Versatile coordination modes of novel hemilabile S-NHC ligands. *Dalton Trans.* **2009**, 2474–2476.
- (12) Liang, Q.; Janes, T.; Gjergji, X.; Song, D. Iron complexes of a bidentate picolyl-NHC ligand: synthesis, structure and reactivity. *Dalton Trans.* **2016**, 45, 13872–13880.
- (13) Mondal, M.; Ranjeesh, T. K.; Gupta, S. K.; Choudhury, J. Labile coordination approach for the modulation of the electronic properties of ruthenium(II) and iridium(III) complexes within an "Nheterocyclic carbene (NHC)–pyridyl" dynamic platform. *Dalton Trans.* **2014**, *43*, 9356–9362.
- (14) Yamauchi, Y.; Hoshimoto, Y.; Kawakita, T.; Kinoshita, T.; Uetake, Y.; Sakurai, H.; Ogoshi, S. Room-Temperature Reversible Chemisorption of Carbon Monoxide on Nickel(0) Complexes. *J. Am. Chem. Soc.* **2022**, *144*, 8818–8826.
- (15) Hoshimoto, Y.; Kinoshita, T.; Ohashi, M.; Ogoshi, S. A Strategy to Control the Reactivation of Frustrated Lewis Pairs from Shelf-Stable Carbene Borane Complexes. *Angew. Chem., Int. Ed.* **2015**, *54*, 11666–11671.
- (16) Hoshimoto, Y.; Ogoshi, S. Development of Metal Complexes Equipped with Structurally Flexible Carbenes. *Bull. Chem. Soc. Jpn.* **2021**, *94*, 327–338.
- (17) For examples on LA-mediated reversible electronic modulation on metal centers, see: (a) Lee, B. Y.; Bazan, G. C.; Vela, J.; Komon, Z. J. A.; Bu, X. α-Iminocarboxamidato-Nickel(II) Ethylene Polymerization Catalysts. J. Am. Chem. Soc. 2001, 123, 5352-5353. (b) Kim, Y. H.; Kim, T. H.; Lee, B. Y.; Woodmansee, D.; Bu, X.; Bazan, G. C. α-Iminoenamido Ligands: A Novel Structure for Transition-Metal Activation. Organometallics 2002, 21, 3082-3084. (c) Hesp, K. D.; McDonald, R.; Ferguson, M. J.; Schatte, G.; Stradiotto, M. (κ^2 -P,S)Pt(benzyl) complexes derived from 1/3-PiPr2-2-StBu-indene: facile synthesis of carbanion- and borate-containing zwitterions. Chem. Commun. 2008, 5645-5647. (d) Hesp, K. D.; McDonald, R.; Ferguson, M. J.; Stradiotto, M. New Cationic and Zwitterionic $Cp*M(\kappa^2-P,S)$ Complexes (M = Rh, Ir): Divergent Reactivity Pathways Arising from Alternative Modes of Ancillary Ligand Participation in Substrate Activation. J. Am. Chem. Soc. 2008, 130, 16394-16406. (e) Azoulay, J. D.; Rojas, R. S.; Serrano, A. V.; Ohtaki, H.; Galland, G. B.; Wu, G.; Bazan, G. C. Nickel α -Keto- β -Diimine Initiators for Olefin Polymerization. Angew. Chem., Int. Ed. 2009, 48, 1089-1092. (f) Azoulay, J. D.; Koretz, Z. A.; Wu, G.; Bazan, G. C. Well-Defined Cationic Methallyl α-Keto-β-Diimine Complexes of Nickel. Angew. Chem., Int. Ed. 2010, 49, 7890-7894. (g) Trofymchuk, O. S.; Gutsulyak, D. V.; Quintero, C.; Parvez, M.; Daniliuc, C. G.; Piers, W. E.; Rojas, R. S. N-Arylcyano- β -diketiminate Methallyl Nickel Complexes: Synthesis, Adduct Formation, and Reactivity toward Ethylene. Organometallics 2013, 32, 7323-7333. (h) Tseng, K.-N. T.; Kampf, J. W.; Szymczak, N. K. Regulation of Iron-Catalyzed Olefin Hydroboration by Ligand Modifications at a Remote Site. ACS Catal. 2015, 5, 411-415. (i) Lambic, N. S.; Brown, C. A.; Sommer, R. D.; Ison, E. A. Dramatic Increase in the Rate of Olefin Insertion by Coordination of Lewis Acids to the Oxo Ligand in Oxorhenium(V) Hydrides. Organometallics 2017, 36, 2042-2051. (j) Geri, J. B.;

- Shanahan, J. P.; Szymczak, N. K. Testing the Push-Pull Hypothesis: Lewis Acid Augmented N₂ Activation at Iron. J. Am. Chem. Soc. 2017, 139, 5952–5956. (k) Goudy, V.; Jaoul, A.; Cordier, M.; Clavaguéra, C.; Nocton, G. Tuning the Stability of Pd(IV) Intermediates Using a Redox Noninnocent Ligand Combined with an Organolanthanide Fragment. J. Am. Chem. Soc. 2017, 139, 10633–10636. (l) Shanahan, J. P.; Szymczak, N. K. Lewis Acid Effects on Calculated Ligand Electronic Parameters. Organometallics 2020, 39, 4297–4306. (m) Schmid, L.; Chábera, O.; Rüter, I.; Prescimone, A.; Meyer, F.; Yartsev, A.; Persson, P.; Wenger, O. S. Borylation in the Second Coordination Sphere of Fe^{II} Cyanido Complexes and Its Impact on Their Electronic Structures and Excited-State Dynamics. Inorg. Chem. 2022, 61, 15853–15863.
- (18) For selected examples on LA-mediated ligand substitution reaction, see: (a) Liberman-Martin, A. L.; Levine, D. S.; Liu, W.; Bergman, R. G.; Tilley, T. D. Biaryl Reductive Elimination Is Dramatically Accelerated by Remote Lewis Acid Binding to a 2,2'-Bipyrimidyl-Platinum Complex: Evidence for a Bidentate Ligand Dissociation Mechanism. *Organometallics* **2016**, *35*, 1064–1069. (b) Warioba, C. S.; Jackson, L. G.; Neal, M. A.; Haines, B. E. Computational Study on the Role of Zn(II) Z-Type Ligands in Facilitating Diaryl Reductive Elimination from Pt(II). *Organometallics* **2023**, *42*, 16–26.
- (19) For an example on proton-mediated electronic mudulation, see: Brennan, C.; Draksharapu, A.; Browne, W. R.; McGarvey, J. J.; Vos, J. G.; Pryce, M. T. Unexpected reversible pyrazine based methylation in a Ru(II) complex bearing a pyrazin-2'-yl-1,2,4-triazolato ligand and its effect on acid/base and photophysical properties. *Dalton Trans.* **2013**, 42, 2546–2555.
- (20) For examples on proton-mediated ligand alternation reaction, see: (a) Himeda, Y.; Onozawa-Komatsuzaki, N.; Sugihara, H.; Kasuga, K. Simultaneous Tuning of Activity and Water Solubility of Complex Catalysts by Acid-Base Equilibrium of Ligands for Conversion of Carbon Dioxide. *Organometallics* **2007**, 26, 702–712. (b) Hashiguchi, B. G.; Young, K. J. H.; Yousufuddin, M.; Goddard, W. A., III; Periana, R. A. Acceleration of Nucleophilic CH Activation by Strongly Basic Solvents. *J. Am. Chem. Soc.* **2010**, 132, 12542–12545. (c) Dixon, N. A.; McQuarters, A. B.; Kraus, J. S.; Soffer, J. B.; Lehnert, N.; Schweitzer-Stenner, R.; Papish, E. T. Dramatic tuning of ligand donor properties in (Ttz)CuCO through remote binding of H⁺ (Ttz = hydrotris(triazolyl)borate). *Chem. Commun.* **2013**, 49, 5571–5573.
- (21) Acosta-Calle, S.; Miller, A. J. M. Tunable and Switchable Catalysis Enabled by Cation-Controlled Gating with Crown Ether Ligands. *Acc. Chem. Res.* **2023**, *56*, 971–981.
- (22) van der Ham, A.; Hansen, T.; Lodder, G.; Codée, J. D. C.; Hamlin, T. A.; Filippov, D. V. Computational and NMR Studies on the Complexation of Lithium Ion to 8-Crown-4. *ChemPhysChem* **2019**, 20, 2103–2109.
- (23) Ouyang, G.-H.; He, Y.-M.; Li, Y.; Xiang, J.-F.; Fan, Q.-H. Cation-Triggered Switchable Asymmetric Catalysis with Chiral Aza-CrownPhos. *Angew. Chem., Int. Ed.* **2015**, *54*, 4334–4337.
- (24) Asada, T.; Hoshimoto, Y.; Ogoshi, S. Rotation-Triggered Transmetalation on a Heterobimetallic Cu/Al N-Phosphine-Oxide-Substituted Imidazolylidene Complex. J. Am. Chem. Soc. 2020, 142, 9772–9784.
- (25) Branzi, L.; Baron, M.; Armelao, L.; Rancan, M.; Sgarbossa, P.; Graiff, C.; Pöthig, A.; Biffis, A. Coordination chemistry of gold with *N*-phosphine oxide-substituted imidazolylidenes (POxIms). *New J. Chem.* **2019**, 43, 17275–17283.
- (26) Dorta, R.; Stevens, E. D.; Hoff, C. D.; Nolan, S. P. Stable, Three-Coordinate $Ni(CO)_2(NHC)$ (NHC = *N*-Heterocyclic Carbene) Complexes Enabling the Determination of Ni-NHC Bond Energies. *J. Am. Chem. Soc.* **2003**, *125*, 10490–10491.
- (27) Dorta, R.; Stevens, E. D.; Scott, N. M.; Costabile, C.; Cavallo, L.; Hoff, C. D.; Nolan, S. P. Steric and Electronic Properties of N-Heterocyclic Carbenes (NHC): A Detailed Study on Their Interaction with Ni(CO)₄. J. Am. Chem. Soc. 2005, 127, 2485–2495. (28) A set of (R_a) and (S_a) atropisomers of 3a were identified in the asymmetric unit of the single crystal, as bulky Al $(C_6F_5)_3$ would inhibit

the rotation of the *N*-phosphinoyl moiety upon complexation and induce N-P axial chirality. For details on complexation-induced N-P axial chirality, see: (a) Kinoshita, T.; Sakuraba, M.; Hoshimoto, Y.; Ogoshi, S. Complexation between MOTf (M=Li and Na) and N-Phosphine Oxide-substituted Imidazolylidenes via Coordination of the N-Phosphoryl Group. *Chem. Lett.* **2019**, 48, 230–233. (b) Asada, T.; Hoshimoto, Y.; Kawakita, T.; Kinoshita, T.; Ogoshi, S. Axial Chirality around N-P Bonds Induced by Complexation between $E(C_6F_5)_3$ (E=B, Al) and an N-Phosphine Oxide-Substituted Imidazolinylidene: A Key Intermediate in the Catalytic Phosphinoylation of CO_2 . *J. Org. Chem.* **2020**, 85, 14333–14341. (c) Hoshimoto, Y.; Yamauchi, Y.; Terada, T.; Ogoshi, S. Complexation-Induced N-P Axial Chirality in Sm(II) N-Phosphine-Oxide-Substituted Imidazolylidene and Imidazolinylidene Complexes. *Can. J. Chem.* **2023**, 101, 429–433.

- (29) Kumar, P S. V.; Raghavendra, V.; Subramanian, V. Bader's Theory of Atoms in Molecules (AIM) and its Applications to Chemical Bonding. *J. Chem. Sci.* **2016**, *128*, 1527–1536.
- (30) Macchi, P.; Sironi, A. Chemical bonding in transition metal carbonyl clusters: complementary analysis of theoretical and experimental electron densities. *Coord. Chem. Rev.* **2003**, 238–239, 383–412.
- (31) He, W.; Beattie, D. D.; Zhou, H.; Bowes, E. G.; Schafer, L. L.; Love, J. A.; Kennepohl, P. Direct metal—carbon bonding in symmetric bis(C—H) agostic nickel(I) complexes. *Chem. Sci.* **2021**, *12*, 15298—15307.
- (32) Bistoni, G.; Rampino, S.; Scafuri, N.; Ciancaleoni, G.; Zuccaccia, D.; Belpassi, L.; Tarantelli, F. How π back-donation quantitatively controls the CO stretching response in classical and nonclassical metal carbonyl complexes. *Chem. Sci.* **2016**, *7*, 1174–1184.
- (33) Gómez-Suárez, A.; Nelson, D. J.; Nolan, S. P. Quantifying and understanding the steric properties of N-heterocyclic carbenes. *Chem. Commun.* **2017**, *53*, 2650–2660.
- (34) Heully, J.-L.; Lindgren, I.; Lindroth, E.; Lundqvist, S.; Mårtensson-Pendrill, A.-M. Diagonalisation of the Dirac Hamiltonian as a basis for a relativistic many-body procedure. *J. Phys. B: At. Mol. Phys.* **1986**, *19*, 2799–2815.
- (35) Chang, C.; Pelissier, M.; Durand, P. Regular Two-Component Pauli-Like Effective Hamiltonians in Dirac Theory. *Phys, Scr.* **1986**, 34, 394–404.
- (36) van Lenthe, E.; Baerends, E. J.; Snijders, J. G. Relativistic regular two-component Hamiltonians. *J. Chem. Phys.* **1993**, 99, 4597–4610.
- (37) DiMucci, I. M.; Lukens, J. T.; Chatterjee, S.; Carsch, K. M.; Titus, C. J.; Lee, S. J.; Nordlund, D.; Betley, T. A.; MacMillan, S. N.; Lancaster, K. M. The Myth of d⁸ Copper(III). *J. Am. Chem. Soc.* **2019**, 141, 18508–18520.
- (38) A bonding interaction between Ni(CO)₄ and Lewis-acidic centers at the surface of γ -Al₂O₃ has already been investigated; for details, see: Rao, K. M.; Spoto, G.; Zecchina, A. IR Investigation of Ni(CO)₄ Interaction with Silicalite, ZSM-5, Zeolite-Y, and γ -Al₂O₃. Langmuir 1989, 5, 319–325.
- (39) For the isolation of a Ni complex including a CO−Li⁺ interaction, see: Hou, H.; Rheingold, A. L.; Kubiak, C. P. An Anionic Zerovalent Nickel Carbonyl Complex Supported by a Triphosphine Borate Ligand: An Ni-C≡O-Li Isocarbonyl. *Organometallics* **2005**, 24, 231–233.
- (40) For the formation of Lewis adducts between M–CO units and Lewis-acidic species, see: (a) Kotz, J. C.; Turnipseed, C. D. Evidence for the Interaction of Trimethylalane with the Oxygen Atom of a Terminal Metal Carbonyl Group. *Chem. Commun.* **1970**, 41–42. (b) Burlitch, J. M.; Petersen, R. B. Variable site of Lewis basicity of π -C₅H₅W(CO)₃⁻ in complexes with (C₆H₅)₃In and (C₆H₅)₃Al. *J. Organomet. Chem.* **1970**, 24, C65–C67. (c) Petersen, R. B.; Stezowski, J. J.; Wan, C.; Burlitch, J. M.; Hughes, R. E. A Novel Metal-Carbonyl-Metal Bonding System. Synthesis and Stereochemistry of Al[W(CO)₃C₅H₅]₃(C₄H₈O)₃. *J. Am. Chem. Soc.* **1971**, 93, 3532–3533. (d) Ulmer, S. W.; Skarstad, P. M.; Burlitch, J. M.;

Hughes, R. E. Synthesis, Stereochemistry, and Bonding of Tetrapyridinemagnesium (II) cyclopentadienyltricarbonylmolybdate(-I)) and Related Compounds. J. Am. Chem. Soc. 1973, 95, 4469-4471. (e) Darensbourg, M. Y.; Jimenez, P.; Sackett, J. R.; Hanckel, J. M.; Kump, R. L. Interactions of $CpM'(CO)_3^-$ (M' = Cr, Mo, W) with Cations: Effects on CO Exchange and RX Addition Reactions. J. Am. Chem. Soc. 1982, 104, 1521-1530. (f) Zhuang, B.; Sun, H.; He, L.; Zhou, Z.; Lin, C.; Wu, K.; Huang, Z. Synthesis, structure and possible formation pathway of a novel cobalt carbonyl compound with new type B-O bond, Co(CO)₃(PPh₃)₂BEt₃, and mixed metal Co-Fe-S cluster with possible nonlinear optical property, [Et₄N][CoFe₂(CO)₈S(PPh₃)]. J. Organomet. Chem. 2002, 655, 233-238. (g) García, F.; Hopkins, A. D.; Kowenicki, R. A.; McPartlin, M.; Rogers, M. C.; Silvia, J. S.; Wright, D. S. Syntheses and Structure of Heterometallic Complexes Containing Tripodal Group 13 Ligands $[RE(2-py)_3]^-$ (E = Al, In). Organometallics 2006, 25, 2561-2568. (h) Fuchs, J.; Irran, E.; Hrobárik, P.; Klare, H. F. T.; Oestreich, M. Si-H Bond Activation with Bullock's Cationic Tungsten(II) Catalyst: CO as Cooperating Ligand. J. Am. Chem. Soc. 2019, 141, 18845-18850. (i) Tinnermann, H.; Sung, S.; Csókás, D.; Toh, Z. H.; Fraser, C.; Young, R. D. Alkali Metal Adducts of an Iron(0) Complex and Their Synergistic FLP-Type Activation of Aliphatic C-X Bonds. J. Am. Chem. Soc. 2021, 143, 10700-10708. (j) Fritz, M.; Demeshko, S.; Würtele, C.; Finger, M.; Schneider, S. Linearly Bridging N2 versus CO: Chemical Bonding and Spin-Controlled Reactivity. Eur. J. Inorg. Chem. 2023, 26, No. e202300011.

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