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Metal-Ligand Cooperation Facilitates Bond Activation and Catalytic Hydrogenation with Zinc Pincer Complexes

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ABSTRACT: A series of PNP zinc pincer complexes capable of bond activation via aromatization/dearomatization metal-ligand cooperation (MLC) were prepared and characterized. Reversible heterolytic N-H and H-H bond activation by MLC is shown, in which hemilability of the phosphorus linkers plays a key role. Utilizing this zinc pincer system, base-free catalytic hydrogenation of imines and ketones is demonstrated. A detailed mechanistic study supported by computation implicates the key role of MLC in facilitating effective catalysis. This approach offers a new strategy for (de)hydrogenation and other catalytic transformations mediated by zinc and other main group metals.

INTRODUCTION

In moving towards a more sustainable future, it is imperative to develop "green" chemical transformations that are both economical and free of waste generation. The most successful metals employed in catalytic reactions have historically been the noble metals, which typically suffer from high price, low abundance and toxicity. Thus, in the past two decades, there has been a notable increase in efforts to develop both organocatalysts and catalysts based on main group metals, which are generally more abundant and cost effective alternatives to the noble metals. Furthermore, significant effort has focused on developing transformations that are atom-efficient or generate byproducts that are benign or useful in their own right (e.g. H:O and H:).

With respect to utilizing molecular main group metal complexes for catalytic transformations, one common impediment is redox incoherence, often nullifying their ability to undergo the fundamental organometallic reactions of oxidative addition and reductive elimination (Figure 1). For example, in homogeneous zinc, magnesium and calcium catalyzed reactions, the metal is almost exclusively limited to the M2+ oxidation state, and thus productive cycles depend on only sigma bond metathesis and/or insertion steps which do not alter the oxidation state of the metal. Nonetheless, utilizing this paradigm, complexes of these metals and other main group metals have demonstrated versatile catalytic applications including hydroamination, polymerization, hydrosilylation, hydroboration and hydrogenation in which metal hydride intermediates are often speculated (and occasionally isolated). Zinc, in particular, has shown activity for all of these transformations, and is a promising metal due to its low cost, relatively high abundance and low toxicity.

Our group is interested in leveraging metal-ligand cooperation (MLC), in which bond activation occurs across both metal and ligand, to effect transformations in an atom-economical, waste-free fashion. The MLC approach has long been demonstrated for advancements with the noble metals, and more recently with more abundant first row transition metals, for reversible bond activation resembling oxidative addition and reductive elimination (Figure 1). Importantly, the metal center does not change oxidation state in this process, and as a result, this activation mode could also be potentially operative with redox-inefficient main group metals.

Figure 1. Top: Bond activation via oxidative addition/reductive elimination compared to bond activation by metal-ligand cooperation. Bottom: Nominal coordination modes of the PNP ligand to a generic metal (M) in which the ligand is an aromatized (La) neutral ligand versus a dearomatized (L:X) anionic ligand.

With respect to aromatization/dearomatization MLC with main group metals, Braunstein, Danopoulos and co-workers characterized a family of dearomatized lithium and potassium complexes, but bond activation with these complexes has not yet been demonstrated. Most notably, Berben and co-workers have shown that aluminum complexes supported by a tridentate bis(imino)pyridine ligand can activate N-H and O-H bonds across the aluminum center and ligand. Other examples with Group 13 metals have also been introduced and explored with similar application. In the case of zinc, we are only aware of reports in which the ligand itself is implicated as a proton or electron reservoir, and very recently,
MLC was proposed to facilitate methanol activation en route to a characterized zinc methylcarbonate species in a novel CO₂ reduction process. Nonetheless, direct observation of reversible H-X bond activation across any main group metal and ligand via aromatization/dearomatization is unknown, and main group metal-ligand cooperation remains an undeveloped area.

Based on our previous efforts, we hypothesized that it should be possible to synthesize pincer complexes of main group metals, such as zinc, that could reversibly activate bonds by MLC in which the oxidation state of the metal remains unchanged. If so, zinc-ligand cooperation could be employed to catalyze useful transformations, such as (de)hydrogenation reactions, which typically occur at a transition metal center. Herein, we establish that both aromatized and dearomatized zinc complexes supported by the PNP ligand (Figure 1) can be prepared and isolated in a straightforward manner. We demonstrate the ability of this system to heterolytically cleave and form H-H and N-H bonds. Finally, this new mode of bond activation by zinc is applied towards the additive-free, catalytic hydrogenation of imines and ketones.

RESULTS AND DISCUSSION

Access to a dearomatized zinc pincer complex is readily achieved via the reaction of the PNP₄⁺⁻⁻ pro-ligand with dimethylzinc at 140°C in toluene to generate PNP₄⁺⁻⁻ZnMe (1) concurrent with the loss of methane (Scheme 1). This simultaneous deprotonation and complexation approach is analogous to the procedure employed to prepare lithium and potassium pincer complexes, and is not uncommon in the synthesis of zinc alkyl and amido species. In contrast to procedures employed in the preparation of traditional dearomatized transition-metal pincer complexes, this approach, the addition of base is obviated by the basic nature of dimethylzinc itself, which is capable of deprotonating the side arm of the ligand. Upon mixing, the colorless ligand and dimethylzinc starting materials form a bright yellow solution, which upon heating evolves gas and exhibits an orange/red color, atypical of Zn²⁺ compounds but quite typical of dearomatized PNP pincer complexes. From monitoring the reaction progress with ¹H and ³¹P{¹H} NMR spectroscopy, we propose that the room temperature yellow intermediate is formed by coordination of the PNP₄⁺⁻⁻ ligand to ZnMe₂, and subsequent heating results in the loss of methane and formation of the orange/red complex 1 (see Supporting Information). The dearomatized compound, PNP₄⁺⁻⁻ZnMe, exhibits several characteristic signals in solution by NMR spectroscopy. For example, in toluene-d₈, multiplets at -0.02 ppm in the ¹H NMR spectrum and 3.47 ppm in the ¹H NMR spectrum indicate the presence of a zinc methyl moiety coupled to two inequivalent phosphorus atoms. In addition, the inequivalent arms of the pincer ligand can be distinguished by both ¹H (doublet at 3.53 ppm and broad singlet at 2.62 ppm) and ¹C{¹H} (doublet at 57.43 ppm and doublet of doublets at 31.19 ppm) NMR spectra, associated with the deprotonated and protonated arms, respectively, as confirmed by HSQC and DEPTQ NMR experiments. Most evidently, the ³¹P{¹H} NMR of 1 exhibits two distinct doublets (40.75 ppm and 5.74 ppm), with J₀ = 18 Hz, indicative of the inequivalent phosphorus atoms, with similar shifts as the known lithium and potassium complexes.

Utilizing the same protocol, a dearomatized zinc ethyl compound, PNP₄⁺⁻⁻ZnEt (2), was also prepared (Scheme 1). The zinc ethyl complex forms via a similar pathway, and its spectroscopic features essentially match that of 1 (see Supporting Information Table S2).

Single crystals of PNP₄⁺⁻⁻ZnR (R = Me, Et) and PNP₄⁺⁻⁻Zn(N(SiMe₃))₂ were grown by cooling saturated pentane solutions to -32°C, and the molecular structures were determined by X-ray crystallography (Figure 2), confirming the formation of a monomeric, dearomatized zinc pincer complex in each case. The zinc center in 1 and 2 is pseudotetrahedral, distorting the PNP ligand quite substantially from its typically planar conformation. Supporting the spectroscopic data, conclusive evidence of the dearomatized nature of the ligand is clear in each case: (i) the hydrogen atoms on the pincer arms were found and refined, and (ii) the C-C distances are indicative of double bond character on one of the arms and single bond character on the other arm (see Figure 2 and Supporting Information Table S1). Also in agreement with the spectroscopic data, the Zn-P distance of the ligand arm that remains protonated increases 1 < 2 < 3 < 4 (2.675Å, 2.965Å and 4.914Å, respectively); in the case of 3, the second phosphorus arm is clearly unbound in the solid state, rendering the zinc center three-coordinate. As such, it is understandable that the ³¹P{¹H} NMR spectrum of 1 and 2 exhibit two doublets (from P-P coupling), whereas that for 4 exhibits two singlets, indicating an open arm in solution. Though a coordination number of four is common for zinc, three-coordinate zinc is less common but has been observed.

While the preparation of organozinc (RZnX) species is formally an oxidative addition of an organic RX to Zn°, these species are typically stoichiometric reagents; thus, in moving towards catalytic applications, the development of molecular and recyclable zinc compounds that can perform bond activation is significant. We previously reported the activation of N-H bonds mediated by MLC with a dearomatized ruthenium complex. We found that electron-deficient anilide derivatives reacted via aromatization/dearomatization MLC to afford ruthenium anilido complexes.
The PNP\textsubscript{tBu} pincer ligand is detached from the metal center, rendering complex S tetrahedral. A tetrahedral geometry is quite common for four-coordinate zinc centers, and the conformation also resembles the structure found for PNP\textsubscript{tBu}ZnCl (see Supporting Information); however, the NMR spectroscopy data indicates an equivalence of the side arms in solution consistent with a 5-coordinate zinc center, hinting that there may be a difference between the solid and solution state behavior for both 3 and S. To probe this difference, a variable temperature NMR experiment was performed for 5. At low temperature, inequivalence of the methylene and tert-butyl groups on the side arms by \textsuperscript{1}H NMR (Figure 2) and the phosphorus linkers by \textsuperscript{31}P\textsubscript{1H} NMR (see Supporting Information) could be observed. Decoalescence of the signals is observed around 253K and 238K in the \textsuperscript{1}H NMR and \textsuperscript{31}P\textsubscript{1H} NMR spectra, respectively. We also performed the analogous experiment with compound 3 and observe the same phenomenon (see Supporting Information). These experiments confirm the labile nature of the phosphorus linkers in solution and are significant in that the hemilability of the side arm of PNP\textsubscript{tBu} complexes has been proposed, but, to our knowledge, has never been demonstrated experimentally. This finding is meaningful in that hemilability in pincer complexes has been extensively studied\textsuperscript{47-51} and is believed to be significant in catalytic mechanisms of related compounds.\textsuperscript{52}

![Figure 2. Molecular structures of complexes (1), (2), (4) and (5). Selected hydrogen atoms omitted for clarity. Some groups displayed as wireframe for clarity. Select bond lengths (Å) and angles (°): (1): Zn1-C24 1.9969(18), Zn1-P1 2.4252(5), Zn1-P2 2.6750(5), P2-C7 1.8646(18), C7-C6 1.510(3), P1-C1 1.7608(18), C1-C2 1.387(2). (2): Zn1-C24 1.9731(14), Zn1-P1 2.4267(4), Zn1-P2 2.965, P1-C1 1.7491(15), C1-C2 1.394(2), P2-C7 1.8684(14), C7-C6 1.5043(19), N1-Zn1-P1 85.36(3). (4): Zn1-N2 1.8849(18), Zn1-P1 2.3369(6), Zn1-P2 4.914, P1-C1 1.746(2), C1-C2 1.402(3), P2-C7 1.866(2), C7-C6 1.508(3), P1-Zn1-N1 87.56(5), N1-Zn1-N2 127.36(7). (5): Zn1-N2 1.9954(14), Zn1-P1 2.4606(4), Zn1-C30 1.9872(18), P1-C1 1.8367(17), C1-C2 1.509(2), P2-C7 1.8648(18), C7-C6 1.510(2). See Supporting Information for molecular structure of (3).]

The nature of ammonia, aniline and other amines, reversible addition was observed. Related palladium and copper chemistry has also been explored.\textsuperscript{43} We began our reactivity studies with the room temperature addition of the electron-poor aniline derivative, H\textsubscript{2}NAr (Ar = 2-chloro-4-nitrophenyl), to 1, which resulted in the immediate formation of the aromatized complex, PNP\textsubscript{tBu}Zn(\textsubscript{Me})NHAr (S), in which N-H activation has occurred across both the metal center and the ligand side-arm (Scheme 1).\textsuperscript{44} Notably, the oxidation state of the zinc center remains +2, in that the pincer ligand itself reassumes a neutral charge, with both the methyl and anilido moieties serving as X-type ligands.\textsuperscript{45} In solution, NMR spectroscopy provides conclusive evidence of the formation of 5, indicating symmetrization of the pincer side arms. Specifically, the \textsuperscript{31}P\textsubscript{1H} NMR changes drastically from two doublets for 4 to a broad singlet for 5 and the \textsuperscript{1}H NMR displays a doublet for four hydrogen atoms at 2.99 ppm, demonstrating solution-state equivalence of the methylene side arms. NMR data suggests that the addition across the side arm is irreversible in the case of this weakly basic amine.

![Figure 3. Select region of the \textsuperscript{1}H NMR spectrum of 5 at variable temperatures. Dashed regions indicate the observable inequivalence of the methylene hydrogen atoms (left) and the tert-butyl hydrogen atoms (right) as temperature decreases (from bottom to top). Peak at 2.08 is toluene; peak below 0 is Zn(\textsubscript{Me}).

Most interestingly, addition of the more basic N-benzylaniline (PhN(\textsubscript{H})Bn) to 1 under ambient conditions resulted in an observable equilibrium. Analysis of a mixture of PNP\textsubscript{tBu}ZnMe (1) and PhN(\textsubscript{H})Bn by \textsuperscript{1}H and \textsuperscript{31}P\textsubscript{1H} NMR spectroscopies indicates the formation of an equilibrium mixture containing three major species: free amine, deaeromated zinc methyl complex with coordinated amine (6),\textsuperscript{43} and the N-H activated compound, PNP\textsubscript{tBu}Zn(\textsubscript{Me})NPhBn (7) (Scheme 2). Compound 7 is characterized as a component of the equilibrium mixture by NMR spectroscopy. Specifically, diagnostic signals at 23.19 ppm in the \textsuperscript{31}P\textsubscript{1H} NMR spectrum for the solution-state equivalent phosphorus atoms and at 4.33 ppm in the \textsuperscript{1}H NMR spectrum for the CH\textsubscript{2}Ph moiety in the amido ligand are observed. In addition, a small amount (<5%) of a species tentatively identified as PNP\textsubscript{tBu}ZnNPPhBn is formed. This species exhibits \textsuperscript{31}P\textsubscript{1H} NMR characteristic signals similar to 1 and...
is presumably formed from protolytic cleavage of the Zn-Me bond. The equilibrium between complexes 1, 6 and free amine and complex 7 could be probed by NMR spectroscopy via a variable temperature NMR study. Specifically, a d₅-toluene solution of a roughly equimolar mixture of PNPMe₂ZnMe and N-benzylideneaniline was heated in increments from room temperature to 120°C, allowing the overall equilibrium between 1 and 7 to be established at each temperature. Kₑq was determined by integration of the corresponding 1H NMR spectra and a Van’t Hoff plot permitted extraction of thermodynamic quantities (see Supporting Information). Kₑq over the temperature range was found to be between 1 and 100, with an experimentally determined Kₑq of 68.8 for the equilibrium mixture at 298K, associated with a ΔG of -2.4 kcal/mol, which agrees reasonably well with our computed results (+1.1 kcal/mol at 393.15K and 1M standard states, vide infra). Observation of this equilibrium provides evidence that (i) free amine can coordinate to and then react with 1 via MLC to rearmatize the ligand and generate 7, (ii) free amine can be released by MLC to regenerate 1 and (iii) that the reactants and product are quite similar energetically, such that the equilibrium is observable in solution by NMR spectroscopy. This unprecedented reversible N-H bond activation by MLC with a main group metal resembles transition-metal-like behavior.

Scheme 2. Reactivity of 1 Towards PhN(H)Bn, H₂, and D₂, and Trapping of a Proposed Zinc Hydride Complex

Subsequently, we turned our attention to the activation of a non-polar, environmentally relevant substrate, H₂. Oxidative addition of H₂ to a transition metal is very important in several transition-metal catalyzed processes, and we have observed reversible H₂ activation by MLC with related transition metal systems with applications in (de)hydrogenation catalysis. Thus, we explored whether PNPMe₂ZnMe could potentially activate hydrogen, a challenging non-polar bond, in the same fashion as the N-H bond. Treatment of a solution of 1 in toluene with 5 bar of H₂ in a high-pressure J. Young tube resulted in no significant change, even upon gradual heating to 80°C. Heating the sample at 90°C, however, resulted in partial decomposition, with grey precipitation presumed to be Zn₃, observed lining the walls of the NMR tube, and both CH₃D and CHD were detected by 1H NMR spectroscopy (see Supporting Information). This result further suggests the activation of H₂ and (D₂) via MLC. Moreover, observable incorporation of deuterium into 1 and the 45% deuterium incorporation into the free ligand at the end of the reaction both imply that the H₂/D₂ activation is reversible. While dihydrogen has been utilized as the hydride source in the preparation of zinc hydride derivatives, we are unaware of any examples of reversible hydrogen activation by molecular zinc species. Nonetheless, it is noteworthy that the relatively low incorporation likely indicates that the decomposition pathway in the absence of imine is facile as compared to the reverse dehydrogenation process. Finally, to confirm the role of the zinc center in the H₂/D₂ activation, a control experiment was performed with the PNPMe₂ free ligand and D₂. No incorporation of deuterium into the side arms was observed when employing the same conditions (see Supporting Information).

Based on the unique activity of the main group metal zinc-based pincer complex towards H₂ cleavage, insertion and reversible activation of N-H bonds (vide supra), we posited that leveraging MLC might permit hydrogenation of polar unsaturated bonds with this system. To our knowledge, zinc catalyzed hydrogenation has been demonstrated for imines under very high H₂ pressures (68-100 bar)13,14 and ketones with limited scope and high H₂ pressure (100 bar). Reductions catalyzed by molecular zinc compounds utilizing other hydrogen sources, specifically hydroxylases and hydroboranes, have been demonstrated, including imines, ketones, aldehydes and nitriles. The lack of reports of hydrogenation, we believe, is related to the energetic cost of sigma bond metathesis to split hydrogen and release product (hence the success of other hydrogen sources with more favorable thermodynamic driving forces). Given the ability of the aromatization/dearomatization MLC approach to facilitate the reversible activation of H₂ and N-H
bonds in this pincer system, we hypothesized that these alternate pathways could potentially bypass other more energetically demanding routes with less favorable thermodynamics, thus allowing hydrogenation at milder conditions compared to previously reported studies.

Indeed, PNP\textsuperscript{tBu*}ZnMe (1) is a moderately active catalyst for the hydrogenation of a model substrate, N-benzylideneaniline (Table 1, Entry 1), at substantially lower pressures (7 bars) than the previous reports (68-80 bars) of zinc-catalyzed imine hydrogenation.\textsuperscript{1,14} Specifically, when 0.5 mmol of N-benzylideneaniline in 1.8 mL toluene was treated with 7 bars of H\textsubscript{2} in the presence of 1 (2.5 mol\%) at 120°C, the corresponding amine was observed by GC-MS and \textsuperscript{1}H NMR spectroscopy in up to 89% yield after 18 hours. Various substituted benzylideneaniline derivatives, with both electron donating and electron withdrawing groups on the phenyl ring, were also hydrogenated to their corresponding amines under similar conditions with high to excellent yields (Entries 2-6). Upon changing the R\textsuperscript{'} group on nitrogen from phenyl to alkyl moieties, higher pressure is needed to obtain substantial conversion (Entries 7-10). Interestingly, N-benzylidene-\textsuperscript{t}Bu\textsubscript{aminine was not amenable to hydrogenation under the reaction conditions, most probably due to the bulky \textsuperscript{t}Bu group (Entry 8). On the other hand, as the R\textsuperscript{'} group was changed to a less bulky alkyl group (Me, Bn), hydrogenation to the corresponding amine products was observed (Entries 7 and 9). R\textsuperscript{'} on carbon is not limited to aromatic substituents; while aldimines with aliphatic R\textsuperscript{'} are relatively elusive, we also found that N-butylbenzylamine can be prepared from the corresponding imine, N-butylidenebenzylamine, in moderate yield with a somewhat longer reaction time (Entry 10). Finally with respect to imines, we demonstrate that 1 also catalyzes hydrogenation of a ketimine, specifically phenyl-(1-phenylethylidene)amine, in good yield (Entry 11).

While several main group metals exhibit activity for imine hydrogenation, homogeneous ketone hydrogenation by main group metals is exceptionally limited. Select examples exist requiring extremely high pressures,\textsuperscript{95,65,66} the use of additives\textsuperscript{96} or that exhibit narrow substrate scope.\textsuperscript{95,65,66} Thus, it is significant that we found that 1 can catalyze the hydrogenation of ketones with aliphatic, aromatic and mixed aromatic/aliphatic R and R\textsuperscript{'} substituents under 65 bar of H\textsubscript{2} at 140°C. Specifically, benzophenone is hydrogenated to diphenylmethanol in up to 88% yield (Entry 12). Additionally, compound 1 catalyzes hydrogenation of 4-heptanone to 4-heptanol and propiophenone to 1-phenyl-1-propanol in good yield (Entries 13-14). With respect to green chemistry, utilizing zinc is inherently significant for these transformations, but also, it is important that (i) there is no need for any base or other additives and (ii) the process is 100% atom economical (as opposed to previously reported zinc reductions with hydrosilanes and hydroboranes).\textsuperscript{1,58,64}

Mechanistically, we propose a plausible cycle (Figure 4) accounting for the observations from the stoichiometric and catalytic experiments. In the first step, the arm of PNP\textsuperscript{tBu*}ZnMe (A), can open (B) allowing for the reversible activation of hydrogen by MLC to generate an aromatized zinc hydride intermediate (C). In the presence of imine or ketone, rather than decomposing, the zinc hydride can be trapped via an insertion pathway, generating the aromatized zinc amido or alkoxy complex (E) (vide supra, Scheme 2). Several pathways seem plausible for the insertion to proceed, but here we propose one concerted path (direct to E) and one stepwise path (via D, see Supporting Information). Insertion of imines or ketones into a zinc hydride bond has been previously proposed and observed.\textsuperscript{11,64,65} The aromatized zinc amido or alkoxy complex, per the stoichiometric experiments, can release product across the side arm

**Table 1. Hydrogenation of Imines and Ketones Catalyzed by 1**

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a) Standard Conditions: 0.5 mmol substrate, 1.8 mL toluene; % catalyst, H\textsubscript{2} pressure, temperature and time as specified b) Product confirmed by GC-MS c) Yield determined by \textsuperscript{1}H NMR spectroscopy of the crude reaction mixture with respect to mesitylene as an internal standard d) 3.6 mL of solvent used to improve substrate solubility e) no enantioselectivity observed f) Range for 3 independent trials
**Figure 4. Proposed Catalytic Cycle and DFT Summary** for imine and ketone hydrogenation in which P = P'Bu₃, X = Me or EC(H)(R)(R') and where E = O or NR'. Computed transition states and Gibbs free energy profile shown for X = Me; EC=(R)R' = PhN=C(H)Ph. Only key hydrogen atoms and only central carbon shown for 'Bu groups in computed structures for clarity. Gibbs free energies are in kcal/mol with respect to the starting materials (I, H₂ and PhN=C(H)Ph) at 393.15 K. See Supporting Information for complete computational details and structures of TSCE, TSCD and D.

DFT calculations at the ωB97M-V/def2-TZVPP/RJICOSX/SMD//M06-L/def2-TZVP/GD3/W06 level of theory were performed on the model system (X = Me; substrate = N-benzylideneaniline, 1M standard states except H₂; 7 atm) to provide insight into the plausibility of the proposed mechanism and to interrogate specifically whether product release via MLC (loss of amine from E to F to B) versus product release by hydrogenolysis (hydrogenation of E to C) is more favorable. In particular, computation indicates that the largest activation energy of the MLC pathway is the initial activation of hydrogen, which is overall 7.4 kcal/mol lower in energy than the highest energy barrier of the sigma bond metathesis pathway, which is the activation of hydrogen at the zinc amido complex (E to C; activation energy: 40.7 kcal/mol). Loss of amine via MLC has a barrier of 28.0 kcal/mol (E to F), which is 12.7 kcal/mol lower in energy than the hydrogenolysis product-forming step of the alternative mechanism. This difference supports the notion that both hydrogen activation and loss of amine through the MLC pathway is substantially more favorable than splitting hydrogen by sigma bond metathesis to generate product (see Supporting Information for full computational details).

To further probe these possible mechanisms, we also performed the hydrogenation experiments with zinc alkyl and amido species without the PNPᵗBu ligand, which would be unable to proceed by a MLC pathway. Dimethylzinc alone shows no activity for imine hydrogenation under the standard conditions. Zinc bis(trimethylsilyl)amide, however, showed some activity across several experimental trials; however, a mercury test suppressed much of its activity for imine hydrogenation, and several trials of the same conditions led to variant results, leading us to believe that this activity is heterogeneous. Supporting this hypothesis, under the ketone hydrogenation conditions, zinc bis(trimethylsilyl)amide alone again gave varying results, and additionally suffers from selectivity problems (see Supporting Information). On the contrary, the presence of mercury had limited to no effect on the activity of I for both imine and ketone hydrogenation. Nonetheless, considering these computational and experimental findings, a contribution of the sigma bond metathesis pathway is unlikely but possible, and in addition, we can-
not rule out other possible outer sphere mechanisms or an FLP-
like mechanism involving the labile arm, which was recently re-
ported for a copper pincer complex. In that respect, it is interest-
ing to note that the proposed pathway does not specifically impli-
cate the labile arm, indicating that PN complexes may also be
suitable for such transformations.

Finally, with respect to the resting state and active catalytic spe-
cies, we also note that during the course of the catalytic reaction,
the zinc methyl moiety (denoted X in Figure 4) may be protonat-
ed with product (supported by the observation of the proposed
complex, PNP^{ax,ax}ZnPPhBn, in the equilibrium described above),
especially as the concentration of amine or alcohol increases dur-
dering the reaction. This substitution may or may not enhance the
catalytic activity of the system. Indeed, monitoring the hydro-
genation of 5 equivalents of N-benzylideneaniline in an NMR
tube with 5 bar of hydrogen, in addition to free amine product,
during the reaction we observed 1 with bound amine (6) and 7 in
equilibrium as the major species, but also a species that may be
the activation of the free imine across the side arm which forms
initially (maximum observed concentration ca 30%) and then
decays as hydrogenation proceeds (depicted as G, but may be
one of several isomers; See Supporting Information) and ob-
servable amounts of PNP^{ax,ax}ZnPPhBn. Therefore, this finding, in
conjunction with the aforementioned equilibrium experiment
and Zn-H trapping experiment (outlined in Scheme 2) suggest
that the formation of PNP^{ax,ax}ZnPPhBn depends on the ratio of
free amine to zinc complex in the system. We hypothesized that
PNP^{ax,ax}ZnPPhBn is likely also catalytically competent. In ac-
cordance, complex 4, prepared in situ from 3 and NaN(SiMe)_3,
is also active for imine and ketone hydrogenation. Regardless of
the X substitution, in agreement with the computation, activation
of hydrogen via MLC is likely the rate-determining step of the
catalytic cycle. Further supporting this notion, we observe an
approximate kinetic isotope effect of 1.71 when the model imine
hydrogenation is performed in a high-pressure NMR tube with a
mixture of H_2 and D_2 (See Supporting Information).

CONCLUSION

In conclusion, a series of dearomatized zinc pincer complexes
supported by the PNP^{ax,ax} ligand were prepared and characterized.
The reactivity of PNP^{ax,ax}ZnMe was explored, allowing for the
unique observation of reversible N-H and H-H bond activation
by MLC by a main group metal complex. Mechanistic studies
indicate that hemilability of the phosphorus arms is key in allow-
ing for this novel bond activation by zinc. Utilizing this zinc pin-
cer system in catalysis, imines and ketones have been hydrogenated
via a mechanism in which aromatization/dearomatization
MLC was found to play a role in key steps of the catalytic cycle:
H_2 activation and product release. Thus, leveraging MLC shows
significant promise in advancing main group metal
(de)hydrogenation catalysis. The study of further bond activa-
tions and catalytic applications of this system is currently under-
way, with the aim of broadening the capabilities of main group
catalysts.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS
Publications website.

General Information, Experimental Details, Computational Details
(PDF)

X-Ray Data for Compounds 1-5 (ZIP/CIFs)

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Notes

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(65) With respect to the insertion, other concerted, stepwise, innersphere and outersphere pathways are plausible. Zn(H)Me could also dissociate from the ligand prior to the insertion. This step is not rate determining, so only some possibilities were explored herein.


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Table of Contents Graphic

[Diagram of catalytic hydrogenation process]

- Aromatization/Dearomatization
- Reversible Bond Activation
- Ligand Hemilability
- Catalytic Hydrogenation

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