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Rare-Earth Supported Nickel Catalysts for Alkyne Semihydrogenation: Chemo- and Regioselectivity Impacted by the Lewis Acidity and Size of the Support

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Abstract: Bimetallic catalysts of nickel(0) with a trivalent rare-earth ion or Ga(III), NiML₃ (where L is [ⁱPr₂PCH₂NPh]⁻, and M is Sc, Y, La, Lu, or Ga), were investigated for the selective hydrogenation of diphenylacetylene (DPA) to (E)-stilbene. Each bimetallic complex features a relatively short Ni-M bond lengths, ranging from 2.3395(12) Å (Ni–Ga) to 2.5732(4) Å (Ni–La). The anodic peak potentials of the NiML₃ complexes vary from -0.48 V to -1.23 V, where the potentials are negatively correlated with the Lewis acidity of the M(III) ion. Three catalysts, Ni-Y, Ni-Lu, and Ni-Ga, showed nearly quantitative conversions in the semihydrogenation of DPA, with NiYL₃ giving the highest selectivity for (E)-stilbene. Initial rate studies were performed on the two tandem catalytic reactions: DPA hydrogenation and (Z)stilbene isomerization. The catalytic activity in DPA hydrogenation follows the order: Ni-Ga > Ni-La >Ni–Y > Ni–Lu > Ni–Sc. The ranking of catalysts by (Z)-stilbene isomerization initial rates is: Ni–Ga >>> Ni–Sc > Ni–Lu > Ni–Y > Ni–La. In operando ³¹P and ¹H NMR studies revealed that in the presence of DPA, the Ni bimetallic complexes supported by Y. Lu, and La form the Ni(η^2 -alkyne) intermediate. (η^2 -PhC = CPh)Ni($^{1}Pr_{2}PCH_{2}NPh$)₂M($\kappa^{2}-^{1}Pr_{2}PCH_{2}NPh$). In contrast, the Ni–Ga resting state is the Ni($\eta^{2}-H_{2}$) species, and Ni-Sc showed no detectable binding of either substrate. Hence, the mechanism of Ni-catalyzed diphenylacetylene semihydrogenation adheres to two different kinetics: an auto-tandem pathway (Ni-Ga, Ni-Sc) versus temporally separated tandem reactions (Ni-Y, Ni-Lu, Ni-La). Collectively, the experimental results demonstrate that modulating a base-metal center via a covalently appended Lewis acidic support is viable for promoting selective alkyne semihydrogenation.

Introduction

The recent emergence of coordination compounds containing metal–metal bonds between transition metals (TMs) and rare earth (RE) elements, that is, the group 3 metals and the lanthanides (Ln), has inspired their use in diverse applications.¹⁻¹⁴ For example, RE–TM bonded complexes are currently being explored for single-molecule magnetism.¹⁵⁻¹⁶ In the realm of catalysis, multimetallic cooperativity between rare earth ions and transition metals can elicit beneficial activity in both heterogeneous¹⁷⁻¹⁸ and cluster¹⁹⁻²² systems by affecting the catalyst stability, electronics, and/or substrate binding affinity. Even so, direct RE–TM bonds, especially those involving lanthanides, is rarely seen in homogeneous catalysis. Relevant precedents are

the rare-earth complexes supported by ferrocene diamide ligands that catalyze ring-opening polymerization, wherein a weak metal-metal interaction was proposed to influence catalytic activity.²³⁻²⁵ Fueled by the rise of bifunctional transition-metal catalysts with an appended Lewis acidic main group or Group 4 metal ion,²⁶⁻⁴⁰ the extension to rare earth metals seemed ripe for exploration.

Our interest in using rare earth ions as a Lewis acidic support for tuning base metal reactivity was realized with the discovery of the catalytically active Ni–Lu complex, NiLu(ⁱPr₂PCH₂NPh)₃ (1). We demonstrated that the Lu(III) promotes Ni(0) activity in olefin hydrogenation. We also showed that the least coordinately saturated (4–coordinate) Lu center, which gave the shortest Ni–Lu bond, also gave the fastest catalytic rates.⁴¹ Because this work was the first example of a directly bound lanthanide ion being used as promoter for molecular base-metal catalysis, we became interested in exploring the catalytic reactivity of other nickel rare-earth bimetallic complexes, and also turned our attention to the semihydrogenation of alkynes.

The semihydrogenation of alkynes to alkenes is an industrially important transformation in bulk and fine chemical manufacturing.⁴²⁻⁴³ However, due to the competitive formation of both (*E*)- and (*Z*)-isomers, as well as the overhydrogenated byproduct, the main challenge is to fully control both the chemo- and stereoselectivity of this reaction.⁴⁴ Classic alkyne semihydrogenation catalysts, such as Lindlar's catalyst⁴⁵ or the Schrock–Osborn catalyst,⁴⁶ can produce (*Z*)-alkenes efficiently, though careful control of reaction conditions is sometimes necessary to avoid overhydrogenation. The stereocomplementary formation of (*E*)- alkenes using H₂ as a reductant is significantly more challenging. As such, well-defined semihydrogenation catalysts that operate under an atmosphere of H₂ are surprisingly limited in number.⁴⁶⁻⁵¹

With rare exceptions,⁵²⁻⁵⁶ (*E*)-alkenes are most commonly formed via an initial hydrogenation of the alkyne to the (*Z*)-alkene, followed by *cis*-to-*trans*-alkene isomerization.⁵⁷⁻⁶⁴ Within a broader catalytic context, the employment of non-precious metal catalysts has attracted intense interest, not only for their affordability and sustainability, but also for their potential to provide unprecedented reactivities and selectivities.⁶⁵⁻⁶⁶ Several (*E*)-selective semihydrogenation catalysts based on Co and Fe have been reported, but common drawbacks include limited substrate scope, the need for a high operating temperature and/or lack of overall selectivity.^{57-59,63} Recently, the first example of Ni-mediated semihydrogenation using H₂ as a reductant was reported using a mixture of Ni(NO₃).6H₂O and triphos.⁶⁰ However, forcing reaction conditions (120°C, 30 bar) were needed to achieve high yields and selectivity.

In this work, we have synthesized and structurally isolated an isostructural series of Ni(0)–RE(III) complexes, NiRE($^{i}Pr_{2}PCH_{2}NPh$)₃ (where RE is Sc, Y, and La). In conjunction with the previously reported Ni–Lu complex (1), and NiGa($^{i}Pr_{2}PCH_{2}NPh$)₃ for comparison with a non-rare earth support, the catalytic utility of the Ni bimetallic complexes toward the (*E*)-selective semihydrogenation of internal alkynes using H₂ at moderate pressure and temperature is presented.

Results

Synthesis and Characterization of ML₃ and NiML₃. Scheme 1 illustrates the two synthetic routes used to prepare the NiML₃ series, where L is [ⁱPr₂PCH₂NPh]⁻. In the first route, the rare-earth metalloligands were isolated prior to metallation with Ni(0). The metalloligands, $ScL_3(2)$ and $YL_3(3)$, were obtained as white powders in good yields (85% and 88% yields, respectively) from the reaction of the corresponding MCl₃ (M = Sc or Y) and 3 equiv of the deprotonated ligand, Li[$^{1}Pr_{2}PCH_{2}NPh$]. The La(III) metalloligand, LaL₃ (4), was obtained from the reaction of La[N(TMS)₂]₃ and 3 equiv of Pr_2PCH_2NHPh in 70% yield. In the second synthetic approach, the metalation sequence was flipped by installing the Ni(0)center first. From the reaction of 3 equiv of $Li[Pr_2PCH_2NPh]$ with Ni(COD)₂, where COD = 1,5cyclooctadiene, the Ni(0) complex, Ni(Li·L)₃ (5), was isolated as a purple solid (90% yield). For complexes 2, 4, and 5, the corresponding ${}^{31}P{}^{1}H$ NMR spectra each display a singlet at -18.8, -12.6, and 20.1 ppm,respectively (Figure S1). The ³¹P NMR spectrum of **3** shows a doublet at -14.1 ppm with a coupling constant (J_{P-Y}) of 36.5 Hz to the $I = \frac{1}{2}$ ⁸⁹Y nucleus. The ¹H{³¹P} NMR spectrum of **2** has 10 peaks, which is consistent with C_3 symmetry: 3 aryl protons, 2 diastereotopic methylene protons, 2 unique methine protons, and 3 methyl peaks that integrate to 18H, 9H, and 9H. The 2:1:1 ratio supports the assignment of P(CHMe₂)(CHMe'Me"), where one of the two ⁱPr substituents displays hindered P-C bond rotation (Figure S4). On the other hand, the ${}^{1}H{}^{31}P{}$ NMR spectra of 3, 4, and 5 each display a singlet for the methylene protons. These complexes are presumably more fluxional, such that the averaged solution structure exhibits C_{3v} symmetry (Figures S6, S8, and S10).



Scheme 1. Synthetic pathways of heterobimetallic NiML₃ complexes 6–9, where M = Sc, Y, La, and Ga.

The heterobimetallic complexes NiScL₃ (6), NiYL₃ (7), and NiLaL₃ (8) were isolated as bright red to orange crystals from the reaction of Ni(COD)₂ with 2, 3, and 4, respectively (crystalline yields 75 to 80%). Further, addition of GaCl₃ to 5 afforded NiGaL₃ (9) after workup as a brown solid (60% yield). The NMR spectra of all these NiML₃ compounds display a single ³¹P{¹H} resonance and a single ¹H{³¹P} resonance for the methylene protons, consistent with C_{3v} symmetry in solution (Figures S12–S15, S18, S19, and S21). Of note, the ³¹P{¹H} NMR shifts in aromatic solvents for the Ni–rare earth series vary subtly (range: –0.8 ppm for 1 to 3.1 ppm for 6, Figure S12), whereas the resonance of Ni–Ga 9 appears significantly more downfield at 31.8 ppm.

Previously, we noted that adding THF to a toluene- d_8 solution of NiLuL₃ (1) generated an equilibrium mixture of 1 and 1–THF, in which a single molecule of THF binds to the Lu center.⁴¹ Complete THF binding resulted in a significant 11-ppm downfield shift of the ³¹P NMR resonance. Herein, a similar behavior was observed for 7 (Δ (³¹P δ) = +10 ppm) and to a smaller extent for 8 (Δ (³¹P δ) = +2.5 ppm at rt, + 6.6 ppm at -83 °C), suggesting that 7–THF and 8–THF are accessible (Figure S13 and S100). By comparison, the change in the ³¹P δ for 6 or 9 is negligible when dissolved in THF- d_8 versus that in toluene- d_8 (Δ (³¹P δ) < 0.3 ppm). These findings indicate that THF binding to a NiML₃ complex can be controlled by the size of M, where a sufficiently large M center (Lu, Y, and La) allows the solvent to bind.

To probe the impact of the M(III) supports on the electronics at Ni, a cyclic voltammetry (CV) study of the NiML₃ series was performed. The cyclic voltammograms were collected at 100 mV/s in 0.1 M [*n*Pr₄N][BAr^F₄]/1,2-difluorobenzene electrolyte solutions (where Ar^F is 3,5-bis(trifluoromethyl)phenyl) and internally referenced to the [FeCp₂]^{+/0} potential. Each NiML₃ complex showed an irreversible Ni(I/0) oxidation process, with the anodic peak potential (E_{pa}) shifting overall by 0.75 V (Figure S27). The E_{pa} values increase in the order: -1.23 V (for 7) < -1.17 V (8) < -1.00 V (1)⁴¹ < -0.57 V (6) < -0.48 V (9). The Ni(I/0) peak potentials in this NiML₃ series correlate well with the Lewis acidity of the M(III) ion, which is parameterized by the p K_a values of the aquo species [M(OH₂)₆]³⁺ (R_{adj}^2 = 0.89, Table S3, Figure S29).^{67.70} This is in accord with the rationale that more Lewis acidic M(III) ions will withdraw electron density from Ni to a greater extent, which leads to more positive Ni(I/0) peak potentials. Of interest to us, the peak potentials showed a worse correlation with size of the M(III) ion using the Shannon M(III) ionic radii (R_{adj}^2 = 0.77, Table S3),^{39,71}which suggests that the flexible nature of L does not accentuate size effects of the M(III) support on the Ni electronics.⁷²

X-ray Crystallography. Single-crystal X-ray diffraction studies were conducted on all monometallic and bimetallic compounds. The solid-state structures and geometrical parameters for monometallic 2 - 5 are provided in the SI (Figure S30–S33, Table S4). The solid-state structures of the bimetallic Ni–M complexes 6-9 are shown in Figure 2, with selected bond distances and other geometrical parameters in Table 1. The

NiML₃ complexes are all isostructural and possess approximate C_3 symmetry. By examining the structures of **1** and **6** – **9**, the impact of M on various bond lengths was assessed. Of note, the Shannon ionic radii of the M(III) ions vary by 0.41 Å, with Ga(III) possessing the smallest ionic radius (0.620 Å) and La(III) the largest (1.032 Å).⁷³ Both the M–N and Ni–M bonds in **1** and **6** – **9** adhere to the same trend: these bonds elongate according to M = Ga < Sc < Lu < Y < La. This accords well with the expected trend based on M(III) radii (Figures S36–S38, Table S7). In comparing Ni–Ga **9** and Ni–La **8**, which represent the two M(III) size extremes, the M–N and Ni–M bond distances differ by 0.51 Å and 0.23 Å, respectively. On the other hand, the Ni–P bond lengths do not substantially change ($\Delta = 0.03$ Å).



Figure 2. Molecular structures of 6 - 9 shown at 50% thermal ellipsoid probability. Hydrogen atoms and lattice solvent molecules have been omitted for clarity. The average bond lengths (Å) are shown. Atom colors: Sc, yellow; Y, light blue; La, purple; Ga, red; Ni, pink; P, orange; N, blue; C, gray.

 Table 1. Geometrical Parameters, Including Bond Lengths (Å)

 and Angles (deg), for 6–9

1	6	7	8	9 ^a	
0.861 (Lu)	0.745 (Sc)	0.900 (Y)	1.032 (La)	0.620 (Ga)	
7.90	5.11	8.43	9.06	2.60	
2.4644(2)	2.3569(7)	2.4823(3)	2.5732(4)	2.3395(8)	
0.91	0.91	0.91	0.89	1.00	
2.2188(7)	2.2184(2)	2.2268(6)	2.2352(5)	2.2027(3)	
2.213(2)	2.102(1)	2.258(1)	2.429(1)	1.9181(5)	
359.40(1)	359.53(1)	354.81(10)	359.74(1)	357.57(3)	
	1 0.861 (Lu) 7.90 2.4644(2) 0.91 2.2188(7) 2.213(2) 359.40(1)	160.861 (Lu)0.745 (Sc)7.905.112.4644(2)2.3569(7)0.910.912.2188(7)2.2184(2)2.213(2)2.102(1)359.40(1)359.53(1)	1670.861 (Lu)0.745 (Sc)0.900 (Y)7.905.118.432.4644(2)2.3569(7)2.4823(3)0.910.910.912.2188(7)2.2184(2)2.2268(6)2.213(2)2.102(1)2.258(1)359.40(1)359.53(1)354.81(10)	16780.861 (Lu)0.745 (Sc)0.900 (Y)1.032 (La)7.905.118.439.062.4644(2)2.3569(7)2.4823(3)2.5732(4)0.910.910.910.892.2188(7)2.2184(2)2.2268(6)2.2352(5)2.213(2)2.102(1)2.258(1)2.429(1)359.40(1)359.53(1)354.81(10)359.74(1)	

Σ N–M–N	352.39(8)	357.68(10)	359.75(4)	359.83(1)	345.74(6)
M to N ₃ plane ^e	0.3559(7)	0.4427(15)	0.2994(10)	-0.06(6)	0.4243(5)
Ni to P ₃ plane ^f	0.0995(3)	0.0881(6)	0.0645(4)	-0.07(2)	0.1991(2)

^{*a*}Two molecules in the asymmetric unit. Values are averages. ^{*b*}Shannon ionic radius of six-coordinate M(III).⁷³ ^{*c*}Formal shortness ratio.⁷⁴ See main text. ^{*d*}Average value. ^{*e*}Positive value indicates M is positioned closer to Ni. ^{*f*}Positive value indicates Ni is positioned away from M.

The intermetal distances in **1** and **6** – **9** are all shorter than the sum of the two metals' covalent radii, signifying a significant Ni–M bonding interaction. To better compare the metal-metal interactions, we calculated the formal shortness ratio (r), or the ratio of the Ni–M bond distance to the sum of their respective covalent radii using Pyykkö covalent radii values.⁷⁴ The r values across the rare-earth supported bimetallic complexes are all similar, 0.89 to 0.91. This suggests that the bonding interaction between Ni and the rare earth ion is independent of the rare earth identity. We further note that the similar r values are not simply a consequence of the bridging ligand L since Ni–Ga **9** has a significantly larger r value of 1.00.

Catalyst Screening for Diphenylacetylene Semihydrogenation. The effect of varying the Lewis acid support on both catalytic activity and selectivity was investigated using the semihydrogenation of diphenylacetylene (DPA) as a model reaction (Table 2, Figures S39-S45). In a standard catalytic run, a J Young NMR tube loaded with the catalyst (2.5 mol%) and substrate (0.45 M DPA) in toluene– d_8 was charged with 4 atm of H₂ and heated to 70 °C for 24 h (4.6 atm H₂ at 70°C). A mixture of Ni(COD)₂ and 3 equiv of the protioligand was tested as a control for a monometallic nickel catalyst (entry 1). This catalyst system gave poor DPA conversion (<40%). In addition, alkyne semihydrogenation proceeded nonselectively to a $\sim 1:1$ mixture of (E)- and (Z)-stilbene with a small amount (6%) of the double alkyne insertion byproduct, 1,2,3,4-tetraphenyl-1,3-butadiene, which will simply be referred to as the (Z,Z)-diene. Mononickel complex 5 performed similarly poorly (entry 2). The catalytic performance of bimetallic 1 and 6-9 are shown in entries 3 to 7. With the single exception of Ni-Sc 6, all the Ni-M catalysts showed nearly complete DPA conversion (\geq 98%) under the standard catalytic conditions. The best performer was Ni-Y 7, which quantitatively converted DPA to (E)-stilbene (entry 4). Both Ni-Lu 1 and Ni-Ga 9 generated (E)-stilbene in high yields (91%); however, both catalysts also generated a significant amount (7-9% yield) of the undesired byproducts: 1,2-diphenylacetylene, which results from double hydrogenation, and (Z,Z)-diene.





ontry	ootolyst	DPA conversion	stilbene	E:Z stilbene	diphenylethane	Z,Z-diene ^c
entry	cataryst	(%)	(%)	ratio	(%)	(%)
1	$Ni(COD)_2 + 3 HL$	38	32	52:48	0	6
2	5 Ni(Li·L) ₃	29	28	7:21	<1	<1
3	1 (Lu)	>99	91	>99:1	7	2
4	6 (Sc)	74	58	40:60	1	15
5	7 (Y)	>99	99	>99:1	<1	<1
6	8 (La)	98	91	20:80	2	5
7	9 (Ga)	>99	92	>99:1	1	7
8^d	7–THF- d_8	>99	78	40:60	21	<1
9 ^e	$7 + Hg^0$	>99	99	>99:1	<1	<1

^{*a*} Catalytic conditions: 2.5 mol % catalyst, 0.45 M DPA, ca. 500 μ L of toluene– d_8 , 4.6 atm H₂ at 70 °C, 24 h. ^{*b*}Values (%) are averages of triplicate runs and were determined by ¹H NMR integration using trimethoxybenzene as an internal standard (error = ± 0.5%). ^{*c*}Z,Z-diene = 1,2,3,4-tetraphenyl-1,3-butadiene. ^{*d*} THF– d_8 solvent. ^{*e*} 300 equiv Hg metal.

Unexpectedly, Ni–La **8** showed an inverted preference for (*Z*)-stilbene over the (*E*)-isomer by a factor of 4. We note, however, that during the catalysis time course of **8**, we observed the gradual growth of different decomposition species, including ${}^{i}Pr_{2}PH$. Hence, a more reasonable explanation is that the *cis*-to*trans* stilbene isomerization was impeded by catalyst death. In support, all NiML₃ complexes, including **8**, catalyze the isomerization of (*Z*)-stilbene to (*E*)-stilbene under the standard catalytic conditions (SI Table S10). Among the bimetallic catalysts, Ni–Sc **6** performed the worst, as it reacted the most sluggishly and produced the most of the (*Z*,*Z*)-diene byproduct (15%).

The choice of solvent was found to be critical for the high catalytic selectivity of 7. Upon changing the solvent to THF– d_8 , the performance of catalyst 7, as the THF adduct 7-THF, suffered greatly, as evidenced by marked decline in both regioselectivity (*E*:*Z* stilbene ratio of 2:3) and chemoselectivity (alkyne versus alkene hydrogenation), as indicated by >20% yield of 1,2-diphenylethane (entry 8). Lastly, catalysis by 7 was uninhibited in the presence of excess Hg, which supports a homogeneous active species (entry 9).

Initial Rate Studies. The (*E*)-selective hydrogenation of disubstituted alkynes proceeds by an autotandem catalytic process, in which a single catalyst executes both the *syn*-hydrogenation of the alkyne and the *cis*-to-*trans* isomerization of the resulting alkene. To disentangle the two catalytic cycles, we separately conducted initial rate studies of DPA hydrogenation and (*Z*)-stilbene isomerization. While H₂ is not consumed in the latter reaction, H₂ was found to be necessary because alkene isomerization either does not proceed, or is very sluggish, in the absence of H₂ (Table S11). The initial rates were determined by monitoring the consumption of the starting material (DPA or (*Z*)-stilbene) under the standard catalytic conditions until either 10% conversion or up to 1 h. The kinetic plots are shown in Figure 3, and the initial rates are listed in Table 3. Of note, catalyst **8** did not decompose during this short time period.



Figure 3. Kinetic plots of substrate concentration (M) versus initial time (s) for: A) the hydrogenation of diphenylacetylene; and B) isomerization of (Z)-stilbene, as catalyzed by 1 and 6–9 under the standard catalytic conditions. For more details, see SI Figures S46–S56.

Table 3. Initial Rate Data for the Syn-Hydrogenation o	f DPA and (Z)-to-(E) Stilbene
Isomerization by NiML ₃ Catalysts ^a	

		Initial Rate	e (×10 ⁻⁵ M/s)
entry catalyst		$-d[DPA]/dt^b$	-d[(Z)-stilbene]/dt
1	1 (Lu)	1.71(8)	1.4(1)
2	6 (Sc)	$0.67(4)^{c}$	3.0(2)
3	7 (Y)	1.9(4)	1.1(1)
4	8 (La)	2.13(1)	0.45(3)
5	9 (Ga)	2.5(4)	40.4(7)

^{*a*}Catalytic conditions: 2.5 mol % catalyst, 0.45 M DPA or (*Z*)-stilbene, ca. 500 µL of toluene– d_8 , 4.6 atm H₂ at 70 °C, ≤ 1 h with an internal standard of 1,3,5-trimethoxybenzene (0.02 mmol). ^{*b*}In all cases except **6**, (*Z*)-stilbene was formed cleanly in the *syn*-hydrogenation, and the rate of the reaction is -d[DPA]/dt. ^{*c*}For **6**, formation of both (*Z*,*Z*)-diene and (*E*)-stilbene were observed, and the corrected rate of *syn*-hydrogenation = -d[DPA]/dt - d[Z,Z-diene]/dt. See SI for details.

In both reactions, Ni–Ga **9** was the fastest of the NiML₃ catalysts. Changing the metal support impacted the catalytic activity, especially in the (*Z*)-stilbene isomerization reaction, where the initial rates varied by nearly 90-fold. On the other hand, the initial rates for DPA hydrogenation only varied by a factor of 3. With the exception of Ni–Ga **9**, the two reactions showed opposite trends in catalytic activity for the different metal supports. For example, catalytic activity in DPA hydrogenation follows the order: **9** (M = Ga) > **8**

(La) > 7 (Y) > 1 (Lu) > 6 (Sc). By comparison, the ranking of catalysts by (*Z*)-stilbene isomerization activity is: 9 (M = Ga) >> 6 (Sc) > 1 (Lu) > 7 (Y) > 8 (La). Upon excluding 9, a *positive* correlation was found between the rate of DPA hydrogenation and the pK_a value of the corresponding aquo species $[M(OH_2)_6]^{3+}$ (Figure 4, slope = +0.37(1) x 10⁻⁵ M/s per pK_a unit; R_{adj}^2 =0.99). This finding suggests that *less* Lewis acidic supports result in greater Ni-based activity in alkyne *syn*-hydrogenation (Figure S57). As an aside, a reasonable correlation also exists between the rate of DPA hydrogenation and E_{pa} for the Ni-rare earth catalysts (R_{adj}^2 =0.94, Table S8). Also excluding 9, the plot of the (*Z*)-stilbene isomerization rate versus the pK_a value of $[M(OH_2)_6]^{3+}$ showed a *negative* correlation (Figure 4, slope = -0.62(5) x 10⁻⁵ M/s per pK_a unit; R_{adj}^2 =0.99), indicating that *more* Lewis acidic supports lead to higher activity in *cis*-to-*trans* alkene isomerization (Figure S58). However, for both reactions, the initial rates for 9 are significantly faster than the correlative values based on Lewis acidity (Figures S59–S60), suggesting the Ga^{III} supporting ion, which is by far the most Lewis acidic, results in a significant change in the nature of the reactive species and/or the mechanism.



Figure 4. Plot of initial rate of *syn*-hydrogenation of diphenylacetylene (black circles, $R_{adj}^2 = 0.99$) and *cis*-to-*trans* isomerization of (*Z*)–stilbene (blue squares, $R_{adj}^2 = 0.99$) versus pK_a of M(OH₂)₆³⁺ for catalysts **1** and **6–8**. The slope, *m*, is provided in the plots.

In operando catalytic studies. To further investigate the impact of the supporting metal, the full reaction time course from DPA to (*E*)-stilbene was monitored by ¹H NMR spectroscopy under the standard catalytic conditions. Three catalysts, Ni–Ga 9, Ni–Sc 6, and Ni–Y 7, were selected because they exhibit markedly different rates and/or selectivity. In the reaction catalyzed by Ni–Ga 9 (Figure 5A), (*Z*)-stilbene is the first intermediate to accumulate, reaching maximum concentration at ~4.5 h, before fully decaying as it converts to (*E*)-stilbene within 8 h. By comparison, the Ni–Sc 6 catalyst required a significantly longer time period (70 h) for the reaction to fully consume (*Z*)-stilbene (Figure 5B). In addition to the (*E*)-stilbene

product, the (Z,Z)-diene byproduct was formed in significant amounts, and primarily at early reaction times when DPA is present. This suggests that after DPA insertion into a Ni–H bond to form a Ni–vinyl intermediate, an off-pathway insertion of a second DPA is competitive with the on-pathway reductive elimination of (Z)-stilbene.



Figure 5. Reaction time course for the semihydrogenation of diphenylacetylene to (*E*)–stilbene under standard conditions for catalysts: A) Ni–Ga 9, B) Ni–Sc 6, and C) Ni–Y 7.

The reaction time course for Ni–Y 7 was remarkably distinct in that a temporal separation of the two catalytic reactions was clearly observed. The *cis*-to-*trans* isomerization of stilbene proceeded only *after* the DPA was almost entirely consumed in the *syn*-hydrogenation reaction (Figure 5C). Temporal separation in auto-tandem catalysis is rare, and to our knowledge, the single well-studied precedent is $[Ru(Cp)(bpy)NCCH_3]^+$, which converts terminal alkynes into the corresponding alcohols through an acetaldehyde intermediate.⁷⁵ The temporal separation of the two catalytic cycles also indicates a greater chemoselectivity for DPA relative to (*Z*)-stilbene by 7. Also of relevance, strong alkyne adsorption and its impact on kinetics had been observed for Cu/SiO₂, a heterogeneous catalyst for (*Z*)-selective semihydrogenation.⁷⁶ Lastly, we note that during the reaction time course shown in Figure 5C, the rate of stilbene isomerization exceeded that of DPA hydrogenation. This was unexpected because the initial rate studies showed the opposite order, where DPA hydrogenation was 1.7 times faster than that of stilbene isomerization. We do not currently understand the origin of the rate enhancement, but we speculate that small amounts of DPA might alter the catalyst speciation to a more reactive species for stilbene isomerization (Figure S75).

Catalyst Resting State. The reaction time course experiments were also monitored by ${}^{31}P{}^{1}H$ NMR spectroscopy to elucidate the catalyst resting state. Throughout the full reaction time course, the catalytic reaction mixture of Ni–Sc **6** only showed the presence of **6** (Figure S65). Hence, no observable binding of either H₂ or DPA was found for the Sc(III)-supported Ni site. During the *in operando* catalysis by Ni–Ga **9**, a single complex with a corresponding ${}^{31}P{}^{1}H$ singlet at 48.5 ppm was observed throughout the full

time course (Figures S71 and S77). This new species also exhibited a broad ¹H resonance at -2.03 ppm, which is typical of a Ni(η^2 -H₂) adduct (Figure S84).⁷⁷⁻⁷⁸ Also, consistent with H₂ binding, the reaction solution color bleached from a dark red-orange (9) to a light yellow-orange (9–H₂).⁷¹ Hence, the strongly Lewis acidic Ga(III) ion promotes H₂ binding at the supported Ni site. Further characterization of the independently synthesized 9–H₂ includes a T₁(min) value of 28(1) ms (500 Hz), which confirms an intact H–H bond (Figure S85).⁷⁷ An H–H distance of 0.86 Å (*c.f.* 0.74 Å in free H₂) was determined from the *J*_{H–D} value of 35 Hz for the corresponding HD adduct (Figure S86).⁷⁸ H₂ binding was reversible as evacuating a solution of 9–H₂ regenerates 9. Of note, the H₂ binding behavior of 9 contrasts that of Ni–Lu 1, where H₂ binding is so weak that it was only perceptible at low temperatures of –80 °C and under 4 atm H₂. The other rare-earth supported Ni complexes herein also show similarly weak H₂ binding akin to 1 (Figures S87–S101).

The *in operando* catalysis of 7 also showed complete formation of a new species that persisted until *t* ~3.5 h. After *t* ~3.5 h, the only species observed was 7. The ³¹P{¹H} spectrum of this new complex consists of a singlet resonance at 30.4 ppm and doublet at –11.7 ppm (J = 48.5 Hz), with an integration ratio of 2:1, respectively (Figure S67). The observed coupling makes evident that one phosphine donor is coordinated to ⁸⁹Y (>99% abundance, $I = \frac{1}{2}$). We further infer that this ligand hemilability allows for strong alkyne binding at Ni. In support, identical spectroscopic features were observed when 5 equiv DPA was added to 7 (Figure S26). Structural elucidation of this species by single-crystal X-ray diffraction confirmed its assignment as the DPA adduct, (η^2 -PhC \equiv CPh)Ni(ⁱPr₂PCH₂NPh)₂Y(κ^2 -ⁱPr₂PCH₂NPh), or 7–DPA (Figure 6). The favorable formation of 7–DPA also rationalizes the greater chemoselectivity of 7 for DPA over stilbene, as well as the temporal separation of the two tandem reactions.



Figure 6. Molecular structure of 7–DPA shown at 50% thermal ellipsoid probability; hydrogen atoms and non-coordinating solvent molecules are omitted for clarity. Only one of the two complexes from the asymmetric unit is shown for clarity. Selected interatomic distances (Å) and angles (deg.) as an average of the two crystallized molecules in the asymmetric unit: Ni–Y, 2.7031(2); Y–P, 2.9898(9); C1–C2, 1.281(5); Ph–C1–C2, 142.48. See Table S6 for more details.

In the solid-state structure of 7–DPA, two molecules crystallized in the asymmetric unit with an average Ni–Y bond length of 2.7031(2) Å, which is longer than that in 7 by 0.22 Å. The Ni–Y bonding interaction remains intact as suggested by the corresponding *r* value of 0.99. The average Y–P distance of 2.9898(9) Å is similar to those in the monometallic YL₃, **3**. Significant π -back-bonding into the alkyne is evidenced by the elongated C1–C2 distance (avg 1.281(5) Å) and the bent Ph–C1–C2 angle (avg 142.48°). The geometric changes in the DPA ligand are similar to those in (η^2 -PhC \equiv CPh)Co(ⁱPr₂PNXyl)₂ZrI(THF), which is also a relevant intermediate in DPA semihydrogenation. ⁵⁹ However, the regioselectivity of the Co-Zr catalyst is poor as stilbene was formed as a mixture of both the *cis* and *trans* isomers.⁵⁹ Lastly, the rearrangement of the [ⁱPr₂PCH₂NPh]⁻ ligand in 7–DPA is reminiscent of the hemilability behavior observed in related heterobimetallic Co-Zr complexes bearing three bridging phosphinoamide ligands.⁷⁹⁻⁸⁰

The combination of ligand hemilability and DPA binding was also observed for Ni–Lu **1** and Ni–La **8** (Figure S61). Also of interest, catalysts **1** and **8** showed a clear temporal separation during catalysis, akin to **7**. These findings suggest that the strong alkyne chemoselectivity is linked to the M(III) rare-earth supports with sufficiently large ionic radii. Hence, we hypothesize that larger supporting metal ions enhance ligand hemilability, which then favors DPA binding. In support, neither Ni–Ga **9** nor Ni–Sc **6** showed any detectable binding of DPA even when DPA was present in large excess (40 equiv) or cooled to –60 °C (Figures S76–S77). To test whether a vacant coordination site at Y(III) in **7** is vital to its catalytic performance, we monitored the catalysis when the Y(III) center is saturated by a coordinating solvent molecule, such as THF. During the catalytic time course in THF, 7–DPA was never observed (40 equiv DPA). Instead, the only observed species was the THF–adduct, 7–THF, which was independently synthesized and crystallographically characterized (Figures S34 and S73, Table S6). The deterioration of the catalyst performance for 7–THF suggests that having an open coordination site on Y to promote ligand hemilability may be critical to the enhanced performance of **7**.

Substrate Scope and Isotope-Labeling Studies. Encouraged by preliminary catalytic studies with DPA, we sought to investigate the utility of 7 toward the semihydrogenation of an assortment of substrates (Table 4, also see Figures S102–S110), using the standard catalytic conditions, unless otherwise noted. *Para*-substituted diarylacetylene featuring a boronate ester proceeded cleanly to the *trans*-diarylethene 10b with excellent regioselectivity (>99:1 *E:Z*) and in 99% yield. 3-Hexyne, a dialkyl-substituted acetylene, gave nearly exclusively *cis*-3-hexene (10c) and some trace hexane. The clean formation of the *cis*-isomer suggests that dialkyl-substituted alkenes may not undergo the critical olefin insertion into the Ni–H species during the alkene-isomerization cycle (*vide infra*). Of interest, 1-phenylpropyne yielded the *trans*-alkene 10d fairly cleanly. Terminal alkynes, on the other hand, are nonviable substrates. An attempted catalytic run with phenylacetylene resulted in catalyst decomposition and no hydrogenation product(s). If instead,

the trimethylsilyl protecting group is employed, such as 1-phenyl-2-trimethylsilylacetylene, then the *trans*alkene **10e** is formed nearly exclusively. Several 1-aryl-2-trimethylsilylacetylene substrates were also investigated, including methoxy or methyl *ortho*-substituted or *para*-substituted fluorine derivatives. For these substrates, the corresponding *trans* products (**10f** - **10h**) were obtained in good yields with excellent regioselectivity, though either higher catalyst loadings (up to 5 mol%) or longer reaction times (up to 60 h) were necessary. Lastly, a substrate bearing two internal alkynes reacted cleanly in a double semihydrogenation with excellent (*E*,*E*)-selectivity (**10i**).





^{*a*}Catalytic Conditions: 2.5 mol % 7, 0.45 M alkyne in ca. 500 μ L of toluene–*d*₈, 4.6 atm H₂, heated to 70 °C, unless otherwise noted. Conversion is reported as an average of triplicate runs (±0.5 error bars) at 24 h (unless otherwise noted), as determined by ¹H NMR integration against an internal trimethoxybenzene standard (0.02 mmol). ^{*b*}After 60h. ^{*c*}3.5 mol % 7. ^{*d*}5 mol % 7, full catalyst decomposition after 24 h, remainder was alkyne. ^{*e*}after 48h.

As previously discussed, the alkene *cis*-to-*trans* isomerization does not consume H_2 , but H_2 is required for the reaction to be kinetically relevant. An isotopic labeling experiment was performed using catalysts **7** and **9**. Using D_2 and otherwise standard catalytic conditions, (*Z*)-stilbene was converted to (*E*)-stilbene with deuterium incorporation. Also, formation of HD, and a minor amount of H_2 , was observed for catalyst **7** (Figure S79–S80). A control experiment showed that **7** does not scramble HD to produce H_2 and D_2 under the same conditions. Hence, the origin of the protons in HD and H_2 is most likely the (*Z*)-stilbene substrate. This supports the alkene isomerization proceeding via alkene insertion into a Ni–H bond and subsequent C–C bond rotation and β -hydride elimination.

Discussion

The mechanistic and kinetic studies provide a comprehensive understanding of how the choice of the M(III) support impacts the Ni–M semihydrogenation catalysis. Both the size and the Lewis acidity of the M(III) ion play significant roles. The larger M(III) ions with an ionic radius ≥ 0.86 Å all favor a catalyst resting state of the alkyne adduct, Ni(η^2 -PhC=CPh), during the first catalytic cycle. This subgroup includes the Ni–Ln catalysts, where M is Lu, Y, and La. During the second catalytic cycle when DPA is depleted, these catalysts change their resting state to the "naked" species, NiML₃, which suggests that both alkene and H₂ binding is weak. Of note, the observation of the Ni(η^2 -PhC=CPh) catalyst resting state is directly coupled to the rare observation of a temporally separated auto-tandem catalysis, which to our knowledge has not been reported for alkyne semihydrogenation. Hence, the temporal separation is a direct consequence of the nearly perfect chemoselectivity of the Ni–Ln catalysts for reacting with alkynes over alkenes when both substrates are present.

The Lewis acidity of the M(III) support can influence both the catalyst resting state and the catalytic rates. The H₂ adduct resting state, Ni(η^2 -H₂), is only observed for Ni–Ga **9**. Previously, increasingly electron-deficient Ni(0) centers were shown to bind H₂ more strongly.^{39,71} The observation of **9**-H₂ as the resting state in both catalytic cycles follows a similar logic in that Ga(III), which is by far the most Lewis acidic ion in this series, would withdraw the most Ni electron density. Catalyst **9** further stands out for its fastest initial rates in (*Z*)-stilbene isomerization in the series, by nearly two orders of magnitude. Even among the remaining Ni–RE catalysts, the *more* Lewis acidic RE ions promote *faster* (*Z*)-stilbene isomerization (Figure 4). By contrast, the *more* Lewis acidic RE ions yield *slower* initial rates in the *syn*-hydrogenation of DPA. We posit that alkyne binding is favored by a more electron-rich Ni center, which can better π -back-bond to the alkyne. Furthermore, we propose that alkyne binding is more critical to the first catalytic cycle, whereas H₂ binding and activation plays a more important role during the second catalytic cycle.





Figure 7. Proposed catalytic cycles for the nickel–catalyzed (*E*)-selective semihydrogenation of diphenylacetylene.

Figure 7 shows the proposed mechanism for the selective hydrogenation of DPA to (*E*)-stillene. Since Ni–Ga **9** and the Ni–Ln (where Ln = Lu, Y, and La) complexes feature different catalyst resting states, their mechanisms are shown separately. The Ni–Ga catalyst first binds H₂ to generate **9**–H₂, the observed resting state. Activation of H₂, followed by phosphine dissociation³⁵ and then DPA binding⁸¹⁻⁸³ leads to the formation of intermediate **I**. The bridging and terminal hydrides, HNi(μ –H)Ga, in **I** is based on the analogous HNi(μ –H)B species reported by Peters,^{28,84} as well as Thomas' HCo(μ –H)Zr intermediate⁷⁹. We note, however, that the mechanistic data does not discern whether alkyne binding or H–H cleavage proceeds first in the cycle. Next, migratory insertion of the alkyne into the Ni–H bond forms the Ni vinyl intermediate, **II**. The first catalytic cycle ends with the reductive elimination of (*Z*)-stilbene and binding of H₂ to regenerate **9**–H₂. In the second catalytic cycle, H₂ activation, followed by phosphine dissociation to allow for (*Z*)-stilbene binding, would provide intermediate **III**, which is the alkene counterpart to **I**. Migratory insertion of the alkene into the Ni–H bind forms the V. Rotation of the ethyl C–C bond followed by β -hydride elimination would generate (*E*)-stilbene, as supported by the isotope labeling studies described above. Alternatively, from **IV**, reductive elimination may proceed to form the 1,2-diphenylethane byproduct.

The Ni–Ln catalysts are proposed to operate through similar intermediates, **I'–IV'**, though their mechanism diverges in a couple key ways. First, by favoring DPA binding, the Ni–Ln catalysts operate via the Ni(η^2 -PhC=CPh) resting state, wherein the dissociated phosphine binds to the Ln ion. Second, owing to

the favorability of the alkyne adduct, (Z)-stilbene cannot enter the *cis*-to-*trans* isomerization cycle until nearly all the DPA is depleted, and Ni(η^2 -PhC=CPh) is no longer the default resting state. The isomerization cycle is similar to that described above for Ni–Ga 9, except that the "naked" NiLnL₃ species is the new resting state. Likewise, H₂ binding is proposed to precede alkene binding since the NiLnL₃ complexes all exhibit weak binding of H₂, whereas alkene binding has not yet been detected for any of these catalysts.

For Ni–Sc **6**, the catalyst resting state throughout the full catalysis is NiScL₃. Presumably, the Sc(III) support is neither sufficiently large (0.75 Å) to accommodate a stable Ni(η^2 -PhC=CPh) adduct, nor particularly Lewis acidic (p K_a 5.1) to promote strong H₂ binding at Ni. The Ni–Sc **6** showed the slowest initial rate in the *syn*-hydrogenation of DPA, which hinders its performance in the overall catalysis. Moreover, Ni–Sc generates the highest percentage of the (*Z*,*Z*)-diene byproduct (15%) in the series. The formation of the *Z*,*Z*-diene points to an alternative path whereby migratory insertion of DPA into the Ni vinyl bond in intermediate **II** occurs prior to reductive elimination.

Conclusion

The Ni-rare earth complexes add to the few first-row transition metal-rare earth complexes with direct metal-metal bonding interactions. More significantly, this study demonstrates the utility of rare earth metal ions as promoters in base-metal catalysis. Both the ionic size and Lewis acidity of the metal support are important tunable parameters for the catalytic activity and selectivity. First, the Ni electronics are significantly perturbed by variation of the M(III) support: the Ni(I/0) peak potentials were tuned overall by 0.75 V and showed a strong correlation with M(III) Lewis acidity. Second, the M(III) support can remotely tune the substrate-binding preference of the active Ni center, and consequently, determine the catalyst resting state. For example, the weak Lewis acidity and the large ionic size of the rare-earth supports are posited to be responsible for the strong Ni-alkyne binding in Ni–Y 7, Ni–Lu 1, and Ni–La 8. Presumably, these more electron-rich Ni centers can better π -back-bond into the alkyne unit, while the large rare-earth supports provide an intramolecular, open coordination site to favor hemilability of the phosphine ligand. On the other hand, strong Lewis acidity appears to be the dominant parameter in priming Ni for strong H_2 binding as only the highly Lewis acidic Ga(III) formed a stable Ni(η^2 -H₂) adduct under ambient conditions. The delineation of size and Lewis acidity in determining substrate binding is corroborated by the fact that Ni-Sc 6, which falls in the middle in terms of both Lewis acidity and size, showed neither alkyne nor H_2 binding under ambient conditions.

The stark differences in substrate binding among the Ni–M catalysts have important mechanistic and kinetic ramifications. When the Ni–M complex binds alkyne strongly, the ensuing auto-tandem catalysis shows a clear temporal separation between the two catalytic cycles. This temporal separation also indicates

a stronger chemoselectivity for alkyne over alkene reactivity. Similarly, for the Ni-rare earth catalysts, the rate of DPA *syn*-hydrogenation appears to be directly influenced by its ability to bind alkyne, such that *less* Lewis acidic RE ions yield faster initial rates in the *syn*-hydrogenation of DPA. The opposite trend was found for the rate of alkene isomerization, where *more* Lewis acidic RE ions yield faster initial rates. This may suggest that H_2 binding and activation is dependent on the Lewis acidity of the support and is the critical step in alkene isomerization for these catalysts.

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/.

X-ray crystallographic data for complexes **2–9**, **7–**THF, **7–**DPA, and **8**' (CIF) Experimental procedures, characterization and spectroscopic data (PDF)

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