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Thermodynamic and kinetic studies of H₂ and N₂ binding to bimetallic nickel-group 13 complexes and neutron structure of a Ni(η^2 -H₂) adduct†

Ryan C. Cammarota,^{‡a} Jing Xie,^{‡abe} Samantha A. Burgess,^{‡c} Matthew V. Vollmer,^a Konstantinos D. Vogiatzis,^{‡abf} Jingyun Ye,^{‡ab} John C. Linehan,^{‡c} Aaron M. Appel,^{‡c} Christina Hoffmann,^{‡d} Xiaoping Wang,^{‡d} Victor G. Young, Jr.^a and Connie C. Lu^{‡*a}

Understanding H₂ binding and activation is important in the context of designing transition metal catalysts for many processes, including hydrogenation and the interconversion of H₂ with protons and electrons. This work reports the first thermodynamic and kinetic H₂ binding studies for an isostructural series of first-row metal complexes: NiML, where M = Al (1), Ga (2), and In (3), and L = [N(o-(NCH₂PiPr₂)C₆H₄)₃]³⁻. Thermodynamic free energies (ΔG°) and free energies of activation (ΔG^\ddagger) for binding equilibria were obtained *via* variable-temperature ³¹P NMR studies and lineshape analysis. The supporting metal exerts a large influence on the thermodynamic favorability of both H₂ and N₂ binding to Ni, with ΔG° values for H₂ binding found to span nearly the entire range of previous reports. The non-classical H₂ adduct, (η^2 -H₂)NiInL (3-H₂), was structurally characterized by single-crystal neutron diffraction—the first such study for a Ni(η^2 -H₂) complex or any d¹⁰ M(η^2 -H₂) complex. UV-Vis studies and TD-DFT calculations identified specific electronic structure perturbations of the supporting metal which poise NiML complexes for small-molecule binding. ETS-NOCV calculations indicate that H₂ binding primarily occurs *via* H–H σ -donation to the Ni 4p_z-based LUMO, which is proposed to become energetically accessible as the Ni(0) → M(III) dative interaction increases for the larger M(III) ions. Linear free-energy relationships are discussed, with the activation barrier for H₂ binding (ΔG^\ddagger) found to decrease proportionally for more thermodynamically favorable equilibria. The ΔG° values for H₂ and N₂ binding to NiML complexes were also found to be more exergonic for the larger M(III) ions.

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Introduction

Developing homogeneous base metal catalysts which can activate H₂ and selectively mediate catalytic hydrogenation, H₂ oxidation, and proton reduction have been active areas of

research in recent years.^{1–5} In such processes, the strength of H₂ binding, the interplay between σ -donation (H₂ $\sigma \rightarrow M$) and π -back-donation (M \rightarrow H₂ σ^*), the resulting extent of H–H activation, and the ease of generating reactive M–H species all can play a critical role in determining catalytic activity and selectivity. H₂ activation is typically initiated by side-on binding of H₂ to form a M(η^2 -H₂) adduct, which precedes the generation of reactive M–H species *via* subsequent oxidative addition or deprotonation events.^{6–8} Despite the ubiquity of H₂ binding as a key fundamental reaction step in catalysis,⁹ limited experimental data has been reported regarding the thermodynamic and kinetic favorability of H₂ binding to transition metals, especially for the first-row transition metals.^{6–8} To the best of our knowledge, thermodynamic H₂-binding data have only been reported for a handful of first-row metal complexes: Cr(PCy₃)₂(CO)₃,^{10,11} [Mn(CO)(dppe)₂][†],¹² [Fe(P₄N₂)][†],¹³ and Co(TPB),¹⁴ where dppe is bis(diphenylphosphino)ethane, P₄N₂ is a tetrachosphine with two pendant amines, and TPB is tris(*o*-diisopropylphosphinophenyl)borane. Moreover, without a series of similar M(η^2 -H₂) adducts with which to compare

^aDepartment of Chemistry, University of Minnesota-Twin Cities, 207 Pleasant Street SE, Minneapolis, Minnesota 55455, USA. E-mail: clu@umn.edu

^bSupercomputing Institute, Chemical Theory Center, University of Minnesota-Twin Cities, 207 Pleasant Street SE, Minneapolis, Minnesota 55455, USA

^cCatalysis Science Group, Pacific Northwest National Laboratory, P. O. Box 999, MS K2-57, Richland, Washington 99352, USA

^dNeutron Scattering Division, Oak Ridge National Laboratory, Oak Ridge, Tennessee 37831, USA

^eKey Laboratory of Cluster Science of Ministry of Education, School of Chemistry and Chemical Engineering, Beijing Institute of Technology, Beijing 100081, China

^fDepartment of Chemistry, University of Tennessee, Knoxville, Tennessee 37996, USA

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‡ These authors contributed equally.



thermodynamic data, the understanding gained by quantifying H₂ binding to a single metal complex are relatively limited.

Recently, the double-decker ligand, [N(*o*-(NCH₂PⁱPr₂)C₆H₄)₃]³⁻ (abbreviated as L), was used to prepare bimetallic (η²-H₂)M_AM_BL complexes in which group 13 Lewis acidic supporting metal ions (M_B) induce H₂ binding at Ni(0) and Co(-I) metal centers (M_A).^{15,16} Figueroa and Gabbaï have independently shown that appending a Lewis acidic σ-acceptor to a d¹⁰ transition metal induces binding of a Lewis base donor *trans* to the σ-acceptor.¹⁷⁻¹⁹ We and others have used a tethered σ-acceptor to promote activity at the transition metal for catalytic CO₂ functionalization.²⁰⁻²² Here, we report the first thermodynamic and kinetic studies of H₂ binding to a Ni center in an isostructural series, where M_B = Al, Ga, and In. In the case of the In(III) supporting ion, strong H₂ binding allowed for structural elucidation *via* single-crystal neutron diffraction. Significant modulation of the thermodynamic favorability of both H₂ and N₂ binding to Ni was enabled by introducing and varying the group 13 supporting metal ion, with H₂ binding free energies (Δ*G*^o) found to span nearly the entire range of previously reported values. The (η²-H₂)NiML complexes also exhibit different extents of H-H activation and kinetic rates of H₂ binding and loss. Through a combination of experimental and theoretical studies on binding and electronic structure, we present a comprehensive understanding of how a supporting group 13 metal ion poises a proximal transition metal for small-molecule binding and influences both the thermodynamics and kinetics of binding equilibria.

Results & discussion

Part I. Neutron diffraction study of (η²-H₂)NiInL and NMR characterization of (η²-H₂)NiAlL

We have previously reported a series of bimetallic NiML complexes featuring Ni(0)→M(III) dative bonds (M = Al (1), Ga (2), and In (3)), where larger group 13 ions, Ga and In, promote H₂ binding to formally d¹⁰ Ni centers.¹⁶ The resulting non-classical H₂ adducts, (η²-H₂)NiGaL (2-H₂) and (η²-H₂)NiInL (3-H₂), were characterized by ¹H NMR spectroscopy techniques. In addition, the remarkable stability of 3-H₂ allowed for its molecular structure to be determined by single-crystal X-ray diffraction at 123 K.¹⁶ We currently report a single-crystal neutron diffraction study of 3-H₂ at 100 K (Fig. 1) and the corresponding X-ray structure at 100 K. To our knowledge, 3-H₂ is the first H₂ adduct of Ni, or of any d¹⁰ metal, to be structurally characterized by neutron diffraction. To date, only two other Ni(0) H₂-adducts have been reported, [O(SiH(*o*-ⁱPr₂PC₆H₄)₂)₂]Ni₂(η²-H₂)₂ and [PhB(*o*-ⁱPr₂PC₆H₄)₂]Ni(η²-H₂), both of which were characterized *in situ* at low *T*.^{23,24}

Table 1 displays selected structural parameters for 3-H₂. Of note, the bond distances for non-hydrogen atoms in the X-ray and neutron structures of 3-H₂ are essentially identical within experimental error (Table S4†). The Ni-H distances between the two structures are also within error, though the H-H bond length differs by 0.12 Å. The neutron structure, which is more reliable for placement of H atoms, validates an intact H₂ molecule that is bound in side-on fashion to the Ni center. Upon

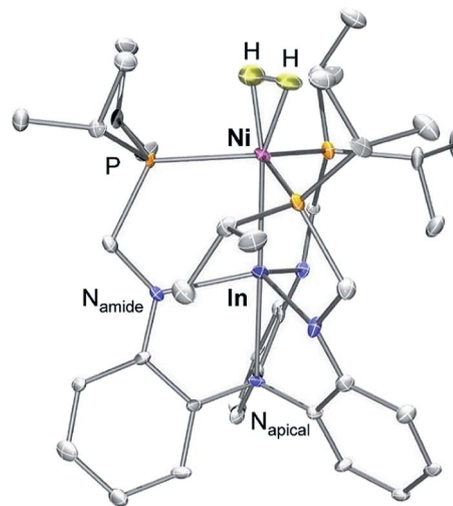


Fig. 1 Neutron structure of 3-H₂. Ellipsoids are shown at the 50% probability level, and H atoms (with the exception of H₂) have been omitted for clarity.

H₂ coordination, the Ni-In bond distance increases slightly from 2.457(1) Å in 3 to 2.4789(2) Å in 3-H₂. The H-H bond length of 0.80(2) Å is slightly elongated relative to that in free H₂ (0.74 Å), but shorter than the solution-state distance of 0.91 Å that was determined based on the *J*_{HD} coupling constant of 3-HD.¹⁶ This discrepancy can be attributed to rapid librational motion of H₂ in 3-H₂, a phenomenon which typically leads to an average contraction of ~0.07 Å in the apparent solid-state H-H bond distance relative to the solution-state distance determined by NMR studies.^{7,25-27} Consistent with this attribution, fast H₂ rotation in solution relative to molecular tumbling is also supported by the previously reported *T*₁ (min) value for the coordinated H₂ ligand of 3-H₂.¹⁶

Notably, the Ni-H bond distance is a parameter that neutron diffraction is uniquely able to experimentally evaluate. The two equivalent Ni-H bond lengths of 1.61(2) Å fall on the longer end of the wide range of the Ni-H distances reported for terminal Ni hydrides (*cf.* 1.32 to 1.65 Å).²⁸ Although direct comparisons are

Table 1 Selected structural metrics for 3-H₂ from X-ray and neutron studies at 100 K^a

	3-H ₂ (neutron)	3-H ₂ (X-ray)
H-H	0.80(2)	0.92(3)
Ni-H	1.61(2)	1.58(2)
	1.61(2)	1.65(2)
Ni-In	2.39(2)	2.4789(2)
Ni-P (avg.)	2.26(1)	2.2618(4)
In-N _{amide} (avg.)	2.13(1)	2.115(1)
In-N _{apical}	2.40(2)	2.366(1)
Ni to P ₃ -plane	0.31	0.29
In to N ₃ -plane	0.52	0.50

^a Values in Å (estimated standard deviations in parentheses). See Table 6 for crystallographic details. See Table S4 for a detailed comparison of the X-ray and neutron structures.



sparse due to the rarity of structurally characterized $M(\eta^2\text{-H}_2)$ complexes, the M–H and H–H distances in **3-H**₂ are both similar to those for $(\eta^2\text{-H}_2)\text{Co}(\text{TPB})$, which has M–H and H–H distances of 1.66/1.67 Å and 0.83(2) Å, respectively.²⁹ Short contact distances between the H₂ unit and the hydrogen atoms of the ligand isopropyl phosphine groups were observed in both **3-H**₂ and $(\eta^2\text{-H}_2)\text{Co}(\text{TPB})$, the closest of which were 2.24 Å and 2.10 Å, respectively. These distances are, within error, close to the intermolecular H⋯H van der Waals distance (2.2 Å).³⁰

Previously, no reaction was observed between NiALL (**1**) and H₂ (1 atm) at room temperature. Subsequent studies have found that either high H₂ pressure or low *T* is needed to observe H₂ binding to **1**. At 34 atm H₂ and 232 K, a new species was observed by ¹H NMR spectroscopy that is assigned as $(\eta^2\text{-H}_2)$ NiALL (**1-H**₂) based on a diagnostic resonance for bound H₂ at –1.5 ppm (Fig. S8 and S9†) and a short *T*₁ (min) value of ≤49(5) ms (500 MHz, THF-*d*₈; Fig. S10†).^{31,32} The ¹H NMR spectrum of the HD isotopomer, $(\eta^2\text{-HD})\text{NiALL}$ (**1-HD**), which was formed at 213 K under 3.8 atm HD, displayed a characteristic 1 : 1 : 1 triplet for the bound HD ligand, with *J*_{HD} = 34.4 Hz (Fig. S11†).

With the addition of **1-H**₂ to complete the isostructural trio of Ni($\eta^2\text{-H}_2$) complexes, it is apparent that H₂ activation increases as the supporting metal is varied down group 13 from Al to Ga to In, as reflected by the decreasing *J*_{HD} values (in Hz): 34.4 for **1-HD** > 33.2 for **2-HD** > 31.7 for **3-HD** (at 213 K; Fig. S11†). Thus, the corresponding estimated solution-state H–H bond distances increase as the supporting metal is varied down group 13, from 0.86 Å for **1-H**₂, to 0.88 Å for **2-H**₂, to 0.91 Å for **3-H**₂.³³ The relative extents of H–H bond activation for the $(\eta^2\text{-H}_2)\text{NiML}$ complexes are also in accord with those predicted by DFT calculations (Tables S13, S19 and S20†). That **1-H**₂ would have the shortest H–H distance of the trio, and the closest to that of free H₂, is suggestive of the weak and labile nature of H₂ binding to **1** relative to H₂ binding to **2** and **3**.^{6,7} We further note that the isostructural mononuclear Ni complex, NiLH₃ (**4**),³⁴ does not bind H₂ even under forcing conditions (193 K, 34 atm H₂),²⁰ which suggests that the supporting metal

plays a pivotal role in inducing H₂ binding and controlling the extent of H₂ activation. Lastly, both **2-H**₂ and **3-H**₂ exhibit a greater extent of H₂ activation compared to the two other Ni(0) H₂-adducts: $[\text{O}(\text{SiH}(\text{o-}^1\text{Pr}_2\text{PC}_6\text{H}_4)_2)_2]\text{Ni}_2(\eta^2\text{-H}_2)_2$ (*J*_{HD} = 34.2 Hz) and $[\text{PhB}(\text{o-}^1\text{Pr}_2\text{PC}_6\text{H}_4)_2]\text{Ni}(\eta^2\text{-H}_2)$ (*J*_{HD} = 36.5 Hz).^{23,24}

Part II. H₂ and N₂ binding energies for the NiML triad

Intrigued by the varied propensities for binding and activating H₂ enabled by changing a single atom, we set out to understand how the group 13 supporting metal influences the thermodynamics and kinetics of Ni–H₂ binding equilibria. Previously, $(\eta^2\text{-H}_2)\text{NiGaL}$ (**2-H**₂) was generated *in situ* under 1 atm H₂, but reverted to **2** upon exposure to vacuum or Ar.¹⁶ Because of its reversible H₂ binding at ambient conditions, **2** was an ideal candidate for initiating binding equilibrium studies. Variable-temperature (VT) ³¹P NMR spectra of **2** in toluene-*d*₈ under 6.8 atm H₂ displayed a single resonance, which shifted downfield from 42.7 to 56.4 ppm as the temperature was decreased from 368 K to 221 K (Fig. 2a). As a control experiment, the VT NMR profile of **2** under Ar showed a minimal change in the ³¹P shift ($\Delta\delta < 0.5$ ppm) over a similar *T* range (Fig. S12 and S13†). Hence, the VT NMR behavior of **2** under an H₂ atmosphere is consistent with an equilibrium between **2** and **2-H**₂ that is governed by fast chemical exchange relative to the ³¹P NMR timescale (202.4 MHz), where the latter is favored at low *T* and both species are three-fold symmetric in solution.

Analogous VT ³¹P NMR experiments were also performed for **2** under various H₂ pressures of 1.0, 13.6, and 34 atm (Fig. S1, S14 and S15†). Of note, for all H₂ pressures examined, the same low *T* convergence of the ³¹P resonance to ~56.9 ppm was observed at 193 K, which corresponds to the ³¹P chemical shift of **2-H**₂ (Fig. 2b). At high *T*, the observed ³¹P chemical shift approaches that of **2**, with closer convergence observed at lower H₂ pressures. These observations are consistent with rapid interconversion between **2** and **2-H**₂, such that the observed chemical shift (³¹P δ) is the population-weighted average of the chemical shifts of these exchanging species.³⁵ Thus, the

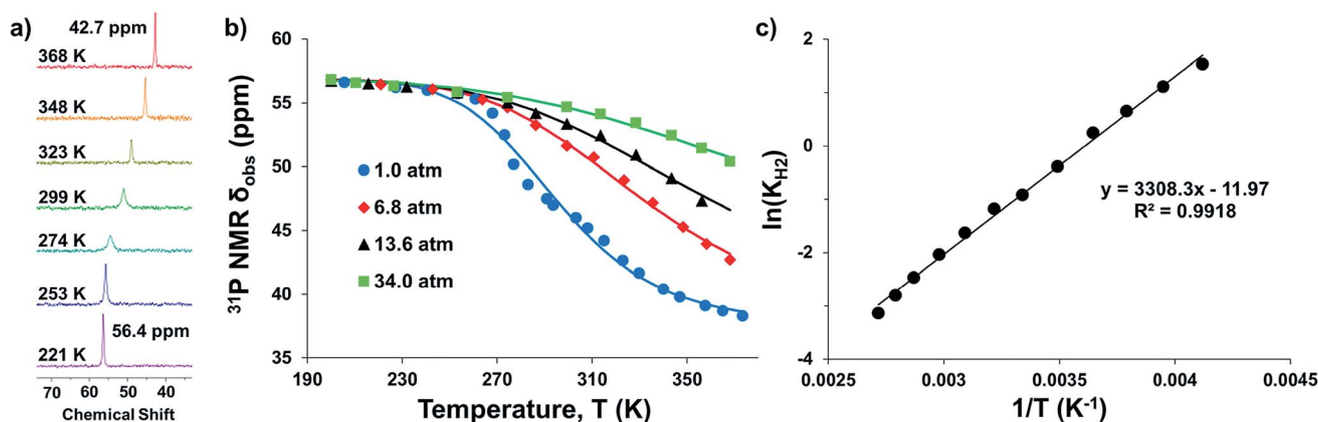


Fig. 2 (a) VT ³¹P NMR spectra of **2** under 6.8 atm H₂ in toluene-*d*₈ from 368 to 221 K. (b) Plots of ³¹P δ vs. *T* obtained for various H₂ pressures. Data are shown as points, and solid traces represent the best-fit curves obtained by varying ΔH° and ΔS° as parameters (see ESI†). (c) Representative van't Hoff plot of $\ln(K_{\text{H}_2})$ vs. $1/T$ at 6.8 atm H₂. The thermodynamic binding parameters shown in Table 2 were determined from van't Hoff plots for data sets collected at 6.8 and 13.6 atm H₂, which exhibit fast chemical exchange.



observed ^{31}P δ can be converted into the equilibrium concentration ratio of **2** and **2-H₂** (see ESI†). The equilibrium constant for H₂ binding, K_{H_2} , was determined according to eqn (1), where P_{H_2} is the H₂ pressure:

$$K_{\text{H}_2} = \frac{[(\eta^2\text{-H}_2)\text{NiML}]}{[\text{NiML}] \times P_{\text{H}_2}} \quad (1)$$

Next, the thermodynamic parameters for H₂ binding, ΔH° , ΔS° , and ΔG° , were determined using two complementary methods: (1) non-linear fitting of the ^{31}P δ vs. T plot in Fig. 2b; and (2) linear regression of the van't Hoff plot of $\ln(K_{\text{H}_2})$ vs. $1/T$, as shown in Fig. 2c. The analyses were performed using the 6.8 and 13.6 atm datasets, where chemical exchange was rigorously fast relative to the ^{31}P NMR timescale and the variability of ^{31}P δ with T was significant enough to reflect the sampling of an adequate portion of the binding equilibrium. The van't Hoff plot gives $\Delta H^\circ = -6.3(2)$ kcal mol⁻¹, $\Delta S^\circ = -23.0(7)$ cal (mol K)⁻¹, and $\Delta G^\circ = +0.6(3)$ kcal mol⁻¹ for H₂ binding to **2**, where the standard state conditions are defined as 298 K, 1 atm H₂, and 1 M concentrations of all other species in toluene-d₈. Non-linear fitting of the ^{31}P δ vs. T plot yields the same thermodynamic values within experimental error (Fig. S16, S17 and Table S2†).

The H₂ binding equilibria for the other NiML complexes, **1** (M = Al) and **3** (M = In), were also investigated by VT ^{31}P NMR spectroscopy. For H₂ binding to **1**, the VT ^{31}P NMR spectra in toluene-d₈ (363 to 210 K, 34 atm H₂) show a single ^{31}P resonance shifting downfield relative to that of **1** (30.7 ppm) with decreasing T , and ultimately converging to a chemical shift of ~44.3 ppm for **1-H₂** (Fig. S18†). The corresponding van't Hoff analysis for H₂ binding to **1** gives $\Delta H^\circ = -6.3(1)$ kcal mol⁻¹, $\Delta S^\circ = -26.4(4)$ cal (mol K)⁻¹, and $\Delta G^\circ = +1.6(2)$ kcal mol⁻¹ (Fig. S19†). Since **3** binds H₂ strongly, sub-ambient H₂ pressure was necessary to establish a measurable equilibrium between **3** and **3-H₂** (Fig. S20–S22†). Under 1 atm of 10% H₂ in Ar, where $P_{\text{H}_2} = 0.1$ atm, a single ^{31}P peak was observed from 299 to 357 K (Fig. S20†). However, the lineshapes of the observed ^{31}P peaks are noticeably broadened, which indicates that chemical

exchange between **3** and **3-H₂** falls into the fast-intermediate regime relative to the ^{31}P NMR timescale.³⁶ This is problematic because non-Lorentzian lineshapes and exchange broadening may diminish the reliability of the thermodynamic parameters determined under fast-intermediate exchange conditions.³⁷ Hence, we caution that the thermodynamic values for H₂ binding to **3**, $\Delta H^\circ = -14.8(6)$ kcal mol⁻¹, $\Delta S^\circ = -37(2)$ cal mol⁻¹ K⁻¹, and $\Delta G^\circ = -3.7(7)$ kcal mol⁻¹ (Fig. S23† and Table 2), should be treated as estimates.

To assess the reliability of this estimated ΔG° value, a control experiment was performed on **2** under identical conditions with 0.1 atm H₂, which showed similarly broad ^{31}P peaks due to fast-intermediate chemical exchange (Fig. S24 and S25†). Comparing the binding parameters obtained for **2** under 0.1 atm H₂ to those obtained under rigorously fast exchange conditions (6.8 and 13.6 atm H₂), we find that the thermodynamic favorability of H₂ binding was overestimated by 0.7 kcal mol⁻¹ using the 0.1 atm H₂ dataset. Thus, by applying this 0.7 kcal mol⁻¹ empirical correction, we propose that a better ΔG° estimate for H₂ binding to **3** is $-3.0(7)$ kcal mol⁻¹. In support, lineshape simulations of VT ^{31}P NMR spectra of **3** under 1 atm H₂ independently gives $\Delta G^\circ = -2.3(2)$ kcal mol⁻¹, which is within experimental error of the corrected value of $-3.0(7)$ kcal mol⁻¹ (Fig. S26†).

Next, we sought to investigate the related equilibria of N₂ binding to the NiML complexes using VT ^{31}P NMR experiments. The equilibrium between **2** and **2-N₂** was monitored at 1 atm N₂ and low T (226 to 193 K). Distinct ^{31}P resonances were observed for both **2** and **2-N₂** (Fig. S30†), which is characteristic of slow chemical exchange relative to the ^{31}P NMR timescale (161.9 MHz). Quantitative integration of the ^{31}P NMR peaks for **2** and **2-N₂** allowed for a straightforward determination of K_{N_2} using eqn (2), where P_{N_2} is the N₂ pressure:

$$K_{\text{N}_2} = \frac{[(\text{N}_2)\text{NiML}]}{[\text{NiML}] \times P_{\text{N}_2}} \quad (2)$$

Conversely, the interconversion of **3** and **3-N₂** is fast relative to the ^{31}P NMR timescale (161.9 MHz) at 1 atm N₂ and $T > 288$ K (Fig. S6 and S31†). As such, the observed chemical shift of the single ^{31}P NMR resonance represents the population-weighted average of the chemical shifts of **3** and **3-N₂**, and the VT NMR data were analyzed as previously described for H₂ binding to **2** (Fig. S31 and S32†). Lastly, the observed equilibrium between **1** and **1-N₂** at 51 atm N₂ switches from slow exchange at low T (≤ 210 K) to fast exchange at higher T (≥ 243 K) (Fig. S33†). Thus, K_{N_2} was evaluated based on the distinct ^{31}P NMR peak integrations for **1** and **1-N₂** at low T , and based on the observed ^{31}P chemical shift at high T . The thermodynamic binding parameters for the H₂ and N₂ binding equilibria of the NiML complexes are compiled in Tables 2 and 3. Additionally, DFT calculations using the M06-L³⁸/bs1 method correctly predict the experimental trends in both the H₂ and N₂ binding free energies for the trio of NiML complexes (Tables 2 and 3).

Across the NiML series, ΔG° for both H₂ and N₂ binding was modulated by ~5 kcal mol⁻¹, with increasing thermodynamic favorability for both H₂ and N₂ binding observed as the

Table 2 Experimental and DFT-calculated H₂ thermodynamic binding parameters for **1**, **2**, and **3**^a

	1	2 ^b	3
ΔH° (kcal mol ⁻¹)	-6.3(1)	-6.3(2)	-14.8(6) ^d
ΔS° (cal mol ⁻¹ K ⁻¹)	-26.4(4)	-23.0(7)	-37(2) ^d
ΔG° (kcal mol ⁻¹)	1.6(2)	0.6(3)	-3.7(7) ^d -3.0(7) ^e
ΔG° (DFT) ^c	2.8	0.9	-1.9

^a Standard deviations obtained from van't Hoff linear regression analyses are given in parentheses. Standard state is defined as 298 K, 1 atm H₂ (or N₂), and 1 M of all other species in toluene-d₈. ^b H₂ binding studies have also been conducted for **2** in THF (Fig. S27–S29): $\Delta G^\circ = +0.1(1)$ kcal mol⁻¹, $\Delta H^\circ = -7.5(1)$ kcal mol⁻¹, and $\Delta S^\circ = -25.4(1)$ cal mol⁻¹ K⁻¹. ^c Units of kcal mol⁻¹, see Computational methods in Experimental section for details. ^d Estimated values extracted from fast-intermediate exchange regime data (Fig. S20–S23). ^e Corrected ΔG° value of $-3.0(7)$ kcal mol⁻¹ is likely a better estimate. See text for explanation of the +0.7 kcal mol⁻¹ correction term.



Table 3 Experimental and DFT-calculated N₂ thermodynamic binding parameters for **1**, **2**, and **3**^a

	1	2	3
ΔH° (kcal mol ⁻¹)	-4.7(2)	-4.7(3)	-14.5(3)
ΔS° (cal mol ⁻¹ K ⁻¹)	-27.5(5)	-23(1)	-45(1)
ΔG° (kcal mol ⁻¹)	3.5(3)	2.1(5)	-1.2(4)
ΔG° (DFT) ^b	6.3	3.6	0.2

^a Same as Table 2 footnote *a*. ^b Same as Table 2 footnote *c*.

supporting metal was varied down group 13 (Tables 2 and 3). Notably, ΔG° values for H₂ binding to the NiML complexes nearly span the entire range of previously reported values (from -2 to +3 kcal mol; Tables S6 and S7†). This is remarkable considering that the full range of literature ΔG° values encompasses H₂ binding to various transition metals (*e.g.* Cr, Mo, W, Re, Fe, Ru, Co, and Ir) in diverse ligand environments.^{10,11,13,14,39–45} In contrast, the NiML trio features an isostructural Ni site within the same ligand framework, where the primary difference is the identity of the group 13 metal. We hypothesize that the strength of the Ni(0)→M(III) dative interaction directly tunes the binding at Ni in the position *trans* to M(III), where greater Ni(0)→M(III) interactions lead to stronger small-molecule binding. This hypothesis is generally consistent with other literature examples wherein the interaction of a σ -acceptor with a d¹⁰ transition metal enhances donor-binding at the *trans* position.^{17–19} DFT calculations predict that supporting Ni with In(III) in **3** increases the favorability of H₂ binding by ~8 kcal mol⁻¹ relative to NiLH₃ (**4**), which are the two limiting extremes in this series (Table S16†).

To the best of our knowledge, (η^2 -H₂)Co(TPB) and [(η^2 -H₂)Re(CN^tBu)₃(PCy₃)₂]⁺ were previously reported to have the most favorable H₂ binding free energies, with $\Delta G_{1M}^\circ = -4.8(9)$ and $-4.8(1.3)$ kcal mol⁻¹, respectively (Table S7†).^{14,39} Notably, **3** binds H₂ even more favorably, with $\Delta G_{1M}^\circ \sim -6.5(7)$ kcal mol⁻¹, which was obtained by converting ΔG° at $P_{H_2} = 1$ atm to ΔG° for [H₂] = 1 M. In addition, the determination of both H₂ and N₂ binding energies for first-row transition metal complexes is rare.^{10,11,13,14,46} Across the NiML series, the binding free energies for H₂ are more favorable than those for N₂ by 1.9(4), 1.5(5), and ~1.8(8) kcal mol⁻¹ for **1**, **2**, and **3**, respectively (Tables 2 and 3). A similar trend was reported for [Fe(P₄N₂)]⁺ and Cr(CO)₃(PCy₃)₂, whereas ΔG_{1M}° energies for H₂ and N₂ binding to Co(TPB) are identical within error (Tables S6–S9†). The finding that N₂ binding is more competitive with H₂ binding for Co(TPB) than it is for NiML complexes is consistent with the greater π -basicity of low-valent Co toward N₂.^{47,48}

In further examining the ΔH° and ΔS° contributions for binding to NiML (**1–3**), the large negative ΔS° values for H₂ and N₂ binding reflect the entropic cost of binding a gas molecule (Tables 2 and 3). In general, a greater entropic cost was observed for N₂ binding than for H₂ binding; this trend can be explained in part by the larger absolute entropy of N₂ relative to H₂.¹¹ It is also striking that both H₂ and N₂ binding to **3** are considerably more exothermic ($\Delta\Delta H^\circ \approx -9$ kcal mol⁻¹) and entropically

costly ($\Delta\Delta S^\circ \approx -13$ and -20 cal mol⁻¹ K⁻¹, respectively) compared with binding to **1** and **2** (Tables 2 and 3). The fact that binding to **3** has the most favorable enthalpic and most unfavorable entropic terms is consistent with tighter and more rigid binding of both H₂ and N₂. Intriguingly, because **1** and **2** have identical ΔH° values for both H₂ and N₂ binding, the differences in their observed binding propensities originate from the differences in ΔS° values.

Part III. Kinetics of self-exchange for NiML and (η^2 -H₂)NiML complexes

To understand the dependence of the kinetics of the H₂ binding process on the supporting metal, VT ³¹P NMR lineshape simulations of NiML under 1 atm H₂ were performed to extract self-exchange rates for the interconversion between NiML and (η^2 -H₂)NiML. Exchange rates were determined *via* least-squares fitting of the VT ³¹P NMR spectra using a two-site, non-mutual exchange model (see Experimental section and ESI† for details).^{49,50} Good agreement was generally achieved between the experimental and simulated spectra, as illustrated in Fig. 3a for the VT ³¹P NMR study of **1** under 1 atm H₂. Similar lineshape analyses were performed for each NiML complex, which allowed exchange rates to be determined at seven or more different temperatures between 213 K and 344 K (Fig. S42–S44†). Notably, the exchange rate decreases by a factor of ~6 at 298 K upon varying the supporting metal from Al to In (Table 4). Exchange rates at 298 K were found to correlate strongly with both H₂ binding free energies ($R^2 = 0.996$, Fig. S47†) and H–H bond distances in (η^2 -H₂)NiML complexes ($R^2 = 0.958$, Fig. S48†), with slower kinetic rates of H₂ exchange for more thermodynamically favorable binding equilibria where H₂ is more activated. The H₂ binding equilibrium for complex **1** is especially dynamic, with interconversion between **1** and **1**-H₂ occurring over 28 000 times per second at 298 K and 1 atm H₂ (Fig. 3a and Table 4).

We propose a self-exchange mechanism comprised of H₂ loss from (η^2 -H₂)NiML to generate NiML and free H₂, and H₂ binding to another NiML complex to form (η^2 -H₂)NiML (Fig. 3c). The rate constants for H₂ loss from (η^2 -H₂)NiML, k_{loss} , were determined at each *T* based on the exchange rates and the known equilibrium concentrations of (η^2 -H₂)NiML (see ESI†). Eyring analyses for k_{loss} allowed for the determination of activation barriers for H₂ loss from (η^2 -H₂)NiML complexes, as shown in Fig. 3b for **1**-H₂. Eyring plots for H₂ loss from **2**-H₂ and **3**-H₂ are shown in Fig. S49 and S50,† respectively. The free energy barriers for H₂ loss ($\Delta G_{\text{loss}}^\ddagger$) were found to be similar for all complexes (9.1 to 9.4 kcal mol⁻¹), with ΔH^\ddagger and ΔS^\ddagger values ranging from 9.3 to 11.3 kcal mol⁻¹ and from -0.3 to +7.5 cal (mol K)⁻¹, respectively (Table 4). The fact that $\Delta S_{\text{loss}}^\ddagger$ values are positive or close to zero in all cases is consistent with the expected gain in H₂ freedom of motion, while positive $\Delta H_{\text{loss}}^\ddagger$ values suggest that partial Ni–H₂ bond breaking is the dominant process involved in reaching the transition state for H₂ loss. Interestingly, both $\Delta H_{\text{loss}}^\ddagger$ and $\Delta S_{\text{loss}}^\ddagger$ values for the (η^2 -H₂)NiML complexes decrease as the supporting metal is varied down group 13 (Al > Ga > In; Table 4).



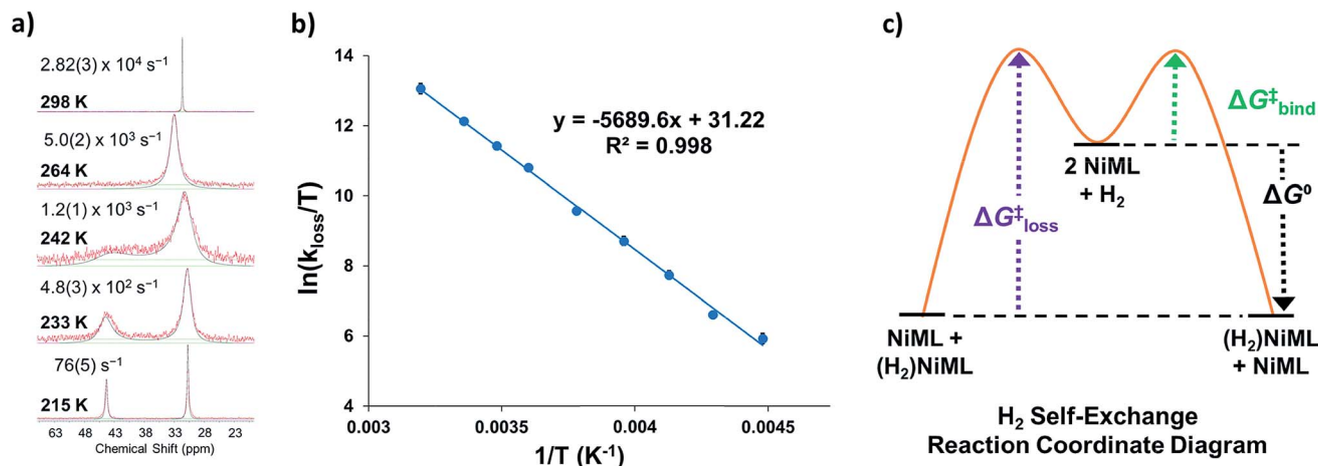


Fig. 3 (a) Selected VT ^{31}P NMR spectra of **1** under 1 atm H_2 from 215 to 298 K (red lines, toluene- d_8 , 162 MHz) and corresponding simulated spectra (black lines), which were utilized to extract exchange rates at each T (standard deviations in parentheses). (b) Eyring plot of $\ln(k_{\text{loss}}/T)$ vs. $1/T$, where k_{loss} is the first-order rate constant for H_2 loss from 1-H_2 . (c) Proposed reaction coordinate diagram for chemical exchange between $(\eta^2\text{-H}_2)\text{NiML}$ and NiML via unimolecular H_2 loss.

Table 4 Comparison of thermodynamic and kinetic binding parameters for H_2 self-exchange, binding, and loss for NiML complexes (**1**–**3**)^a

Complex		1	2	3
Self-exchange ^b	Rate _{ex} ($\times 10^4$)	2.82(3)	2.2(2)	0.437(5)
	Rate _{rel}	6.4	4.9	1.0
H_2 loss ^c	k_{loss} ($\times 10^6$)	55(3)	6.5(5)	0.60(3)
	$\Delta H_{\text{loss}}^\ddagger$	11.3(2)	10.1(4)	9.3(4)
	$\Delta S_{\text{loss}}^\ddagger$	7.5(1)	3.5(1)	-0.3(1)
	$\Delta G_{\text{loss}}^\ddagger$	9.1(2)	9.1(4)	9.4(4)
H_2 binding	$\Delta G_{1\text{M}}^\ddagger$ ^d	-1.9(2)	-2.9(2)	-6.5(7)
	$\Delta G_{\text{bind}}^\ddagger$	7.2(3)	6.2(5)	2.9(8)

^a All values determined for a standard state of 1 M H_2 , 1 M for all other species in toluene- d_8 , and 298 K. $\Delta G_{1\text{M}}^\ddagger$, ΔG^\ddagger , and ΔH^\ddagger values are in units of kcal mol⁻¹, and ΔS^\ddagger values are in units of cal (mol K)⁻¹. Rate_{ex} and k_{loss} are in s⁻¹, and rate_{rel} is the relative rate. ^b $[\text{Ni}]_{\text{total}} = 7.5$ mM. ^c $\Delta G_{\text{loss}}^\ddagger \approx \Delta G_{\text{exchange}}^\ddagger$ for unimolecular H_2 loss. ^d $\Delta G_{1\text{M}}^\ddagger$ are directly derived from values in Table 2 by converting the standard state for H_2 from 1 atm H_2 to 1 M. See ESI for details.

Furthermore, $\Delta S_{\text{loss}}^\ddagger$ values (~ 0 or >0) are consistent with unimolecular H_2 loss from $(\eta^2\text{-H}_2)\text{NiML}$, as opposed to H_2 self-exchange via an intermolecular $\text{LMNi}\cdots\text{H}_2\cdots\text{NiML}$ transition state. Such a transition state also seems unlikely because of the steric clash between the diisopropyl ligand groups surrounding each Ni site. Using the proposed reaction coordinate diagram, one can further extract the free energy barrier for H_2 binding ($\Delta G_{\text{bind}}^\ddagger$) by using the thermodynamic relationship: $\Delta G_{\text{loss}}^\ddagger = \Delta G_{\text{bind}}^\ddagger - \Delta G_{1\text{M}}^\circ$. Of note, $\Delta G_{1\text{M}}^\circ$ is the free energy of H_2 binding after converting the H_2 standard state from 1 atm (as given in Table 2) to 1 M. Hence, the $\Delta G_{\text{bind}}^\ddagger$ values for **1**, **2**, and **3** are 7.2(3), 6.2(5), and ~ 2.9 (8) kcal mol⁻¹, respectively, where the activation barriers for H_2 binding are lower for more thermodynamically favorable binding equilibria (Table 4).

Activation barriers and rate constants for H_2 binding and loss have seldom been reported despite their relevance for

many catalytic processes. $\Delta H_{\text{loss}}^\ddagger$ values for $(\eta^2\text{-H}_2)\text{NiML}$ (9.3 to 11.3 kcal mol⁻¹) are comparable to those reported (in kcal mol⁻¹) for $[\text{Ru}(\eta^2\text{-H}_2)\text{H}_3(\text{PPh}_3)_3]^+$ (8.8),³¹ $\text{Cr}(\eta^2\text{-H}_2)(\text{CO})_3(\text{PCy}_3)_2$ (12.1),¹⁰ $\text{Ir}(\eta^2\text{-H}_2)(\text{H})_2\text{X}(\text{P}^t\text{Bu}_2\text{Me})_2$ (9.4 to 11.3 for X = Cl, Br, I),⁴⁰ and $\text{Ir}(\eta^2\text{-H}_2)(\text{H})_2\text{X}(\text{P}^i\text{Pr}_3)_2$ (10.1 to 11.4 for X = Cl, Br, I),⁵¹ and are significantly lower than those reported for $\text{W}(\eta^2\text{-H}_2)(\text{CO})_3(\text{PCy}_3)_2$ (16.9)⁴⁴ and $\text{Ru}(\eta^2\text{-H}_2)(\text{H})_2(\text{PPh}_3)_3$ (17.9).³¹ Perhaps the most thoroughly studied H_2 binding equilibrium is that of $\text{W}(\eta^2\text{-H}_2)(\text{CO})_3(\text{PCy}_3)_2$, for which rate constants for both H_2 binding (k_{bind}) and loss (k_{loss}) have been directly measured to be $\sim 2 \times 10^6$ M⁻¹ s⁻¹ and 469 s⁻¹ at 298 K, respectively.^{44,52} In comparison, the k_{loss} values for $(\eta^2\text{-H}_2)\text{NiML}$ complexes are 3 to 5 orders of magnitude greater than that for $\text{W}(\eta^2\text{-H}_2)(\text{CO})_3(\text{PCy}_3)_2$ (Table 4). This can be rationalized by the fact that H_2 is much more activated in $\text{W}(\eta^2\text{-H}_2)(\text{CO})_3(\text{PCy}_3)_2$, which co-exists with its dihydride species at 298 K ($K \sim 0.25$).^{7,44,45} In contrast, $\text{H}(\eta^2\text{-H})\text{ML}$ dihydride species, which have some precedent in the literature,^{24,53,54} have not been observed.^{16,55} DFT calculations predict such a species to be unstable relative to $(\eta^2\text{-H}_2)\text{NiML}$ by 12 to 19 kcal mol⁻¹ under 1 atm H_2 (Table S25, S26 and Fig. S55[†]).

Part IV. Insights from quantum chemical calculations and UV-Vis spectra

Complexes **1**–**3** and NiLH_3 (**4**), as well as their H_2 and N_2 adducts, were investigated by density functional theory (DFT) calculations using several different functionals and basis sets (see Experimental section and Table S3[†]). The M06-D3^{56,57}/bs4 method gave the best agreement between the optimized and experimental geometries for the three Ni–In complexes: **3**, 3-N_2 , and 3-H_2 (Tables S13–S15[†]).¹⁶ On the other hand, the relative free energies (ΔG°) for H_2 and N_2 binding to **1**–**3** were best matched by M06-L³⁸/bs1 (Tables S16 and S17[†]), which correctly predicted the trends of stronger H_2 binding than N_2 and increasing binding favorability for $\text{M} = \text{In} > \text{Ga} > \text{Al}$ (Tables 2 and 3).



To better understand chemical bonding between H₂ and Ni in the (η²-H₂)NiML complexes, we conducted energy decomposition analysis (EDA).⁵⁸ EDA allows for the total interaction energy (ΔE_{int}) between the H₂ and NiML fragments to be partitioned into the following terms: electrostatic energy (ΔE_{elstat}), Pauli repulsive interaction (ΔE_{Pauli}), dispersion (ΔE_{disp}), and covalent interaction (ΔE_{orb}).⁵⁸ The ΔE_{orb} term can be further divided into individual energy contributions associated with specific orbital interactions, and thereby allows for differentiation of σ, π, and δ bonding interactions. The extended transition-state method coupled with natural orbitals for chemical valence theory (ETS-NOCV⁵⁹) was then utilized to analyze the bonding interactions between Ni and H₂. Fig. 4 illustrates the two most important NOCV pairs for the Ni-H₂ interaction in 3-H₂, in which chemical bonding is indicated by deformation in the electron densities (Δρ). Overall, two orbital interactions are important: (1) σ-donation from the H₂ σ-bond to the empty Ni 4p_z orbital, which accounts for 56% of ΔE_{orb} and is comprised of Δρ₁ (38%; Fig. 4) and Δρ₃ (18%; Fig. S53†); and (2) π-back-donation from a Ni 3d_π orbital to the empty H₂ σ* orbital, which accounts for 35% of ΔE_{orb}. Similarly, greater contributions of σ-donation (57–59% of ΔE_{orb}) compared to π-back-donation (34–35% of ΔE_{orb}) were also observed for Ni-H₂ orbital interactions in 1-H₂ and 2-H₂ (Table S22†).

The total interaction energy (ΔE_{int}) between the H₂ and Ni fragments becomes increasingly favorable in the order, (η²-H₂)NiLH₃ < 1-H₂ < 2-H₂ < 3-H₂ (Table S21†). Of interest, the less favorable ΔE_{int} predicted for (η²-H₂)NiLH₃ arises from a large, unfavorable ΔE_{Pauli} term, which is the repulsive interaction energy between like spins in the H₂ and NiLH₃ fragments. This makes sense as NiLH₃ lacks a Ni→M interaction that would result in attenuation of Ni electron density. Within the (η²-H₂)NiML series, both the ΔE_{elstat} and ΔE_{orb} terms become more favorable as M is varied down the group 13 triad (Table S21†). Furthermore, the greater relative importance of σ-donation to H₂ binding than π-back-donation is consistent with the trend that H₂ binding favorability increases as the Ni center becomes

more electron-deficient, as judged by the Ni(I/0) redox potentials (Fig. S71 and Table S39†).¹⁶ Hence, we propose that increased electron-withdrawal *via* Ni→M dative bonding as M is varied down group 13 induces stronger electrostatic and covalent interactions between H₂ and NiML, with a concomitant decrease in unfavorable Pauli repulsion between the two fragments.

Given the rarity of H₂ and N₂ binding to a Ni(0) metal center, we next sought to understand the specific electronic perturbations of the supporting group 13 metal ion that poise NiML to bind small molecules. To lend insight, UV-Vis spectroscopy studies in conjunction with time-dependent density functional theory (TD-DFT) calculations were conducted to investigate the electronic structure of complexes 1–4 and their H₂ and N₂ adducts. The UV-Vis spectra of NiLH₃ and 1–3 are shown in Fig. 5a (THF, 298 K), along with the corresponding TD-DFT predicted spectra, where M06-D3/bs4 showed the best agreement with experiment (Fig. 5b and Table S29†). We also sought to understand how the binding of H₂ and N₂ to the NiML complexes further impacts electronic structure. In this regard, it is notable that small-molecule binding to NiML often manifests in vibrant color changes. For example, exposure of a THF solution of 3 under Ar to an N₂ or H₂ atmosphere resulted in a color change from a deep red-purple color to a lighter red (3-N₂) or yellow-brown (3-H₂), respectively (Fig. 5c).

The UV-Vis spectra of 1–3 share the same pattern of three peaks between 400 and 800 nm, which are marked with asterisks in Fig. 5a and listed in Table 5 as peaks I, II, and III. These peaks all red-shift upon varying the supporting metal from Al to Ga to In. The TD-DFT calculated transition energies for 1–3 agree reasonably well with experiment, with the predicted excitations generally blue-shifted by 0.04 to 0.3 eV for peaks I to III (Table 5). The TD-DFT results also correctly predicted the red-shift of all peaks for 2 relative to 1, but the subtle spectral differences between 2 and 3 were not discerned by TD-DFT despite testing several methods (Table S29, Fig. S57 and S58†). For NiLH₃, a broad absorption feature is observed at ~500 nm, which was fit with two overlapping peaks with maxima at 491 and 533 nm, along with a low-intensity shoulder at 663 nm (Table S27, Fig. S56†).

UV-Vis peaks I–III for NiLH₃ and 1–3 were assigned based on the TD-DFT results (Table 5). Each peak corresponds to an electronic excitation from either a Ni 3d orbital or a ligand-based molecular orbital (MO) to the lowest unoccupied molecular orbital (LUMO), the latter of which is highly similar for all complexes. For NiLH₃, the LUMO has both Ni 4p_z and P 4p character, whereas the LUMO for complexes 1–3 also has additional contributions from Ni (3d, 4s) and M (s, p_z) atomic orbitals (Fig. S59 and Table S31†). For all complexes, peaks I and II arise from electronic excitations to the LUMO from the Ni d_{xy}/d_{x²-y²} and d_{xz}/d_{yz} orbitals, respectively. Peak III for NiLH₃ is a transition from a pure d_{z²} orbital to the LUMO, while peak III for 1–3 is a more complex transition from a mixed ligand-based arene π* MO with partial Ni d_{z²} character to the LUMO (Table S28†).

Based on these transition assignments, semi-quantitative MO diagrams for NiLH₃ and 1–3 can be constructed (Fig. 6).

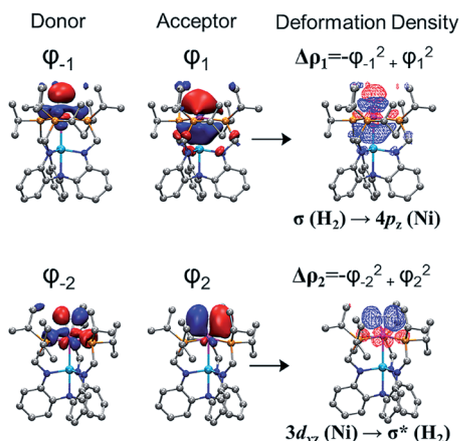


Fig. 4 The two dominant NOCV pairs in 3-H₂ and their associated deformation densities (Δρ₁, top right; Δρ₂, bottom right; contour iso-value = 0.04 a.u.). The colors of the deformation densities indicate the flow of electrons, from red to blue, involved in the Ni-H₂ interaction.



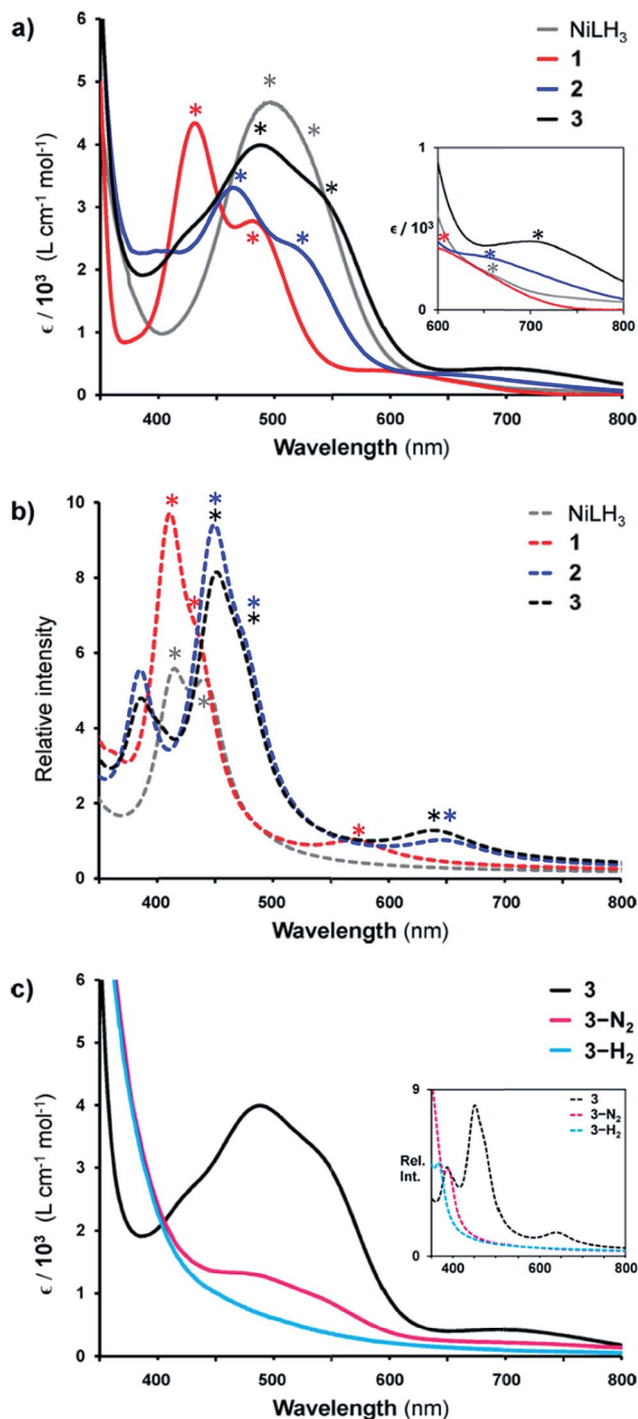


Fig. 5 (a) UV-Vis spectra of NiLH₃ (grey), 1 (red), 2 (blue), and 3 (black) obtained in THF at room temperature. Inset is a close-up of the 600–800 nm range. (b) TD-DFT predicted spectra of NiLH₃ and 1–3 (MO6-D3/bs4, SMD/THF) shown as dotted lines. (c) UV-Vis spectra of 3 (under Ar, black), 3-N₂ (under 1 atm N₂, pink), and 3-H₂ (under 1 atm H₂, cyan) in THF at 298 K, with TD-DFT plots shown in the inset. Note that small features corresponding to 3 can be seen in the UV-Vis spectrum of 3-N₂ due to the equilibrium binding of N₂ under these conditions. Asterisks in spectra correspond to bands I, II, or III, as listed in Table 5.

Table 5 UV-Vis peaks (nm) for NiLH₃ and 1–3, with TD-DFT data (in parentheses) and transition assignments

Peak	Transition	NiLH ₃	1	2	3
I	$d_{xy}/d_{x^2-y^2} \rightarrow \text{LUMO}$	663 (490)	600 (573)	638 (650)	699 (642)
II	$d_{yz}/d_{xz} \rightarrow \text{LUMO}$	491 (413)	490 (437)	508 (478)	530 (475)
III	$\pi\text{-ligand} + d_{z^2} \rightarrow \text{LUMO}$	533 (443) ^a	430 (410)	464 (448)	488 (449)

^a Pure Ni $d_{z^2} \rightarrow \text{LUMO}$ transition.

For NiLH₃, the d-orbital manifold is consistent with that expected for a trigonal-planar metal center with three σ -donors: the degenerate $d_{xy}/d_{x^2-y^2}$ orbitals are the most destabilized, followed by d_{z^2} , and then the d_{xz}/d_{yz} set. For 1–3, the presence of the supporting group 13 metal ion results in the stabilization of the Ni $3d_{z^2}$ orbital *via* Ni \rightarrow M(p_z/s) dative bonding, which is consistent with the blue shift in the predicted pure $3d_{z^2} \rightarrow \text{LUMO}$ transitions in the bimetallic NiML complexes (363 to 385 nm; Table S30[†]) relative to NiLH₃ (443 nm). The Ni d_{xz}/d_{yz} orbitals, on the other hand, are destabilized upon the introduction of the supporting metal and its variation down group 13. Presumably, the π -back-bonding interaction, Ni $d_{xz}/d_{yz} \rightarrow$ P–C σ^* , becomes weaker as the Ni center becomes more electron-deficient due to stronger electron withdrawal by the supporting metal (In > Ga > Al > no support).¹⁶ In support, both the Ni–P bonds elongate and the ³¹P NMR signal shifts downfield from NiLH₃ to 1 to 2 to 3.

Another notable MO trend involves the energy difference between the Ni d_{xz}/d_{yz} orbitals and the LUMO, as reflected in the peak II energies, which decrease upon the introduction and variation of the supporting metal down group 13. This particular energy gap is important because the LUMO and the Ni d_{xz}/d_{yz} orbitals are the frontier Ni-based MOs that participate in small-molecule binding, with the Ni-based LUMO accepting σ -donation from either the H₂ σ -bond or the N₂ lone pair, and the Ni d_{xz}/d_{yz} orbitals participating in π -back-bonding to either the H₂ σ^* or N₂ π^* LUMOs. Overall, the Ni $d_{xz}/d_{yz} \rightarrow \text{LUMO}$ excitation energy decreases by ~ 0.2 eV across the NiML complexes, from 2.53 eV in 1 to 2.44 eV in 2 to 2.34 eV in 3 (Table S28[†]). Thus, it is reasonable to propose that varying the supporting metal from Al to Ga to In results in the destabilization of the Ni d_{xz}/d_{yz} orbital set and the stabilization of the Ni-based LUMO, such that both become more energetically accessible to interact with small-molecule substrates.

The involvement of the LUMO in small-molecule binding is further supported by examining the changes in the UV-Vis spectra for 3 upon binding H₂ and N₂ (Fig. 5c). Notably, both the experimental and theoretical spectra of 3-H₂ and 3-N₂ lack any intense features between 400 and 700 nm. Instead, the lowest energy transitions for 3-H₂ and 3-N₂ are predicted at 375 and 391 nm, respectively. The shift to higher energy excitations for the H₂ and N₂ adducts can be qualitatively explained by the nature of their acceptor MOs, which we define as the lowest-energy unoccupied MO with significant Ni character, rather



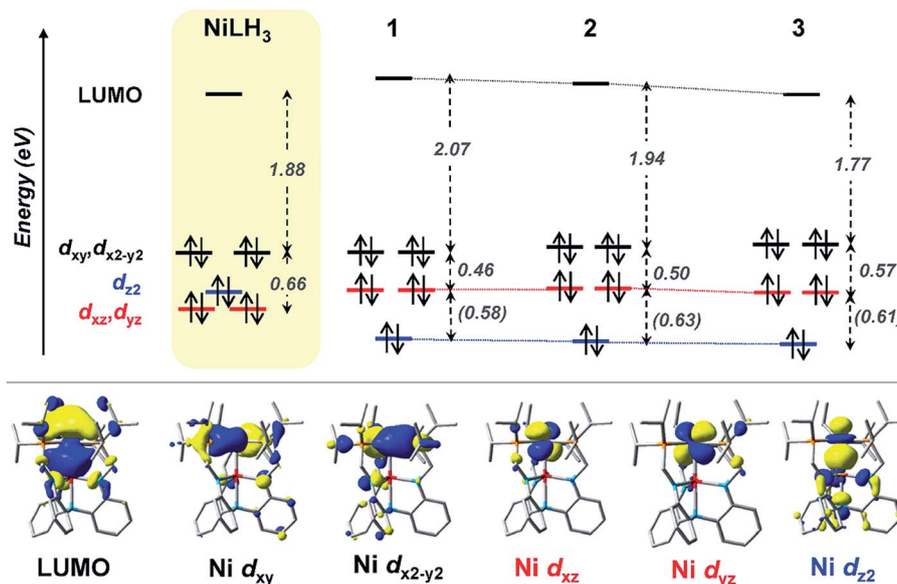


Fig. 6 (Top) MO diagrams of NiLH₃ and complexes 1–3. The ligand field energies (in eV) are based on UV-Vis data (Table 5). TD-DFT calculated energies are shown (in eV, italicized within parentheses) when experimental values were not obtainable. (Bottom) MOs for complex 2, which are representative of those for the series. Note that MO energies are drawn to scale, but energy comparisons across the complexes are qualitative.

than those that are ligand based (Fig. S60–S63 and Tables S33–S38†). For both 3-H₂ and 3-N₂, the acceptor MO appears to be an anti-bonding combination of the LUMO in 3 and the donor MO of the small molecule (Fig. S64†). Thus, upon small-molecule binding *via* the dominant σ -donation interaction with the LUMO of NiML, the resulting acceptor orbitals in (L')NiML adducts (L' = H₂, N₂) to which electrons can be excited are of higher energy relative to the Ni 3d manifold, and thereby give rise to higher energy electronic transitions.

Part V. Linear free-energy relationships

We examined linear free-energy relationships across the NiML series to quantify the effect of the supporting group 13 metal on the thermodynamics and kinetics of small-molecule binding. For H₂ binding, an excellent correlation was observed between the thermodynamic free energies (ΔG_{1M}°) and the free energies

of activation ($\Delta G_{\text{bind}}^\ddagger$), where more exergonic binding equilibria have smaller activation barriers ($R^2 = 0.999$, Fig. 7a). The slope of +0.93 in the $\Delta G_{\text{bind}}^\ddagger$ versus ΔG_{1M}° plot indicates that a decrease of ~ 1 kcal mol⁻¹ in ΔG_{1M}° for H₂ binding corresponds to a commensurate lowering of the free energy barrier. Reasonably strong correlations were also found between the Ni 3d_{xz}/d_{yz} → LUMO excitation energy and ΔG° for H₂ ($R^2 = 0.924$) and N₂ ($R^2 = 0.963$) binding (Fig. 7b). This correlation makes sense since the Ni 3d_{xz}/d_{yz} orbitals and the LUMO are the Ni-based frontier orbitals that participate in small-molecule binding (*vide supra*). The slopes of the plots of ΔG° for H₂ and N₂ binding *versus* the Ni 3d_{xz}/d_{yz} → LUMO excitation energy (in kcal mol⁻¹) are close to unity, where a decrease of 1 kcal mol⁻¹ in the Ni 3d_{xz}/d_{yz} → LUMO excitation energy is associated with a nearly equivalent increase in the favorability of H₂/N₂ binding.

The isostructural nature of the (η^2 -H₂)NiML series allows us to further interrogate which intrinsic properties of the group 13

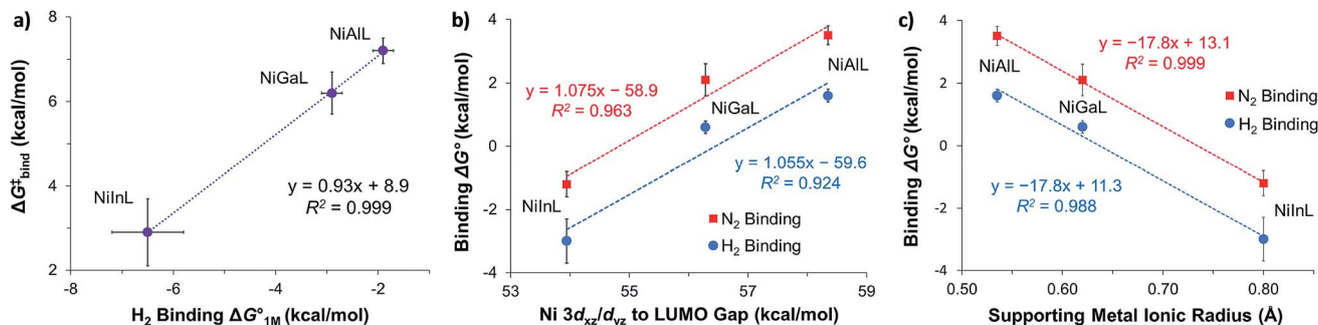


Fig. 7 (a) Plot of free energy barrier for H₂ binding ($\Delta G_{\text{bind}}^\ddagger$) vs. thermodynamic free energy for H₂ binding (ΔG_{1M}°), with the standard deviation in all values shown by error bars and all standard states defined to be [H₂] = 1 M in toluene-*d*₈. (b–c) Plots of ΔG° for H₂ (blue circles) and N₂ binding (red squares) vs. Ni 3d_{xz}/d_{yz} to LUMO energy gap (b) and vs. supporting metal Shannon ionic radii (c). Ni 3d_{xz}/d_{yz} to LUMO energy gaps were measured experimentally by UV-Vis spectroscopy with transition assignments from TD-DFT calculations (see ESI† and Table 5).



support (M) dictate the favorability of H₂ and N₂ binding. A strong negative correlation was found between the size of the supporting metal, as represented by Shannon's M(III) ionic radii,⁶⁰ and ΔG° for both H₂ ($R^2 = 0.988$) and N₂ ($R^2 = 0.999$) binding (Fig. 7c), where larger supporting ions induce more exergonic binding. The finding that larger supporting metals better poise Ni for small-molecule binding is likely a composite effect of both electronic and steric factors. Larger group 13 supporting metals show stronger Ni→M bonding interactions (Table S24b†) and shift the Ni(0/I) oxidation to more positive potentials,¹⁶ both of which should lower the Pauli repulsion associated with binding a donor (Fig. S71†). Also, larger group 13 ions force Ni to move further above the P₃-plane which should minimize the structural reorganization energy associated with small-molecule binding (Fig. S72†).

The confluence of steric and electronic effects is also supported by the fact that neither effect can solely account for the observed trends. For example, the position of Ni above the P₃-plane is identical for both Al and Ga (0.13 Å), despite their distinct differences in ΔG° for H₂/N₂ binding. Also, correlations between ΔG° for H₂/N₂ binding and Ni redox potentials are comparatively poor, whether one considers the Ni(0/I) oxidation or the Ni(-I/0) reduction potentials ($R^2 = 0.726$ to 0.891 , Fig. S68 and S69†).^{16,61} Typically Lewis acidity is expected to correlate with σ -accepting ability, yet no linear relationship exists between ΔG° for H₂/N₂ binding and the relative Lewis acidities of group 13 metals, as given by the pK_a values of the corresponding M(H₂O)₆³⁺ complexes ($R^2 = 0.02$ – 0.05 , Fig. S66†).^{62,63} It is plausible that the Lewis acidity scale based on M–OH₂ bonds, where H₂O is the Lewis base, is an inappropriate benchmark for a Ni(0) Lewis base. You and Gabbaï have recently proposed that the double-decker ligand framework, which imposes spatial constraints, may “accentuate” the influence of the group 13 ion's size.⁶⁴ It is noteworthy that ΔG° values for H₂ ($R^2 = 0.945$) and N₂ ($R^2 = 0.977$) binding correlate reasonably well with the degree of Ni→M dative bonding, as quantified by the ratio of the solid-state Ni–M bond distance to the sum of the covalent radii of Ni and M (Fig. S67†).^{65,66} This finding is consistent with the hypothesis that group 13 metals can significantly tune the reactivity of a proximal Ni center *via* a direct Ni(0)→M(III) dative interaction.

Conclusion

H₂ and N₂ binding is atypical for Ni complexes and facilitating H₂ activation to form reactive Ni(η^2 -H₂) and/or Ni–H species poses a difficult hurdle for developing homogeneous Ni catalysts for H₂ oxidation, proton reduction, and related processes. Excitingly, we find that supporting Ni with group 13 metals induces the binding of H₂ and N₂ to Ni, with H₂ binding found to be ~ 2 kcal mol⁻¹ more favorable than N₂ binding in each case. The pivotal role of the supporting metal in promoting binding to the NiML complexes is highlighted by the inability of NiLH₃, a similarly ligated mononuclear Ni center, to bind H₂ or N₂ under any conditions examined. The dramatic tuning effect of the supporting group 13 metal is illustrated by the wide range of ΔG° values for H₂ binding, which span ~ 8 kcal mol⁻¹

(including NiLH₃) and nearly cover the entire range of previously reported values. H₂ binding to NiInL (3) is more exergonic than any such equilibrium reported, allowing for solid-state characterization of (η^2 -H₂)NiInL (3-H₂) *via* neutron diffraction, which is unprecedented for an H₂ adduct of Ni or any d¹⁰ metal.

Theoretical calculations have provided important insights into the role of the supporting metal in promoting small-molecule binding. Specifically, complementary σ -interactions are proposed based on ETS-NOCV calculations, where H₂ → Ni σ -donation to the energetically-accessible Ni(4p_z)-based LUMO is the dominant binding interaction, which is likely induced by electron withdrawal from Ni *via* the dative Ni(0)→M(III) bond. The involvement of the LUMO in small-molecule binding is supported by TD-DFT and UV-Vis studies, along with the strong correlation between ΔG° values for H₂/N₂ binding and the Ni 3d_{xz}/d_{yz} → LUMO excitation energy. Notably, the size of the supporting M(III) ion correlates best with ΔG° values for H₂ and N₂ binding, with larger group 13 supporting metals inducing more favorable binding by: rendering Ni more electron-deficient, favoring stronger Ni→M interaction, and minimizing the structural reorganization energy. Moreover, thermodynamically favorable H₂ binding equilibria (In > Ga > Al) have proportionally smaller free energies of activation.

Overall, a thorough understanding of H₂ and N₂ binding to NiML complexes, and the integral role of the supporting metal therein, has been presented. Future work will explore the impact of the thermodynamics and kinetics of H₂ binding equilibria on catalytic CO₂ hydrogenation reactivity, where H₂ binding to displace formate has been found to be the rate-determining step in catalysis.^{20,67} Additionally, the generalizability of the strategy of favorably altering base-metal reactivity *via* interactions with group 13 supporting metals will be assessed, and efforts are currently underway to extend our studies to other ligand frameworks and transition metals.

Experimental section

Additional information is provided in the ESI.†

General considerations

Unless otherwise stated, all manipulations were performed under an Ar or N₂ atmosphere inside a glovebox or using standard Schlenk techniques. Standard solvents were deoxygenated by sparging with N₂ and dried by passing through activated alumina columns of a SG water solvent purification system. Deuterated solvents and HD gas (97% D content) were purchased from Cambridge Isotope Laboratories, Inc. Deuterated solvents were degassed *via* freeze–pump–thaw cycles, and either stored over activated 4 Å molecular sieves or stirred with Na–K alloy and distilled. N₂ and H₂ gases were purchased from Matheson Trigas, Inc., and a gas mixture of 10% H₂/90% Ar was purchased from Praxair, Inc. All other reagents were purchased from commercial vendors and used without purification unless otherwise noted. The ligand N(*o*-(NHCH₂PⁱPr₂)C₆H₄)₃ (abbreviated as LH₃), NiAlL (1), NiGaL (2), NiInL (3), (η^2 -H₂)NiGaL (2-H₂), (N₂)NiInL (3-N₂), (η^2 -H₂)NiInL (3-H₂), and NiLH₃ (4) were



synthesized according to the literature.^{16,34,68} ^1H and ^{31}P NMR spectra were recorded on Bruker (400 or 500 MHz) or Varian (500 MHz) spectrometers and referenced to internal residual solvent (or H_3PO_4 for ^{31}P NMR spectra). For VT NMR experiments, the temperature was calibrated using a methanol (≤ 298 K) or an ethylene glycol (> 298 K) standard. UV-Vis spectra were collected on a Cary-14 instrument. Cyclic voltammetry experiments were performed using a CHI Instruments 620D potentiostat. The one-cell setup utilized a glassy carbon working electrode, platinum wire counter electrode, and Ag/AgNO_3 reference electrode in CH_3CN .

In situ generation of $(\eta^2\text{-H}_2)\text{NiAlL}$ (abbreviated as 1-H_2)

A solution of **1** (15 mg, 19.6 μmol) in THF-d_8 (~ 0.30 mL) was added to a PEEK NMR cell and pressurized to 34 atm H_2 . The following NMR data are reported at 34 atm H_2 ; an equilibrium between **1** and 1-H_2 can also be observed under 1 atm H_2 at low T . $^1\text{H}\{^{31}\text{P}\}$ NMR (ppm, THF-d_8 , 232 K, 500 MHz): 7.34 (br, 3H, ArH), 6.87 (br, 3H, ArH), 6.34 (br, 6H, ArH), 3.00 (br, 3H, CHH'), 2.87 (br, 3H, CHH'), 2.10 (m, 6H, $\text{CH}(\text{CH}_3)_2$), 1.27 to 0.90 (36H, $\text{CH}(\text{CH}_3)_2$), -1.5 (br, 2H, $(\text{H}_2)\text{Ni}$, T_1 (min) $\leq 0.49(5)$ s at 200 K). ^{31}P NMR (ppm, 200 K, 202.4 MHz): 44.9 (in THF-d_8), or ~ 44.3 (in toluene- d_8). The lability of H_2 precluded elemental analysis from being obtained.

In situ observation of $(\text{N}_2)\text{NiAlL}$ (1-N_2) and $(\text{N}_2)\text{NiGaL}$ (2-N_2)

A solution of **1** (3.7 mg, 4.8 μmol) in toluene- d_8 (0.30 mL) was added to a PEEK NMR cell and pressurized to 51 atm N_2 . Similarly, a solution of **2** (5.0 mg, 6.2 μmol) in toluene- d_8 (0.41 mL) was added to a J. Young NMR tube under 1 atm N_2 . Neither 1-N_2 nor 2-N_2 was isolable due to lability of the N_2 ligand. Both 1-N_2 and 2-N_2 were observed in the presence of **1** and **2**, respectively, which limited ^1H NMR characterization due to overlapping resonances. For 1-N_2 : ^{31}P NMR (ppm, toluene- d_8 , 190 K, 51 atm N_2 , 202.4 MHz): ~ 32.2 . For 2-N_2 : ^{31}P NMR (ppm, toluene- d_8 , 193 K, 1 atm N_2 , 162 MHz): ~ 43.5 .

X-ray and neutron diffraction crystallographic and structure refinement details

X-ray diffraction. A gold block of $(\eta^2\text{-H}_2)\text{NiInL}$ (3-H_2) was placed onto the tip of a MiTeGen Dual-Thickness Micro-LoopTM and mounted on a Bruker Photon II CMOS diffractometer for data collection at 100(2) K (Table 6). The data collections were carried out using Mo $K\alpha$ radiation (graphite monochromator), and the data intensity was corrected for absorption and decay (SADABS).⁶⁹ Final cell constants were obtained from least-squares fits of all measured reflections. The structure was solved using SHELXT-16 and refined using SHELXL-16, which were executed from the ShelXle graphical user interface.⁷⁰ A direct-methods solution was calculated which provided most non-hydrogen atoms from the E-map. Full-matrix least-squares/difference Fourier cycles were performed to locate the remaining non-hydrogen atoms. All non-hydrogen atoms were refined with anisotropic displacement parameters. Hydrogen atoms were placed in ideal positions and refined as riding atoms with relative isotropic

Table 6 Crystallographic details for 3-H_2 (X-ray and neutron structures)

Radiation type	X-ray	Neutron
Chemical formula	$\text{C}_{39}\text{H}_{62}\text{N}_4\text{P}_3\text{InNi}$	$\text{C}_{39}\text{H}_{62}\text{N}_4\text{P}_3\text{InNi}$
F_w	853.36	853.36
Cryst syst	Orthorhombic	Orthorhombic
Space group	$P2_12_12_1$	$P2_12_12_1$
a (\AA)	12.2127(4)	12.2010(10)
b (\AA)	14.5402(5)	14.5638(12)
c (\AA)	22.5601(8)	22.547(2)
α (deg)	90	90
β (deg)	90	90
γ (deg)	90	90
V (\AA^3)	4006.1(2)	4006.4(6)
Z	4	4
λ (\AA), μ (mm^{-1})	0.71073, 1.198	0.60–3.36, 0.1570 + 0.1306 λ
T (K)	100(2)	100(2)
Θ	2.285 to 36.348	7.352 to 78.740
Reflns collected	237 431	14 303
Unique reflns	19 473	4718
Data/restraint/parameters	19 473/0/451	4718/1068/991
$R1$, $wR2$ ($I > 2\sigma(I)$)	0.0181, 0.0392	0.0620, 0.1262

displacement parameters,⁷¹ with the exception of the apical H_2 ligand in 3-H_2 , for which the H atoms were sufficiently resolved in the Fourier difference map to allow tentative placement. Images were rendered using POV-ray.⁷²

Neutron diffraction. Neutron diffraction data were collected using the TOPAZ single-crystal time-of-flight (TOF) Laue diffractometer at the Spallation Neutron Source (SNS), at Oak Ridge National Laboratory (Oak Ridge, TN).⁷³ A block-shaped crystal of 3-H_2 , with dimensions of $0.35 \times 0.30 \times 0.30$ mm, was grown from a concentrated toluene solution under 1 atm H_2 at ~ 280 K for several weeks. The crystal was mounted on the tip of a polyimide capillary using fluorinated grease and transferred to the TOPAZ goniometer for data collection at 100 K (Table 6). To ensure good coverage and redundancy, data were collected using crystal orientations optimized with CrystalPlan software⁷⁴ for optimal coverage of symmetry-equivalent reflections of the orthorhombic cell. The integrated raw Bragg intensities were obtained using the 3-D ellipsoidal Q-space integration in accordance with previously reported methods.⁷⁵ Data reduction, including neutron TOF spectrum, Lorentz, and detector efficiency corrections, was carried out with the ANVRED3 program.⁷⁶ A spherical absorption correction was applied with $\mu = 0.1570 + 0.1306 \lambda \text{ cm}^{-1}$. The reduced data were saved as SHELX HKLF2 format, in which the wavelength is recorded separately for each reflection, and data were not merged. Starting with the X-ray structure at 100 K as an input model (where all the H atoms were placed except for the H_2 ligand), the neutron crystal structure was refined using the SHELXL-14/7 program^{70,71} with RIGU restraints for the H-atoms' anisotropic displacement parameters.⁷⁷

General procedure for H_2/N_2 binding studies

A toluene- d_8 solution of NiML (15 mM) was filtered and transferred to either a J. Young NMR tube (≤ 3.8 atm) or a PEEK NMR



cell (≥ 6.8 atm).^{78,79} For high-pressure studies, the PEEK cell was sealed and connected to a high-pressure line equipped with a vacuum pump and an ISCO syringe pump. The line was purged with H₂ or N₂ gas three times. Next, the headspace was degassed by opening the PEEK cell to static vacuum (3×30 s), and H₂ or N₂ gas was then delivered to the cell from an ISCO syringe pump running constantly at the desired pressure (*i.e.* continuous gas feed). The contents of the PEEK cell were mixed using a vortex mixer for ~ 15 min prior to NMR data collection to allow for pressure stabilization and equilibration. For low-pressure studies, J. Young NMR tubes were pressurized after freeze–pump–thaw cycles. VT ³¹P NMR spectra were acquired at several different T (193 to 368 K) in all studies, with an adequate number of scans collected (typically 64 to 160 scans) to quantify the observed peak position (for fast chemical exchange) and/or the relative peak integrations (for slow chemical exchange). ³¹P spectra were typically collected with a recycle delay time of 2 s and an acquisition time of 1.68 s. In cases where quantitative integration of multiple peaks was desired, a longer delay time of 10 s was used.

CAUTION: H₂ is a highly flammable gas. Pressurized vessels must be handled with care using proper personal protective equipment. PEEK cells were employed for the high-pressure NMR studies, as described previously.^{78,79}

General procedure for kinetic studies of H₂ self-exchange

A solution of NiML (7.5 mM in 0.70 mL) was prepared in toluene-*d*₈ and transferred to a J. Young NMR tube. VT ³¹P NMR spectra were obtained at various T (≥ 7 data points, 214 K to 344 K) for samples under 1 atm Ar to determine intrinsic linewidths and chemical shifts for NiML complexes. Subsequently, after freeze–pump–thaw cycles, the VT ³¹P NMR profile was obtained under 1 atm H₂. NMR lineshape analysis was performed using the gNMR (version 5.0) program⁸⁰ to extract H₂ self-exchange rates at each T (see ESI for details[†]). ³¹P NMR spectra were typically collected with a delay time of 2 s, an acquisition time of 1.68 s, and with 160 scans.

Computational methods

Density functional theory (DFT). DFT calculations were performed using the Gaussian 09 program package.⁸¹ Four functionals (M06-L,³⁸ M06-D3,^{56,57} PBE0,⁸² and PBE0-D3^{56,82}) and five basis sets (denoted as bs0 to bs4, Table S3[†]) were evaluated. The M06-L/bs1 method (def2-TZVPP for H₂, N₂, Ni, Al, Ga, and In, with the SDD effective core potential⁸³ for In; def2-TZVP basis set for N, P; and def2-SVP for C and H atoms)⁸⁴ gave the best match to the experimental binding energies, and had been used in related prior studies.⁶⁷ Geometric structures were optimized in the gas phase at 0 K, using the crystal structure atomic coordinates as the initial geometries when available (1–3, 3-H₂, and 3-N₂). Vibrational frequency analyses were performed with the harmonic approximation to confirm the nature of all species (0 and 1 imaginary frequency for ground-state and transition-state structures, respectively). Unless otherwise noted, Gibbs free energies at 298.15 K and 1 atm were computed by adding zero-point vibrational energies and thermal corrections. Solvation effects

were considered by performing single-point calculations for all stationary points using the SMD solvation model.⁸⁵ The H₂ and N₂ binding energies for NiML were calculated according to the following equation:



In eqn (3), X_(g) is H₂ or N₂, and the standard state is defined as 1 atm for X_(g). The overall energy for eqn (3) was determined based on the gas-phase free energies for H₂ or N₂ and the Gibbs free energies with solvation (in toluene) for NiML_(solv) and X-NiML_(solv).

Energy decomposition analysis (EDA). The EDA method⁵⁸ was implemented in the Amsterdam Density Functional (ADF 2016)^{86,87} program package to study chemical bonding between H₂ and Ni in the (η^2 -H₂)NiML complexes. Single-point energy calculations were performed using PBE0-D3/TZ2P⁸⁸ on the M06-L/bs1-optimized geometries. Relativistic effects for In were included by applying the zeroth-order regular approximation (ZORA).⁸⁹

EDA breaks the total interaction energy (ΔE_{int}) between the H₂ and NiML fragments into four components, as shown in eqn (4):

$$\Delta E_{\text{int}}(\text{EDA}) = \Delta E_{\text{elstat}} + \Delta E_{\text{Pauli}} + \Delta E_{\text{orb}} + \Delta E_{\text{disp}} \quad (4)$$

ΔE_{elstat} is the attractive, quasi-classical electrostatic interaction between the electrons and the nuclei; ΔE_{Pauli} is the repulsive interaction between the occupied orbitals of the fragments; ΔE_{orb} is the interaction between the occupied molecular orbitals of one fragment and the unoccupied molecular orbitals of the other fragment; and, ΔE_{disp} corresponds to the dispersive effects between the two fragments. The extended transition state—natural orbitals for chemical valence (ETS-NOCV) method,⁵⁹ in combination with the energy decomposition scheme, was utilized to break down the orbital interaction component (ΔE_{orb}) into contributions from specific NOCV pairs (see ESI for further details[†]).

Time-dependent (TD) DFT calculations. TD-DFT calculations (M06-D3/bs4, Gaussian 09) with solvent considerations (SMD, THF) were performed to aid in assigning electronic transitions in the absorption spectra of NiLH₃ (4), NiML (1–3), (η^2 -H₂) NiML, and (N₂)NiML complexes (M = Al, Ga, In). The basis sets denoted by bs4 were used: LANL2DZ⁹⁰ for In and 6-311G(d,p)⁹¹ for all other atoms.

Conflicts of interest

The authors declare no competing financial interests.

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References

- 1 L. Alig, M. Fritz and S. Schneider, *Chem. Rev.*, 2019, **119**, 2681–2751.
- 2 S. J. C. Robinson and D. M. Heinekey, *Chem. Commun.*, 2017, **53**, 669–676.
- 3 R. H. Morris, *Acc. Chem. Res.*, 2015, **48**, 1494–1502.
- 4 P. J. Chirik, *Acc. Chem. Res.*, 2015, **48**, 1687–1695.
- 5 R. M. Bullock and M. L. Helm, *Acc. Chem. Res.*, 2015, **48**, 2017–2026.
- 6 R. H. Crabtree, *Chem. Rev.*, 2016, **116**, 8750–8769.
- 7 G. J. Kubas, *Chem. Rev.*, 2007, **107**, 4152–4205.
- 8 P. G. Jessop and R. H. Morris, *Coord. Chem. Rev.*, 1992, **121**, 155–284.
- 9 G. J. Kubas, *Catal. Lett.*, 2005, **104**, 79–101.
- 10 J. M. Millar, R. V. Kastrup, M. T. Melchior, I. T. Horvath, C. D. Hoff and R. H. Crabtree, *J. Am. Chem. Soc.*, 1990, **112**, 9643–9645.
- 11 A. A. Gonzalez and C. D. Hoff, *Inorg. Chem.*, 1989, **28**, 4295–4297.
- 12 D. G. Abrecht and B. Fultz, *J. Phys. Chem. C*, 2012, **116**, 22245–22252.
- 13 D. E. Prokopchuk, G. M. Chambers, E. D. Walter, M. T. Mock and R. M. Bullock, *J. Am. Chem. Soc.*, 2019, **141**, 1871–1876.
- 14 D. L. M. Suess, C. Tsay and J. C. Peters, *J. Am. Chem. Soc.*, 2012, **134**, 14158–14164.
- 15 M. V. Vollmer, J. Xie and C. C. Lu, *J. Am. Chem. Soc.*, 2017, **139**, 6570–6573.
- 16 R. C. Cammarota and C. C. Lu, *J. Am. Chem. Soc.*, 2015, **137**, 12486–12489.
- 17 B. R. Barnett, C. E. Moore, P. Chandrasekaran, S. Sproules, A. L. Rheingold, S. DeBeer and J. S. Figueroa, *Chem. Sci.*, 2015, **6**, 7169–7178.
- 18 T.-P. Lin, C. R. Wade, L. M. Pérez and F. P. Gabbaï, *Angew. Chem., Int. Ed.*, 2010, **49**, 6357–6360.
- 19 T.-P. Lin, R. C. Nelson, T. Wu, J. T. Miller and F. P. Gabbaï, *Chem. Sci.*, 2012, **3**, 1128–1136.
- 20 R. C. Cammarota, M. V. Vollmer, J. Xie, J. Ye, J. C. Linehan, S. A. Burgess, A. M. Appel, L. Gagliardi and C. C. Lu, *J. Am. Chem. Soc.*, 2017, **139**, 14244–14250.
- 21 J. Takaya and N. Iwasawa, *J. Am. Chem. Soc.*, 2017, **139**, 6074–6077.
- 22 P. Steinhoff, M. Paul, J. P. Schroers and M. E. Tauchert, *Dalton Trans.*, 2019, **48**, 1017–1022.
- 23 B. A. Connor, J. Rittle, D. VanderVelde and J. C. Peters, *Organometallics*, 2016, **35**, 686–690.
- 24 W. H. Harman, T. P. Lin and J. C. Peters, *Angew. Chem., Int. Ed.*, 2014, **53**, 1081–1086.
- 25 R. H. Morris and R. J. Wittebort, *Magn. Reson. Chem.*, 1997, **35**, 243–250.
- 26 P. A. Maltby, M. Schlaf, M. Steinbeck, A. J. Lough, R. H. Morris, W. T. Klooster, T. F. Koetzle and R. C. Srivastava, *J. Am. Chem. Soc.*, 1996, **118**, 5396–5407.
- 27 G. J. Kubas, C. J. Burns, J. Eckert, S. W. Johnson, A. C. Larson, P. J. Vergamini, C. J. Unkefer, G. R. K. Khalsa, S. A. Jackson and O. Eisenstein, *J. Am. Chem. Soc.*, 1993, **115**, 569–581.
- 28 N. A. Eberhardt and H. Guan, *Chem. Rev.*, 2016, **116**, 8373–8426.
- 29 W. A. Gunderson, D. L. M. Suess, H. Fong, X. Wang, C. M. Hoffmann, G. E. Cutsail, J. C. Peters and B. M. Hoffman, *J. Am. Chem. Soc.*, 2014, **136**, 14998–15009.
- 30 R. S. Rowland and R. Taylor, *J. Phys. Chem.*, 1996, **100**, 7384–7391.
- 31 J. Halpern, L. Cai, P. J. Desrosiers and Z. Lin, *J. Chem. Soc., Dalton Trans.*, 1991, 717–723.
- 32 R. H. Crabtree and D. G. Hamilton, *J. Am. Chem. Soc.*, 1986, **108**, 3124–3125.
- 33 T. A. Luther and D. M. Heinekey, *Inorg. Chem.*, 1998, **37**, 127–132.
- 34 L. J. Clouston, R. B. Siedschlag, P. A. Rudd, N. Planas, S. Hu, A. D. Miller, L. Gagliardi and C. C. Lu, *J. Am. Chem. Soc.*, 2013, **135**, 13142–13148.
- 35 M. P. Williamson, *Prog. Nucl. Magn. Reson. Spectrosc.*, 2013, **73**, 1–16.
- 36 R. G. Bryant, *J. Chem. Educ.*, 1983, **60**, 933–935.
- 37 J. Feeney, J. G. Batchelor, J. P. Albrand and G. C. K. Roberts, *J. Magn. Reson.*, 1979, **33**, 519–529.
- 38 Y. Zhao and D. G. Truhlar, *J. Chem. Phys.*, 2006, **125**, 194101.
- 39 D. M. Heinekey, M. H. Voges and D. M. Barnhart, *J. Am. Chem. Soc.*, 1996, **118**, 10792–10802.
- 40 B. E. Hauger, D. Gusev and K. G. Caulton, *J. Am. Chem. Soc.*, 1994, **116**, 208–214.
- 41 D. G. Gusev, A. B. Vymenits and V. I. Bakhmutov, *Inorg. Chem.*, 1992, **31**, 1–2.
- 42 M. Mediati, G. N. Tachibana and C. M. Jensen, *Inorg. Chem.*, 1990, **29**, 3–5.
- 43 D. G. Gusev, V. I. Bakhmutov, V. V. Grushin and M. E. Vol'pin, *Inorg. Chim. Acta*, 1990, **177**, 115–120.
- 44 K. Zhang, A. A. Gonzalez and C. D. Hoff, *J. Am. Chem. Soc.*, 1989, **111**, 3627–3632.
- 45 A. A. Gonzalez, K. Zhang, S. P. Nolan, R. Lopez de la Vega, S. L. Mukerjee, C. D. Hoff and G. J. Kubas, *Organometallics*, 1988, **7**, 2429–2435.
- 46 D. E. Prokopchuk, E. S. Wiedner, E. D. Walter, C. V. Popescu, N. A. Piro, W. S. Kassel, R. M. Bullock and M. T. Mock, *J. Am. Chem. Soc.*, 2017, **139**, 9291–9301.
- 47 P. L. Holland, *Dalton Trans.*, 2010, **39**, 5415–5425.



- 48 C. Tsay and J. C. Peters, *Chem. Sci.*, 2012, **3**, 1313–1318.
- 49 J. L. Sudmeier, J. L. Evelhoch and N. B. H. Jonsson, *J. Magn. Reson.*, 1980, **40**, 377–390.
- 50 D. S. Stephenson and G. Binsch, *J. Magn. Reson.*, 1978, **32**, 145–152.
- 51 T. Le-Husebo and C. M. Jensen, *Inorg. Chem.*, 1993, **32**, 3797–3798.
- 52 D. C. Grills, R. van Eldik, J. T. Muckerman and E. Fujita, *J. Am. Chem. Soc.*, 2006, **128**, 15728–15741.
- 53 K. M. Gramigna, D. A. Dickie, B. M. Foxman and C. M. Thomas, *ACS Catal.*, 2019, **9**, 3153–3164.
- 54 W. H. Harman and J. C. Peters, *J. Am. Chem. Soc.*, 2012, **134**, 5080–5082.
- 55 R. C. Cammarota, L. J. Clouston and C. C. Lu, *Coord. Chem. Rev.*, 2017, **334**, 100–111.
- 56 S. Grimme, J. Antony, S. Ehrlich and H. Krieg, *J. Chem. Phys.*, 2010, **132**, 154104–154122.
- 57 Y. Zhao and D. G. Truhlar, *Theor. Chem. Acc.*, 2008, **120**, 215–241.
- 58 L. Zhao, M. von Hopffgarten, D. M. Andrada and G. Frenking, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.*, 2018, **8**, e1345.
- 59 M. P. Mitoraj, A. Michalak and T. Ziegler, *J. Chem. Theory Comput.*, 2009, **5**, 962–975.
- 60 R. D. Shannon, *Acta Crystallogr., Sect. A: Cryst. Phys., Diffr., Theor. Gen. Crystallogr.*, 1976, **32**, 751–767.
- 61 M. V. Vollmer, J. Xie, R. C. Cammarota, V. G. Young, E. Bill, L. Gagliardi and C. C. Lu, *Angew. Chem., Int. Ed.*, 2018, **57**, 7815–7819.
- 62 D. D. Perrin, *Ionisation Constants of Inorganic Acids and Bases in Aqueous Solution*, Pergamon Press, Elmsford, NY, 2nd edn, 1982.
- 63 E. Y. Tsui, R. Tran, J. Yano and T. Agapie, *Nat. Chem.*, 2013, **5**, 293–299.
- 64 D. You and F. P. Gabbaï, *Trends in Chemistry*, 2019, DOI: 10.1016/j.trechm.2019.03.011.
- 65 A. Amgoune and D. Bourissou, *Chem. Commun.*, 2011, **47**, 859–871.
- 66 B. Cordero, V. Gómez, A. E. Platero-Prats, M. Revés, J. Echeverría, E. Cremades, F. Barragán and S. Alvarez, *Dalton Trans.*, 2008, **21**, 2832–2838.
- 67 J. Ye, R. C. Cammarota, J. Xie, M. V. Vollmer, D. G. Truhlar, C. J. Cramer, C. C. Lu and L. Gagliardi, *ACS Catal.*, 2018, **8**, 4955–4968.
- 68 P. A. Rudd, S. Liu, L. Gagliardi, V. G. Young and C. C. Lu, *J. Am. Chem. Soc.*, 2011, **133**, 20724–20727.
- 69 L. Krause, R. Herbst-Irmer, G. M. Sheldrick and D. Stalke, *J. Appl. Crystallogr.*, 2015, **48**, 3–10.
- 70 G. M. Sheldrick, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2008, **64**, 112–122.
- 71 G. M. Sheldrick, *Acta Crystallogr., Sect. C: Struct. Chem.*, 2015, **71**, 3–8.
- 72 *Persistence of Vision (TM) Raytracer*, Persistence of Vision Pty. Ltd., Williamstown, Victoria, Australia, 2004.
- 73 Z.-L. Xue, A. J. Ramirez-Cuesta, C. M. Brown, S. Calder, H. Cao, B. C. Chakoumakos, L. L. Daemen, A. Huq, A. I. Kolesnikov, E. Mamontov, A. A. Podlesnyak and X. Wang, *Eur. J. Inorg. Chem.*, 2019, 1065–1089.
- 74 J. Zikovsky, P. F. Peterson, X. P. P. Wang, M. Frost and C. Hoffmann, *J. Appl. Crystallogr.*, 2011, **44**, 418–423.
- 75 A. J. Schultz, M. R. V. Jorgensen, X. Wang, R. L. Mikkelsen, D. J. Mikkelsen, V. E. Lynch, P. F. Peterson, M. L. Green and C. M. Hoffmann, *J. Appl. Crystallogr.*, 2014, **47**, 915–921.
- 76 A. J. Schultz, K. Srinivasan, R. G. Teller, J. M. Williams and C. M. Lukehart, *J. Am. Chem. Soc.*, 1984, **106**, 999–1003.
- 77 A. Thorn, B. Dittrich and G. M. Sheldrick, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2012, **68**, 448–451.
- 78 C. R. Yonker and J. C. Linehan, *Prog. Nucl. Magn. Reson. Spectrosc.*, 2005, **47**, 95–109.
- 79 C. R. Yonker and J. C. Linehan, *J. Organomet. Chem.*, 2002, **650**, 249–257.
- 80 P. H. M. Budzelaar, *gNMR User Manual, 5.0*, IvorySoft, 2006.
- 81 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, *Gaussian 09, Revision E. 01*, Gaussian, Inc., Wallingford CT, 2009.
- 82 C. Adamo and V. Barone, *J. Chem. Phys.*, 1999, **110**, 6158–6170.
- 83 D. Andrae, U. Haeussermann, M. Dolg, H. Stoll and H. Preuss, *Theor. Chim. Acta*, 1990, **77**, 123–141.
- 84 F. Weigend and R. Ahlrichs, *Phys. Chem. Chem. Phys.*, 2005, **7**, 3297–3305.
- 85 A. V. Marenich, C. J. Cramer and D. G. Truhlar, *J. Phys. Chem. B*, 2009, **113**, 6378–6396.
- 86 *ADF2016, SCM, Theoretical Chemistry*, Vrije Universiteit, Amsterdam, The Netherlands, <http://www.scm.com>.
- 87 G. te Velde, F. M. Bickelhaupt, E. J. Baerends, C. Fonseca Guerra, S. J. A. van Gisbergen, J. G. Snijders and T. Ziegler, *J. Comput. Chem.*, 2001, **22**, 931–967.
- 88 E. Van Lenthe and E. J. Baerends, *J. Comput. Chem.*, 2003, **24**, 1142–1156.
- 89 E. Van Lenthe, J. Snijders and E. Baerends, *J. Chem. Phys.*, 1996, **105**, 6505–6516.
- 90 P. J. Hay and W. R. Wadt, *J. Chem. Phys.*, 1985, **82**, 299–310.
- 91 R. Krishnan, J. S. Binkley, R. Seeger and J. A. Pople, *J. Chem. Phys.*, 1980, **72**, 650–654.

