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Facile H/D Exchange at (Hetero)Aromatic Hydrocarbons Catalyzed by a Stable Trans-Dihydride N-Heterocyclic Carbene (NHC) Iron Complex

Subhash Garhwal,[†] Alexander Kaushansky,[†] Natalia Fridman,[†] Linda J. W. Shimon,[∥] and Graham de Ruiter^{†*}

[†]Schulich Faculty of Chemistry, Technion – Israel Institute of Technology, Technion City, 3200008 Haifa, Israel; ^{II}Department of Chemical Research Support, Weizmann Institute of Science, Rehovot, 7610001, Israel

Supporting Information Placeholder

ABSTRACT: Earth-abundant metal pincer complexes have played an important role in homogenous catalysis during the last ten years. Yet, despite intense research efforts, the synthesis of iron PC_{carbene}P pincer complexes has so far remained elusive. Here we report the synthesis of the first $PC_{NHC}P$ functionalized iron complex [($PC_{NHC}P$)FeCl₂] (1), and the reactivity of the corresponding transdihydride iron(II) dinitrogen complex $[(PC_{NHC}P)Fe(H)_2N_2)]$ (2). Complex 2 is stable under an atmosphere of N_2 and is highly active for hydrogen isotope exchange at (hetero)aromatic substrates under mild conditions (50 °C, N₂). With benzene- d_6 as deuterium source, easily reducible functional groups such as esters and amides are well tolerated, contributing to the overall wide substrate scope (e.g. halides, ethers, amines). DFT studies suggest a complex assisted σ -bond metathesis pathway for $C(sp^2)$ –H bond activation, which is further discussed in this study.

INTRODUCTION

Driven by their pre-eminent two-electron chemistry, the predictable reactivity and selectivity of precious metals have made them the premier choice as catalysts in many synthetic processes.¹ The growing environmental, economic, and geopolitical concerns associated with using precious metals, in conjunction with their limited availability, are strong incentives to rethink current strategies and establish more environmental friendly alternatives.² One such methodology relies on using earth-abundant metals such as cobalt,³ manganese⁴ and in particular iron,⁵ as catalyst for a variety of organic transformations.⁶ Indeed, during the past decade we have seen a resurgence of using earth-abundant metals in homogenous catalysis. One of the main reasons behind this resurgence is our ability to utilize the unique properties of earth-abundant metals (e.g. spin-state reactivity) via elaborate ligand designs.⁷

From the wide variety of available ligand architectures, pincer-type ligands have contributed tremendously to the development of earth-abundant metal catalysis.8 The most commonly encountered structural motifs within this set of ligands are those featuring an XNX (X = NR, PR₂, P(OR)₂) pincer type geometry with a central amino or pyridine donor.8 In contrast, earthabundant metal pincer complexes presenting a carbene as central donor are virtually absent from the literature,⁹ in particular those containing a PC_{NHC}P type geometry (Figure 1).¹⁰ Their absence is quite surprising as carbenes often impart distinct electronic and steric properties to the metal center.¹¹ For instance, when comparing PCP versus PNP pincer complexes of the 2nd and 3rd row transition metals, the PCP complexes featuring a central Nheterocyclic carbene (NHC) typically bind stronger to the metal center,¹² while simultaneously increasing its electron density,13 benefitting catalyst stability and reactivity.12a



Figure 1. Selected examples of first-row transition-metal $PC_{carbene}P$ pincer complexes, and the herein reported reactivity of $[(PC_{NHC}P)Fe(H)_2N_2)]$ (2).

Yet, despite these advantages no catalytic activity of iron, cobalt or manganese $PC_{NHC}P$ pincer complexes have been reported,¹⁴ while synthetic methodologies to access

these complexes for iron are currently lacking.^{10b} Considering these limitations, developing NHC-centered phosphine-functionalized pincer complexes could hold great benefits for iron-based catalysis, especially when containing metal-hydrides.¹⁵

Here we report the synthesis and characterization of novel PC_{NHC}P pincer complex [(PC_{NHC}P)FeCl₂)] (1), that upon exposure to 2.2 equiv. of NaBHEt₃ generates the first example of a stable *trans*-dihydride iron(II) dinitrogen complex $[(PC_{NHC}P)Fe(H)_2N_2)]$. (Scheme 1; 2). Iron complex 2 is highly stable at room temperature and does not readily reductively eliminate H₂ upon exposure to N_2 , which is a commonly observed deactivation pathway for other *trans*-dihydride iron(II) complexes lacking π -acidic CO ligands.^{15b, 16} The equatorial N₂ ligand in 2 is readily displaced under catalytic conditions enabling hydrogen/deuterium (H/D) exchange at (hetero)aromatic hydrocarbons with benzene- d_6 as deuterium source. Generally, the reaction occurs under mild conditions (N₂, 50 °C), and is tolerant of a variety of functional groups including, ethers, esters, amides, halides, and heterocycles. To the best of our knowledge, complex 2 is one of the very few iron-based catalysts capable of catalytic H/D exchange at heteroaromatics using a readily available deuterium source.

RESULTS AND DISCUSSION

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Synthesis of iron PC_{NHC}P pincer complexes

Realizing the lack of current synthetic methodologies for preparing earth-abundant metal PC_{NHC}P pincer complexes, we became interested in a ligand platform known as dipyrido[1,2-c;2',1'-e]imidazolin-6-ylidenes,¹⁷ whose rigid framework might allow for strong binding of earth-abundant metals. We commenced our studies by synthesizing azolium salt **A1** via a modification of a known literature procedure.^{17a} Subsequent deprotonation of **A1** with potassium *tert*-butoxide (KO^tBu) in THF, resulted in the formation of free carbene **A2** (Scheme 1).

Scheme 1. Synthesis of $PC_{NHC}P$ iron complexes 1 and 2, and the observed H/D exchange in benzene- d_6 .

tΒι ?([/]Pr)₂ ?(′Pr)₂ ?(′́Pr)₂ KO^tBu FeCl₂.1.5 THF THF, N₂, RT THF, N₂, RT 15 h Ŕ([′]Pr)₂ ^tBu ^tBu 1:88% A2: 90% Paramagnetic NaBHEt₃ THF, -110 °C - BT N₂, 2 h Hydrogen Isotope Exchange ^tBu ^tBu P(′Pr)2 P([/]Pr)₂ C₆H₆ $C_6 D_6$ (Pr)₂ ([/]Pr)₂ 3 2:70% tRu diamagnetic diam netic Addition of $FeCl_2 \cdot 1.5THF$ (1.1 equiv.) to a stirred solution of A2 in THF (15 mL), resulted in the formation of a new paramagnetic species (1) as judged by ¹H NMR spectroscopy (Figure S12). High-resolution mass spectrometry (HRMS) is consistent with the assignment of 1 as [(PC_{NHC}P)FeCl₂)] (Figure S13), which was also confirmed by X-ray crystallography.

The solid-state structure of **1** is shown in **Figure 2**, and features an iron metal center in a distorted trigonal bipyramidal geometry. The axial phosphine donors are only weakly bound to the iron metal center, which is evident from the long iron phosphine distances of 2.782(2) Å (Fe–P1) and 2.765(2) Å (Fe–P2). The iron carbene (Fe–C1) distance of 2.062(6) and the NCN angle of 102.3(5)° are typical for other iron NHCs reported in the literature.¹⁸



Figure 2. Solid state structures of $[(PC_{NHC}P)FeCl_2]$ (1, left) and $[(PC_{NHC}P)Fe(H)_2N_2]$ (2, right). Thermal ellipsoids are shown at the 30% probability level. Hydrogen atoms and co-crystallized solvent molecules are omitted for clarity.

With iron $PC_{NHC}P$ pincer complex 1 in hand, we investigated its reactivity towards a variety of hydride donors, because of the wide applicability of transitionmetal hydrides in catalysis.^{8b, 19} Addition, of two equiv. of NaBHEt₃ to a THF solution of complex 1 at -110° C, afforded a new diamagnetic species (2) that is stable for several days in solution at room temperature (Scheme 1). The ¹H NMR spectrum of complex 2, exhibits a single characteristic triplet at -8.79 ppm (2H), whose ${}^{2}J_{P-H}$ values of 43.0 Hz are consistent with a tentative assignment of 2 as the *trans*-dihydride iron complex $[(PC_{NHC}P)Fe(H_2)N_2)]$. Additional T₁ measurements (at 298K) support such a trans-dihydride assignment where the decay time of 420 ms is similar to those reported for other transition-metal trans-dihydride complexes.^{15b, 16} Definite structural assignment of 2 was provided by Xray crystallography (Figure 2). Although the crystals were not of sufficient quality to allow comparison of the bond metrics, it does allow for identification of complex **2** as $[(PC_{NHC}P)Fe(H)_2N_2)]$ with the coordinated N₂ opposite to the PC_{NHC}P carbene, forcing the two hydrides in a trans geometry. As a result, the combined spectroscopic (NMR) and crystallographic data confirm the formation of a stable classical iron(II) transdihydride, which is unprecedented.²⁰

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Hydrogen isotope exchange (HIE) at (hetero)aromatics

The ability to selectively exchange hydrogen for either deuterium or tritium is important for understanding many fundamental processes in organometallic,²¹ medicinal,²² and biological chemistry.²³ Classically, HIE is catalyzed by noble metals such as ruthenium,²⁴ rhodium,²⁵ and iridium.²⁶ In contrast, only a few studies report on the HIE with earth-abundant metals.²⁷ Given the importance of earth-abundant metal-hydride species in hydrogen isotope exchange (HIE) reactions,²⁸ we reasoned that iron complex **2** could be a straightforward entry towards facile H/D exchange at aromatic hydrocarbons.

Our studies into H/D exchange started with the observation that the hydride resonance at -8.79 ppm (t, ${}^{2}J_{P.H} = 43.0$ Hz) in complex 2 slowly disappeared after dissolving 2 in benzene- d_6 . No other changes in the ¹H NMR spectrum of 2 were observed. Analysis of the solution by ²H and ³¹P NMR spectroscopy revealed the appearance of a triplet (²H) at -8.68 ppm (${}^{2}J_{P.D} = 6.6$ Hz), and a quintet (³¹P) at 133.27 ppm (${}^{2}J_{D.P} = 6.2$ Hz), which is consistent with the formation of the di-deuteride iron(II) complex 3 (Figure S3) Similarly, dissolving 3 in benzene results to the reformation of complex 2 in quantitative yields.

The reversible H/D exchange between the solvent and complex 2 indicates reversible $C(sp^2)$ -H activation,

which is promising for allowing catalytic HIE with other hydrocarbons.^{25b, 29} As evident from Table 1, complex 2 indeed efficiently catalyzes the H/D exchange between the solvent (benzene- d_6) and a variety of (hetero)aromatic hydrocarbons. The reaction occurs under mild conditions (N₂, 50-80 °C) and typically requires less than three hours for high levels of deuterium incorporation. For example, toluene is exclusively deuterated at the meta-, and paraposition (>95%), which are the sterically most accessible positions. (Table 1; [d₃]-4). Likewise, for *m*-xylene, the *meta*-position was preferentially deuterated (98%). In both substrates, deuteration of the ortho-position was not observed. These data suggest that for toluene and mxylene, the observed regioselectivity is primarily dictated by steric factors. Computational studies (vide infra) corroborated these findings and showed that for toluene C-H bond activation at the *meta*- and *para*-positions is energetically more favorable than at the ortho-position (Table 2). Note however, that by using elevated temperatures and longer reaction times different regioselectivities can be observed (Figure S20 and S21). Notwithstanding, these results are akin to those obtained by Chirik,^{27f} and Leitner,^{24a} whose iron and ruthenium $[H_4-^{iPr}CNC)Fe(N_2)_2]$ complexes; pincer and [(PNP)Ru(H)₂(H₂)] showed comparable regioselectivity.





^aSee supporting information for experimental details. ^bYields were determined by ¹H NMR spectroscopy in the presence of an internal standard (tetraethylsilane).

Encouraged by these initial results, we sought to increase the substrate scope to include a variety of electronically and sterically differentiated substrates (**Table 1**). For example, fluorobenzene was completely

deuterated within three hours (Table 1, $[d_5]$ -6). Monitoring the reaction by ¹H and ¹⁹F NMR spectroscopy revealed that the *ortho-* and *meta*-position are preferentially deuterated (Figures S71 and S72). Only

after 30 minutes deuteration of the para-position is observed, while complete deuteration of the parapositions takes nearly three hours. These results reflect the level of deuteration in the order of *ortho* (98%) > meta (95%) > para (90%) as shown in **Table 1**. Interestingly, for the related chlorobenzene deuterium incorporation was only observed at the para-position, whereas for 4fluoroanisole incorporation was observed at the orthoposition (Table 1; $[d_1]$ -7 and $[d_2]$ -9). Clearly, besides the established steric effects, electronic effects are important contributors to the observed regioselectivity (vide infra). More detailed computational studies regarding the observed regioselectivity and mechanism for H/D exchange are presented in Figure 3 and Table 2, and are further discussed in the computational section of this manuscript. Nonetheless, the ability of 2 to efficiently deuterate chlorobenzene demonstrates that the complex 2 is stable towards reductive elimination of H₂ and subsequent oxidative addition of the aryl halide.

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Besides aryl halides, arenes bearing other electron withdrawing substituents were efficiently deuterated as well, albeit primarily at sterically accessible $C(sp^2)$ –H bonds (**Table 1**; [**d**₅]-**10**). In a similar manner, electron rich arenes were also efficiently deuterated. Dimethylaniline showed near quantitative incorporation of deuterium (>98%) at the *meta-* and *para-*positions, whereas for anisole complete deuteration was observed to yield anisole-*d*₈ within the course of 3 hours (**Table 1**).

Substrates containing reducible substituents such as esters $([d_2]-12)$ and amides $([d_3]-13)$ are tolerated as well. For instance, ethyl 4-fluorobenzoate is selectively deuterated at the ortho position (60%), while N,Ndimethylbenzamide is selectively deuterated at the metaand para position (Table 1). The difference in regioselectivity between $[d_2]$ -12 and $[d_3]$ -13 is due to different steric requirements of the directing group, although electronic effects from the fluorine atom cannot be excluded (vide infra). Unfortunately, substrates containing ketones (e.g. acetophenone, benzophenone, and/or cyclohexyl phenyl ketone) are not susceptible for catalytic H/D exchange. It is known that iron-dihydride complexes are good ketone hydrogenation catalysts.³⁰ In the absence of a suitable proton donor, the iron-hydride is nucleophilic enough to attack the ketone to generate an alkoxide intermediate,³¹ incapable of performing HIE. However, catalyst decomposition to form inactive Fe(I) or Fe(0) species cannot be excluded.³² Nonetheless, these results do not negate the fact that complex 2, tolerates several functional groups (e.g. halides, ethers, esters, amides, and/or heterocycles).

In addition to the various substituted aromatic hydrocarbons, heteroaromatic compounds are also good substrates for HIE with catalyst **2**. For instance, in 5membered heterocyclic compounds such as, N-methylpyrrole, 1-methylimidazole, 2-methylfuran, and 2,5dimethylfuran, most aromatic protons were deuterated with incorporation levels ranging from 80 to 98% (**Table 1**, **[d₂]-16–[d₂]-19**). In addition, for 2-methylfuran

deuteration of the methyl substituent was also observed to a large extend (50%). Because of the vastly different physicochemical and electronic properties of pyrroles, imidazoles, and furans, it is difficult to rationalize the regioselectivity for each substrate individually. In general, H/D exchange of hydrogen atoms ortho to a heteroatom is highly favorable and is most-likely directed by pre-coordination of the heteroatom to the metal center. Pre-coordination also explains the deuteration of the $C(sp^3)$ -H bonds in 2-methyl furan (Table 1; [d₃]-18). However, for 2,5-dimethylfuran $([d_2]-19)$ and 2,6dimethylpyridine ($[d_1]$ -15) coordination to the metal center is hampered due to steric crowding and deuteration of the $C(sp^3)$ -H bonds is not observed. Furthermore, for 5-membered heterocycles, deuteration of C-H bonds adjacent to methyl substituents is more feasible because they are sterically more accessible compared to their 6membered counterparts (e.g. compare [d₃]-16-[d₂]-19 with $([d_1]-15)$. The exception appears to be Nmethylpyrrole (Table 1), whose lack of *ortho* reactivity we are not able to explain at this stage. For 6-membered aromatic heterocycles such as 2,6-lutidine deuteration was regioselective for the sterically most accessible para proton (Table 1, [d₁]-15). For guinoline all aromatic C-H bonds were deuterated to yield quinoline- d_7 (Table 1, [d₇]-21). The different degree of deuteration in quinoline reflects the different bond dissociation energies of the various C(sp²)-H bonds.33

Comparing these results to the *state*-of-the-art, it was realized that despite the tremendous progress in homogeneous catalysis, earth-abundant metal catalyzed HIE at aromatic $C(sp^2)$ -H bonds remains extremely rare.²⁷ For example, the cobalt catalyzed H/D exchange reported by Zou and co-workers is only applicable to indoles,^{27b} while the cobalt catalysts developed by Chirik and coworkers primarily catalyze H/D exchange at (benzylic) C(sp³)-H bonds.^{27e} For iron related studies, the seminal work of Chirik and co-workers is important as it describes the first example of the iron catalyzed HIE at aromatic C(sp²)-H bonds including those of pharmaceuticals.^{27f} However, due to the instability of the *in-situ* formed dihydride, hydrogen or deuterium gas is always required to ensure catalyst stability and efficiency as shown in a recent study.^{27a} The herein reported results, however, demonstrate for the first time that such *trans*-dihydrides can be stable, ultimately leading to the deuteration of wide variety of aromatic hydrocarbons at lower temperatures with shorter reaction times and greater efficiencies,^{27f} even when compared to PNP ruthenium dihydride complexes reported by Milstein and Leitner.^{24a} However, it must be mentioned that comparison to the state-of-the-art is challenging due to the different nature of (i) the used solvents (e.g. benzene vs. THF), (ii) the deuterium source (D_2 , D_2O_2 , or C_6D_6), and (iii) the used catalyst loadings.

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Computational mechanistic investigations.

To further understand the regioselectivity, and to gain more insight into the mechanism of the H/D exchange between benzene- d_6 and (hetero)aromatic hydrocarbons, we used density functional theory (DFT) to obtain more detailed information about the relevant intermediates and transition states (**Figures 3**). To reduce the computational load, calculations were performed on a model system with R¹=Me (**Figure 3**). Additional computational details are reported in the supporting information.

Before going into the details of the herein reported H/D exchange and C–H bond activation, there are several mechanisms by which C-H bonds can be activated that include (i) oxidative addition, (ii) σ -bond metathesis, or other types of mechanisms.³⁴ Low valent (late) transition metals generally activate $C(sp^2)$ -H bonds via oxidative addition. High valent (early) transition metals - on the other hand – prefer a σ -bond metathesis pathway, because of their general lack of available d-electrons. A variant of σ -bond metathesis, σ -complex-assisted metathesis or σ -CAM, has also been developed for late transition metals to explain the facile hydrogen exchange at bound E-H (E = B, C, or Si) σ -complexes.³⁵ For example, Leitner,^{24a} Lau,^{24b} and others,³⁶ have used σ -CAM to explain metalcatalyzed H/D exchange at aliphatic and aromatic hydrocarbons.

For iron, oxidative addition of $C(sp^2)$ -H bonds is well known and has been established for nearly half a

century.³⁷ On the other hand, Eisenstein and co-workers have shown that, for iron, σ -CAM is also a plausible mechanistic pathway for H/D exchange at aromatic hydrocarbons.³⁸ Our computational studies also indicates that H/D exchange with benzene- d_6 most likely occurs via a σ -CAM based mechanism (Figure 3). Starting from 3, labeled A in Figure 3, formation of the σ -complex C is a two-step process and energetically uphill by 18.4 kcal mol⁻¹. The structure of intermediate C is shown in **Figure 3**, and features in an η^2_{C-H} interaction of benzene with the iron metal center resulting in elongation of the C–H bond by 0.032 Å. Such intermediates have also been implied for aromatic $C(sp^2)$ –H activation with ruthenium and iridium.³⁹ The C(sp²)-carbon atom engaged in the η^2_{C-H} interaction is located 0.125 Å above the plane of the PC_{NHC}P pincer ligand, while the C_b-H_b bond is rotated upward by 27.0° and located 0.420 Å above the PC_{NHC}P plane. The slight upward rotation, might indicate an attractive interaction between the axial deuteride (D1) and $H_{\rm b}$ as their calculated distance of 2.098 Å is slightly shorter than the sum of their van der Waals radii (2.200 Å).³⁸ The attractive interaction between D1 and the $C_{\rm b}$ -H_b bond is somewhat reminiscent of the hydride "cis effect" proposed by Eisenstein and Caulton,40 which describes the interaction between a metal-hydride and a coordinated dihydrogen molecule en-route towards dynamic hydrogen atom exchange. As such the "ciseffect" demonstrates important concepts that are also observed in the herein described σ -CAM (vide infra).³⁵



Figure 3. Calculated free energy profiles ($\Delta H/\Delta G$) in kcal mol⁻¹ and plausible mechanistic pathways for hydrogen isotope exchange (HIE) of 3 with benzene. Hydrogen atoms (except H_b) are omitted for clarity. For computational details, see the supporting information

The critical C-H activation step occurs via a σ -complex-assisted metathesis pathway and leads to the formation of the non-classical hydride isotopologue (**D**) with an overall energy barrier (ΔG) of 25.9 kcal mol⁻¹. The transition state structure (TS1) is shown in Figure 3 (bottom), and features an iron complex with a distorted pentagonal bipyramidal geometry. Obviously, starting from C, further distortion of the $Fe \cdots C$ axis is necessary to access the planar four-center transition state in TS1 that is typically observed for σ -CAM based mechanisms. In **TS1**, the calculated $D1 \cdots H_b$ and $Fe \cdots C_b$ distances of 1.193 and 2.106 Å are significantly shorter than those found by Leitner and Milstein,^{24a} and are indicative of a σ -CAM based mechanism with a late transition-state structure exhibiting very little oxidative addition character as shown by Eisenstein and co-workers.⁴¹ Overall, the barrier going from C to D is 7.5 kcal mol^{-1} , indicating that H/D exchange occurs quite readily once the σ -complex is formed and that the slightly elevated temperatures (50 °C) are necessary to facilitate dissociation of the N₂ ligand (Figure 3).

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The formation of non-classical hydride isotopologue **D** is also supported by NMR studies. Monitoring a solution of **2** in benzene- d_6 shows the formation of a triplet at -8.79 ppm (${}^{1}J_{H-D} = 21.1$ Hz) in the ${}^{1}H\{{}^{31}P\}$ NMR spectrum after 15 min (**Figures S1–S4**). The large value of the H–D coupling constant is indicative for the formation of a non-classical hydride intermediate, as *cis/trans* HD couplings are general small or negligible.⁴² In this process, oxidative addition from an Fe(0)/Fe(II) complex was not observed experimentally and computationally.

Rotation around the iron H–D bond (Figure 3, $D \rightarrow E$) is essentially barrierless and the transition state structure TS2 is shown in Figure 3 (bottom). The calculated bond lengths in TS2 do not differ significantly from those calculated for **D** and the complex remains essentially octahedral. After rotation, the first stage of the H/D exchange is complete (Figure 3, $A \rightarrow E$). Hereafter, a nearly identical, but reverse, pathway follows that start with σ -CAM to yield the σ -bonded deuterated benzene adduct (F). Final two-step ligand exchange between the substrate and N₂ regenerates an isotopologue of the starting catalyst (H). In general, the herein presented mechanism is similar to that proposed by Leitner and Milstein, whose ruthenium PNP complex also catalyzes H/D exchange between aromatic hydrocarbons and benzene-d₆.^{24a} Besides the mechanism outlined in **Figure** 3, a modified version is also energetically accessible (Figure S76; Path B). In this mechanism, instead from the *trans*-dideuteride, σ -CAM is initiated from the iron(II) cis-dideuteride with a very similar transition state that is only 5.2 kcal mol^{-1} higher in energy.

We also used computational studies to gain more insight into the observed regioselectivity. For a series of substituted benzenes, we calculated the energy barriers of **TS1-R** (**Table 2**; R = H, Me, NMe₂, OMe, and F). For benzene (R = H) the calculated barrier is 25.2 kcal mol⁻¹ which is slightly lower than for TS1, due to isotope effects.⁴³ Overall, as our calculations will show, the observed regioselectivity is combination of steric and electronic effects (*vide infra*). As expected strong steric effects are observed for toluene (R = Me; **Table 2**). The energy barrier for *ortho* C–H bond activation (30.4 kcal mol⁻¹) is significantly higher than for activation at the *meta*- and *para*-positions ($\Delta G = 25.9/25.2$ kcal mol⁻¹). Similar effects are observed for dimethylaniline, where C–H bond activation at the *ortho*-position has an energy barrier of 31.3 kcal mol⁻¹ (**Table 2**). These results are inline with our experimental results (**Table 1**)

Because for anisole no strong steric effects are expected and pre-coordination of the heteroatom is commonly invoked in C–H bond activation strategies, there are no clear differences in the energies for *ortho*, *meta*, or *para* C–H bond activation. As a result, complete deuteration of all C–H bonds is observed, including those present in the –OMe substituent (**Tables 1** and **2**).

Table 2. Relative energies (ΔG) in kcal mol⁻¹ of the transition state (TS1-R) for C–H exchange at the *ortho*, *meta*, and *para* positions of toluene, fluorobenzene, anisole, and dimethylaniline.^{a,b}

$L_{3}F_{e}^{P}-N_{2} \xrightarrow{H} L_{3}F_{e}^{P}-N_{2} \xrightarrow{H} L_{3} \xrightarrow{H}$			
Substituent	Position		
	Ortho	Meta	Para
R = H	25.2	25.2	25.2
R = Me	30.4	25.9	25.2
$R = NMe_2$	31.3	25.2	27.5
R = OMe	25.6	24.5	25.7
R = F	23.9	23.6	25.1

^aEnergies are reported relative to structure **A** (Figure 3). ^bSee supporting information for computational details.

Besides the obvious steric effects, electronic effects are also important. This is particularly evident in fluorinated substrates $[d_5]$ -6 and $[d_2]$ -9. We begin by noting that the overall transition barriers TS1-F for C(sp²)-H activation in fluorinated substrates are lower than those calculated for their non-fluorinated counterparts (Table 2). Despite having the strongest C-H bond, the more facile activation of fluoroarenes is wellknown and is due to the larger increase of the M-CAr bond strength relative to that of the CAr-H bond.44, 45 This is particularly true when fluorine substituents are introduced at the ortho-position as demonstrated by Jones and Perutz.⁴⁶ This pronounced *ortho*-fluorine effect is clearly seen in $[d_2]-9$, which shown nearly exclusive deuteration (~85%) at ortho-position. Similarly, for fluorobenzene ([d₅]-6), the highest level of deuteration is observed at ortho-position (98%). In contrast, the para-

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position shows a lower level of deuteration (~90%), consistent with a transition state barrier that is 1.2 kcal mol⁻¹ higher than that calculated for the *ortho*-position (**Table 2**). Besides the electronic of fluorine, other electronic effects are observed as well. To illustrate, we calculated the natural charges on the aromatic carbons and hydrogen atom H_b (**Figure S79**). As expected upon C–H bond activation, additional negative charges develop on aromatic carbon atoms (incl. C_b), while a positive charge develops on H_b. Consequently, π -donation of the coplanar NMe₂ substituent effectively destabilizes this negative charge, resulting in an overall higher activation energy for C–H bond activation (**Table 2**).

Overall, these computational studies show that for the majority of substrates the observed regioselectivity can be explained by a combination of steric and electronic factors (**Table 2**) and that a σ -CAM based mechanism is able to explain the observed HIE (**Figure 3**).

Conclusions

In summary, we have reported the synthesis and characterization of a rare *trans*-dihydride iron(II) dinitrogen complex (2) that is based on a novel $PC_{NHC}P$ pincer type motif. This compound is stable under N₂ for several days and can be used for selective H/D exchange at (hetero)aromatic hydrocarbons with benzene- d_6 as deuterium source. Deuterium incorporation typically exceeds 90% and is generally regioselective for sterically accessible $C(sp^2)$ -H bonds unless overriding electronic effects are present. Computational studies indicate that a σ -bond metathesis pathway is mainly responsible for the observed hydrogen isotope exchange (HIE), which is accessible under mild conditions (N₂, 50 °C). Overall, the herein reported results convey a convenient and straightforward method for H/D exchange that displays a wide functional group tolerance (e.g. esters, amides, halides, ethers, etc.). Furthermore, the robustness and stability of the herein reported catalyst holds great potential for other organic transformations that rely on hydride transfer or those that focus on small molecule activation. Current efforts are directed towards synthesizing the corresponding manganese, cobalt, and nickel complexes and exploring their reactivity in a variety of organic transformations.

ASSOCIATED CONTENT

Supporting Information

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

Crystal data (CIF)

xyz-coordinate files (ZIP)

Synthetic procedures, characterization data, catalysis, and computational studies (PDF)

AUTHOR INFORMATION

Corresponding Author

* Email: graham@technion.ac.il

Notes

The authors declare no competing financial interests.

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