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Experimental and Computational Studies of Hydrogen Bonding and Proton Transfer to [Cp*Fe(dppe)H].

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The present contribution reports experimental and computational investigations of the interaction between Cp*Fe(dppe)H and different proton donors (HA), focusing on the structure of the proton transfer intermediates and on the potential energy surface of the proton transfer leading to the dihydrogen complex [Cp*Fe(dppe)(H₂)]⁺. With p-nitrophenol (PNP) a UV-visible study evidences the formation of the ion pair stabilized by hydrogen bond between the non-classical cation [Cp*Fe(dppe)(H₂)]⁺ and homoconjugated anion ([AHA]⁻). With trifluoroacetic acid (TFA), the hydrogen bonded ion pair containing simple conjugate base (A-), in equilibrium with the free ions, is observed by IR spectroscopy when using a deficit of the proton donor, whereas an excess leads to the formation of the homoconjugated anion. The interaction with hexafluoroisopropanol (HFIP) was investigated quantitatively by IR spectroscopy and by ¹H and ³¹P NMR spectrometry at lowtemperatures (200-260 K) and by stopped-flow kinetics near room temperature (288-308 K). The hydrogen bond formation to give Cp*Fe(dppe)H···HA is characterized by ΔH° = -6.5±0.4 kcal mol⁻¹ and ΔS° = -18.6±1.7 cal mol⁻¹ K⁻¹ and the activation barrier for the proton transfer step, which occurs only upon intervention of a second HFIP molecule, is $\Delta H^{\dagger}_{1} = 2.6\pm0.3$ kcal mol⁻¹ and ΔS^{\ddagger}_1 = -44.5±1.1 cal mol⁻¹ K⁻¹. The computational investigation (at the DFT/B3LYP level with inclusion of

solvent effects by the polarizable continuum model) reproduces all the qualitative findings, provided the correct number of proton donor molecules are used in the model, but the proton transfer process is computed as less exothermic relative to the experiment.

Keywords: Iron, Hydride Ligand, Dihydrogen Bonding, Proton Transfer Mechanism, DFT Calculations

Introduction

Proton transfer processes to and from transition metal centers and hydride ligand sites are of fundamental importance for catalysis and are also relevant to the biochemical genesis of dihydrogen.^[1] It has now been quite firmly established that, when both a hydride ligand and a metal-based electron pair are present in the same complex, proton donors show a kinetic preference for the hydride site, [2-8] though exceptions have recently been reported from studies carried out in our laboratories.^[9, 10] It has also been quite firmly established that hydrogen bonded adducts are well defined intermediates along the proton transfer pathway.^[11-13] Thus, referring to the general Scheme 1, species II or III are usually generated faster than species V or VI, a phenomenon that is not necessarily related to the thermodynamic preference for hydrogen bond formation at the hydride (I) or metal (IV) site. Whether indeed there is a general correlation between the strength of the hydrogen bonding at a particular site and the rate of proton transfer at the same site is not yet known.

<Scheme 1>

The proton transfer kinetics to the hydride ligand has been studied for several iron subgroup complexes. A number of different acids (HBF4, CF_3CO_2H , CF_3SO_3H , HCl, HBr) were used by

Basallote et al., [14-18] but they were too strong to identify hydrogen bonded intermediates, and species like I or II have been only anticipated as transition states. More recent work, on the other hand, has provided kinetic evidenced for the importance of species **II** in the reverse deprotonation reaction for complex trans-[FeH(η^2 -H₂)(dppe)₂]⁺.^[19] Some of us have studied the kinetics of transformation of dihydrogen bonded Ι nonclassical complex II in complex into of case CpRuH(CO)(PCy₃) using fluorinated alcohols as proton donors.^[20] The activation enthalpy and entropy of the proton transfer process as well as the thermodynamic parameters for the dihydrogen bond formation step have been obtained for the protonation by perfluoro-tert-butanol (PFTB) in hexane, but low solubility of the ion-paired complex in hexane the prevented us from determining the equilibrium thermodynamic parameters of proton transfer step. The enthalpies and entropies of both the dihydrogen bond and the molecular hydrogen complex formation steps have been obtained for triphosRe(CO)₂H/PFTB,^[21] triphosRu(CO)H₂/HFIP,^[22] PP₃OsH₂/TFE,^[23] and RuH₂(dppm)₂/HFIP^[24] systems [triphos = CH₃C(CH₂PPh₂)₃; PP₃ = P(CH₂CH₂PPh₂)₃; dppm = Ph₂PCH₂PPh₂; HFIP = hexafluoroisopropanol, TFE = 2, 2, 2-trifluoroethanol]. The protonation of these hydrides is exothermic and exoentropic, the equilibrium shifting toward cationic dihydrogen complexes upon cooling. However, the proton transfer step was too fast to be studied by conventional spectroscopic methods (IR, NMR) and no

5

activation data were derived. To date, there is no example in the literature where both activation and equilibrium thermodynamic data have been determined for the same system and where a quantitative energy profile can be presented for the proton transfer process to a transition metal hydride.

Some of us have recently reported an experimental study of the protonation of complex Cp*FeH(dppe) in CH₂Cl₂ with a variety of proton donors of different acid strength (2monofluoroethanol, MFE; TFE; HFIP; PFTB; and trifluoroacetic acid, TFA).^[25] Using weaker acids, we have not only confirmed the results of Hamon et al. ^[5, 6] for the protonation with HBF₄ that the proton transfer is faster at the hydride site to give intermediate dihydrogen complex, [Cp*Fe(H₂)(dppe)]⁺, but an also found the spectroscopic signatures for the establishment of a hydrogen bond at the hydride site and have determined the thermodynamic characteristics of this interaction, whereas no evidence was obtained for the establishment of a hydrogen bond with the metal center. We have also experimentally determined that the fluorinated alcohols are able to transfer the proton only upon assistance by a second alcohol molecule. No information of this kind could be obtained in the case of trifluoroacetic acid (TFA), because of the faster rate of the proton transfer process. In addition, we have established that the isomerization yielding the final classical dihydride product [Cp*FeH₂(dppe)]⁺ occurs via an internal rearrangement process rather than via a reversible deprotonation at the

hydride site followed by a slower protonation at the metal site.^[25] This conclusion followed the key observation that the rate of isomerization is independent from the nature of the proton donor, whereas the rate of proton transfer at the hydride site increases with the hydrogen bond strength.

The main aim of the present study is to obtain more detailed information on the structure of the proton transfer intermediates and on the proton transfer potential energy surface, specifically: (i) the role of the second proton donor molecule; (ii) the hydrogen bonding status in the intermediate dihydrogen complex (i.e. as II or III) and final product (i.e. as **V** or **VI**); (iii) the energetic cost of the proton transfer process and of the subsequent internal rearrangement leading to the final dihydride product; (iv) the intimate mechanism of this rearrangement. In order to address these questions, we have carried out new experimental studies with additional proton donors that contain UV-visible and IR spectroscopic probes that are more sensitive to the chemical environment for carried the deprotonated acid. We have out careful equilibrium investigations of the proton transfer step for the Cp*Fe(dppe)H/HFIP system. We have carried out a variable temperature stopped-flow kinetic study of the same proton transfer process in order to gain information on the activation parameters for the proton transfer step. Finally, we have completed our study with a computational investigation of the energetics of hydrogen bonding at the hydride and metal

sites, as well as the potential energy surface for the proton transfer process at both sites. In this contribution, we present the new spectroscopic results, the variable temperature stopped-flow results, and the computational work on the proton transfer step and on the intimate nature of the conjugate base. An accompanying computational study of the rearrangement mechanism will be presented separately.^[26]

Experimental Section

All manipulations were carried out under an argon atmosphere by standard Schlenk techniques. The Cp*Fe(dppe)H hydride was synthesized according to the literature.^[27]

IR and UV-visible investigations

The IR measurements were performed on a "Specord M82" spectrometer (IR) on 0.1-0.15 M (for the v(OH) measurements) or 0.02-0.025 M (for the v(MH), v(CO) and v^{as} (OCO) measurements) hydride solutions in CH₂Cl₂ (0.12 cm path length) using CaF₂ cells. UV measurements were performed on Specord M-40 and Varian Cary 5 spectrophotometers in CH₂Cl₂. Low temperature IR and UV measurements were carried out by use of a cryostat (Carl Zeiss Jena) in the 200-290 K temperature range. The accuracy of the temperature adjustment was ± 0.5 K. The reagents

were mixed at low temperature and the cold mixtures were subsequently transferred into the pre-cooled cryostat.

Variable temperature UV-visible spectra of homoconjugated PNP anion were obtained for the equimolar mixture of PNP and potassium p-nitrophenolate in the presence of excess 18-crownshown by independent measurements, 6. As was PNP forms hydrogen bonded adduct with the crown-ether with band at 330-346 nm. Subtraction of this band from the spectra gives the band of homoconjugated PNP anion presented on



Figure 1.

9



Figure 1>

NMR investigations

The NMR studies were carried out in standard 5 mm-NMR tubes containing solutions of the complexes in CD₂Cl₂. The ¹H and ³¹P NMR data were collected with a Bruker AMX 400 spectrometer operating at 400.13 and 161.98 MHz respectively. The spectra were calibrated with the residual protonated solvent resonance (¹H) and with external 85% H_3PO_4 (³¹P). The conventional inversion-recovery method (180- τ -90) was used to determine the variable-temperature longitudinal relaxation time T₁. Low temperature experiments were carried out in the 180-260 K temperature range using a TV-3000 Bruker temperature unit. The apparatus was calibrated with a methanol standard. The accuracy and stability of temperature was ±1 K. All samples were allowed to equilibrate at every temperature for at least 10 min. All mixings between the alcohols and the hydride complexes were performed at low temperature.

Proton transfer equilibrium constants from NMR and UVvisible data

For the calculation of the proton transfer equilibrium constant leading from Cp*Fe(dppe)H···HOR and ROH to $[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^- [R = (CF_3)_2CH] (K_1 = k_1/k_{-1},$ see Scheme 2), the concentration of $[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^-$, and sum of Cp*Fe(dppe)H on one side, the and Cp*Fe(dppe)H···HOR, on the other side, were derived from the relative intensities measured of the significant NMR resonances (hydride ligand resonance in the ¹H NMR spectrum and phosphine resonance in the ³¹P NMR spectrum) or from the measured intensity of the UV band in comparison with the spectra of pure Cp*Fe(dppe)H and [Cp*Fe(dppe)(H₂)]⁺. The UVvisible spectrum of pure Cp*Fe(dppe)H does not change upon formation of the hydrogen bonded adduct Cp*Fe(dppe)H···HOR as verified by the stopped-flow experiment. The spectrum of pure $[Cp*Fe(dppe)(H_2)]^+$ was obtained by low-temperature (200 K) protonation with HBF4 and it is also assumed to be independent from the hydrogen bonding with the [ROHOR] - homoconjugated anion.

<Scheme 2>

The sum of Cp*Fe(dppe)H and Cp*Fe(dppe)H····HOR equilibrium concentrations could be partitioned to the individual species, given the independent knowledge of $K_{\rm H0}$ (see Results section) and the calculation of [ROH] from Equation 1. Using $K_{\rm H0}$ to express [Cp*Fe(dppe)H···HOR] as a function of [ROH] and ([Cp*Fe(dppe)H] + [Cp*Fe(dppe)H···HOR]) and insertion into Equation 1 gives a quadratic equation from which [ROH] can be calculated. From that, the calculation of K_1 is straightforward.

$$[ROH] = C_{ROH} - [Cp*Fe(dppe)H \cdots HOR] - 2$$
$$[[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^-]$$
(1)

Stopped-flow investigations

The stopped-flow kinetic runs were carried out in the temperature range $15-35^{\circ}$ C with a Hitech SF-61-DX2 apparatus coupled to a Hitech diode-array UV-visible spectrophotometer. Given the extreme air-sensitivity of the hydride compound, unacceptable results were obtained at the low concentrations required for work in a suitable absorbance range when using a regular 1 cm cell (ca. 5×10^{-4} M). This phenomenon is attributed to oxidation by oxygen diffusion through the instrument transfer lines, as confirmed by the observation of small and irreproducible signal evolutions when shooting the same

hydride solution from both syringes. Switching to a tenfold concentration and to a smaller path-length (1.5 mm) reduced the oxidation problem below acceptable noise levels. Only the data that were collected within the first 2 seconds were analyzed, yielding reproducible results. The temperature range was limited by the fact that the instruments shows leaks at $T < 15^{\circ}$ C, while it is too tight at $T > 35^{\circ}$ C. All experiments were run with the same stock solutions during the After each temperature change, the thermal same dav. equilibrium was reached within ca. 15 min and was checked by the reproducibility of the first few shots. At least 3 shots were used at each temperature to obtain the averaged results reported in the Results section. The rate data were obtained by a global fit using the SPECFIT software.^[28]

Computational details

Quantum computations were performed with the Gaussian98^[29] package at the DFT/B3LYP level.^[30-32] Core electrons of the Fe and of the P atoms were described using the effective core pseudopotentials of Hay-Wadt^[33, 34] and valence electrons were described with a standard double- ζ basis set.^[29] In the case of the P atoms, a set of d type functions was added to the standard basis functions.^[35] Carbon atoms and hydrogen atoms non-bonded to the metal together with atoms of proton donors (C, F, H) not involved in hydrogen bonds were described with a 6-31G basis set.^[36] The hydrogen atom directly bonded to the Fe atom together with hydrogen and oxygen atoms of the proton donors involved in hydrogen bonding were then described with a 6-31G(d,p) set of basis function.^[37] Solvent effects were taken into account by means of polarized continuum model (PCM) calculations^[38, 39] using standard options.^[29] Then, the free energies of solvation were computed in dichloromethane $(\Box\Box8.93)$ solvent at the geometries optimized in gas phase. The complexation energies in gas phase were also corrected from the basis set superposition errors according to the counterpoise method of Boys and Bernardi.^[40] Test calculations on the real complex Cp*Fe(dppe)H were carried out by using the IMOMM method, [41] with a program built from modified versions of Gaussian98^[29] for the quantum mechanics part and mm3(92)^[42] for the molecular mechanics part. The Cp*Fe(dhpe)H (dhpe = PH₂CH₂CH₂PH₂) plus the proton donor constituted the quantum mechanical part (QM) of the system while the four phenyl ligands were described by molecular mechanics (MM). The QM part of the calculations was done at the B3LYP level using the same basis defined previously. The MM part calculations used the mm3(92) force field.^[43] Torsional contributions involving dihedral angles with the metal center were set to zero. All of the geometrical parameters were optimized except the bond distances of atoms involved in the QM-MM frontier. The frozen values were 1.41 Å for the P-H bonds of the Cp*Fe(dhpe) in the quantum part and the crystallographic values for the P-C in the MM part.

14

a. Interaction of Cp*FeH(dppe) with PNP: UV-visible study.

The absorptions of substituted phenols in the visible spectrum are highly characteristic of their protonation and hydrogen bonding state.^[44-46] Therefore, we used *p*-nitrophenol (PNP) to distinguish between all species in equilibrium. While the visible bands of the phenol (ArOH), of its conjugate base ([ArO]⁻), and of the homoconjugated anion ([ArOHOAr]⁻) are centered at different positions as shown in Table 1, the bands of hydrogen bonded complexes $ArOH \cdots [X]^-$ and $[ArO]^- \cdots HX$, appear at intermediate positions.^[47] The exact band positions are dependent on the solvent^[46] and temperature. The values obtained in this work for CH_2Cl_2 solutions (Table 1) are in good agreement with the literature data.



Figure 1 shows the spectra of the key species obtained.

Table 1. Visible bands (λ_{\max}/nm) and their molar absorption coefficients $(\varepsilon/L mol^{-1} cm^{-1})$ of the free p-nitrophenol and its conjugate base in CH₂Cl₂.

T/K	ArOH	[ArOHOAr] ⁻	[Ar0] ⁻	
280	305	398	423	
	(8050)	(19750)	(21250)	
200	312	400	430	
	(9970)	(25250)	(27605)	

Visible spectra were recorded for CH_2Cl_2 solutions of PNP (0.001-0.003 M) in the presence of Cp*FeH(dppe) at different concentrations (PNP/Fe ratios from 1:0.1 to 1:2) and at temperatures between 200 K and room temperature. The spectra show wide bands with a complex shape, resulting from the

overlap of both the phenol in its various forms and the iron hydride complexes (both free and dihydrogen bonded). The latter bands are wide (for Cp*FeH(dppe) the half-height band width $\Delta \lambda_{\lambda_2}$ is ca. 130 nm)^[48] and have lower molar absorption coefficients (for Cp*FeH(dppe) ε (λ_{max} 388 nm) = 2370 L mol⁻¹ cm⁻ ¹ at 200K). Therefore, they contribute only in a minor way to the total absorption affecting mainly quantitative results. An analysis of the spectral changes indicates the overlap of three bands, with maxima centered respectively at 312, 340 and 380 nm. On the basis of literature precedents, the following assignments could be made: on the acid side, the band centered at 312 nm is assigned to free PNP and the band at 340 nm is the nonclassical dihydrogen bonded complex assigned to $Cp^*(dppe)Fe-H\cdots$ HOAr (Ar = $p-C_6H_4NO_2$). The 28 nm red shift is caused by the effect of hydrogen bonding on the electronic absorption by the phenol chromophore. On the conjugate base side, the band at 380 nm is attributed to a hydrogen bonded phenolate ion, since this is blue-shifted from the free phenolate band by ca. 50 nm. The absence of free phenolate is signaled by the absence of a band at 430 nm. There are two possibilities for this species: the hydrogen-bonded ion pair $[Cp*(dppe)Fe(H_2)]^+ \cdots [OAr]^-$, and the homoconjugated anion [ArOHOAr]-, and the latter could either be free or further hydrogen bonded to the cationic dihydrogen complex.

Upon increasing the Cp*Fe(dppe)H amount at constant initial phenol concentration and at constant temperature, the bands at





Figure 2. The plot of the intensity changes at 380 nm vs the Cp*(dppe)FeH mole fraction (Equation 2) gives a break point for a mole fraction of (or near) 0.3, indicating a 1:2 binding stoichiometry for the ionic species,



Figure 3.^[49] The blue-shift of this band relative to free $[ArOHOAr]^-$, see Table 1, suggests further hydrogen bonding of homoconjugated anion with the cationic dihydrogen complex, $[Cp^*(dppe)Fe(H_2)]^+...[ArOHOAr]^-$.

see



Figure 2>



Figure 3>

Mole fraction = $C_{FeH} / [C_{FeH} + C_{PNP}]$ (2)

The spectral changes are fully reversible in the 200-260 K temperature range, showing that no significant isomerization to the classical dihydride complex occurs below 260 K in agreement with our previous report.^[25] Upon lowering the temperature, the band of the free phenol decreases and those of hydrogen-bonded phenol (340 nm) and hydrogen bonded ion





Figure 4. Note that neither the free phenolate band, expected at 420-430 nm, nor the band of homoconjugated anion [ArOHOAr]⁻, expected at ca. 400 nm, is observed, suggesting that hydrogen bonded ion pair does not essentially dissociate below 260 K. These qualitative observations are in agreement with the equilibria depicted in Scheme 2 ($R = p-O_2N-C_6H_4$) and with the exothermicity for both the hydrogen bond formation and for the proton transfer steps.



Figure 4>

The exothermicity of the hydrogen-bond formation falls in line with the H-bond strength previously determined with the MFE and proton donors (-4.6 and -5.9 kcal mol⁻¹, TFE respectively) and with the fact that PNP has a proton donor ability and acidity ($P_i = 1.27^{[50]}$ and $pK_a = 10.8^{[51]}$ or $10.4^{[52]}$ in DMSO) comparable to those of PFTB (P_i = 1.33 and pK_a = 10.7^[51] in DMSO). Therefore, the strength of the H-bonding interaction with PNP is expected to be even greater. The exothermicity of the proton transfer step is also in qualitative agreement with the thermodynamic parameters

previously determined for the Cp*Fe(dppe)H/HFIP system (see also later).

b. Interaction of Cp*FeH(dppe) with the TFA: IR study.

The interaction of Cp*FeH(dppe) with TFA was previously investigated kinetically, but the initial proton transfer is too fast to measure within the stopped-flow time constraints. Only the rate of isomerization of the intermediate dihydrogen complex was kinetically accessible, while the initial proton transfer is quantitative within the minimum time lapse for the first measurement (ca. 1 ms). Trifluoroacetate is a weaker base than *p*-nitrophenolate or $[(CF_3)_2CHO]^-$ and gives weaker bonding with the dihydrogen cation.^[20] Although the trifluoroacetate anion is colorless preventing the use of UVvisible spectroscopy, IR spectroscopy can be conveniently used in this case because the $v^{as}(OCO)$ bands are sufficiently diagnostic of the hydrogen bonding state of the TFA anion. The IR spectrum in the carbonyl stretching region for a 2:1

in



Figure 5(a). Two new bands at 1692 and 1713 $\rm cm^{-1}$, assigned respectively to the free and hydrogen-bonded [CF3COO] - anion,

are observed. At the same time, the acid band at 1780 cm⁻¹ is completely consumed and the v(FeH) band at 1840 cm⁻¹ has half the intensity of a stock solution with the same concentration. When using a 5-fold excess of the acid, no bands attributable to the free anion or to the hydrogen-bonded ion pair are



Figure 5(b), whereas a wide and low intensity band is observed at 1620 $\rm cm^{-1}.$ This corresponds to the free

homoconjugated ion, in which $[CF_3COO]^-$ anion is bonded with two TFA molecules.^[53] Evidently, since TFA is a much stronger acid, the initial hydrogen bond between TFA and the hydride complex is not observed, because proton transfer occurs very rapidly and its equilibrium position leaves no measurable amounts of the hydride precursor in solution. On the other hand, the trifluoroacetate anion is a relatively weak base, being energetically stabilized by resonance. Thus, it is a weak proton acceptor for hydrogen bonding and consequently it is present in large proportions as the free base in solution, in equilibrium with the hydrogen bonded base. In the presence of excess acid, the only carbonyl species present in solution are CF₃COO(H and $[CF_3COO(HOOCCF_3)_2]^-$.



c. Interaction of Cp*FeH(dppe) with HFIP: thermodynamics of the hydrogen-bond formation.

Our previous studies of hydrogen-bonding were limited to the proton donors MFE and TFE, for which proton transfer does not occur at all or very slowly at room temperature.^[25] By carrying out IR studies in the v_{OH} region at low temperatures (200-280 K) according to our established protocol, we have now obtained the thermodynamic parameters for the Cp*Fe(dppe)H-HFIP hydrogen-bonding interaction ($\Delta H^{\circ} = -6.5\pm0.4$ kcal mol⁻¹, $\Delta S^{\circ} = -18.6\pm1.7$ cal mol⁻¹ K⁻¹), see



Figure 6. These parameters indicate stronger bonding for HFIP relative to TFE ($\Delta H^{\circ} = -5.4\pm0.3$ kcal mol⁻¹ and $\Delta S^{\circ} = -13.6\pm0.9$ cal mol⁻¹ K⁻¹),^[25] as expected.

<



Figure 6>

The same investigation could not be carried out for the hydrogen-bonding interaction with PNP, because this phenol (whose acidity and proton donor strength are close to those of PFTB as shown above) yields proton transfer processes, as indeed is evident from the low temperature UV-visible spectra (e.g.



Figure 2).

d. Interaction of Cp*FeH(dppe) with the HFIP: thermodynamics of the proton transfer equilibrium.

We have previously analyzed the proton transfer equilibrium for the Cp*FeH(dppe) with the HFIP system by UV-visible spectroscopy, but the analysis did not take into account the hydrogen bonding equilibrium [$K_{\rm H0}$ in Scheme 2, R = (CF₃)₂CH] for the starting material. The knowledge that is now available for this equilibrium (see previous section), allows us to recalculate the equilibrium and thermodynamic parameters for

see

the proton transfer step. However, knowledge is still lacking about the hydrogen bonding equilibrium for the ionic dihydrogen complex product [$K_{\rm H1}$ in Scheme 3, R = (CF₃)₂CH]. The hydrogen bond in the product ion pair is expected to be stronger relative to that of the neutral precursor, because of the stronger Coulombic component for the charge separated species. Moreover [(CF₃)₂CHO]⁻ is stronger base than pnitrophenolate, for which no dissociation of hydrogen bonded ion pair was observed (see above). Thus, the product is likely to remain as a hydrogen bonded ion pair in the relatively nonpolar dichloromethane solvent, especially in the low temperature range used for the equilibrium measurements (200-Under this hypothesis, the equilibrium $K_{\rm H1}$ may be 260K). neglected, yielding K_1 (= k_1/k_{-1}).

A van't Hoff analysis of the K_1 constants derived from the previously published^[25] UV-visible data yields $\Delta H = -5.5\pm0.3$ kcal mol⁻¹ and $\Delta S = -13.0\pm0.6$ cal mol⁻¹ K⁻¹. It is necessary to point out that, besides the neglect of the $K_{\rm H1}$ equilibrium, this analysis also makes use of the hypothesis of a temperature- and anion-independent UV-visible spectrum for species [Cp*(dppe)Fe(H₂)]⁺···[ROHOR]⁻ (see Experimental section and Supporting Information).

< Scheme 3 >

alternative way to analyse the proton transfer An equilibrium is provided by NMR spectrometry. Our previous contribution included ¹H and ³¹P NMR investigations of the interaction between Cp*FeH(dppe) and ROH (TFE, HFIP and PFTB), [25] but did not include the accurate measurement of the proton transfer equilibrium position as a function of temperature. The formation of the hydrogen bond occurs without energy barrier, thus the NMR resonances of free and an hydrogen-bonded hydride complexes are in the fast exchange regime at all temperatures, as are those of the dihydrogen complex $[Cp*Fe(dppe)(H_2)]^+$ and that of its hydrogen bonded adduct [Cp*Fe(dppe)(H₂)]^{+...} [ROHOR]⁻. These are seen at δ ca. -17.3 and ca. -12.5 in the $^{1}\mathrm{H}$ NMR spectrum and at δ ca. 93.8 and ca. 108 in the ³¹P NMR spectrum, respectively. The distinction of these two resonances, on the other hand, shows that the proton transfer equilibrium is in the slow exchange positions of these peaks regime. The do not shift significantly with temperature and with the nature or concentration of the proton donor. Thus, the chemical shifts cannot be used as indicators for the presence and extent of hydrogen bonding, neither for the starting neutral hydride complex, nor for the cationic nonclassical complex.

The measured relative intensities of the significant resonances (hydride ligand resonance in the ¹H NMR spectrum and phosphine resonance in the ³¹P NMR spectrum) yield the equilibrium concentration of the various species reported in

34

Table 2, as detailed in the Experimental part. The experiment cannot give equilibrium data at temperatures above 270 K, because of the incipient isomerization to the classical dihydride product. We should mention here that the resonances of the neutral hydride complex were previously reported to broaden extensively as temperature was raised, a phenomenon attributed to a fast self-exhange with the that was adventitious product of one-electron oxidation.^[25] The present experiment was carried out on a recrystallized, pure sample and under rigorous conditions. The measured ¹H NMR resonance for the neutral hydride complex was sharp, with a well resolved P coupling and much longer T_1 values relative to the previous report.

Table 2. Concentrations of different species as a function of temperature from the ¹H and ³¹P NMR study of the reaction between Cp*FeH(dppe) and HFIP.^a

T	¹ H spectra			³¹ P spectra					
	[FeH] ^b	[FeH···H] c	[Fe(H ₂)] ^d	K_1	[FeH] ^b	[FeH···H] ^c	[Fe(H ₂)] ^d	K_1	
220	0.0323	0.0224	0.0127	125.17	0.0343	0.0204	0.0141	171.33	
240	0.0373	0.0174	0.0120	72.58	0.0379	0.0168	0.0126	82.28	
250	0.0393	0.0154	0.0109	55.10	0.0397	0.0150	0.0114	60.16	
260	0.0411	0.0136	0.0095	41.31	0.0411	0.0136	0.0096	41.80	
270	0.0430	0.0117	0.0082	32.82	0.0431	0.0116	0.0084	34.30	
$^{a}C_{Fe} = C_{HFIP} = 0.0547 \text{ M}.$ $^{b}[FeH] = [Cp*Fe(dppe)H].$ $^{c}[FeH \cdot \cdot \cdot H] = -$									

 $[[]Cp*Fe(dppe)H \cdots HOR]. \quad [Fe(H_2)] = ([Cp*Fe(dppe)(H_2)]^+] + [[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^-]).$

As Table 2 shows, the integration of both the ¹H and the ³¹P data leads to very similar concentration data. Again, under the assumption that the $K_{\rm H1}$ equilibrium favors the hydrogen
bonded adduct quantitatively (i.e. $[Fe(H_2)]$ in Table 2 corresponds to the concentration of the hydrogen bonded species, $[[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^-]$), the derived values of K_1 in Table 2 correspond to the true equilibrium constant for the proton transfer process, k_1/k_{-1} . The van't Hoff analysis on K_1 yields $\Delta H_1 = -3.2\pm0.1$ kcal mol⁻¹ and $\Delta S_1 =$ -4.8 ± 0.4 cal mol⁻¹ K⁻¹ from the ¹H NMR data; $\Delta H_1 = -3.9\pm0.1$ kcal mol⁻¹ and $\Delta S_1 = -7.4\pm0.4$ cal mol⁻¹ K⁻¹ from the ³¹P NMR data. These values are rather close to those established from the low-temperature UV-visible data. Averages of the values obtained with the three different methods are: $\Delta H_1 = -4.2\pm1.2$ kcal mol⁻¹; $\Delta S_1 = -8\pm4$ cal mol⁻¹ K⁻¹.

e. Interaction of Cp*FeH(dppe) with the HFIP: proton transfer activation barrier.

The proton transfer from HFIP to Cp*FeH(dppe) has been investigated by the stopped-flow method in the temperature range 288-308 K. Our previous kinetic investigation of this process was limited to room temperature.^[25] The transformation consists in the establishment of the proton transfer equilibrium, followed by the rearrangement to the classical dihydride product. As already established by the previous study, the hydrogen bond formations ($K_{\rm H0}$ and $K_{\rm H1}$) are instantaneous and equilibrated, the proton transfer step (k_1 , k_{-1} ; first measurable rate) is equilibrated and [R_FOH]- dependent, whereas the internal rearrangement from dihydrogen complex to classical dihydride (k_2 , second measurable rate) is irreversible and [R_FOH]-independent.

The values of k_1 and k_{-1} were previously obtained from the slope and the intercept of the k_{obs} vs. [R_FOH] plot for each alcohol. We have now taken a more rigorous approach, consisting of the use of the complete model $A \rightarrow$ B (k_{1obs}), $B \rightarrow$ A (k_{-1obs}) and $B \rightarrow C$ (k_2) . The use of SPECFIT^[28] allows in principle the independent determination of the values of the three rate constants for each experiment. The A kinetic species is the equilibrium mixture of free and hydrogen-bonded Cp*Fe(dppe)H, whereas B and C are the equilibrium mixtures of free and hydrogen-bonded intermediate dihydrogen complex and final dihydride product, respectively. The expression relating the observed rate constant k_{1obs} to the true rate constants k_1 and the equilibrium constant $K_{\rm H0}$ is given in Equation 3. The value of k_{-1obs} corresponds to k_{-1} under the above approximation of quantitative formation of the $[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^-$ hydrogen bond.

$$k_{1obs} = k_1 K_{H0} [ROH]^2 / (1 + K_{H0} [ROH])$$
 (3)

This analysis method has the advantage of not requiring the use of different ROH concentrations at each temperature to determine the individual rate constants. However, the analysis requires precise knowledge of the spectrum of the

intermediate species B. In fact, the three rate constants are highly correlated with each other and with the position of the proton transfer equilibrium, thus equally excellent data fits result from analyses carried out with different B spectra that reflect different equilibrium situations. The correct fit is only obtained if the B spectrum is that of the pure dihydrogen Of course, this intermediate species cannot be complex. generated in solution in a pure state, in presence of the same counterion and under the same temperature conditions of the kinetic measurement. In order to circumvent this problem, we different solutions of the "pure" have generated several dihydrogen complex under different conditions and recorded their spectra, and we also constructed spectra of the pure complex with the same counterion as described in the Supporting Information. Consistent results were obtained by using five different spectra, but only limited to the activation parameters related to the forward rate constant k_1 . The most reliable results are believed to be ΔH^{\ddagger}_1 = 2.6±0.3 kcal mol⁻¹ and ΔS^{\ddagger}_1 = -44.5±1.1 cal mol⁻¹ K⁻¹.^[54] The full set of results is available as Supporting Information. The activation parameters for the k_{-1} and k_2 steps turned out to be extremely sensitive to the nature of the B spectrum. Their discussion is not warranted here (see Supporting Information). The activation parameters for the reverse proton transfer step (k_{-1}) may be more appropriately estimated from the independent knowledge of the activation parameters of the forward step plus the equilibrium parameters obtained from the UV-visible or NMR studies (previous section). The activation parameter for the isomerization step (k_2) will be measured independently and more accurately using a stronger proton donor, and the results will be reported in a separate contribution.^[26]

In conclusion, the combination of the various techniques used to investigate the interaction between Cp*Fe(dppe)H and HFIP, to yield $[Cp*Fe(dppe)(H_2)]^+$ and $[ROHOR]^-$ (R = $(CF_3)_2CH)$, yields the following information: (i) the equilibrated proton transfer process occurs starting from the hydrogen-bonded adduct Cp*Fe(dppe)H···HOR only after intervention of a second ROH molecule and leads to the hydrogen-bonded adduct $[Cp*Fe(dppe)(H_2)]^+ \cdots [ROHOR]^-;$ (*ii*) this product then equilibrates with the free ions but this equilibrium most probably lies on the side of the hydrogen bonded ion pair, at least in dichloromethane at low temperatures; (iii) the proton transfer process is exothermic (ΔH° between -3 and -5 kcal mol⁻ ¹) and with a negative entropy (ΔS° between -5 and -13 cal mol⁻ ¹ K⁻¹); (*iv*) the proton transfer has a low activation enthalpy $(2.6\pm0.3 \text{ kcal mol}^{-1})$ and a very negative activation entropy (-44.5 \pm 1.1 cal mol⁻¹ K⁻¹).

f. Theoretical Study.

Theoretical calculations have been carried out on the reaction between models of the Cp*Fe(dppe)H complex and

different proton donors HA in order to highlight some of the governing factors of each step of the protonation reaction.^[20]. The presentation of the theoretical results is divided in two parts. The first one will be devoted to the thermodynamics of the formation of the initial hydrogen bonded complexes, both in gas phase and in dichloromethane (DCM). In the second part, we will focus on the thermodynamics and the kinetics of the proton transfer process converting the hydrogen bonded complexed pairing effects are also discussed in this section.

f.1. Determination of the hydrogen bonding site.

We have considered the interaction of the Cp*Fe(dhpe)H(dhpe = $H_2PCH_2CH_2PH_2$) model complex with the following set of proton donors with increasing acidity: MFE < TFE < HFIP < PFTB < TFA, i.e. the proton donors that were used in our previous experimental investigation.^[25] The formation of the hydrogen bond was investigated at both the hydride and metal sites. The



optimized geometries of the hydrogen bonded adducts are







80

2.688 Å

0





presented in

Figure 7 with the associated parameters in Table 3.





Table	3. Parameters		describing the		he hyd	hydrogen		bonding	
	interac	tion at	the	hydride	and i	metal	sites	for	complex
	Cp*Fe(d	hpe)H.							

HA		$d(H \cdots X)^{a}$	Δd	Δd	$\Delta v (0 - 1) b$	ΔE	ΔE	ΔG	$\Delta H^{\circ f}$
		21)	(Fe-H)5	(O-H) ^D	H) ²	Gase	Gas BSSE ^d	DCMe	
		Å	Å	Å	Cm^{-1}	Kcal mol ⁻¹	kcal mol ⁻¹	kcal mol ⁻¹	kcal mol ⁻¹
MFE	Ha H · · ·	1.617	0.007	0.016	-330	-10.6	-6.1	-3.7	-5.7
	${\tt H}\cdot\cdot\cdot {\tt Fe^h}$	2.949	-0.004	0.006	-122	-6.8	0.2	-0.7	-2.6
TFE	Ha Ha	1.584	0.012	0.021	-432	-12.2	-6.1	-4.1	-6.8
	$\texttt{H}\cdot\cdot\cdot$ Fe ^h	2.688	-0.004	0.012	-257	-7.2	-1.0	0.7	-4.8
HFIP	На Н · · ·	1.513	0.016	0.027	-571	-13.8	-7.4	-4.2	-8.0
	${\tt H}\cdot\cdot\cdot{\tt Fe^h}$	2.593	-0.005	0.015	-361	-8.1	-0.9	1.2	-6.0
PFTB	На Н · · ·	1.381	0.014	0.045	-898	-14.3	-7.6	-3.7	_ 10.0
	${\tt H}\cdot\cdot\cdot{\tt Fe^h}$	2.675	-0.006	0.017	-353	-7.2	0.6	3.6	-7.0
TFA	Ha H · · ·	1.374	0.015	0.048	-954	-14.5	-10.7	-5.9	_ 10.4
	${\tt H}\cdot\cdot\cdot {\tt Fe^h}$	2.418	-0.007	0.021	-701	-9.1	-4.3	-0.1	-8.9

a X = (H, Fe)

 $^{\rm b}$ Difference between the value in the free compounds and in the hydrogen bonded adduct

 $^{\rm c}$ Complexation energy in the gas phase.

 $^{\rm d}$ Complexation energy in the gas phase corrected by the basis set superposition error.

^e Complexation free energy in dichloromethane.

 $^{\rm f}$ Enthalpy of the hydrogen bond formation from Iogansen's empirical rule. $^{\rm [55]}$

^g Hydride site

 $^{\rm h}$ Metal site

The geometrical parameters that describe the hydrogen bond are quite similar to those computed $previously^{[20]}$ for CpRuH(CO)(PH₃), as a model of the CpRuH(CO)(PCy)₃ complex, interacting with moderate proton donors. The computed dihydrogen bonds $(AH \cdots HFe)$ are slightly shorter than those reported for the CpRuH(CO)(PH₃) model complex due to the higher basicity of Cp*Fe(dppe) (E_j = 1.35 for Cp*Fe(dppe)H^[25] compared to E_j = 1.02 for CpRuH(CO)(PCy₃).^[12] The formation of the hydrogen bonds is reflected in the lengthening of the OH bond distances (Δd (OH)) from their values in the isolated proton donors and consequently by the shift of v(O-H), see Table 3. Note that Δd (OH) is greater for the hydride site, signaling stronger bonding. The binding of the proton donors at the hydride site also produces a Fe-H bond lengthening, which follows the order of the acidic strength of the proton donor. On the other hand, the Fe-H bond is slightly shortened by the binding at metal site.

The energy changes associated with the hydrogen bond formation are more negative for the hydride site than for the metal site for all the proton donors (Table 3). This trend is observed both in gas phase and in DCM solvent. ΔG values at the metal site are positive for TFE, HFIP and PFTB, and very slightly negative for MFE and TFA, meaning that the corresponding hydrogen bonded complexes presumably do not exist in solution. The gas phase complexation energies of the different proton donors at the hydride site follow the acidic strength order. The same trend is found at the Fe site, except for the PFTB molecule. In comparison with the other proton donors, PFTB is a quite bulky molecule due to the three CF_3 groups, preventing the PFTB to reach easily the Fe site. The complexation energies are in the same range as those obtained for the same kind of interactions in other hydride systems.^[56-58]

The hydrogen bond formation enthalpies, ΔH° , reported in Table 3 are calculated by means of the Iogansen's empirical rule,^[55] using the computed OH frequency shifts. There is rather good agreement between the computed hydrogen bonding enthalpies and those obtained from the IR experiments (-4.6 kcal mol⁻¹ for MFE, -5.9 kcal mol⁻¹ for TFE).^[25] Our computed ΔH° values are slightly more negative, possibly because the experimental values are obtained in the dichloromethane solvent whereas the frequency calculations were carried out in the gas phase, where the interaction between the proton donor and the Cp*Fe(dhpe)H complex is stronger. As expected, the more acidic the proton donor, the stronger the interaction (as reflected by more negative values of ΔH°). This is in agreement with the experimental trends of ΔH° for MFE, TFE and In addition, bonding energies corrected by the basis HFIP. set superposition error (BSSE) have been also calculated because previous results concerning M-H···H-OR hydrogen bonds showed that the BSSE can be very important in this type of Indeed, the BSSE energy can be up to 50% of the systems.^[59] interaction energy both at the hydride and the metal site. Consequently, the BSSE-corrected energies at the hydride site

are close to the enthalpies of formation of the hydrogen bond. At the metal site, the BSSE causes the complexation energies to be close to zero or even positive, meaning that the BSSE is the main component of the complexation energies. These first computational results identify the hydride site of the Cp*Fe(dhpe)H complex as the initial bonding site for all proton donors prior to the proton transfer process, in agreement with the experimental results on the Cp*Fe(dppe)H complex.^[25]

We have checked the suitability of the model used above by carrying out additional calculations on different models of the iron complex with TFE as proton donor, see Table 4. Whatever the model used, the complexation energy is always more negative at the hydride site than at the metal site. At the hydride site, going from CpFe(dhpe)H to Cp*Fe(dhpe)H, the complexation energy is slighly more negative because the electron richer Cp* ligand enhances the basicity of the hydride ligand. Adding the phenyl on the P atoms at а molecular mechanics level does not change the energy in a significant way, meaning that there is no steric hindrance at the hydride site caused by the phenyl groups. In fact, the Cp* is slightly tilted away from the hydride site and the access to the proton acceptor site is consequently easier at the



Figure 7). In order to introduce the electronic effects of the phenyl substituents while keeping the computation

affordable, B3LYP interaction energies on the real Cp*Fe(dppe)H system have been calculated by means of full quantum mechanical single point calculations at the IMOMM optimized geometries. The reliability of such approach has been previously discussed.^[60-62] The interaction energy at the hydride site remains almost unchanged, showing that the electronic effect of the phenyl substituents is also small. The results at the metal site contrast with those at the hydride site. CpFe(dhpe)H and Cp*Fe(dhpe)H model complexes lead to comparable energies, but adding phenyl ligands at the IMOMM level decreases considerably the interaction energy. This highlights the steric hindrance caused by the phenyl substituents of the dppe ligand when the proton donor comes close to the metal site. The electronic effects of the phenyl groups, on the other hand, seem less important as for the case of the hydride site, as shown by the similar values of the IMOMM and IMOMM/B3LYP interaction energies.

Table 4. Gas phase hydrogen bond energies (in kcal mol⁻¹) at hydride and metal sites with TFE for different models of the real Cp*Fe(dppe)H complex.

	CpFe (dhpe) H B3LYP	Cp*Fe(dhp e)H B3LYP	Cp*Fe(dpp e)H IMOMM	Cp*Fe(dppe)H IMOMM/B3LYP ^a
ΔE Gas	-10.6	-12.2	-12.8	-12.3
$h \cdots h$ ΔE Gas	-7.5	-7.2	-4.4	-3.2

^a Interaction energies computed at the B3LYP level on the IMOMM optimized geometries.

As a conclusion of this calibration study, Cp*Fe(dhpe)H seems a good model of the real system for what concerns the interaction energies at the hydride site, whereas it overestimates the interaction energies at the metal site, mainly due to the neglect of the steric effects introduced by the phenyl substituents.

f.2. Proton transfer for TFA and HFIP.

In some of the proton transfer investigations we shall use a system comprising two proton donor molecules, in line with the experimental results of the kinetics study and with previous computations on other systems.^[20] This considerably enlarges the size of the system. Therefore, for computational limitations and in order to keep the computations affordable, we have only considered the CpFe(dhpe)H model in this part. However, we have first carried out test computations in order to check how the computational level and the chosen model affect the gas phase hydride proton affinity (PA), i.e. the energy change involved in Equation 4, for the monohydride iron complex. The results are reported in Table 5.

 $[Fe]-H + H^{+} \rightarrow [[Fe](\eta^{2}-H_{2})]^{+}$ (4)

Table 5. Proton affinity of the monohydride complex

	Proton affinity (kcal mol ⁻¹)
CpFe(dhpe)H/B3LYP	250.1 (251.1ª)
CpFe(dhpe) $H/CCSDT^{b}$	255.1
Cp*Fe(dhpe)/B3LYP	258.7 (259.4ª)
Cp*Fe(dppe)H/IMOMM	256.5
Cp*Fe(dppe)H/B3LYP	267.1

^aComputed adding polarization functions on the carbon atoms of the Cp ring. ^b Single point CCSDT computation on DFT/B3LYP optimized geometries.

The B3LYP level of computation with the basis set used appears suitable for our purpose. CCSDT proton affinities are close to the B3LYP results. The effect of adding polarization functions on the carbon atoms of the Cp can be considered as not important, because the values only change by about 1.0 kcal mol⁻¹. The proton affinity increases on going from the simplest to the largest models. Substitution of Cp by the more basic Cp* ligand increase the PA by 8.6 kcal mol⁻¹. The electronic effect of the phenyl substituents further increases the PA by a similar amount (8.6 kcal mol⁻¹). The model system adopted presents a slightly lower proton affinity than the real system, but the difference amounts to only 7%.



(a) TFA. Starting from the hydrogen bonded adductsresented in the previous section

(b) Figure 7) we have studied the proton transfer process both to the hydride and to the metal site. We started by optimizing the geometry of the final products: the ion pair comprising the dihydrogen complex (protonation at the hydride site) or dihydride complex (protonation at the metal site) and $[CF_3COO]^-$.

In the gas phase, both ion pairs are found as minima (



Figure 8). In the optimized structures the O····H distances are 2.053 Å and 1.849 Å at hydride and metal site, respectively. So, the proton transfer seems more accomplished at the hydride site, highlighting the stronger basicity of the hydride ligand relative to the metal. One must notice that ion pairs were previously reported as stable structure along the pathway to the protonation of metal hydrides only when a second molecule of a moderate proton donor was involved in the transfer, by homoconjugated pairing effect.^[20] Here, a second proton donor molecule is not required to obtain a stable ion pair. This is in good agreement with the experimental result when protonation of the Cp*Fe(dppe)H complex is carried out with low concentration of TFA (section b).



Figure 8>



Figure 9 reports the energetic profile for the proton transfer from one molecule of TFA to CpFe(dhpe)H. The O-H bond length of the TFA donor has been chosen as the reaction coordinate. All the other geometrical parameters were optimized for each fixed value of the reaction coordinate. The profiles in DCM solvent were obtained from single point calculations on each point of the gas phase profiles with the PCM continuum model of the solvent. The energy of the dihydrogen bonded adduct is taken as the zero of energy.



Figure 9>

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The gas phase energy barrier to form the ion pair at the hydride site and at the metal site are respectively 10.9 kcal mol^{-1} and 16.0 kcal mol^{-1} . In DCM solvent the corresponding values are 4.6 kcal mol⁻¹ and 11.3 kcal mol⁻¹. The kinetic effects of the solvent is to lower the energy barriers in comparison to the gas phase. The ion pairs resulting from the protonation at hydride and metal site are found respectively at 10.2 kcal mol⁻¹ and 15.8 kcal mol⁻¹ above the hydrogen bonded complexes in the gas phase, and at 2.2 kcal mol⁻¹ and 10.8 kcal mol⁻¹ in DCM. We can see that there is also a thermodynamic effect of the solvent, stabilizing the charged species to a greater extent than the initial neutral hydrogen bonded ones. Both solvent effects (kinetic and thermodynamic) are more pronounced for the protonation at the hydride site, in good agreement with the experimental evidence that proton transfer takes place at hydride site to form the ion pair with the nonclassical [Cp*Fe(dppe)(η^2-H_2)]⁺ cation prior to the dihydrogendihydride isomerization.

We have also computed the energy profile of the proton transfer with a second TFA molecule assisting protonation by homoconjugated pairing. This situation can be related to experiments with an excess of TFA. The resulting $[Fe(H_2)]^+ \cdots [OC(CF_3)O \cdots HOOCCF_3]^-$ ion pairs are also found as minima. The ion pairs together with the initial dihydrogen



Figure 10.



Figure 10>

In the initial hydrogen bonded adducts with two TFA molecules, the latter are joined by a $0 \cdots H$ hydrogen bond, with O-H distances of 1.735 Å and 1.773 Å for the hydride and Fe bonded adducts, respectively. In the two corresponding ion pairs there is a strong hydrogen bond in the homoconjugated pair, with similar intermolecular hydrogen bond length (1.351

Å and 1.347 Å, respectively). However, the O···H distance between the homoconjugated anion and the cationic $[CpFe(dhpe)H_2]^+$ is shorter at the hydride site (2.215 Å) than at the metal site (2.439 Å) reflecting the greater basicity of the former. Moreover, these values are longer than those computed with a single molecule of CF₃COOH, showing that the second molecule enhances the acidity and favors the proton release.



Figure 11 reports the energetic profile of the proton transfer process to CpFe(dhpe)H involving two hydrogen bonded TFA molecules. The role of the second TFA molecule is to provide extra stabilization, through a strong hydrogen bond, to the $[CF_3COO]^-$ base left over after the proton release, thereby reducing its basicity. The same effect was reported for the protonation of CpRuH(CO)(PCy₃) ^[20] and Cp*Fe(dppe)H^[25] by weaker proton donors at the experimental and computational levels. There is a low energy barrier of about 3.0 kcal mol⁻¹ to overcome to form the protonated dihydrogen complex in the gas phase, and this completely vanishes in CH₂Cl₂, meaning that this is a quite easy process. This result is in good agreement with the experiment, because the proton transfer is very fast for TFA and the associated rate constant could not Moreover, as the solvent stabilizes the be estimated.^[25] charge-separated proton transfer product to a greater extent than the neutral hydrogen bonded complex, the process is more exothermic in dichloromethane than in gas phase.



Figure 11>

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situation is quite different when considering The the protonation at the Fe site. The potential energy curve for protonation exhibits a barrier of ca. 4.2 kcal mol^{-1} in the gas phase, slightly higher than that for the protonation at the hydride site. However, a barrier of $3.1 \text{ kcal mol}^{-1}$ still remains when adding the solvent effect. The kinetics of the transfer consequently enhanced proton are by the dichloromethane solvent only at the hydride site as we have found previously for the protonation with a single molecule of TFA. On the whole, our theoretical results confirms that the protonation is favored at the hydride site. In conclusion, shown that there is a correlation between they the thermodynamics of the hydrogen bond and the proton transfer kinetics for the two sites.

We have performed the (C) HFIP. same computations reported above for the protonation process with HFIP because both kinetic and thermodynamic data are experimentally available. Attempts to optimize ion pairs related to proton transfer (at hydride and at metal sites) always failed when considering only one molecule of HFIP, the system unfailingly going back to the initial hydrogen bonded complexes. As HFIP is a weaker acid than TFA, its conjugated base is consequently stronger than [CF₃COO]⁻. The [(CF₃)₂CHO]⁻ anion is a stronger base than the CpFe(dhpe)H complex preventing the loss of one proton. For HFIP, a second molecule is needed to localize the ion pair as a minimum, by homoconjugate pairing effect. One

may remark that, for HFIP, the same oxygen atom of one conjugate base is involved in two hydrogen bonds (



(d) Figure 12) in comparison to TFA where one oxygen atom is involved in the hydrogen bonding with the homoconjugate acid and the other one with the protonated complex



(e) Figure 10).. The $0 \cdots H$ bond lengths between the homoconjugated anion and the protonated complex are respectively 1.807 Å and 2.031 Å at the hydride and the metal sites reflecting a stronger interaction in the former case. Conversely, hydrogen bonding with the proton of the second alcohol molecule is stronger in the latter case (1.423 Å and 1.381 Å, respectively).



Figure 12>

The potential energy curves of the proton transfer with the two HFIP molecules



Figure 13) point out the same general solvent effects as found with TFA. At the hydride site, the energy barrier to

form the ion pair in dichloromethane (5.6 kcal mol⁻¹) is lower than in the gas phase (8.7 kcal mol⁻¹), illustrating the kinetic effect of the dichloromethane solvent. The proton transfer step with two HFIP molecules appears as an endothermic process by 4.2 kcal mol⁻¹. Concerning the proton transfer at the metal site, the solvent lowers the energy barrier found in gas phase. However, a barrier of 12.9 kcal mol⁻¹, more than twice higher than that for the protonation at the hydride site (5.7 kcal mol⁻¹) still remains.



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We have also considered that, after the formation of the dihydrogen bonded complex at the hydride site, the second HFIP molecule may form a hydrogen bond at the metal site. In other words, the metal and the oxygen atom of the first HFIP molecule may compete as basic centers for the incoming HFIP proton. The optimized geometry of this adduct is represented in



Figure 14. This complex lies 5 kcal mol⁻¹ above the adduct of the HFIP dimer hydrogen bonded at the hydride site


Figure 12, top left). Consequently, it can be stated that in the presence of an excess of HFIP the initial complex involves the participation of an HFIP dimer forming a dihydrogen bond at the hydride site. As for the TFA case, we find a correlation between the thermodynamics of hydrogen bonding and the kinetics of proton transfer, in favor of the hydride site.



Figure 14>

Although the calculations reproduce the main features of the proton transfer process, there is a discrepancy between the calculated and experimental thermodynamics, The experiment shows that the proton transfer from a single TFA molecule is possible, whereas the calculation shows this as endothermic. The experiment also shows exothermicity for proton transfer with two HFIP molecules, whereas the calculation also shows this as slightly endothermic. In both cases, the error has the same origin: the relative stability of the protonation product (the dihydrogen complex) is underestimated with respect to that of the reactant (the monohydride + the proton donor). The cause of this discrepancy is evident from the calibration of the proton affinity discussed above (Equation 4 and Table 5). The model system adopted, with Cp and dhpe instead of the actual Cp* and dppe ligands, presents a lower proton affinity than the real system. Thus the protonation process in the model system will be less exothermic than in the real one.

Discussion

a. Hydrogen bonding site

The calculations agree with the experimental study in terms of the identification of the hydride ligand as the preferred hydrogen bonding site. According to the calculations, the interaction with the metal is much less exothermic than the interaction with the hydride. Steric effects probably play the most important role favoring $M-H\cdots H$ over $M\cdots H$ hydrogen bonds for the Cp*FeH(dppe) system. For this system, hydrogen bonding at the hydride site involves only the hydride ligand, and hydrogen bonding at the metal site only the metal centre, since the two sites are mutually trans. The proton donor substituents get much closer to the metal coordination sphere the metal when binding occurs at site. Indeed, our computational work shows that the interaction energy at the metal site decreases considerably when moving from dhpe to dppe, whereas a much smaller effect is observed for the interaction energy at the hydride site. For all other previously investigated hydride complexes, hydrogen bonding to the metal site occurs in a *cis* position relative to the hydride site. The cutoff between metal site and hydride site is not so clear in those cases because even for $M \cdots H$ distances of about 2.6 Å (the usual distance for this kind of interactions) there will also be relatively short $H \cdots H$ distances. These hydrogen bonds could be more appropriately described as bifurcated,^[59, 63] and the steric effects may play more similar roles for the two types of interactions.

There does not appear to be a clear correlation between the hydrogen bonding site preference and the basicity factor, which reflects the electronic effects. The basicity factors of hydride ligands reported up to date vary from 0.7 to $1.6.^{[13]}$ Relative to complex Cp*FeH(dppe) ($E_j = 1.36\pm0.02$),^[25] which prefers to use the hydride site, hydrogen bonding has been shown to occur at the metal site for compounds with both higher basicity factors, e.g. (NP₃)ReH₃ ($E_j = 1.46\pm0.01$) [NP₃ = N(CH₂CH₂PPh₂)₃]^[10] and (PP₃)RhH ($E_j = 1.40$),^[64] and lower ones, e.g. WH₄(dppe)₂ ($E_j = 1.20$).^[65] Thus, the reasons for the preference of the metal or hydride site for hydrogen bonding are not clear at the moment. This topic definitely needs further work, both at the experimental and at the theoretical levels, before attaining greater understanding and predictive power.

The experimentally determined hydrogen bonding energies (-6.5 \pm 0.4 kcal mol⁻¹ for HFIP, *cf.* -5.9 \pm 0.4 kcal mol⁻¹ for TFE and -4.6 \pm 0.4 kcal mol⁻¹ for MFE)^[25] follow the acidity strength and are of the same order than those reported for other M-H··H interactions.^[56-58, 66, 67] The computed hydrogen bond formation

75

enthalpy values are slightly more negative than those obtained experimentally, possibly because the frequency calculations were carried out in the gas phase, where the interaction between the proton donor and the Cp*Fe(dhpe)H complex is stronger than in the dichloromethane solvent. The basicity of complex Cp*FeH(dppe) is sufficiently high to allow proton transfer even in the presence of a proton donor as weak as TFE. The dihydrogen bond formed as the first stage of the protonation reaction determines the direction of the proton transfer process yielding the non-classical dihydrogen complex.

b. Proton transfer activation barrier and equilibrium

The dihydrogen complex formation is reversible below 260 K, the equilibrium shifts to the right with a temperature decrease or with an acid strength increase. For the Cp*FeH(dppe)/HFIP system the negative enthalpy ($\Delta H_1 = -3.2 \div -$ 5.5 kcal mol⁻¹) and entropy ($\Delta S_1 = -4.8 \div -13.0$ cal mol⁻¹ K⁻¹) values for proton transfer step were estimated from the low temperature NMR and UV-visible studies. The enthalpic profile

76









Figure 15>

Our previous kinetic investigation of the Cp*FeH(dppe)/HFIP, though limited to a single temperature (298

K), had allowed us to establish that the proton transfer process is equilibrated and that the forward rate law has a first order dependence on the alcohol concentration. In combination with the kinetics study involving the proton donors TFE (weaker), PFTB and TFA (stronger), the study also revealed that the proton transfer rate constant increases with proton donor strength.^[25] The variable temperature the investigation reported here has provided the activation parameters for the proton transfer process, ΔH^{\ddagger}_1 = 2.6±0.3 kcal mol⁻¹ and ΔS^{\ddagger}_1 = -44.5±1.1 cal mol⁻¹ K⁻¹. Even though the activation enthalpy is relatively small, the entropic term is such that the activation free energy is in the 12-16 kcal mol⁻¹ range between 200 and 300 K, in agreement with the observation that the NMR resonances of the two equilibrating species are observed separately (slow exchange limit). The proton transfer equilibrium, however, is characterized by a smaller negative entropy (-4 to -7 e.u. from the NMR study; -13 e.u. from the UV-visible study), indicating that the transition state is much more ordered than both reagents and products. This is so because the two entities that constitute the starting and final systems are rather loose, the reagents having a hydrogen bond of moderate strength (H···H distance of 1.394 Å for the Cp*(dppe)FeH·2HFIP adduct) and the product featuring a hydrogen bonded ion pair with a relatively long H_2 anion distance (H···O distance of 1.807 Å). On the contrary, in the TS there is a strong interaction between the two tightly bonded units. In fact, the transition state can be regarded as an H₂ entity (H-H distance of 0.915 Å) simultaneously bonded to both the $[Cp*Fe(dppe)]^+$ and the $[AHA]^$ units (distances Fe-H and H-O of 1.568 and 1.40 Å, respectively). Therefore, the two units have lost degrees of freedom in the TS leading to a negative activation entropy.

The barrier is particularly sensitive to the strength of the proton donor as shown by the computational study. The hydrogen bonds have been described as incipient protontransfer reactions.^[68] Thus it can be expected that a correlation exists between the strength of the М-Н • • Н interaction and the proton transfer activation barrier. Indeed we have found such correlation in the two proton transfer reactions studied, with TFA and HFIP as a proton donors. Τn the limit, increasing the dihydrogen bond strength should lead to a complete transfer of the proton on the hydride and result in the formation of a η^2 -dihydrogen ligand.

In this respect, it is interesting to analyze the role played by the intervention of a second proton donor molecule, leading to the dihydrogen complex $[Fe(H_2)]^+\cdots[AHA]^-$. This intervention is crucial as shown both experimentally and theoretically. During the proton transfer from H-A···HA to the hydride site, several processes are simultaneously taking place: (A) weakening of the H-A bond; (B) weakening of the Fe-H bond; (C) establishment of the Fe-(H₂) three-center-twoelectron bond; (D) strengthening of the hydrogen bond between

79

A- (originating from the proton donor) and the second HA molecule, leading to the homoconjugated anion AHA-. Only contributions A and D depend on the nature of the proton donor. The calculated parameters (see Results section) clearly show that the second HA molecule increases the strength of the primary dihydrogen bonding interaction (socalled cooperative effect in hydrogen bonding)^[69] and therefore facilitates proton transfer. It is also possible to imagine that the slightly acidic protons of the dichloromethane solvent molecules interact in the same manner with the proton donor molecule, thereby assisting the proton transfer process.^[70] On the other side of the equilibrium process, the anion stabilization provided by the second proton donor molecule also contributes to lower the activation barrier and

81









c. Nature of the conjugate base

Our previous spectroscopic study (see Introduction) could not provide detailed information on the nature of the proton donor conjugate base,^[25] i.e. whether it is free or hydrogen bonded (as in II or ∇), by itself or associated with the proton donor in a homoconjugated anion. The reason for this lack of information is the fact that the conjugate bases of the fluorinated alcohols used in that study are colorless species and do not exhibit any typical IR absorptions that could signal their hydrogen bonding with the product dihydrogen complex. The p-nitrophenol used in the current study show a diagnostic change of the absorption maximum as a function of their protonation status, whereas the anion of TFA exhibits IR absorptions that are also quite sensitive to hydrogen bonding.

The experimental evidence, backed up by the computational work, show that the stronger proton donor TFA is capable to transfer the proton without assistance by a second TFA molecule (equation (a) in Scheme 4). The ion pair $[Cp^*(dppe)Fe(H_2)]^+ \cdots [OCOCF_3]$, in equilibrium with the free ions, forms under conditions of a deficit of TFA. Excess acid then quantitatively produces the free homoconjugated anion, $[CF_3COO(HOOCCF_3)_2]^-$. On the other hand, the phenol, like the fluorinated alcohols, is a much weaker proton donor than TFA (in DMSO, $pK_a = 3.45$ for TFA, 10.7 PFTB, 17.9 HFIP and 23.5 TFE)^[51] and therefore behaves in the same fashion as the fluorinated alcohols, for which the intervention of a second proton donor molecule was proven by the kinetic investigation (equation (b) in Scheme 4).

< Scheme 4 >

d. Solvent effects

The protonation of a neutral hydride is a reaction involving the generation of charged species from neutral reactants. In this type of process solvent effect are very important. Polar solvents will assist the charge generation process. This behavior can be observed comparing the computed energy profiles for the proton transfer in gas phase and

for

instance



 f_{i} f_{i

Figure 13). The stabilization caused by the solvent is greater on the products side (ion pair) than at any other

point along the pathway. As a result, the monotonically growing energy profile obtained in gas phase is turned in a double well in DCM. The polarity of the solvent will also play a major role in the charge separation step which leads from the hydrogen bonded ion pair to the free ions.

Conclusion

The present study has allowed us to zoom in closer on the energy profile and mechanism of the proton transfer process to compound Cp*Fe(dppe)H. The general features of this process are identical to those established for the protonation of many other hydride complexes: faster proton transfer to the hydride site, followed by isomerization. The combined experimental and theoretical work has addressed the strength of the hydrogen bonding interaction, the energetic barrier to the proton transfer step, and the thermodynamics of the proton transfer equilibrium. The preferred site of hydrogen bonding and the strength of the resulting interaction are shown to correlate with the site of the kinetic protonation and with proton transfer barrier. Finally, important the new information on the proton transfer assistance by a second proton donor molecule was revealed by the computational study. The second molecule enhances the ability of the proton donor and stabilizes the transition state and the final product by homoconjugated anion formation. However, sufficiently strong proton donors (in our case, TFA) are capable to transfer the proton without the need of such assistance.

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Figure 1. UV-visible spectra at 200 K in CH₂Cl₂ of (a) PNP (0.001 M); (b) potassium *p*-nitrophenolate (0.001 M in the presence of 18-crown-6); (c) homoconjugated PNP anion (band derived from the spectrum of a 1:1 mixture of (a) and (b)); (d) PNP (0.001 M) in the presence of Cp*FeH(dppe) (0.0005 M).



Figure 2. UV-visible spectra for a CH₂Cl₂ solution containing PNP (0.001 M) and Cp*FeH(dppe) at 200 K. The Fe/PNP molar ratios are, respectively: 0.1, 0.2, 0.3, 0.5, 1 and 2.



Figure 3. Intensity changes at 380 nm vs the Cp*(dppe)FeH mole fraction. The data are from the UV-visible spectra

shown in



Figure **2**.



Figure 4 UV-visible spectra taken at each 10 K interval between 200 and 260 K for a CH₂Cl₂ solution containing PNP (0.001 M) and Cp*FeH(dppe) (0.00033M).



Figure 5. IR spectra at 200 K of CH₂Cl₂ solutions containing Cp*FeH(dppe) and TFA. (a) [FeH] = 0.06M, [TFA] = 0.03M. (b) [FeH] = 0.006 M, [TFA] = 0.03 M. The spectra of the solution containing only the free acid is also shown for comparison in both (a) and (b).



Figure 6. Equilibrium constant at different temperatures for the hydrogen bond formation between Cp*Fe(dppe)H and HFIP.



Figure 7. Optimized geometries of the hydrogen bonded adducts at hydride and metal sites with the set of proton donors. Hydrogen atoms (except for those involved in the hydrogen bond) are omitted for clarity.





Figure 9. Potential energy curves for the proton transfer from one TFA molecule to complex CpFe(dhpe)H at the hydride (top) and metal (bottom) sites. The O-H distance of the transferring proton is the reaction coordinate. Energies are in kcal mol⁻¹.



Figure 10. Top view of the optimized geometries of hydrogen bonded complexes (top) and of the ion pair (bottom) with two TFA molecules. Left: hydride site protonation; right: metal site protonation. Hydrogen atoms of Cp and PH₂CH₂CH₂PH₂ are not displayed.



Figure 11. Potential energy curves for the proton transfer from two TFA molecules to the Fe complex at the hydride (top) and metal (bottom) sites. The O-H distance of the transferring proton has been taken as the reaction coordinate. Energies are in kcal mol⁻¹.



Figure 12. Optimized geometries of hydrogen bonded complexes (top) and of the ion pair (bottom) with two HFIP molecules. Left: hydride site protonation; right: metal site protonation. Hydrogen atoms of Cp and PH₂CH₂CH₂PH₂ are not displayed.



Figure 13. Potential energy curves for the proton transfer from two HFIP molecules to the Fe complex at the hydride (top) and metal (bottom) sites. The O-H distance of the transferring proton has been taken as the reaction coordinate. Energies are in kcal mol⁻¹.



Figure 14 Optimized geometry of a CpFe(dhpe)H·2(HFIP) isomer with one HFIP molecule hydrogen bonded at the hydride site and the second one at the metal site. Hydrogen atoms of Cp and PH₂CH₂CH₂PH₂ are not displayed.



Figure 15 Enthalpic profile for the reaction between Cp*Fe(dppe)H and HFIP.



Scheme 1



Scheme 2

 $Cp*(dppe)Fe(H_2)^+\cdots[ROHOR]^- \rightleftharpoons Cp*(dppe)Fe(H_2)^+ + [ROHOR]^-$ Scheme 3

 $Cp^*(dppe)Fe-H\cdots HA \longrightarrow Cp^*(dppe)Fe(H_2)^+\cdots A^-$ (a)

 $Cp*(dppe)Fe-H\cdots HA + HA \implies Cp*(dppe)Fe(H_2)^+\cdots AHA^-$ (b) Scheme 4

