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Hydrogen-Bond Acceptance of Bifunctional Ligands in an Alkyne–Metal π Complex

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Typically, organometallic catalyst selectivity and reactivity have been controlled by variations in the metal and ancilliary ligands used, focusing on steric and electronic properties of the latter.^{1,2} In contrast, nature's metalloenzymes use secondary interactions such as hydrogen bonding or proton transfer in the active site.³ An increasing number of synthetic catalysts and related systems show the benefits of secondary interactions,⁴ which are generally difficult to control. Using hydrogen bonding or proton transfer from or to imidazolyl5 and pyridylphosphines,4d,6,7 our group has reported rate enhancements in anti-Markovnikov alkyne hydration and alkene isomerization by factors of up to 10000. In studying the origin of remarkable rate enhancements in alkyne hydration, we have discovered hydrogen bonding between the terminal alkyne proton and a heterocyclic nitrogen. To probe this unusual structural feature, we report the first use of scalar coupling across a hydrocarbon C-H bond $({}^{2h}J_{CN})$, a technique used to characterize hydrogen bonds in biological samples.8

Initial DFT calculations on the alkyne hydration pathway first suggested the importance of structures like **5** (Scheme 1).⁹ However, in **5** the orientation of the alkyne alone does not allow us to conclude that there is C–H–N hydrogen bonding: in **5c**, a complex lacking the nitrogens, the same alkyne orientation is seen in both an X-ray crystal structure (with BF_4^- counterion)¹⁰ and our calculations. It appeared from the literature¹⁰ that to make an alkyne complex like **5** and study its structure before it isomerized to a vinylidene complex, we would need the smallest alkyne, phosphine, and ancilliary Cp ligand possible, making **5a,b** with ligands **1a** and **1b** our synthetic targets, with **1c** as a control.

Thus, ligands 1a,¹¹ 1b,⁹ and 1c were each converted to 2 using CpRu(COD)Cl. In the case of 2c, ionization of the chloride (Scheme 1) was done in the presence of HCCH to give 5c, which could be isolated and stored cold, but it converted smoothly to 6c after 3 h at 50 °C (Scheme 2).

In contrast, early experiments made it very clear that extraordinary measures would be needed to isolate or even observe **5a** or **5b**. First (Scheme 1), the chlorides in **2a** and **2b** were ionized in the presence of water¹² to give aquo complexes **3a** or **3b**. Storage of **3a** or **3b** under vacuum led to loss of water and mixtures of **3** and **4**, with a greater amount of **4** in the case of **4b**. When cooled below -40 °C in CD₂Cl₂, chelate complex **4a** did not react at an appreciable rate with acetylene (Scheme 2), but near -40 °C was converted to symmetrical π complex **5a**, which isomerized smoothly to vinylidene **6a** at higher temperatures (0 °C, 3 h). In the imidazole case, **4b** was inert to acetylene until ca. 10 °C, where unsymmetrical addition product **7b** was the only product detected. Complex **7b** reasonably arises from unseen vinylidene **6b**. Interestingly, the NMR spectrum of **7b** (e.g., two sets of sharp resonances for the imidazoles at -40 °C) shows coalescence behavior at higher Scheme 1. Precursor Synthesis



temperatures consistent with an equilibrium between **7b** and **6b** with $E_a = 15.9$ kcal mol⁻¹.

In summary, when imidazole derivative **4b** reacts with acetylene, alkyne complex **5b** is not seen, because of rapid conversion to **7b**, whereas, when starting with pyridyl analogue **4a**, alkyne complex **5a** can be observed at -40 °C, but at temperatures closer to 0 °C it evolves to **6a**. In contrast, control alkyne complex **5c** requires temperatures 50–90 °C higher for conversion to vinylidene **6c**, all showing the profound effects and assistance of the pendant bases on these transformations.

Bonding in vinylidenes **6a** and **6c** was compared using isotopomers derived from H¹³C¹³CH. Significantly, nearly identical values for ${}^{1}J_{CC}$ (Scheme 2) are consistent with the same C–C bond length and degree of backbonding in each case.¹³

To clarify the structure of 5a, various unsuccessful attempts were made to grow crystals at -40 °C. Instead, spectroscopic characterization¹⁴ of alkyne bonding in **5a** and **5c** was made using H¹³C¹³-CH, resulting in 5a- and 5c-(13C)2. At -40 °C the ten-line AA'XX' pattern could be analyzed¹³ to show that there were significant differences in couplings involving the hydrogens (Scheme 2). Data above for ${}^{1}J_{CC}$ in **6a** and **6c** show that the electronic effects of the two ligands are identical, consistent with almost identical ${}^{1}J_{CC}$ values in the π complexes. We could find no experimental data on effects of hydrogen bonding on alkyne NMR coupling constants and only a single theoretical paper (on HCCH-OH₂).¹⁵ Comparing alkyne and hydrogen-bonded alkyne, the changes in ${}^{1}J_{CH}$ and ${}^{2}J_{CH}$ seen by us experimentally (+1.5 and +1.9 Hz) resemble those predicted¹⁵ for HCCH-OH₂ (+2.55 and +1.65 Hz).¹⁶ In short, the differences in NMR couplings are for those couplings to hydrogen, and we attribute this to effects of hydrogen bonding in 5a. Attempts were made to engage the CH bonds of π complex 5c in hydrogen bonding. Instead, addition of imidazole 8 led to addition to the alkyne (9).

Scheme 2. Alkyne Hydrogen Bonding and Enhanced Reactivity of **Bifunctional Complexes**



Because NMR coupling constants are a novel tool for studying alkyne hydrogen bonding, we wanted to provide additional evidence for structure 5a. The remarkable discovery of scalar couplings across hydrogen bonds has stimulated a great deal of experimental work on proteins, DNA, and supramolecular interactions⁸ and theoretical investigations of simple systems, but there appears to have been no use in coordination or organometallic chemistry. Ligand 1a- ^{15}N was made 9 and converted to $3a \cdot (^{15}N)_2$ and $4a \cdot (^{15}N)_2$. Addition of acetylene at low temperature led to 5a-(15N)2, whose ¹³C NMR spectrum acquired at -50 °C with decoupling of both ¹H and ³¹P allowed observation of a somewhat broadened¹⁷ doublet ($^{2h}J_{CN} =$ 3 ± 0.5 Hz) as expected for coupling of one natural abundance acetylene 13C to the nearest 15N nucleus, whereas a similar spectrum of 5a showed a singlet.

Previous theoretical work by Del Bene et al. on a simple system¹⁸ showed that when the C-H-N angle is near 135°, ${}^{2h}J_{CN}$ would be approximately one-third the magnitude when the hydrogen bond is linear. Indeed, DFT calculations on 5a itself show a slightly unsymmetrical structure with two C-H-N angles (124.5 and 135.7°).¹⁹ Determining the strength of the hydrogen bonds in 5a remains a subject for future study, but as an estimate, we note that calculations on conversion of **5c** to **6c** indicate that $\Delta G = -17.9$ kcal mol⁻¹ whereas for similar conversion of **5a** to **6a**, $\Delta G = -12.7$ kcal mol⁻¹. The difference may be attributed to thermodynamic stabilization of 5a by two hydrogen bonds. Despite this effect,

experiments show a pronounced kinetic acceleration by the heterocycles on isomerization of alkyne to vinylidene (5 to 6), and importantly, **3a** is a very competent catalyst (>99% yield of hexanal from 1-hexyne after 5 h at 70 °C using 2 mol % 3a).

In summary, the presence of C-H-N hydrogen bonding in an alkyne π complex was revealed using NMR coupling information, from both data within the alkyne ligand as well as between alkyne and pyridine $({}^{2h}J_{CN})$. Ongoing experimental and theoretical studies are designed to elucidate further the favorable effects of bifunctional ligands in alkyne hydration and related reactions, and these will be reported in due course.

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Supporting Information Available: Details of compound preparation and characterization and calculations. This material is available free of charge via the Internet at http://pubs.acs.org.

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