Cooperative C–H Bond Activation by a Low-Spin d⁶ Iron–Aluminium Complex

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Abstract: The reactions of transition metal complexes underpin numerous synthetic processes and catalytic transformations. Typically, this reactivity involves the participation of empty and filled molecular orbitals centred on the transition metal. Kinetically stabilised species, such as octahedral low-spin d⁶ transition metal complexes, are not expected to participate directly in these reactions. However, novel approaches that exploit metal ligand-cooperativity offer an opportunity to challenge these preconceptions. Here we show that inclusion of an aluminium-based ligand into the coordination sphere of neutral low-spin d⁶ iron complex leads to unexpected reactivity. Complexes featuring an unsupported Fe–Al bond are capable of the intermolecular C–H bond activation of pyridines. Mechanistic analysis suggests that C–H activation proceeds through a reductive deprotonation in which the two metal centres (Fe and Al) act like a frustrated Lewis-pair. Key to this behaviour is a ground state destabilisation of the d⁶ iron complex, brought about by the inclusion of the electropositive aluminium-based ligand. These findings have immediate implications for the design of reagents and catalysts based on 1st row transition metals.

Introduction.

The concept of metal-ligand cooperativity has greatly enriched the chemistry and catalytic applications of transition metal complexes.^{1,2} Cooperative strategies that take advantage of the Lewis-acidic nature of a transition metal centre in combination with a Lewis basic ligand have become established features, frequently employed in the design of novel catalytic systems. Equally attractive but less common are transition metal complexes that bear a Lewis acidic functionality in the ligand.^{3–5} These new design principles offer an opportunity to overturn existing paradigms in transition metal chemistry. This is particularly important for applications of inexpensive and sustainable 1st row transition metals (e.g. Fe).^{6–9} Despite the exciting opportunities in this field, many of the emerging reactions that involve metal-ligand cooperativity still proceed through established mechanisms, such as oxidative addition. For example, transition metal complexes bearing Lewis-acidic ligands (based on B or Al) can activate the *ortho* C–H bond of pyridine substrates (Figure 1).^{10–13} C– H activation is believed to take place by an oxidative addition mechanism at the transition metal centre, leading to products in which the pyridyl group is directly bonded to this metal.^{13–15} The main group ligand plays

a role in substrate coordination and determining the *ortho* selectivity, but itself does not lead to new types of reactivity.

In this paper, we report the synthesis and characterisation of bimetallic complexes in which an unsupported¹⁶ aluminium-based ligand is bound to an iron(II) dihydride fragment. These complexes are neutral low spin d⁶ species based on an octahedral parent geometry and as such are expected to be chemically inert.¹⁷ Textbook examples of such compounds with more classical ligand systems possess a stable 18-electron configuration and a large HOMO (t_{2g}) – LUMO (e_g) gap, limiting direct reactivity at the metal site.¹⁸ Inclusion of the aluminium-based ligand results in a significant distortion of geometry away from octahedral due to favourable, but weak, interactions in the secondary coordination sphere. This distortion leads to an unusual ground state destabilisation and raises the energy of the HOMO of the iron centre. This effect exposes an entirely new type of reactivity of the neutral low spin d⁶ centre. These complexes were found to selectively break the *ortho* C–H bond in pyridine. Mechanistic studies reveal that the two metal centres act as a frustrated Fe–Al Lewis pair (FLP)^{19–23} leading to a deprotonation rather than oxidative addition pathway.²⁴

The findings both compliment and expand upon the known nucleophilicity of anionic analogues such as $[Fe(\eta^5-C_5H_5)(CO)_2]^-$. More broadly, these results suggest that the shape and electronic structure of transition metal centres can be modulated through incorporation of Lewis-acidic ligands, leading to perturbation of the electronic structure, and exposing new types of reactivity.



Figure 1. Well-defined bimetallic systems for the selective *ortho* C–H activation of pyridines. Current systems that require ligand dissociation prior to bond activation versus a novel pathway occurring at a coordinatively saturate iron aluminylene system.

Synthesis and characterisation.

Complexes **2a** and **2b** can be readily prepared by reacting FeBr₂, PMe₃ and the respective β -diketiminate aluminium hydrides **1a** (R = Mesityl or Mes) or **1b** (R = 2,6-Diisopropylphenyl or Dipp) in toluene or benzene and were isolated in 80-90% yield (Figure 2). Both complexes exhibit a mutually coupled spin system comprised of one triplet and one doublet resonance in the ³¹P{¹H} NMR spectra with an integration ratio of 1:2. These resonances are consistent with the magnetic non-equivalence of the axial and equatorial phosphine ligands. The most characteristic features in the ¹H NMR spectra are broad signals at $\delta_{\rm H}$ = -13.08 (**2a**) or -13.20 (**2b**) ppm assigned to the bridging hydrides which are coupled to the quadrupolar I = 5/2 ²⁷Al nucleus.

Reduction of these precursors using magnesium turnings in THF affords complexes **3a** and **3b** in almost quantitative NMR yield. Both **3a** and **3b** can be isolated as dark red/orange crystalline solids. In contrast to complexes **2a-b**, **3a-b** both show only one singlet resonance in the ³¹P{¹H} and a well resolved quartet hydride resonance in the ¹H NMR spectra. The data are consistent with a highly symmetric structure in solution – an observation that could suggest fast ligand exchange on the NMR timescale.

a) preparation of the low-valent complexes 3a-b





Figure 2. a) Preparation of complexes 2a-b and 3a-b. b) X-ray structure of 2b. c) X-ray and calculated structure of 3b.

Structure and bonding.

Single crystals suitable for X-ray diffraction could be obtained for all four compounds. Data for **2b** and **3b** are depicted in Figure 2. The position of the hydride ligands in **3b** could not be refined but their presence is evident from the corresponding ¹H NMR spectrum. When comparing these structures, it becomes apparent that drastic changes in the Fe---Al intermetallic distances occur upon reduction. For example, a decrease of the Fe---Al distance from 2.459(1) Å in **2b** to 2.194(1) Å in **3b** becomes visible in the solid-state structures. While the Fe---Al separation of **2b** almost matches the sum of their covalent radii²⁵ (formal shortness ratio²⁶ (FSR) = 1.02) it is clearly below this value in **3b** (FSR = 0.91). Such short Fe---Al bond lengths appear to be diagnostic for an aluminylene²⁷ metalloligand covalently bound to iron.²⁸

Based on the experimental data it is evident that **2a-b** and **3a-b** are low-spin and diamagnetic 18 valence electron complexes. While **2a-b** can be confidently described as σ -alane²⁹ complexes, three extreme bonding scenarios¹⁶ may be considered for the bimetallic species **3a-b** (Scheme 3a): (**A**) a four-electron, $\eta^2:\eta^2$ - coordination of H–Al–H to the 14 electron fragment [Fe(PMe₃)₃], (**B**) two electron coordination of an aluminylene metalloligand to a 16 electron Fe(II) dihydride fragment, (**C**) strong polarisation of the TM–M bond resulting in a cationic AI metalloligand and an anionic iron dihydride fragment due to the electronegativity difference between the two metals ($\Delta \chi_P = 0.22$).^{30,31}



Figure 3. Analysis of the bonding in **3a-b**. a) extreme bonding descriptions. b) calculated Fe---Al distances and NPA charges of **2a** and **3a** (ω B97X // 6-31g** (H,C,N,P) / SDDAll (Fe,Al)). c) QTAIM analysis and WBIs of **2a** and **3a**.

More insight into the structure and bonding in these complexes was gained by DFT calculations and the optimised structure of **3b** is depicted in Figure 2c. In complexes **3a-b**, the iron fragment adopts a pseudo-octahedral coordination geometry in an all-*cis* configuration of the hydride and phosphine ligands, respectively (**3b**: L-Fe-L = 83.1-101.6°; L = H, PMe₃). Only the Al metalloligand deviates from this geometry being bent away from its axial position by about 45°. The calculated structures also reflect the short Fe---Al distances observed in the solid-state structure of **3b** (Figure 3b). The Al–H bonds in these species appear to be significantly longer in comparison to the parent dibromide complexes (e.g., 1.838 vs. 2.045 Å). The Wiberg bond indices (WBI) for the Fe---Al bond drastically increase from 0.14 in **2b** to 0.55 in **3b** while the WBIs for the Al–H bonds are slightly lower in **3b** (0.23, 0.23) than in **2b** (0.27, 0.26). These data suggest the presence of a direct metal–metal bond due to a preceding double Al–H bond activation at the transition metal centre.^{30,32-35}



Figure 4. a) Energy profile for the relaxed scan of the AI-Fe-P_{ax} angle in the simplified model complex **3c** $(\omega B97X // 6-31g^{**} (H,C,N,P) / SDDAII (Fe,AI))$. b) energies of the occupied frontier orbitals in **3c** as function of the AI-Fe-P_{ax} bending angle. c) Orbital isosurfaces (isovalue = 0.05) at 0° (equilibrium structure) and 40° (metalloligand in the axial position) bending angles.

These findings are further underpinned by quantum theory atoms-in-molecules (QTAIM) calculations (Figure 3c).³⁶ In **2b**, bond critical points (BCPs) are found between Fe and H, Al and H but not between Fe and Al. In **3b**, the QTAIM analysis reveals BCPs between Fe and Al as well as Fe and H whereas no BCPs are found between Al and H.

Analysis of the NPA charges in **3b** reveals significant polarisation of the Fe–Al bond showing values of -0.97 for Fe and 1.25 for Al while negative charges on the hydrides are low (-0.18/0.18). The large negative charge accumulation on Fe likely results from the strong electron donating nature of both the two hydrides and aluminylene ligand. This assumption is supported by ETS-NOCV calculations³⁷ on **3b** revealing that the donation from the hydrides and the aluminylene ligand to iron accounts for more than 64 % ($\Delta \rho_1 = -74.7$ kcal/mol) of the total orbital interaction energy ($\Delta E_{orb} = -117.1$ kcal/mol) whereas backdonation from the Fe–H bonds to the empty Al p-orbitals were identified as the second ($\Delta \rho_2 = -16.6$ kcal/mol, 14 %) and third ($\Delta \rho_3 = -14.2$ kcal/mol, 12 %) largest contributions to ΔE_{orb} (see table S5 in the SI). These secondary interactions are likely responsible for the bent position of the aluminylene metalloligand.

NBO calculations were used to gain further insight into the bonding in these complexes. For the sake of clarity, we considered a simplified model complex **3c** (P' = PH₃, R = Me) which, however, gives rise to highly similar frontier MOs as in **3a-b** (see table S6 in the SI). The NBO analysis identifies a σ -bond between Fe (4s, 40.8%, 3d 57.8%) and Al (3s 73.6%, 3p 26.4%) as main contributor to the HOMO, while the LUMO+2 comprises an empty p-orbital on aluminium and possesses antibonding character with respect to the Fe–Al σ -bond. The HOMO is significantly higher (about 1.0 eV) in energy than the lower MOs which are predominantly non-bonding and possess largely 3d-character (HOMO-1 to HOMO-3). This difference appears to be a consequence of the distorted coordination geometry and vanishes when the metalloligand is moved towards the axial position (Figure 4).

Intramolecular C–H activation.

The complex **3a** was found to undergo an intramolecular C–H activation affording the cyclometalated complex **4a** (Figure 5a). Heating a toluene- d_8 solution of **3a** to 80°C results in slow formation of a new species as revealed by a growing new singlet resonance at $\delta_P = 30.0$ ppm in the ³¹P NMR spectrum. In the ¹H NMR, **4a** gives rise to a new hydride signal (broadened quartet) that integrates to 3H. The reaction was monitored over time and complete consumption of the starting material observed after 24 h. The new species was formed in 71% NMR yield. Crystals suitable for X-ray diffraction could be grown by diffusion of tetramethylsilane into a saturated solution of *n*-pentane and the solid-state structure of **4a** is depicted in Figure 5b. Heating a solution of **3b** in toluene- d_8 to 80 °C only results in the slow decomposition of this complex and the formation untraceable species.



Figure 5. a) Intramolecular C-H activation of **3a**, b) X-ray structure of **4a**, c) calculated free energy profile for the conversion of **3a** to **4a** (B3PW91 / PCM (toluene) / D3 // 6-31g** (H,C,N,P) / SDDAll (Fe,Al)). Energies are given in kcal/mol. d) acceptor (LUMO+6) and donor (HOMO) orbitals in intermediate **1-B**.

More insight into the conversion of **3a** to **4a** was gained from a combination of kinetic experiments and DFT calculations.¹³ Experimentally, the reaction was found to be first order with respect to **3a**. An Eyring analysis over a temperature range of 60-100 °C gave the activation parameters $\Delta H^{\dagger} = 23.4 \pm 0.5$ kcal mol⁻¹ and $\Delta S^{\dagger} = 12.5 \pm 1.3$ cal mol⁻¹ K⁻¹ which correspond to an associated $\Delta G_{298K}^{\dagger} = 27.1 \pm 0.8$ kcal mol⁻¹. DFT calculations were benchmarked against the experimental data. Best results were obtained by using the B3PW91 functional and a

6-31g** (H,C,N,P) / SDDAII (Fe,AI) basis set. Solvent (PCM, Benzene) and dispersion corrections (D3) were directly included in the optimisation of the stationary points.

The calculated free energy profile for the intramolecular C–H activation is shown in Figure 5c. The reaction sequence is initiated by a slight rotation of the aluminylene ligand around the Fe---Al axes (**1-A** to **1-B**) followed by a deprotonation of the mesityl CH₃ group by the iron metal centre (**1-TS**_{BC}). Intermediate **1-C** features the Fe-(μ -H₃)-Al bridging motif and the deprotonated mesityl residue. Another slight conformational change initiates the Al–C bond formation (**1-TS**_{CD}) and leads to the cyclometalated product (**1-D**). Formation of the product is exergonic by -13.3 kcal mol⁻¹ with an overall barrier of 27.9 kcal mol⁻¹ being in good agreement to the experimental value (27.1 ± 0.8 kcal mol⁻¹).

Analysis of the frontier MOs reveals that the HOMO in the ground state of **3a** (Figure 5d) acts as electron donor for the reductive deprotonation of the CH_3 group in **1-TS**_{BC} while an empty p-orbital at Al (LUMO+6, Figure 5d) acts as electron acceptor for the deprotonated CH_2 group (**1-TS**_{CD}) to form the Al–C bond in the final product (**1-D**).

Intermolecular C–H Activation

Complexes **3a** and **3b** are also capable of promoting intermolecular C–H activation. For example, **3a** readily reacts with pyridine (1 equiv., C₆D₆, room temperature) resulting in the selective C–H activation in the 2-position of the heterocycle (> 95% NMR yield, Figure 6a). Like **4a**, **5a** exhibits a sharp singlet resonance in the ³¹P{¹H} NMR spectrum at δ_P = 29.6 ppm as well as a broadened quartet hydride signal at δ_H = -15.33 ppm in the ¹H NMR spectrum. The reaction of **2b** with pyridine required more forcing reaction conditions (40 °C, 10 equiv. of pyridine, 18 h) but led to the analogue product **5b** in about 80 % NMR yield. Crystals suitable for X-ray diffraction could be obtained for **5b** and the solid-state structure confirms the *ortho* C–H activation and C–Al bond formation in 2-position of the pyridine heterocycle (Figure 6b).

A small array of substrates was investigated. Employing pyridines with different substituents in the 4-postion (R = Me, Ph, NMe₂) under the same conditions did not show any change in the reactivity or selectivity of the C– H activation with **3a**, in all cases giving products that arise from substitution at the 2-position (Figure 6c). On the other hand, 2-phenylpyridine did not react with **3a**, presumably due to steric interference of the phenyl substituent with the ligand system on aluminium.

DFT-calculations on the *ortho* C–H activation of pyridine suggest a similar reaction pathway as for the intramolecular C–H activation of **3a** (Figure 6c). Again, a two-step mechanism has been identified that proceeds *via* the deprotonation of pyridine substrate. Coordination of the pyridine nitrogen to the aluminium centre controls the site selectivity of the reaction. NBO calculations reveal that the coordination of pyridine also facilitates bond breaking. In **2-A**, the Fe–Al bond becomes more polarised in comparison to **1-B** and the Wiberg bond index drops (see tables S7 and S8 in the SI). The following transition state **2-TS_{AB}** is significantly lower in energy than for the intramolecular deprotonation of the mesityl CH₃ group in **3a** (14.7 vs. 27.9

kcal/mol) being in line with a fast reaction at room temperature. Both, the intermediates after coordination (**2**-**A**, 0.0 kcal/mol) and deprotonation (**2**-**B**, 0.8 kcal/mol) of the pyridine appear to be almost thermoneutral in comparison to the starting materials. However, a final switch from N- to C-coordination of the deprotonated pyridine is facile (**2**-**TS**_{BC}, barrier of 5.5 kcal/mol) and highly exergonic due to formation of the Al–C bond in **2**-**C** (-20.7 kcal/mol).

A comparison of the reaction rates using an excess of pyridine or pyridine- d_5 in two independent reactions gave an unusually large kinetic isotope effect (KIE) of 14.0 ± 0.2 at 297 K. The experimentally observed $k_{\rm H}/k_{\rm D}$ value likely results from a large classical KIE including a significant contribution of quantum tunnelling.³⁸ The rate determining elementary step (**2-A** to **2-B**) is nearly thermoneutral with a centred transition state along the reaction coordinate and thus a maximum primary KIE (up to 7-8) may be expected.^{38,39} On the other hand, KIEs greater than 10 are generally considered to be caused by quantum tunneling and are diagnostic for proton transfer reactions.^{40–43} Both assumptions are in agreement with the calculated mechanism in Figure 6c and support a non-oxidative addition pathway.





Figure 6. a) Pyridine *ortho* C–H activation with **3a-b** (NMR Yields are given in parentheses), b) X-ray structure of **5b**, c) calculated free energy profile for the conversion of **3a** to **5a** (B3PW91 / PCM (benzene) / D3 // 6-31g** (H,C,N,P) / SDDAII (Fe,AI)). Energies are given in kcal/mol.

Conclusions.

In summary, we described two well-defined bimetallic Fe–Al complexes that possess a distorted octahedral geometry. This distortion leads to an unusual ground state destabilisation and raises the energy of the HOMO (the Fe–Al bond). Consequently, the HOMO becomes accessible for an encounter substrate and leads to an enhanced reactivity of these complexes including intramolecular sp³ C–H activation as well as in the selective *ortho* C–H activation of pyridine substrates. These reactions were found to follow a novel cooperative mechanism in which a reductive deprotonation of the C–H bond was identified as key elementary step. We could show that the Fe–Al bonds these complexes are highly polarised and react as a bimetallic frustrated Lewis pair in which the destabilised HOMO acts as Lewis donor orbital while empty p-orbitals on Al serve as Lewis acceptors. The reaction leads to the formation of a Fe–H and Al–C bonds. These results may lay the foundation for the rational design of future catalytic systems with prospects in the field of base-metal catalysis.

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