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Introduction

In the past decade there has been significant interest in transition metal (TM) free systems which activate H_2 .¹ Two main strategies have emerged to facilitate this reactivity: the use of lowvalent main group (MG) compounds,² and so-called 'frustrated Lewis pairs' (FLPs).³ In both cases, reactivity arises from simultaneously having access to a high-lying HOMO and low-lying LUMO (Fig. 1). Various low-valent MG compounds containing multiple E–E bonds (E = Al, Si, Ga, Ge, Sn),^{4,5} or single-site low-valent centres such as carbenes and heavier tetrylene analogues, have been shown to react with $\mathrm{H_2}$.⁶ The scope of Lewis bases (LBs) and, to a lesser extent, Lewis acids (LAs), which can be used in H_2 -activating FLPs has expanded to include a number of elements from across the periodic table. This is principally due to the readily tuneable steric and electronic profiles of the individual LA and LB sites.⁷⁻⁹ Many FLP systems display reversible H_2 cleavage, which has facilitated their rapid expansion into the field of catalytic hydrogenation.¹⁰ The same is not true for low-valent MG compounds; examples of reversible $H₂$ activation are very rare and limited to antiaromatic boracycles,¹¹ a phosphorusbased singlet biradicaloid,¹² and only one low-valent group 14 compound: a dinuclear $Sn(i)$ distannyne.¹³ The design of singlesite MG systems which are ergoneutral for H_2 activation requires fine-tuning of thermodynamic (e.g. weak E-H bond strengths promoting an accessible formal E^{n+2}/E^n couple) and kinetic factors, both of which are constrained to a mononuclear species, and is hence especially challenging. **EDGE ARTICLE**
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Base-induced reversible H_2 addition to a single Sn(II) centret

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A range of amines catalyse the oxidative addition (OA) of H_2 to $[(Me_3Si)_2CH]_2Sn$ (1), forming [(Me_{3Si)2}CH]₂SnH₂ (2). Experimental and computational studies point to 'frustrated Lewis pair' mechanisms in which 1 acts as a Lewis acid and involve unusual late transition states; this is supported by the observation of a kinetic isotope effect (KIE; $k'_{(H_2)}/k'_{(D_2)} = 1.51 \pm 0.04$) for Et₃N. When DBU is used the energetics of H_2 activation are altered, allowing an equilibrium between 1, 2 and adduct [1. DBU] to be established, thus demonstrating reversible oxidative addition/reductive elimination (RE) of $H₂$ at a single main group centre.

> The ability of $L_2Sn(\pi)$ compounds to undergo OA has been inversely correlated with the size of the singlet–triplet (HOMO– LUMO) gap, which may be diminished through the use of extremely strong σ -donor ligands. Aldridge *et al.* have employed a bis(boryl)tin(π) system to achieve the only example of direct OA of H_2 to a mononuclear Sn(π) centre, irreversibly forming the $Sn(v)$ dihydride; boryl ligands are even stronger σ -donors than hydride or alkyl ligands, permitting a successful reaction outcome.^{6d}

Fig. 1 Representative orbital interactions between H_2 and main group compounds: (a) unsaturated E–E compounds e.g. distannynes (Ar $=$ C_6H_2 -2,6- $(C_6H_3$ -2,6- Pr_2)₂-4-X; X = H, SiMe₃, F); for X = H, the reaction is reversible at 80 °C;^{5a,13} (b) single site low-valent centres e.g. carbenes;^{6a} (c) sterically hindered LAs and LBs (FLPs); (d) this work.

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Conversely, the irreversible base-induced RE of H_2 from organostannanes is well-known.¹⁴ Wesemann and others have studied RE from ArSnH₃ and $[(Me₃Si)₂CH]SnH₃$ compounds to yield various mononuclear Sn and Sn-Sn bound species ($Ar =$ terphenyl).¹⁵ Nevertheless, there has yet to be a report of reversible OA and RE occurring on a single $Sn(II)$ scaffold. Lappert's stannylene $[(Me₃Si)₂CH]₂Sn (1)$, which can act as both Lewis acid (LA) and base (LB), is a paradigmatic system for investigating OA to low-valent MG centres, yet to date its reactivity with H_2 has been unexplored.¹⁶ Herein we report the use of FLP methodology to promote formal OA of $H₂$ to this simple dialkylstannylene. Furthermore we document the first example of reversible H_2 addition to a single-site MG complex, which accesses an FLP via reversible dissociation of a classical 1 LB adduct; formation of the latter renders OA of $H₂$ to 1 energetically less favourable, enabling RE to occur from the $Sn(w)$ dihydride and reform 1, which is in equilibrium with $1 \cdot$ LB.¹⁷

Results and discussion

1 is in a rapid solution-phase equilibrium with its dimer $[1]_2$, which has been crystallographically characterised and contains a formal Sn=Sn double bond.¹⁸ When a d_8 -toluene solution of $1/[1]_2$ was placed under an atmosphere of H₂ (4 bar) in a sealed NMR tube, no change was observed in the ${}^{1}\mathrm{H}$ NMR spectrum, even after prolonged periods $($ >48 h), confirming that neither 1 nor $[1]_2$ can react with H_2 alone. Separately, addition of Et₃N (20 mol%) to a solution of 1 resulted in no perturbation of their ¹H NMR resonances, suggesting no interaction between the components; *i.e.* the formation of an FLP.¹⁹ Placing this new mixture under H_2 (4 bar, RT) resulted in the solution turning from deep red to colourless over the course of 24 h, with the $^1\mathrm{H}$ NMR spectrum revealing complete consumption of 1 and a new Sn–H triplet resonance at $\delta = 5.10$ ppm $[{}^{3}J({}^{1}H-{}^{1}H) = 2.2$ Hz] with attendant satellites $[{}^{1}J({}^{117}Sn-{}^{1}H) = 1704 \text{ Hz}; {}^{1}J({}^{119}Sn-{}^{1}H) =$ 1784 Hz], in addition to signals for the $Si(CH₃)₃$ and methine protons $\left[\delta/\text{ppm} = 0.17 \right]$ (s) and -0.42 (t, $\left[\frac{3}{1}H^{-1}\text{H}\right] = 2.2 \text{ Hz}$), respectively] (see Fig. 2). ¹¹⁹Sn NMR spectroscopy showed only a triplet of triplets at -196 ppm $\left[\frac{1}{1}\right]^{119} \text{Sn}^{-1} \text{H}$ = 1784 Hz,
 $\frac{27}{119} \text{Sn}^{-1} \text{H}$ = 87 Hz, which collapsed to a singlet upon $\frac{1}{11}$ $J(^{119}Sn-^{1}H) = 87 Hz$] which collapsed to a singlet upon ¹H decoupling. Collectively these data correspond to the previously unreported dihydride $[(Me₃Si)₂CH]₂SnH₂ (2), which was$ confirmed by comparison with an authentic sample prepared by the reaction of LiAlH₄ and $[(Me₃Si)₂CH]₂SnCl₂$ (see ESI[†] for details).

Isotopic investigation

When D_2 was used in place of H_2 , the methine peak present in the 1 H NMR spectrum of the product mixture resolved as a singlet, while the Sn–H signal was absent and replaced by a Sn–D signal at $\delta = 5.11$ ppm $\left[\frac{1}{117} \text{Sn}^{-2} \text{H}\right] = 262 \text{ Hz}, \frac{1}{119} \text{Sn}^{-2} \text{H}\right] = 274 \text{ Hz}$ in the ²H NMR spectrum. These results demonstrate the formation of dideuteride $2-D_2$,²⁰ and that the Sn-bound protons in 2 must originate from the hydrogen atmosphere.

In order to probe the mechanism further, a d_8 -toluene solution of $1/[1]_2$ and Et₃N was reacted with a 1 : 1 mixture of

Fig. 2 $^{-1}$ H/²H NMR spectra from the reaction of 1 and 20 mol% Et₃N: (a) CH(SiMe₃)₂ region under H₂ (left); H₂/D₂ (1 : 1) (middle); HD (right). (b) SnH region under H_2 (left); H_2/D_2 (1 : 1) (middle; ²H NMR); HD (right). \bullet denotes trace formation of 2 from H_2 in commercial HD gas.

 H_2/D_2 . The resultant ¹H NMR spectrum was very similar in appearance to that of 2, with two exceptions: the relative integration of the Sn–H peak did not match that of the methine signal (1.2 : 2; consistent with the faster rate of reaction with H_2 vs. D_2 – vide infra), and the C–H resonance was composed of overlapping peaks commensurate with a mixture of 2 and $2-D_2$. No spectroscopic evidence was seen for the formation of 2-HD, which was independently and selectively obtained by analogous reaction of $1/[1]_2$ under an HD atmosphere. These observations provide strong evidence that delivery of both atoms from $H_2/D_2/$ HD to a single Sn centre occurs either simultaneously, or in a near-concerted fashion.

Kinetic analysis

By analogy with established FLP systems, and the microscopic reverse of the polar mechanism by which dehydrogenation of $ArSnH₃$ species is proposed to occur,^{15a} we envisaged a reaction mechanism in which 1 and Et_3N form a weakly associated 'encounter complex' which subsequently reacts with H_2 (Scheme 1).²¹ Assuming that encounter complex formation is a rapid pre-equilibrium prior to rate-limiting H_2 activation gives the expected rate law: rate = $k'[1][Et_3N][H_2]$, where $k' = (k_1k_2)$ / k_{-1} . Calorimetric studies on H₂ activation by the FLP Mes₃P/ $B(C_6F_5)_3$ (Mes = 2,4,6-C₆Me₃H₂) found the rate to be very accurately modelled as a single, termolecular step, which formally gives the same rate law.²²

To confirm the order of catalytic Et_3N , the method of time (t) scale normalisation was used;²³ normalisation to the scale of t . $[Et₃N]^x$ resulted in the superposition of all reactant traces only when $x = 1$, confirming the rate to be first order with respect to the amine (Fig. 3a). Determination of reaction order with respect to 1 requires its concentration to be known accurately at any given time in a reaction mixture. However, since the

Scheme 1 Proposed reaction mechanism for H_2 heterolysis by 1, catalysed by $Et₅N$

observed ¹H NMR resonances are a weighted average of the signals from 1 and $[1]_2$ ($\Delta G_{293K} = 3.1$ kcal mol⁻¹), with both species present at significant concentrations under reaction

Fig. 3 (a) Solutions of 1 (0.03 mmol) in d_8 -toluene (0.5 mL) under H₂ (4 bar) containing various base concentrations were prepared. When the stannylene concentration, [Tot], is plotted against the normalised timescale t \cdot [Et₃N]¹, all traces overlap, confirming the order in base to be one. (b) Linearised rate data for a similar solution of 1 (0.03 mmol) in d_8 -toluene (0.5 mL) under H₂ (4 bar) containing Et₃N (0.6 M; 10 eq.).

conditions, simple observation of the concentration of 1 is not directly possible by ¹H NMR spectroscopy.^{18a} The concentration of 1 can, however, be calculated from the total concentration of " R_2 Sn" species in solution, [Tot], present as either monomer or dimer, which are related to the concentrations of 1 and $[1]_2$ by:

$$
[Tot] = [1] + 2[(1)_2]
$$
 (1)

The dimerisation equilibrium of 1 can be expressed as:

$$
K_0 = \frac{[1]^2}{[(1)_2]}
$$
 (2)

Combining eqn (1) and (2) and solving for $[1]$ yields:

$$
[1] = \frac{1}{4} \left(\sqrt{K_0} \sqrt{8[\text{Tot}] + K_0} - K_0 \right) \tag{3}
$$

Inserting eqn (3) into the expected rate law (vide supra) gives:

$$
-\frac{d[Tot]}{dt} = \frac{k^*}{4} \left(\sqrt{K_0} \sqrt{8[Tot] + K_0} - K_0 \right) \tag{4}
$$

where, if the amount of H_2 is sufficiently high that its concentration remains approximately constant:

$$
k^* = \frac{k_1 k_2}{k_{-1}} [\mathbf{B}][\mathbf{H}_2]
$$
 (5)

Rearrangement and integration by substitution of eqn (4) (see ESI†) gives:

$$
\sqrt{\frac{8[\text{Tot}]}{K_0} + 1} + \ln\left(\frac{\sqrt{K_0 + 8[\text{Tot}]} - \sqrt{K_0}}{\sqrt{K_0 + 8[\text{Tot}]_0} - \sqrt{K_0}}\right) - \sqrt{\frac{8[\text{Tot}]_0}{K_0} + 1}
$$

= -k*t (6)

Therefore, plotting the variable portion of the LHS of this expression against t gives a straight line of gradient $-k^*$, confirming the proposed first-order dependence on 1 (Fig. 3b).

Using the known value of $[H_2]$ in toluene at 4 bar (293 K)²⁴ provides a value of $k'_{(Et_3N)} = 0.47 \pm 0.03 \text{ M}^{-1} \text{ s}^{-1}$.²⁵ As well as Et₃N, 2-tert-butyl-1,1,3,3-tetramethylguanidine (Barton's base, TBTMG) and 1,2,2,6,6-pentamethylpiperidine (PMP), were also found to form FLPs with $1/[1]_2$, with corresponding rates of H₂ cleavage: $k'_{(TBTMG)} = 5.0 \pm 0.3 \text{ M}^{-1} \text{ s}^{-1}$, $k'_{\text{[PMP]}} = 0.0266 \pm 0.0018 \text{ M}^{-1} \text{ s}^{-1}$.²⁶ Despite the similar basicity to Et₃N, the bulkier Hünig's base (iPr₂EtN; $pK_{a(MeCN)}$): 18.0)²⁷ was ineffective for $\rm H_2$ heterolysis, as was the weaker base 2,4,6-collidine $(pK_{a(MeCN)}$: 14.98).²⁸ Clearly H₂ activation requires that the LB be sufficiently basic and not too sterically encumbered, in line with observations of other FLP systems.²⁹

A kinetic analysis of the isotopic systems permitted quanti fication of the KIE: $k'_{(H_2)}/k'_{(D_2)} = 1.51 \pm 0.04$ when Et₃N was used as the base. In addition, the acceleration in rate from a more polar solvent could also be quantified: $k_{\rm (THF)}^\prime / k_{\rm (toluene)}^\prime = 1.97~\pm~0.04$ (when Et₃N was used).

Coordinating bases

When the less sterically bulky 1,8-diazabicyclo[5.4.0]undec-7ene (DBU) is used, an interaction with 1 can be clearly seen in the $\mathrm{^{13}C^{11}_{1}}H\}$ NMR spectrum: upon gradual addition of DBU to 1/ $[1]_2$, the methine resonance undergoes a substantial upfield shift, reaching a limiting value of $\delta = 18.5$ ppm (10-fold excess of DBU). Using the established 13 C NMR chemical shift values for 1 and $[1]_2$ (60.0 ppm and 28.7 ppm, respectively),^{18a} this is consistent with a fast equilibrium between 1 DBU, 1 and $[1]_2$ (Scheme 2; see ESI† for full details). A value of $\Delta G = -3.7 \pm$ 0.2 kcal mol⁻¹ for the formation of 1 DBU from $[1]_2$ was obtained from a van't Hoff analysis of variable temperature UV-Vis spectra.

While the reaction of 1/DBU mixtures (containing 0.1–10 equivalents of DBU) with H_2 proceed rapidly, they do not reach completion, indicative of a reversible process (see Fig. S7 in ESI†).

The reversibility can be explicitly demonstrated by the $(CH_3)_3$ Si region of the ¹H NMR spectrum, whereby addition of DBU to a solution of 2 led to the appearance of a signal corresponding to the dehydrogenated mixture 1 · DBU \leftrightarrow 1 \leftrightarrow [1]₂; this increased in intensity at the expense of the $(CH_3)_3S$ peak of 2 (Fig. 4a–c). No \rm{H}_{2} is observed in the $^{1}\rm{H}$ NMR spectrum as the solution was degassed multiple times in order to accelerate the reaction – however, the very small amount of H_2 generated (approx. 0.3 bar) would likely hamper detection. Furthermore, the methine resonance of the 1 \cdot DBU \leftrightarrow 1 \leftrightarrow [1]₂ mixture is subject to a significant upfield shift compared to $[1]/[1]_2$ (dependent upon the DBU concentration), and so is obscured beneath the relatively intense $(CH₃)₃Si$ region. Upon charging this reaction with H_2 , restoration of 2 was rapidly observed (Fig. 4d). For the equilibrium involving $H₂$ (Scheme 2), an equilibrium constant, $K_{\rm eq} = 164\,\pm\,5,$ in favour of 2 can be calculated from the relative intensities of the $(CH₃)₃Si$ resonances, providing $\Delta G = -3.0$ kcal mol⁻¹ (1 bar H₂).

Using the similarly unhindered but less basic 4-(dimethyl amino)pyridine (DMAP) also gave an adduct 1 DMAP, but no reaction with H_2 at room temperature. However, heating a solution of 1 with excess DMAP (4 bar H₂, 2 h, 100 °C) yielded 2 in 31% conversion.

Computational investigation

To gain further insight into the mechanism of $H₂$ activation, DFT calculations were performed for various $1/LB$ pairs;³¹ the computed reaction profiles for both the $Et₃N-$ and DBU-

Scheme 2 Equilibrium between product 2 + DBU and the dehydrogenated mixture 1 · DBU \leftrightarrow 1 \leftrightarrow [1]₂.

Fig. 4 The reversibility of the reaction between $1/[1]_2$, DBU and H₂ can be shown explicitly by a series of ${}^{1}H$ NMR spectra depicting: (a) a solution of 2 (0.03 mmol) in d_8 -toluene (0.5 mL) (b) the same solution with added DBU (0.03 mmol, 1 equivalent) and mesitylene (2%) as an internal standard; (c) after degassing three times over the course of one week, showing the formation of $1 \cdot DBU \leftrightarrow 1 \leftrightarrow [1]_2$; (d) reformation of 2 after the addition of H_2 (4 bar). *Small amount of silicone grease from the independent synthesis of 2.

mediated reactions are depicted in Fig. 5. When $LB = Et_3N$, the reaction was found to proceed via initial H_2 heterolysis leading to a tight ion pair intermediate $[1H]^{-}[Et_{3}NH]^{+}$ (int₁). Facile rearrangement to int_2 and subsequent delivery of the H⁺ to the lone pair on the $[1H]$ ⁻ moiety furnishes 2 (Fig. 6a); a very similar mechanism was found when $LB = DBU$. In support of this polar mechanism, the rate using $Et₃N$ as the LB was found to be faster in THF $(k'_{\text{(THF)}}/k'_{\text{(toluene)}} = 1.97 \pm 0.04)$. The low

Fig. 5 Computed free energy profile for Et₃N- and DBU-assisted H₂ activation with 1. Relative free energies (in kcal mol $^{-1}$) are with respect to $0.5 \cdot [1]_2 + LB + H_2$.

Fig. 6 (a) Structural representations of the computed transition states for the heterolysis of H₂ by 1 and Et₃N. H–H distances are given for TS_1 and int₁. (b) The computed adduct formed between 1 and DBU. All energies (in kcal mol $^{-1}$) are relative to 0.5 \cdot [1] $_2$ + LB + H $_2$. Si–CH $_3$ and C–H groups omitted for clarity.

barriers to rearrangement of the intermediates also offer an explanation as to why H/D exchange is not observed upon reaction with an H_2/D_2 mixture or HD: collapse of the ion pairs is likely much faster than solvent cage escape.

Although the located transition states (TSs) are energetically close-lying, the overall reaction barrier appears to be determined by the H_2 splitting step, which is in line with kinetic measurements. Free energy data computed for the $H₂$ splitting step for reactions with different bases are compiled in Table 1 alongside other properties. For $Et₃N$, TBTMG and PMP, no favourable adduct formation was found with 1, and the ΔG^{\ddagger} values follow the order TBTMG \leq Et₃N \leq PMP, which is consistent with experimental reaction rates. For the coordinating bases DBU (Fig. 6b) and DMAP, adducts favourable relative to free $[1]_2$ and base were computationally determined. This reduces the absolute value of $\Delta G_{\text{reaction}}$ such that an

Table 1 Computational and pK_a data for reactions of a series of bases with 1 and H_2^a

Property	Et ₃ N	TBTMG	PMP	DBU	DMAP
1.1				-3.4	-3.6
TS_1^a	20.4	18.3	21.4	17.0	20.1
int_1^a	18.1	5.4	16.1	6.6	16.8
$\Delta G^{\ddagger a}$	20.4	18.3	21.4	20.4	23.7
$\Delta G_{\rm reaction}$	-7.7	-7.7	-7.7	-4.3	-4.1
$PA^{a,b}$	-270.1	-286.0	-272.7	-283.5	-272.2
pK_a^c	18.8	23.6	18.7	24.3	18.0
$d(HH)^d/\AA$	0.99	0.87	0.96	0.88	0.99

 a Free energy data relative to $0.5\cdot [\mathbf{1}]_2 + \text{base} + \text{H}_2$ (kcal mol $^{-1}$); ΔG^{\ddagger} is activation free energy. b Proton affinity is defined as the free energy of</sup> base + H⁺ \rightarrow baseH⁺. ^c Measured in MeCN.³⁰ ^d H–H distance in TS₁ $(0.76 \text{ Å} \text{ in free H}_2).$

equilibrium is experimentally observed in the case of DBU. For DMAP, the activation barrier is found to be much higher, paralleling results seen by experiment where elevated temperatures are required to obtain product 2.

The energies of all intermediates int_1 are computed to be well above the reference state, which follows from the weak Lewis acidity of 1. The stabilities of int_1 species correlate very well with the general trend in PA and pK_a , but this is not strictly true for the TSs, where steric factors are more important. Unstable int_1 intermediates imply late TSs for the H₂ activation step, which is shown by signicantly elongated H–H distances in the TS structures. The experimentally observed KIE (1.51 \pm 0.04) supports this finding, which is commensurate with ratelimiting H_2/D_2 activation involving considerable H–H/D–D bond breaking.³²

Conclusions

In conclusion, we have demonstrated the ability of FLPmediated reactivity to enable the formal oxidative addition of $H₂$ to an otherwise inert MG centre, and in doing so have also observed the first example of reversible H_2 addition to a singlesite MG complex. We have utilised experimental and computational means to comprehensively explore the mechanism of this transformation and found that H_2 activation in this system differs from those based on more typical FLPs, due to the highenergy nature of the immediate H_2 splitting products, resulting in rare examples of late TSs. The development of methods to harness this FLP-promoted OA/RE H_2 reactivity for hydrogenation catalysis is currently underway.

Conflicts of interest

There are no conflicts to declare.

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the uB97X-D/Def2SVP level. See ESI† for full computational details.

32 In FLPs with early TSs, such as $\text{Mes}_3\text{P/B}(C_6F_5)_3$, a KIE of only 1.1 has been obtained (see ref. 22).