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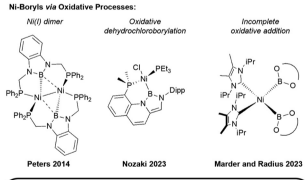
Contrasting reactivity of B-Cl and B-H bonds at [Ni(IMes)₂] to form unsupported Ni-boryls†

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[Ni(IMes)₂] reacts with chloroboranes via oxidative addition to form rare unsupported Ni-boryls. In contrast, the oxidative addition of hydridoboranes is not observed and products from competing reaction pathways are identified. Computational studies relate these differences to the mechanism of oxidative addition: B-Cl activation proceeds via nucleophilic displacement of Cl-, while B-H activation would entail high energy concerted bond cleavage.

The transition metal-catalysed borylation of organic substrates is an efficient method for the preparation of organoboronate esters and boronic acids, 1,2 which serve as important substrates for C-C coupling reactions in the synthesis of complex molecules.^{3,4} Transition metal-boryls (M-BX₂)^{5,6} are key intermediates in these borylation reactions^{7–10} and whilst many of these reactions rely on precious metals, the economic and environmental credentials of the 3d metals have seen a focus on their applications in borylation catalysis. 11 There have been a number of reports of nickelcatalysed borylations¹² but despite Ni-boryls often being proposed as intermediates in these cycles, examples of characterised Ni-boryls remain rare. 13-19 This reflects both the reactivity of the Ni-BX₂ bond and the challenges associated with its formation.

The formation of Pt-boryls via the oxidative addition of B-B and B-X bonds (X = halogen or H) is well established²⁰⁻²⁴ and B-halogen oxidative addition has also been used for the formation of Pd-boryls.²⁵⁻²⁸ In contrast, examples of Ni-boryls formed by oxidative addition are limited to the bis-boryls cis-[Ni(iPr₂Im^{Me})₂- $(B(OR)_2)_2$ (iPr₂Im^{Me} = 1,3-diisopropyl-imidazolin-2-ylidene; OR = $\frac{1}{2}$ catecholato, pinacolato or ethylene glycolato) formed from the



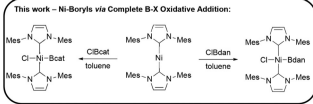


Fig. 1 (top) Examples of Ni-boryls formed by oxidative processes; (bottom) Ni-boryl synthesis via oxidative addition (cat = catecholato; dan = 1,8-diaminonaphthalato).

reaction between the corresponding diboranes and [Ni₂(iPr₂- Im^{Me})₄(μ -(η^2 : η^2)-COD)]/[Ni(i Pr₂ Im^{Me})₂(η^4 -COD)] (Fig. 1).¹⁷ However, detailed analysis of the bonding in these bis-boryls shows oxidative addition to be incomplete with the structure lying between a Ni(II) bis-boryl and a non-classical Ni(0) diborane(4) complex. Other examples of Ni-boryls formed via oxidative processes are the [(PhPBP)Ni]₂ Ni(I) dimer reported by Peters, ¹⁴ resulting from one electron reactivity between the PhPBP(H) pincer ligand and [Ni(COD)2], and the related [(PhPBP)NiCl] and chelating phosphine/boryl formed by oxidative dehydrochloroborylation reported by Nozaki (Fig. 1).19 Boryl formation via simple oxidative addition of a B-X bond (X = halogen or H) at a Ni(0) centre, in a manner analogous to precious metals, has not been reported.

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The reactivity of Ni-boryls is receiving increased attention and the nucleophilicity of the boryl ligand has been demonstrated by the borylation of bromobenzene by [(PNP)Ni(Bcat)]¹³ and the reactivity of cis-[Ni(iPr₂ImMe)₂(Bcat)₂] towards alkyl halides.¹⁸ Incorporation of the boryl into a PBP-pincer ligand has allowed exploitation of its strong σ -donor and potential π acceptor capabilities for the activation of H2, 15,16 dimerization of ethylene²⁹ and cleavage of CO₂.³⁰ With increased interest in Ni-boryls and their applications in bond activation and catalysis we now report the ready formation of stable unsupported Ni-boryls by the oxidative addition of the B-Cl bond at the Ni(0) precursor [Ni(IMes)₂]. Interestingly, the analogous oxidative addition of B-H bonds does not occur, and this stark difference between B-H and B-Cl activation is explored computationally. This further highlights the complexity of the oxidative addition of catalytically relevant substrates at Ni(0) precursors.³¹

The addition of a toluene solution of either ClBcat or ClBdan to a toluene solution of [Ni(IMes)₂], 1, results in a progressive colour change from the dark purple colour of the Ni(0) precursor to a dark yellow or orange solution respectively. This colour change is accompanied by the loss of the ¹¹B NMR resonance associated with the haloborane and the appearance of a new broad resonance at $\delta_{\rm B}$ = 37.3 or 41.6 ppm for the corresponding products of oxidative addition trans-[Ni(IMes)2(Bcat)Cl] (2) and trans-[Ni(IMes)₂(Bdan)Cl] (3) respectively. Crystallisation through the storage of saturated solutions of 2 and 3 at -20 °C allowed for their characterisation by single crystal X-ray diffraction confirming their identity as new unsupported Ni-boryls (Fig. 2).

The Ni-B bond distances of trans-[Ni(IMes)₂(Bcat)Cl] (2) and trans-[Ni(IMes)₂(Bdan)Cl] (3) are 1.908(4) Å and 1.911(3) Å respectively. The ligand trans to the boryl clearly influences the Ni-B bond distance as those observed for 2 and 3 are similar to that of [(PNP)Ni(Bcat)] (1.9091(18) Å), ¹³ in which the boryl ligand is trans to an amide, but within the range defined by being trans to another boryl [(PBP)Ni(Bcat)] (2.015(2) Å)¹⁶ and the heavier halogens in trans-[(iPr2ImMe)2Ni(Bcat)X] (X = Br, 1.872(4) Å; X = I, 1.864(4) Å). 18 Comparison to the Ni-B bond distances observed in cis-[(iPr₂Im^{Me})₂Ni(Bcat)₂] (1.9231(19) Å and 1.9092(18) Å), ¹⁷ for which a B-B interaction is still present and the overall bonding picture is more complex, is also pertinent.

Scheme 1 Reactivity of [Ni(IMes)2(Bdan)Cl] (3) with Na[Et3BH] and MeLi.

Characterisation by multinuclear NMR spectroscopy (see ESI†) showed that clean samples of both 2 and 3 are stable in solution for hours at room temperature. However, the isolation of clean material from the preparation of 2 was hampered by the redistribution of the catechol scaffold as has been observed previously. 32-34 Furthermore, 2 was shown to be unstable in the presence of ClBcat with its addition to samples initiating the decomposition of the Ni-boryl and the further redistribution of the catechol ligand (see ESI†). The Bdan scaffold proved to be far more resistant to redistribution providing a cleaner overall reaction and allowing for more facile isolation of the Ni-boryl 3 in up to 79% of clean material.

To establish both if the chloride ligand of 3 could be substituted and whether the hydride-containing species trans-[Ni(IMes)₂(Bdan)H], the product of oxidative addition of the corresponding hydridoborane, would be a viable target, reactions with Na[Et₃BH] and MeLi were undertaken (Scheme 1). Addition of Na[Et3BH] or MeLi to a toluene solution of 3 at low temperature resulted in a shift in the ¹¹B NMR resonance from $\delta_{\rm B}$ = 41.6 ppm to $\delta_{\rm B}$ = 48.1 and 49.3 ppm, consistent with the substitution of the chloride ligand and formation of the new Ni-boryl complexes trans-[Ni(IMes)₂(Bdan)H] (4) and trans-[Ni(IMes)₂(Bdan)-Me] (5). For 5, filtration followed by removal of the solvent and storage of a concentrated ether solution at -20 °C resulted in the formation of single crystals suitable for X-ray diffraction. The structure of 5 (Fig. 2) confirms the presence of the boryl ligand trans to a Me group and substitution of the chloride ligand. Furthermore, the Ni-B bond distance of 1.9688(17) Å is longer than that in 3 (cf. 1.911(3) Å), consistent with the greater trans influence of the alkyl ligand. Although 5 is thermally unstable in solution (see ESI†), showing significant decomposition in 2 hours at room temperature, it has been fully characterised

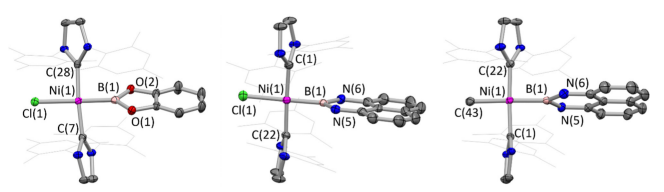


Fig. 2 Molecular structures of 2 (left), 3 (centre) and 5 (right) with thermal ellipsoids at the 50% probability level. Mesityl substituents are shown in wireframe, solvents of crystallisation and hydrogen atoms omitted for clarity

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by multinuclear NMR and elemental analysis. 4 is also thermally unstable but spectroscopic data of isolated material are consistent with its assignment as trans-[Ni(IMes)2(Bdan)H] (see ESI†) with the Ni-H resonance readily identified in the ¹H NMR spectrum at $\delta_{\rm H}$ = -4.62 ppm.

The spectroscopic characterisation of 4 suggests that oxidative addition of hydridoboranes may also be possible at [Ni(IMes)2], similar to well-established Rh-catalysed hydroboration chemistry. 9,35 However, the reactivity of [Ni(IMes)₂] towards the hydridoboranes HBcat and HBdan contrasts strongly to the chloroboranes, with addition of either of these species to a solution of [Ni(IMes)₂] in C₆D₆ providing no evidence for the oxidative addition of either B-H bond and formation of the corresponding Ni-boryls. In the case of HBcat, products resulting from competing one electron pathways were identified and the Ni(1)-containing species [Ni(IMes)₂][Bcat₂] (6, see ESI†) and [Ni₂(IMes)₂][Bcat₂] (7, see ESI†) were both characterised by single crystal X-ray diffraction. The Ni(1) cation 6 has previously been characterised, 36,37 but the mixed Ni(0)-Ni(1) dimer 7, in which each IMes ligand binds through the carbenic carbon to one Ni centre and an \(\eta^6\)-arene interaction to the other Ni, has not been reported and is the IMes analogue of [Ni₂(IPr)₂][OTf] previously reported by Sadighi.38

The addition of HBdan to a C₆D₆ solution of [Ni(IMes)₂] resulted in a dark reaction mixture from which colourless crystals were formed over 1 week. Characterisation of these by single crystal X-ray diffraction (see ESI†) showed these to be the product of B-N dehydrocoupling between two HBdan molecules with an IMes ligand bound to one of the boron centres. This contrasting reactivity of [Ni(IMes)₂] towards both XBcat and XBdan (X = Cl or H) precursors, despite the product of HBdan oxidative addition 4 being prepared via an alternative route, indicates a difference in the mechanism of oxidative addition for B-Cl and B-H bonds at [Ni(IMes)2]. This was therefore investigated computationally, focussing on XBdan for which the experimental outcomes are more clearly defined.

The lowest energy computed reaction profile for the reaction of ClBdan at [Ni(IMes)2] is shown in Fig. 3(a). ClBdan initially forms a B-bound adduct, Int(1-3)1, at +2.6 kcal mol^{-1} (Ni-B = 2.20 Å). The elongated B-Cl bond (1.94 Å), and displacement of the Cl away from the Ni centre (Ni···Cl = 2.77 Å) set up this species for B-Cl activation via nucleophilic displacement. This proceeds via TS(1-3) with a barrier of only 7.8 kcal mol⁻¹ to form ion-pair Int(1-3)2 featuring a Ni-boryl bond (1.93 Å) and an outersphere Cl⁻ held primarily via a Cl⁻···H-N contact (2.25 Å). Attempts to locate a transition state for concerted oxidative addition were unsuccessful, and generally converged on **TS(1-3)**. ³⁹ Ni–Cl bond formation then gives 3 at -22.5 kcal mol⁻¹. B-Cl activation is therefore very favourable thermodynamically and proceeds with an overall barrier of 10.4 kcal mol⁻¹ via nucleophilic displacement of Cl⁻ by [Ni(IMes)₂].

In contrast, HBdan is computed to form a σ -borane complex with $[Ni(IMes)_2]$ at -4.1 kcal mol⁻¹ (Fig. 3(b), Int(1-4)1: Ni-H = 1.67 Å; Ni-B = 2.13 Å; B-H = 1.26 Å). A transition state for concerted oxidative addition on the singlet surface, ¹TS(1-4), was characterised but was prohibitively high in energy (+56.3 kcal mol⁻¹). The corresponding transition state on the (a) B-Cl activation: Nucleophilic Displacement

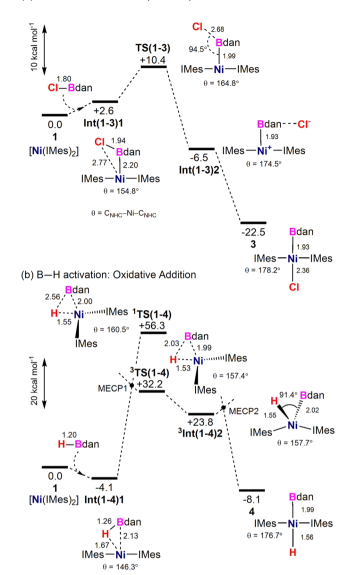


Fig. 3 Computed free energy reaction profiles (kcal mol⁻¹) for (a) B-Cl activation and (b) B-H activation of XBdan at [Ni(IMes)₂]. Note the different scales of the two profiles. Minimum energy crossing points (MECPs) between singlet and triplet surfaces were not located. Method: BP86 (Toluene, B3DJ)/ Def2TZVP//BP86(Toluene)/6-31G(d,p),SDD (Ni, Cl $\{\zeta = 0.640\}$).

triplet surface proved more stable (3TS(1-4), +32.2 kcal mol⁻¹) and could be accessed via spin-crossover MECP1. 3TS(1-4) links to ³Int(1-4)2 at +23.8 kcal mol⁻¹ with a 'see-saw' structure, from which isomerisation and spin-crossover via MECP2 would form 4 at -8.1 kcal mol⁻¹. Overall, while 4 is a viable target thermodynamically, it is kinetically inaccessible via concerted B-H activation as this incurs a prohibitively high energy span of at least 36.3 kcal mol⁻¹ via ³TS(1-4), and possibly higher if either MECP were to lie above this.

The greater kinetic accessibility of B-Cl over B-H activation primarily reflects the ability of Cl⁻ to act as a leaving group: nucleophilic displacement is therefore possible with ClBdan whereas concerted B-H cleavage is the only possibility for Communication ChemComm

HBdan. 40 In addition, concerted B-H oxidative addition may be disfavoured by the combination of the bulky IMes and Bdan ligands; 31,36,41,42 in particular in TS(1-4) the Bdan moiety is forced to be coplanar with the developing Ni···H bond, resulting in the extremely high energy of +56.3 kcal mol⁻¹ (see ESI†).

Finally, alternative reactivity of the stable σ -borane complex Int(1-4)1 was considered computationally.⁴³ One possibility involves IMes transfer to B which proceeds with a barrier of 14.8 kcal mol⁻¹ to form a σ-B-H-bound adduct of HBdan-IMes at {Ni(IMes)} with an energy of +9.6 kcal mol⁻¹. From here reaction with a second equivalent of HBdan to give the dehydrocoupled product seen experimentally along with H2 and the [Ni(IMes)]₂ dimer is computed to be thermodynamically favourable $(G = -9.8 \text{ kcal mol}^{-1}, \text{ see Fig. S25, ESI}^{\dagger})$.

In summary, we report stark differences in the B-X bond activation chemistry of XBR₂ species (X = H or Cl, BR₂ = Bdan or Bcat) at [Ni(IMes)₂], with the facile oxidative addition of B-Cl bonds contrasting the absence of any B-H activation. Computational studies relate these differences to the mechanism involved: B-Cl activation readily proceeds via nucleophilic displacement of Cl-, while B-H activation would entail high energy - and hence inaccessible - concerted bond cleavage.

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Conflicts of interest

There are no conflicts to declare.

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