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Low coordinate NHC-Zinc-Hydride Complexes Catalyze Alkyne C-H Borylation and Hydroboration using Pinacolborane

Richard. J. Procter§†, Marina Uzelac§†, Jessica Cid†, Philip J. Rushworth†, Michael J. Ingleson§*†

§ School of Chemistry, University of Edinburgh, Edinburgh, EH9 3FJ, UK
† School of Chemistry, University of Manchester, Manchester, M13 9PL, UK.
‡ Research and Development, GlaxoSmithKline, Gunnels Wood Road, Stevenage SG1 2NY, U.K.

ABSTRACT: Organozinc compounds containing sp, sp² and sp³ C-Zn moieties undergo metathesis with pinacolborane (HBPin) to produce Zn-H species and organoborate esters (RBPin). This Zn-C-H metathesis step is key to enabling zinc catalyzed borylation reactions, with it used in this work to develop both terminal alkyne C-H borylation and internal alkyne hydroboration. These two conversions can be combined in one-pot to achieve the zinc catalyzed conversion of terminal alkynes to 1,1-diborylated alkenes without isolation of the sensitive (to protodeboronation) alkynyl boronate ester intermediates. Mechanistic studies involving the isolation of intermediates, stoichiometric experiments and DFT calculations all support mechanisms involving organozinc species that undergo metathesis with HBPin. Furthermore, zinc catalyzed hydroboration can proceed via a hydrozincation step, which does not require any exogenous catalyst in contrast to all previously reported alkyne hydroborylations. Bulky N-heterocyclic carbenes (NHCs) are key for effective catalysis as the NHC steric bulk enhances the stability of NHC-Zn species present during catalysis and provides access to low coordinate (NHC)Zn-H cations that are electrophilic yet Brønsted basic. This work provides an alternative approach to access synthetically desirable pinacol-organoboronate esters using earth abundant metal based borylation catalysts.

Keywords: borylation; zinc; hydrides; N-heterocyclic carbene; homogeneous catalysis

Introduction
The considerable utility of organoboranes has made them ubiquitous in synthesis and this provides a continued impetus to develop new routes to form C-B bonds. Hydroboration and C-H borylation of hydrocarbons using commercial boranes, particularly HBPin / B3Pin, represent efficient routes to C-B bonds. Noble metal based catalysts have dominated this area and borylation methodologies now exist that are extremely powerful. The replacement of noble metals with base metals that are earth abundant (e.g. Fe, Cu, Zn, Al), is desirable. However, catalysts based on base metals often react via distinct mechanisms to noble metals due to different propensities to undergo two electron redox steps. For base metal catalyzed borylation metathesis steps at a single redox state are often crucial (Figure 1A and B). While recent progress has been made in developing earth abundant metal based catalysts for alkyne hydroboration (Figure 1A), there are fewer examples of earth abundant metal based catalysts for C-H borylation. Notable exceptions include NHC stabilized Fe and Cu catalysts for arene and alkyne C-H borylation, respectively (Figure 1B). However, most catalysts can only effect one step, hydroboration or C-H borylation, therefore a single earth abundant metal based catalyst that enables both alkyn C-H borylation and hydroboration would be highly desirable. For example, this would represent a useful catalytic conversion for converting terminal alkynes into diborylated alkenes. This conversion is of topical interest as the products are highly useful, however it should be noted that forming 1,1-diborylated alkene from terminal alkynes is rare and much less common than the formation of the 1,2-isomers. Indeed, to the best of our knowledge, it is limited to two reports using either an iridium or a cobalt based catalyst. While highly notable work, Ir and Co have drawbacks with both having extremely low permitted

Figure 1 A-D select relevant previous work. E, inset this work demonstrating Zn-R-H-BPin transmetallation and zinc catalyzed C-H borylation and hydroboration for formation of 1,1-diborylated alkenes.
daily exposures (PDE) as defined by the European Medicines Agency,\textsuperscript{9} while Ir is also of low abundance. Thus the development of a catalyst based on a metal with a high PDE\textsuperscript{10} and high natural abundance that performs C-H borylation and hydroboronation enabling formation of 1,1-diborylated alkenes from terminal alkynes would be significant.

As part of our ongoing interest in catalyzed C-H borylation\textsuperscript{11} and using zinc complexes for \( \sigma \) bond formation we targeted zinc catalyzed C-B bond formation.\textsuperscript{12} In previously reported base metal borylation catalysis a metal hydride is often on an cycle species generated concurrently with the desired organoborane via a \( \sigma \)-bond metathesis step between HBPin and an organometallic species (e.g. Figure 1A and B). Notably to the best of our knowledge, the analogous metathesis reaction of organozinc and hydroboronated species to form a zinc hydride and an organoborane has not been reported. Indeed, the only reported zinc catalyzed C-H borylation, that we are aware of,\textsuperscript{13} is of terminal alkynes using 1,8-naphthalene-diaminato-borane (HBDAn), to afford alkynyl-BDAn. (Figure 1C). Notably, the more mainstream reagent, HBPin, does not generate alkynylBPin compounds under the reported conditions. Furthermore, this report does not invoke the intermediacy of Zn-C species or metathesis of Zn-C with H-B, potentially due to the current lack of precedence for such conversions. Therefore determining the feasibility of Zn-C/H-B metathesis is vital to establish the viability of borylation catalytic cycles mediated by molecular zinc hydrides. While molecular zinc hydrides have attracted significant attention in recent years, this is predominantly for the catalytic hydroduction of polar unsaturated bonds (e.g. C=O, C=N).\textsuperscript{14} In these systems C=O=C=N hydrosilation generates a zinc alkoxide or zinc amide which reacts with a hydroborane to regenerate the zinc-hydride and B-N or B-O containing products (Figure 1D). B-N and B-O bond formation is a significant driving force (due to the partial multiple bond character present in B-N/B-O bonds) which will be absent in putative Zn-C/H-B metathesis reactions. It is however notable that Grignard reagents react with HBPin to form Mg-H and organoBPin species,\textsuperscript{15} thus suggesting the feasibility of an analogous metathesis reaction with organozinc compounds given the commonalities often observed between Mg and Zn chemistry.

In this work we demonstrate that a range of organozinc species do undergo transmetallation with HBPin to form organoBPin species and zinc hydrides. The latter when ligated with bulky N-heterocyclic carbenes (NHC) are effective for both catalytic terminal alkyny C-H borylation and internal alkyny hydroboration. These two conversions can be combined using a single zinc catalyst to achieve the one-pot transformation of terminal alkynes into 1,1-diborylated alkynes using HBPin. 7-DIPP ligated Zn cations (inset Figure 1E, 7-DIPP = 1,3-bis(2,6-diisopropylphenyl)-4,5,6,7-tetrahydro-1H-1,3-diazepin-3-ium-2-ide) are essential for effective catalysis as this NHC affords enhanced thermal stability during catalysis (relative to zinc cations ligated by bulky five membered NHCs) and provides access to low coordinate zinc-hydride cations that are electrophilic yet Brønsted basic.

**Results and Discussion**

**Zn-C / H-B Metathesis Studies**

Commercial diphenylzinc was selected as the initial organozinc reagent for transmetallation studies as it is devoid of the equilibria present using organoazinc-halides and it does not contain zincates which are often present when diorganozinc compounds are made from Grignard reagents and ZnX\textsubscript{2}. On mixing ZnPh\textsubscript{2} with two equiv. of HBPin in THF rapid transmetallation proceeded leading to formation of PhBPin as the major boron containing product (by \( ^1 \)H and \( ^11 \)B NMR spectroscopy). Insoluble material also was observed to form, presumably (ZnH\textsubscript{2})\textsubscript{2}. To confirm that THF coordination to any zinc species is not significantly affecting the transmetallation outcome the reaction was repeated in the more weakly coordinating solvent ortho-dichlorobenzene (o-DCB). From this reaction PhBPin again was observed as the major boron product along with insoluble material (again presumably (ZnH\textsubscript{2})\textsubscript{2}). Similar outcomes were observed when combining commercial ZnEt\textsubscript{2} and HBPin (with EtBPin observed as the major boron containing product) confirming that Zn-C/H-BPin transmetallation proceeds with both dialkyl- and diaryl-zinc compounds.

\[
\begin{align*}
\text{ZnPh}_2 + 2 \text{HBPin} & \rightarrow \text{PhBPin} + \text{ZnH}_2 \\
\text{ZnEt}_2 + 2 \text{HBPin} & \rightarrow \text{EtBPin} + \text{ZnH}_2 \\
\text{THF or o-DCB} &
\end{align*}
\]

**Scheme 1**: Zn-R / HBPin transmetallation reactions.

Solubilizing the zinc hydride product formed post Zn-C/H-B transmetallation is essential to preclude precipitation shifting Zn-C/H-B metathesis equilibrium reactions to completion. StERICally demanding NHCs were selected as solubilizing ligands as low coordinate NHC-ligated zinc hydrides have been previously reported to be thermally stable at 20°C in solution for days while also being useful in a range of catalytic applications.\textsuperscript{14} The two NHCs selected were N,N-bis(2,6-diisopropylphenyl)imidazol-2-ylidene (IDIPP) and 7-DIPP as these both have large percentage buried volumes (%V\textsubscript{bur}) which enforce low coordination numbers at metal centres.\textsuperscript{16} (7-DIPP)ZnCl\textsubscript{2}, 1, and (7-DIPP)ZnPh\textsubscript{2}, 2, were synthesized by addition of the free carbene to ZnCl\textsubscript{2} and ZnPh\textsubscript{2}, respectively. Notably, the solid state structures of both 1 and 2 on crystallization were monomeric and three coordinate at zinc. For the dihalide analogue 1 (which was crystallized from THF) this is in contrast to (IDIPP)ZnCl\textsubscript{2} which coordinated THF, forming (IDIPP)ZnCl\textsubscript{3}(THF) which is four coordinate at Zn.\textsuperscript{17} Comparison of the buried volumes of 1 (49.7%) and 2 (44.7%) to a range of (IDIPP)Zn analogues confirms the greater steric impact of 7-DIPP relative to IDIPP. While the structure of 2 is comparable to previously reported (7-Mes)ZnMe\textsubscript{2},\textsuperscript{18} it is notable that there is a significant angle between the N-C-N plane and the C-Zn-C and C-Zn-C\textsubscript{2} planes (64.6°) in 2 presumably reducing steric interactions between the DIPP and Ph moieties.

![Figure 2](image-url)  
**Figure 2.** Solid state structures of 1 (left) and 2 (right) ellipsoids at 50 % probability and hydrogens omitted for clarity. Selected bond distances (Å) and angles (°) for 2: \( \text{NHC-C-Zn} = 2.098(2); \text{NHC-C-Zn} = 1.989(2) \) and 2.011(2); \( \text{nC-C-Zn-C\text{\_NsH}} = 118.74(7); \text{pC-C-Zn-C\text{\_NsH}} = 123.75(7) \) and 116.90(7).
With a well-defined (NHC)ZnR₂ species in hand transmetalation with HBPin was assessed. Compound 2 was combined with two equivalents of HBPin in THF which led to the formation of PhBPin as the major new boron containing product (by ¹H and ¹¹B NMR spectroscopy) along with other products, including the NHC-derived species, 3 (Scheme 2). In our hands a putative (7-DIPP)ZnH₂ complex could not be isolated from this mixture. Furthermore, other routes to form (7-DIPP)ZnH₂ species (commencing from 1 or 2) were also unsuccessful in our hands. For full characterization the independent synthesis of 3 was achieved by combining 7-DIPP and 2 equiv. of HBPin. Compound 3 is a NHC ring opened product derived from two H transfers from B to C and is related to previously reported products from the reaction of 1,3-bis(2,6-di-isopropylphenyl)-imidazolidin-2-ylidene (SIDIPP) with HBCat or HBPin.¹⁹ The formation of 3 from reactions commencing from 2 / HBPin would potentially also lead to (ZnH₂)₃ formation, precipitation of which could still affect the metathesis equilibrium. To provide support for Zn-H formation by Zn-C/H-B metathesis being thermodynamically favored (IDIPP)ZnPh₂, 4, was synthesized by the combination of IDIPP and ZnPh₂. Addition of two equivalents of HBPin to 4 in o-DCB or THF led to formation of PhBPin as the major new boron containing species along with the previously reported complex ([(IDIPP)ZnH₂]₂(μ-H)₂).²⁰ This confirmed that in this case Zn-C/H-B metathesis is thermodynamically favored and not driven by the insolvability of one of the products.

Scheme 2: Top and bottom, transmetalation between (NHC)ZnPh₂ and HBPin. Middle, the formation of 3. As zinc-catalyzed terminal alkyne C-H borylation (Scheme 1C) is the only zinc catalyzed C-H borylation reaction reported to date to the best of our knowledge,¹⁵ we explored the transmetalation between Zn-alkynyl species and HBPin. Zn-bis alkynyl complex 5 (Figure 3), was synthesized and crystallized from THF, with the solid-state structure indicating it is also a coordinate at zinc. Compound 5 has a shorter bond distance for NHC-Zn (2.037(3) Å) compared to 2 (2.098(2) Å) which contributes to the higher %Vₕob determined for 7-DIPP in 5 (47.3°). There is also a large area in 5 between the N-C-N and the alkynylC-Zn-Calkynyl planes (89.2°) presumed to reduce steric interactions. The reaction of 5 with excess HBPin in THF led to formation of alkynylBPin 6a and compound 3 (Figure 3, by ¹H and ¹¹B NMR spectroscopy after 18 h). Inspection of the in-situ NMR spectra at shorter reaction times revealed that the formation of NHC-ring opened product 3 occurs prior to formation of 6a, indicating, in contrast to 2 and 4 that dissociation of 7-DIPP from zinc in compound 5 and formation of 3 occurs prior to the reaction of the resultant Zn(alkynyl)(THF), species reacting with HBPin to form 6a. Dissociation of bulky ring expanded NHCs from diorganozinc species has been previously observed.¹⁸ Finally, for comparison the IDIPP congener, compound 7 (Figure 3 bottom), was synthesized in-situ from the reaction of 4 with two equiv. of 4-ethylnyltoluene in THF (with evolution of benzene observed by NMR spectroscopy). This Zn-alkynyl species underwent transmetalation with excess HBPin to form only one equivalent of 6a, with the NHC remaining coordinated to zinc. The zinc containing product displayed NMR spectra consistent with the formulation [(IDIPP)Zn(alkynyl)(μ-H)]₂. With only one equivalent of HBPin being consumed in the reaction with 7 (the second zinc-alkynyl moiety does not undergo b-bond metathesis even after long reaction times and with additional HBPin added) this suggests that Zn-Calkynyl / H-BPin metathesis is less energetically favorable than metathesis with Zn-Csp³ analogues (e.g. 4) where two metathesis steps take place. Combined, the above reactions clearly demonstrate that Zn-C / H-B transmetalation between ZnR₂ / (NHC)ZnR₂ species and HBPin proceeds for sp² and sp³ carbon centres and to some extent for C-sp centres. With the viability of metathesis confirmed and the reactivity of molecular zinc hydrides with terminal alkenes to form zinc alkynyls and H₂ previously documented, the feasibility of combining these steps to achieve zinc catalyzed alkyne borylation with HBPin was explored.

Catalytic Terminal Alkyne C-H Borylation

Catalytic studies started with compounds 2 and 4 as these are readily accessible catalyst precursors with the proposed on-cycle (NHC)Zn-hydride species generated by transmetalation with HBPin (forming PhBPin as the by-product). However, using 2 or 4 as the (pre)catalyst in o-DCB, MeCN or THF under a range of conditions led to only partial consumption of 4-ethylnyltoluene, incomplete (<50% in all cases) formation of the desired compound 6a and formation of multiple other products (by in-situ multinuclear NMR spectroscopy) some of which were consistent with products from the reaction of the NHC with HBPin (e.g. 3). During these reactions using 2 or 4 insoluble material also was observed to form concomitant with catalyst deactivation. Catalyst decomposition (as indicated by precipitation of insoluble material) occurred more readily on heating to 60°C with no significant improvement in the conversion to 6a using 2 or 4.
Scheme 3: Protonolysis of 2 and 4 to form complexes 8 and 9.

With catalyst deactivation potentially precluding high conversions to 6a using 2 or 4 cationic analogues of general formula (NHC)ZnPh(anion) were targeted, as hydrides of the general formula, (NHC)Zn(H)(anion), have been reported to be significantly more thermally stable than neutral (NHC)ZnH₂ analogues.¹⁴ε Compound 2 was combined with one equivalent of the strong Bronsted acid HNTf₂ which led to benzene evolution and formation of (7-DIPP)ZnPh(NTf₂), 8. (IDIPP)ZnPh(NTf₂), 9, was synthesized via an analogous route (Scheme 3).

Table 1. Catalyst optimization for alkyne dehydroboration

<table>
<thead>
<tr>
<th>Entry</th>
<th>Catalyst</th>
<th>T (°C)</th>
<th>Time/h</th>
<th>6a (%)</th>
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<tr>
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<td>20</td>
<td>72</td>
<td>94</td>
</tr>
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<td>8⁺</td>
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<td>99</td>
</tr>
<tr>
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<td>9⁺</td>
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</tr>
<tr>
<td>6</td>
<td>9⁺</td>
<td>60</td>
<td>21</td>
<td>11</td>
</tr>
<tr>
<td>7</td>
<td>ZnPh₂</td>
<td>60</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>8</td>
<td>PhZnNTf₂</td>
<td>60</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>9⁺</td>
<td>8 + base</td>
<td>20</td>
<td>18</td>
<td>41</td>
</tr>
</tbody>
</table>

Reaction conditions as shown above unless otherwise indicated. a = using 5 mol% at [0.8 M] 8 b = made in-situ from ZnPh₂ / HNTf₂. c = base = 2,6-di-tert-butylpyridine. d = Yields by in-situ 1H NMR spectroscopy based on the ratio of the product Vs. mesitylene added as internal standard.

In o-DCB or benzene the use of 8 as a borylation catalyst did proceed but led to competitive terminal alkyne C-H borylation andhydroborination in o-DCB (approx. 1:1 mixtures of products were formed) while in benzene the C-H borylation product dominated but lower yields were observed (ca. 75%). In contrast, in THF only alkyne C-H borylation to form 6a was observed over a range of reaction temperatures. While alkyne C-H borylation in THF was slow at 20°C reactions with 8 could be heated in THF (table 1, entries 1-4) without significant catalyst deactivation, thus enabling high conversion to 6a. Notably, under a range of conditions the (IDIPP)Zn analogue 9 gave very low conversions (e.g. entries 5 and 6). Reactions using 9 led to the formation of a black precipitate on heating at 60°C, consistent with the formation of zinc metal and indicating the lower thermal stability under these catalytic conditions for (IDIPP)Zn congeners versus (7-DIPP)Zn congeners. This is consistent with the absence of any additional turnover with longer reaction times (entries 5 Vs 6). In the absence of any NHC, using ZnPh₂ or PhZnNTf₂ as catalysts, extremely low conversions were observed (entries 7-8) indicating the importance of NHC ligation.

Finally, in an attempt to use a base to catalyze any alkyne deprotonation steps (as proposed previously for zinc catalyzed alkyne C-H borylation with HBDan, see Figure 1C)¹³a compound 8 was utilized in combination with a hindered pyridyl base (to prevent pyridyl coordination to zinc and catalyst deactivation as observed previously in carbonyl hydrosilylation using related zinc systems)¹⁴ε. However, this resulted in no significant increase in conversion (entry 9) relative to borylation reactions in the absence of this base, indicating under these conditions that this base is not involved in any deprotonation step.

With effective zinc catalyzed alkyne C-H borylation conditions in hand, the borylation of a range of terminal alkyynes was investigated. Aryl, heteroaryl, vinyl and alkyld substituted alkyynes were all viable substrates with 1.1 equiv. of HBpin sufficient for high yields of 6 in each case (Figure 4). This methodology was also applicable to a range of terminal arylalkynes bearing electron donating and withdrawing functionalities such as OMe, F, Cl, NMe₂ and CF₃. However, substrates containing nitrile and nitro-substituents were not amenable as these were reduced in preference to alkyne C-H borylation. Given the large steric demand of 7-DIPP it is notable that ortho-ClC₆H₄ and mesityl substituted alkyynes, 6g and 6i, were successfully borylated under these conditions.

Figure 4: Scope of (7-DIPP)Zn catalyzed alkyne C-H borylation. Yields by in-situ 1H NMR spectroscopy based on the ratio of product Vs. mesitylene added as internal standard. a = 22 h.

Formation of 1,1-Diborylated Alkenes

While alkyne boronate esters such as 6a-m are useful in their own right²² they are highly prone to protodeborylation. Therefore we sought to perform a subsequent transformation of the alkyne boronate esters in-situ using the same zinc catalyst targeting more robust to protodeborylation products that are also highly useful. Guided by the observation of alkyne hydroboration as a by-product during the formation of 6a in aromatic solvents we targeted the hydroboration of 6a. This reaction would proceed via initial alkyne hydrozincation with a subsequent transmetalation of the Zn-alkenyl intermediate with HBpin furnishing diborylated alkenes (e.g. 10a, Figure 5). Notably, while Zn-B and Zn-Si bonds have been reported to add to alkyynes in...
the absence of catalysts, to the best of our knowledge, catalyst free alkyne hydrozincation has not been reported prior to this work. The hydroboronation of alkynyl boronate esters could produce 1,1 or 1,2 diborylated alkenes, both useful products. However, catalytic routes to form the 1,1-isomer from non-activated alkenes (e.g. alkynes that are not Michael acceptors) are much more limited with those reported to date requiring noble metal or cobalt catalysts.

![Figure 5](image)

**Figure 5:** Hydroboration of 6a to 10a using precatalyst 8.

The hydroboration of 6a using HBPin and 10 mol% 8 as a catalyst precursor was achieved in good yield on heating in a range of hydrocarbon solvents (toluene, benzene or methyl-cyclohexane). In hydrocarbon solvents the 1,1-diborylated alkene, 10a, was formed in high yield (> 95% after heating for 28 h at 90°C) with no 1,2-isomer observed by NMR spectroscopy. To provide support that the catalysis is zinc mediated and that B-H species formed by borane substituent redistribution (minor 1H resonances centered at 22 ppm are observed during this process consistent with formation of (RO)2B products) are not responsible for the observed hydroboration of 6a, control experiments using BH3:THF were carried out. Using 10 mol% BH3:THF (1M in THF) only 25% hydroboration of 6a to 10a was observed after 33 hours at 90°C. This is drastically less than the outcome observed using 10 mol% of 8 as a precatalyst which gives complete conversion of 6a to 10a under identical conditions. Hydroboration conversion using 8 remains high (> 98% conversion after 28 h) even when 5 equivalents of THF was added to the reaction mixture (disfavoring THF being a catalyst poison). These results suggest that (7-DIPP)Zn based species are responsible for the hydroboration reactivity, which proceeds via a highly selective hydrozincation of 6a and subsequent transmetalation with HBPin.

A combined C-H borylation / hydroboration protocol next was developed to form 1,1-diborylated alkenes in one-pot from terminal alkenes operating all in hydrocarbon solvent (C6D6 is used in sealed tubes for NMR monitoring purposes but the transformation also proceeds effectively in toluene). The one-pot conversion of the terminal alkyne to 10a using 8 as the catalyst precursor proceeded in good yield at 90°C with no solvent switch required. Notably, performing the reaction in C6D6 at the lower temperature of 60°C led to 6a being the major product with minimal formation of 10a observed. The higher temperature required to form 10a indicates a greater kinetic barrier (relative to formation of 6a) and that C-H borylation precedes hydroboration under these conditions. With conditions in hand using 8 as a catalyst precursor a number of terminal alkenes were converted in one-pot to the 1,1-diborylated alkenes, 10a-h, in moderate to good yield without isolation of the intermediate alkynyl boronate esters 6 (Figure 6). Performing the tandem functionalization for longer duration or increasing the number of equivalents of HBPin used (from 4 to 5) increases the conversion to the 1,1-diborylated alkenes, indicating that there is still significant catalyst remaining (the mass balance in these reactions is predominantly 6x). This one-pot diborylation process was amenable to a range of terminal alkenes (including aryl groups substituted with electron withdrawing and donating groups and ortho, meta and para substituents). It should be noted that electron withdrawing groups required longer for comparable conversion, suggesting a build-up of cationic character on the alkyne moiety in the transition state of the hydrozincation process. This transformation complements the recently reported cobalt catalyzed conversion of terminal alkynes into 1,1-diborylated alkenes (which focused predominantly on alkyl alkynes). Furthermore, this zinc catalyzed process proceeds with no enyne by-products from homodimerization observed in contrast to the cobalt catalyzed process. As the diborylated alkyne products, 10, are much more resistant to protodeborylation than the respective alkynyl boronate esters they are readily isolated by column chromatography in moderate to good yields. 1,1-diborylated alkenes have been previously utilized in a wide range of transformations (e.g. Suzuki–Miyaura cross coupling, halogenation to the 1,1-dihaloalkenes and oxidation to carboxylic acids) thus are extremely useful.

**Mechanistic Studies**

**Precatalyst Activation and Synthesis of Zn-H Species:**

Compound 8 is a readily accessed catalyst precursor, with a plausible mechanism (Figure 7) commencing by 8 undergoing transmetalation with HBPin to form a low coordinate ([7-DIPP]ZnH)2 species (presumably solvent and/or anion coordinated). Alternatively, 8 can react with a terminal alkyne by protonolysis to evolve benzene and form ([7-DIPP]Zn(alkynyl)]+ species (again the zinc centre will be solvent / anion coordinated). These two cations represent the key proposed on-cycles species that would enable the first process, terminal alkyne C-H borylation. Inspection of the catalytic C-H borylation reactions (in THF and in benzene) by in-situ 1H and 11B NMR spectroscopy indicated the presence of PhBPin in all cases formed by ZnPh / H-B transmetalation is kinetically more facile under these conditions than Zn-Ph protonation and thus metathesis is the key step for accessing zinc-hydride species in this system.
Isolation and structural characterization of the various (7-DIPP)Zn complexes shown in Figure 7, particularly the proposed zinc hydride was targeted. Compound 8, (7-DIPP)ZnPh(NTf$_2$), was isolated as a crystalline solid from THF with the [NTf$_2$] anion bound to the zinc center in a bidentate fashion (Figure 8, left) with no THF bound to zinc in the solid state structure. The majority of the structural metrics for 8 are comparable to the related four coordinate complex (NHC)Zn(C$_2$F$_5$)$_2$O($	ext{CR}$), A (where the carboxylate is also bidentate, R = Cp)*, with similar C-Zn distances. However, the Zn-O distances in 8 (in the range 2.31(4) to 2.33(2) Å) are much larger than that in A (Zn-O = 2.086(2) Å)* suggesting that [NTf$_2$]$^-$ in 8 is more weakly interacting with the zinc centre.

To confirm that Zn-C/H-B transmetalation does proceed from 8, 8 was reacted with HBPin in THF at 60°C. Transmetalation between 8/HBPin in THF does form PhBPin, however, in the absence of terminal alkyn, to react with the zinc-hydride product) subsequent reaction with more HBPin occurs in THF to form 3 and other currently unidentified products, preventing isolation of a zinc-hydride. However, in o-DCB the transmetalation between 8/HBPin proceeds cleanly and alongside PhBPin a single new zinc product that contained a resonance consistent with a terminal Zn-H (δ$_H$ = 3.65 ppm 1 H integral relative to the new NHC resonances) was observed as the major new (7-DIPP)Zn species. This species was isolated by recrystallization from THF coordination to zinc was forthcoming from the solid state structural analysis of crystals obtained from a THF solution of 8. This revealed displacement of NTf$_2$ by one molecule of THF and formation of a three coordinate Zn-h cation, [(7-DIPP)ZnH(THF)]$^+$, termed [11-THF]$^+$. The closest anion--zinc contacts are all long, with 5.41 Å the shortest Zn--F and 5.93 Å the shortest Zn---O contact to NTf$_2$. The geometry at zinc in [11-THF]$^+$ is approximately trigonal planar (Σangles) at Zn = 359.2$^o$) with a relatively (compared to B) large C-Zn-H angle again observed (133(2)$^o$). Other bulky NHCs coordinated to cationic zinc-hydrides lead to ≥ four coordinate zinc centres due to solvent/anion coordination or oligomerization. While 2 and 3-coordinate neutral zinc hydride complexes are known, [11-THF]$^+$ is the first example of a low (< 4) coordinate cationic zinc hydride, to the best of our knowledge, which is accessible due to the large steric demand of the 7-DIPP ligand.

**Alkyne C-H-Borylation:**

The solution state reactivity of 11 relevant to the catalytic cycle proposed in Figure 7 next was explored. Firstly, using 11 in place of 8 as catalyst in the C-H borylation to form 6a resulted in a greater conversion to 6a after 90 minutes relative to the conversion when using 8. Furthermore, no induction period is observed for the formation of 6a when using 11. This is consistent with 11 (or [11-THF]$^+$) being an on-cycle species, or having a low barrier to form an on-cycle species. Next, 11 was reacted with 4-ethyltoluene, which led to concomitant consumption of both the Zn-H and alkynyl C=H resonances in
the \(^1\)H NMR spectrum and the formation of a single major (7-DIPP)Zn containing product. This new species was not isolable in our hands, however, the same compound was obtained by reaction of 5 with one equivalent of HNTf\(_2\) (by \(^1\)H NMR spectroscopy), with the expected terminal alkyne by-product from protonolysis observed (Scheme 4). This supports the identification of this new complex as [(7-DIPP)Zn(alkynyl)NTE], 12, which again maybe anion and/or solvent coordinated.

Scheme 4: Formation of compound 12 (shown as anion coordinated, in THF solution THF could also be bound to zinc, displacing NTE; to produce [12-THF]).

The use of 12 in the C-H borylation catalysis in THF also led to effective formation of 6a with a greater conversion after 90 minutes relative to that using 8, indicating 12 is also an on-cycle species, or has a low barrier to form an on-cycle species. The addition of HBPin to 12 (in the presence of one equiv. of 4-ethyl toluene) resulted in formation of 6a (the zinc-hydride, 11 or [11-THF]), is not observed by NMR spectroscopy as it is consumed by the rapid reaction with 4-ethyl toluene to reform 12) confirming the feasibility of this Zn-alkynyl / H-BPin metathesis step. Repeating the reaction between 12 and excess HBPin in THF in the absence of any additional terminal alkyne still led to formation of 6a, but the ring opened NHC derived product 3 was observed instead of the zinc-hydride (11 or [11-THF]), consistent with 11 not being stable in THF in the presence of excess HBPin. Thus, all of the steps of the proposed C-H borylation cycle shown in Figure 7 are viable, but in THF the zinc hydride 11 can react with additional HBPin (leading to catalyst deactivation) or with a terminal alkyne (leading to turnover), with the latter reaction dominating. Notably, the combination of stoichiometric 11 and 6a in CD\(_2\) (a solvent in which combinations of 11/HBPin do not result in decomposition to form 3) leads to formation of HBPin as the major boron containing species (Figure 9) and resonances consistent with 12. However, not all 11 and 6a are consumed (even on heating) in this reaction. These observations suggest that formation of zinc hydride 11 and 6a from 12 and HBPin is an equilibrium process favouring 12 and HBPin and that consumption of 11 in a subsequent step (e.g. reaction with a terminal alkyne to form Hz and reform 12) is required to drive the reaction to completion.

Figure 9: Reversible sigma bond metathesis between zinc alkynyl 12 and HBPin.

Alkynе Hydroboration:

Regarding the second cycle in figure 7, the alkynyl boronate esters are proposed to undergo hydroboration and then metathesis with HBPin to form the diborylated alkenes. Hydroboration of 6a is more effective in hydrocarbon solvents than in THF, an observation attributable to the decomposition of zinc-hydride 11 in the presence of excess HBPin in THF (and in the absence of terminal alkyne). In contrast, in benzene and other weakly coordinating solvents (e.g. o-DCB) 11 is more robust towards decomposition from reaction with HBPin, permitting prolonged heating which is essential for the transformation of 6 into 10. To the best of our knowledge uncatalyzed hydrozincation of alkynes has not been previously documented. However, direct observation of hydrozincation products by combining 11 and 6a is complicated by the kinetic product being the zinc-alkynyl 12 and HBPin. It is only on prolonged heating that the diborylated products 10 are observed, indicating they are the thermodynamic products from this reaction and suggesting that the hydrozincation of 6a with 11 has a higher barrier than the \(\sigma\)-bond metathesis to form 12 and HBPin.

Figure 10: Hydrozincation of alkyne 13 with 11 and subsequent metathesis with HBPin. Inset, complex C.

To probe the feasibility of the unprecedented catalyst free alkyne hydrozincation step the internal alkyne 1-phenyl-1-propyne, 13, was reacted with stoichiometric 11 in CD\(_2\). While no reaction was observed at room temperature, on heating complete consumption of both 11 and 13 was observed, producing two new species in an approximately 1:1 ratio. These were spectroscopically consistent with the hydrozincation products 14a and 14b (Figure 10). Key diagnostic resonances from the \(^1\)H NMR spectrum included a new alkynyl singlet at 5.32 ppm (for 14a) and a quartet at 4.82 ppm (for 14b), the latter is coupled to the alkynyl-Me group. In the HSQC spectrum these resonances were found to correlate to \(\delta_{13C}\) alkynyl resonances at 142.6 and 140.3 ppm. Crucially, both of these species undergo \(\sigma\)-bond resonance significantly further downfield. While the selectivity in the hydrozincation/hydroboration of 13 is low in contrast to that observed in the hydroboration of 6. We attribute this disparity to the cationic nature of the zinc centre in 11, leading to build-up of positive charge on the alkyne during hydrozincation which is significantly more stabilized by the aryl moiety than the BP\(^n\) in 6 leading to the observed selective formation of 1,1-diborylated products. In contrast with alkyne 13 the electronic disparity between Ph / Me is less (than Ph/BP\(^n\)) resulting in lower selectivity.
DFT Calculations
Catalyst Speciation / Key Properties

The mechanistic studies discussed above indicate the involvement of zinc-hydride and zinc-alkynyl complexes in alkyne C-H borylation, with both potentially being three coordinate mono solvated (NHC)Zn cations in THF. To determine the degree of solvation in THF for subsequent computational investigations the energy change on binding of a second molecule of THF to [11-THF]+ was explored at the M06-2x/cc-pVTZ level. This was performed as regards to the solid state structure in solution it is feasible that an additional THF molecule binds to zinc in [11-THF]+. At this level the calculated structure of [11-THF]+ revealed similar metrics to the solid state structure. For [11-THF]+ the binding of a second molecule of THF to zinc was found to be effectively thermoneutral at 298 K (Figure 11), therefore the three coordinate at zinc species, [11-THF]+, will dominate at raised temperatures and thus mono-solvated species are used in all the subsequent DFT calculations (including the hydrozincation process).12

Analysis of the electronic structure of [11-THF]+ revealed that despite the low coordination number at zinc and the unit positive charge the hydride has significant negative charge (-0.544) based on natural bond orbital (NBO) calculations. This is comparable in magnitude to that previously calculated (at the same level) for the hydride in the four coordinate (at zinc) complex ([DIPP]ZnH(THF)(OTf)).3e Another notable point is that the LUMO of [11-THF]+ has significant zinc p orbital contribution, in contrast to the four coordinate at zinc bis-solvate complex [11-THF]+ (Figure 11, inset) which has negligible zinc contribution to the LUMO. Finally, the energy of the LUMO of [11-THF]+ is significantly lower than for [11-THF]+ as expected based on their respective coordination numbers. Thus [11-THF]+ contains a highly electrophilic zinc centre and an hydridic (thus Brønsted basic) Zn-H moiety.

In contrast to the reactivity of the (7-DIPP)Zn-alkynyl complex 12 with HBPin (an endergonic reaction), the reaction of the phenyl derivative, 8, with HBPin furnishes the zinc-hydride complex 11, indicating an exergonic reaction. Examining this transformation confirmed the transmetallation of [8-THF]+ with HBPin to form [11-THF]+ and PhBPin is exergonic (by 9.0 kcal mol⁻¹, Figure 13). Inspection of the electronic structure of [8-THF]+ and [12'-THF]+ for any significant differences reveals closely comparable LUMOs for the LUMO of [8-THF]+ see Figure 13), but distinct occupied frontier orbitals. For [8-THF]+, the HOMO has significant Zn-C sigma bond character; however, for the alkynyl analogue, [12'-THF]+, the HOMO has Ph=C-C π character. It is lower in energy HOMO-8 of [12'-THF]+ that is the highest energy occupied orbital that contains significant Zn-C sigma bond character, which indicates a reduced nucleophilicity of the zinc-alkynyl moiety relative to the zinc-phenyl. Regarding the thermodynamics of σ-bond metatheses, recent work has determined that the bond dissociation energy for the C-B bond in alkynyl-boronic acids is significantly (ca. 25 kcal mol⁻¹) greater than the C-B bonds in phenyl boronic acids.13 This indicates that the zinc-alkynyl bond in (7-DIPP)Zn compounds has to be significantly stronger than the Zn-Ph bond for the reaction of zinc-hydride 11 and 6a to form HBPin / 12 to be thermodynamically favored. This difference is further supported by the isodesmic reaction (bottom inset, Figure 13) between PhBPin/[12'-THF]+ and 6e/[8-THF]+ which is found to be significantly exergonic. We attribute this to the greater C 2s orbital character in the Zn-(alkynyl) bond relative to that in the

Figure 12: Calculations at the M06-2x/lan12dz/6-311G/PCM(THF) level for: top, the C-H borylation process.

In contrast to the reactivity of the (7-DIPP)Zn-alkynyl complex 12 with HBPin (an endergonic reaction), the reaction of the phenyl derivative, 8, with HBPin furnishes the zinc-hydride complex 11, indicating an exergonic reaction. Examining this transformation confirmed the transmetallation of [8-THF]+ with HBPin to form [11-THF]+ and PhBPin is exergonic (by 9.0 kcal mol⁻¹, Figure 13). Inspection of the electronic structure of [8-THF]+ and [12'-THF]+ for any significant differences reveals closely comparable LUMOs for the LUMO of [8-THF]+ see Figure 13), but distinct occupied frontier orbitals. For [8-THF]+, the HOMO has significant Zn-C sigma bond character; however, for the alkynyl analogue, [12'-THF]+, the HOMO has Ph=C-C π character. It is lower in energy HOMO-8 of [12'-THF]+ that is the highest energy occupied orbital that contains significant Zn-C sigma bond character, which indicates a reduced nucleophilicity of the zinc-alkynyl moiety relative to the zinc-phenyl. Regarding the thermodynamics of σ-bond metatheses, recent work has determined that the bond dissociation energy for the C-B bond in alkynyl-boronic acids is significantly (ca. 25 kcal mol⁻¹) greater than the C-B bonds in phenyl boronic acids.13 This indicates that the zinc-alkynyl bond in (7-DIPP)Zn compounds has to be significantly stronger than the Zn-Ph bond for the reaction of zinc-hydride 11 and 6a to form HBPin / 12 to be thermodynamically favored. This difference is further supported by the isodesmic reaction (bottom inset, Figure 13) between PhBPin/[12'-THF]+ and 6e/[8-THF]+ which is found to be significantly exergonic. We attribute this to the greater C 2s orbital character in the Zn-(alkynyl) bond relative to that in the

Figure 11: Top, energy change on binding a molecule of THF to [11-THF]+. Inset left, LUMO of [11-THF]+ and inset right LUMO of [11-THF]2+ (isoelectronic species) also calculated. NBO charges for [11-THF]+: C-NHC= +0.090, Zn= +1.155, Zn-H= -0.544, O= -0.661. 

C-H Borylation:

Regarding the feasibility of the proposed C-H borylation mechanism outlined in Figure 7, the σ bond metathesis step was explored at the M06-2x/cc-pVTZ level and at the computationally less demanding M06-2x/lan12dz/6-311G/PCM(THF) level. At both these levels the calculated ΔG values were all within 2 kcal mol⁻¹, thus further calculations were performed at the M06-2x/lan12dz/6-311G/PCM(THF) level, and only values at this level are reported from here on unless otherwise stated. Starting from the zinc-alkynyl species [12'-THF]+ (12' designates it contains a phenylacetylene unit not 4-ethynyltoluene), the transmetallation with HBPin to form [11-THF]+ is endergonic, but only by +4.1 kcal mol⁻¹ (Figure 12). The subsequent protoanalysis of [11-THF]+ is exergonic, with the formation of H₂ presumably playing a key role in the overall favorable energetics of the conversion due to the high bond energy of H₂. The overall conversion of phenylacetylene / HBPin to H₂ and 6e is exergonic by 6.1 kcal mol⁻¹. The transition state (TS1) for the key σ-bond metathesis step from the spectroscopically observed resting state (the zinc-alkynyl, [12'-THF]+) to the zinc-hydride [11-THF]+ was relatively low in energy, ΔG = +19.3 kcal mol⁻¹ (relative to [12'-THF]+ / HBPin). These results are consistent with the observed reactivity, with the presence of additional terminal alkyne required to react with the hydride [11-THF]+ whose formation is slightly endergonic from [12'-THF]+ and HBPin.

Figure 12: Calculations at the M06-2x/lan12dz/6-311G/PCM(THF) level for: top, the C-H borylation process.
This will lead to a greater degree of $^6$Zn-C$_6$ polarization for the alkyne derivative and thus a greater electrostatic contribution enhancing the Zn-C bond strength relative to the phenyl derivative as discussed previously for other series of organometallic complexes. Due to the difference in the respective valence orbital energies of Zn and B this effect will be greater for organozinc species than organoboron species. Consistent with this hypothesis is the observation that the magnitude of positive charge on zinc is greater in the alkyne derivative $[12'$-THF]$^+$ relative to the phenyl derivative $[8$-THF]$^+$ (+1.391 compared to +1.366) based on NBO calculations.

The energetic feasibility of internal alkyne hydrozincation was next explored. The highly exergonic nature of positive charge on zinc is greater in the alkynyl derivative and thus a greater electrostatic contribution enhancing the Zn-C bond strength relative to the phenyl derivative as discussed previously for other series of organometallic complexes. The binding of a second molecule of THF to [(IDIPP)Zn(THF)$_2$] and [(IDIPP)Zn(THF)$_2$] to form $[16$-THF]$^+$ (Figure 15, top) is exergonic by $7.6$ kcal mol$^{-1}$ (calculated at the M06-2x/cc-pVTZ level for comparison with the 7-DIPPP analogue see Figure 11). The exergonic nature of this step is in contrast to the energetically neutral nature of THF binding for the 7-DIPPP analogue $[11$-THF]$^+$. As observed for $[11$-THF]$^+$ binding of two THF molecules to zinc increases the energy and alters the character of the LUMO which in both $[11$-THF]$^+$ (Figure 11, bottom) and $[16$-THF]$^+$ (Figure 15, bottom) is higher in energy and has no significant character on zinc in contrast to the three coordinate mono-THF congener $[11$-THF]$^-$ and $[16$-THF]$^+$ (the LUMO for $[16$-THF]$^+$ is closely comparable to that calculated for $[11$-THF]$^+$; see figure 11).

**NHC Dependency:**

Finally, the dramatic disparity in catalytic performance observed between 7-DIPPP and IDIPP was investigated. One major difference between the two series of (NHC)Zn compounds is the relative accessibility of the respective three coordinate zinc complexes. The binding of a second molecule of THF to [(IDIPP)Zn(THF)$_2$] and [(IDIPP)Zn(THF)$_2$] to form $[16$-THF]$^+$ (Figure 15, top) is exergonic by $7.6$ kcal mol$^{-1}$ (calculated at the M06-2x/cc-pVTZ level for comparison with the 7-DIPPP analogue see Figure 11). The exergonic nature of this step is in contrast to the energetically neutral nature of THF binding for the 7-DIPPP analogue $[11$-THF]$^+$. As observed for $[11$-THF]$^+$ binding of two THF molecules to zinc increases the energy and alters the character of the LUMO which in both $[11$-THF]$^+$ (Figure 11, bottom) and $[16$-THF]$^+$ (Figure 15, bottom) is higher in energy and has no significant character on zinc in contrast to the three coordinate mono-THF congener $[11$-THF]$^-$ and $[16$-THF]$^+$ (the LUMO for $[16$-THF]$^+$ is closely comparable to that calculated for $[11$-THF]$^+$; see figure 11).

**Alkyne Hydrozincation:**

The energetic feasibility of internal alkyne hydrozincation was next explored. The fact that hydroboronation is not observed in THF or in benzene at lower temperatures ($\leq 60^\circ$C) suggested a significant barrier (as the Zn-C/H-B metathesis steps explored in this work have all been found to be facile). Hydrozincation calculations utilized the internal alkene 13, where the Zn-alkenyl products, 14, are observed spectroscopically. Calculations indicated that while the hydrozincation of alkene 13 using $[11$-THF]$^+$ was significantly exergonic the transition states (TS$_2$ and TS$_3$, Figure 14) were found to be relatively high in energy (compared to TS$_1$) in both cases. These barriers are consistent with the requirement for heating for internal alkyne hydrozincation/hydroboration to occur. The highly exergonic nature of alkene hydrozincation indicates it will proceed irreversibly, consistent with it being the thermodynamic pathway. Inspection of TS$_2$ (TS$_3$ is extremely similar in character) reveals a considerable deviation from linearity of the alkene moiety (C-C-C angles are 143° and 153°), elongation of the C=C bond (to 1.25 Å) and the Zn-H bond (from 1.65 Å in $[11$-THF]$^+$ to 1.74 Å in TS$_2$) and contraction of the C-H distance (to 1.79 Å). Thus these calculations support the feasibility of the mechanism proposed in figure 7.

**Figure 13:** The exergonic metathesis of $[8$-THF]$^+$ and HBPin. Inset, key select frontier orbitals of $[12'$-THF]$^+$ and $[8$-THF]$^+$ (iso surface value = 0.04). Bottom, the isodesmic reaction between Zn-C/B/C species.

**Figure 14:** The calculated transition state and product energies (in kcal mol$^{-1}$) for the hydrozincation of 13 with $[11$-THF]$^+$. The energetic feasibility of internal alkyne hydrozincation was next explored. The fact that hydroboronation is not observed in THF or in benzene at lower temperatures ($\leq 60^\circ$C) suggested a significant barrier (as the Zn-C/H-B metathesis steps explored in this work have all been found to be facile). Hydrozincation calculations utilized the internal alkene 13, where the Zn-alkenyl products, 14, are observed spectroscopically. Calculations indicated that while the hydrozincation of alkene 13 using $[11$-THF]$^+$ was significantly exergonic the transition states (TS$_2$ and TS$_3$, Figure 14) were found to be relatively high in energy (compared to TS$_1$) in both cases. These barriers are consistent with the requirement for heating for internal alkyne hydrozincation/hydroboration to occur. The highly exergonic nature of alkene hydrozincation indicates it will proceed irreversibly, consistent with it being the thermodynamic pathway. Inspection of TS$_2$ (TS$_3$ is extremely similar in character) reveals a considerable deviation from linearity of the alkene moiety (C-C-C angles are 143° and 153°), elongation of the C=C bond (to 1.25 Å) and the Zn-H bond (from 1.65 Å in $[11$-THF]$^+$ to 1.74 Å in TS$_2$) and contraction of the C-H distance (to 1.79 Å). Thus these calculations support the feasibility of the mechanism proposed in figure 7.

**Figure 15:** Top, exergonic binding of a second molecule of THF at 298 K (in kcal mol$^{-1}$) to $[16$-THF]$^+$. Inset left, LUMO of [(IDIPP)Zn(THF)$_2$] and [(IDIPP)Zn(THF)$_2$] at isosurface value = 0.04 (most hydrogens omitted for clarity), inset right solid state structure of $[9$-THF$_2$], select bond distances (Å) = Zn-C$_{Ph}$ = 1.982(3), Zn-C$_{NHC}$ = 2.044(3), Zn-O = 2.134(2) and 2.044(2).

To confirm that (IDIPP)Zn(NTf$_2$)$_2$ (Y = H, Ph or alkynyl) complexes can indeed coordinate two molecules of THF we attempted to crystallize a number of (IDIPP)Zn(NTf$_2$)$_2$ complexes from THF. Crystals of 9 formed from THF/hexane, and analysis of these by X-ray diffraction revealed that 9 did indeed crystallize with two molecules of THF bound to the zinc center with the NTf$_2$ anion well separated (the cationic portion of this salt is termed $[9$-THF$_2]^+$). The disparity between the observed solid state structures of 7-DIPPP (3 coordinate with only one THF molecule binding) and IDIPP (4 coordinate at Zn with two
THF molecules binding) analogues deposited from THF is consistent with the DFT calculations on the relative energetics of a second molecule of THF binding to zinc. The enhanced steric impact of the 7-DIPP ligand relative to IDIPP is demonstrated by the lower \( \%V_{\text{bun}} \) of IDIPP in [9-THF] \( \text{Zn} \) (35.6\%) relative to 7-DIPPNZn analogues (7-DIPP analogue 8 which is also four coordinate due to NTF \( _2 \) binding bidentate has a \( \%V_{\text{bun}} = 47.2\%). Therefore, we attribute the observed disparity in catalytic reactivity between the 7-DIPP and IDIPP congeners to: (i) the preference for different coordination numbers at zinc when ligated by IDIPP (coordination number 4) Vs 7-DIPP (coordination number of 3); (ii) the greater thermal stability of the 7-DIPP species, such as 8, 11 and 12, relative to their IDIPP congeners (the IDIPP congeners are observed to more rapidly form insoluble species in the presence of HBPin, presumably ZnH \( _2 \) and/or metallic zinc, during catalytic reactions) enabling superior catalyst longevity.

Conclusions
The viability of Zn-C / H-BPin σ-bond metathesis for sp, sp\(^3\) and sp\(^3\) C centers has been demonstrated. This step allows for zinc hydride complexes to be utilized in catalytic borylation reactions. This has been exemplified herein by developing zinc hydrides that enable both the C-H borylation of terminal alkynes and the hydroboration of internal alkynes using the commercial borane HBPin. The latter reaction can proceed via an uncatalyzed alkynie hydrazinocilation. The two borylation processes can be combined to generate a one-pot zinc catalyzed transformation of terminal alkynes that selectively produces the desirable 1,1-diborylated alkene products. This process represents an earth abundant base metal alternative to the established catalytic routes to 1,1-diborylated alkynes (these all currently rely on noble or Co transition metal catalysis both metals that have extremely low permitted daily exposure values in contrast to zinc). Mechanistic studies and DFT calculations support the intermediacy of organozinc species, zinc-hydrides and Zn-C/H-B σ-bond metathesis in both these catalytic processes. Finally, extremely bulky N-heterocyclic carbenes are vital for these borylation reactions, a phenomenon attributable to the larger NHC providing enhanced thermal stability and access to low (three) coordinate (NHC)Zn cations. Highly electrophilic yet Bronsted basic low coordinate, (NHC)Zn-H catalysts offer significant potential for developing new catalytic transformations a number of which are currently under development in our laboratory.

ASSOCIATED CONTENT
Supporting Information. The Supporting Information listed below is available free of charge on the ACS Publications website at DOI: Synthetical methods and characterization details (.PDF), Crystallographic data (.cif).

AUTHOR INFORMATION

Corresponding Author
*E-mail for M.J.I.: michael.ingleson@ed.ac.uk

ORCID
Michael J. Ingleson: 0000-0001-9975-8302
Marina Uzelac: 0000-0002-5060-7017

Author Contributions
All authors have given approval to the final version of the manuscript. §RJP and MU contributed equally to the experimental sections of this work while JC performed the DFT calculations.

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(32) The mono-THF solvate is used throughout for DFT calculations, including for the hydrozincation reactions. For reactions performed in benzene a THF solvate is clearly not feasible, however, the THF solvate still is used to keep the model constant throughout and to simplify the calculations (if THF is replaced by NTF2 mono and bidentate NTF2 binding modes need to be considered for all steps). It should be noted that all the energies obtained using this model correlate well with the observed solution reactivity.


