

PERSPECTIVE

View Article Online



Cite this: New J. Chem., 2021, **45**, 14855

Received 9th June 2020, Accepted 7th July 2020

DOI: 10.1039/d0nj02908d

rsc.li/njc

Haloboration: scope, mechanism and utility

Sven Kirschner, (1) † Kang Yuan† and Michael J. Ingleson (1) *

Haloboration, the addition of B-X (X = Cl, Br, I) across an unsaturated moiety e.g., C = Y or C = Y (Y = C, N, etc.), is dramatically less utilised than the ubiquitous hydroboration reaction. However, haloboration of alkynes in particular is a useful tool to access ambiphilic 1,2-disubstituted alkenes. The stereochemical outcome of the reaction is easily controlled and the resulting products have proven to be valuable building blocks in organic synthesis and materials chemistry. This review aims at providing the reader with a brief summary of the historic development and of the current mechanistic understanding of this transformation. Recent developments are discussed and select examples demonstrating the use of haloboration products are given with a focus on the major areas, specifically, natural product synthesis and the development of boron-doped polycyclic aromatic hydrocarbons (B-PAHs).

1. Introduction

Since the end of the last millennium, the principle of sustainability and atom economy increasingly has impacted the way scientific research is done. A chemical transformation that is 100% atom efficient is the addition reaction, in which a reagent is added across a multiple bond (e.g., $C \equiv C$, $C \equiv C$, or $C \equiv E$, E: O, NR). One very important addition reaction is the Nobel Prize winning hydroboration reaction.² Initially, it was almost exclusively used to access alcohols from alkenes by oxidative B-C bond cleavage. However, organic transformations like the Matteson homologation,³ Petasis variant of the Mannich reaction,⁴ Chan-Lam coupling,⁵ and, of course, the Suzuki-Miyaura reaction⁶ all utilise substrates that can be accessed by hydroboration reactions, rendering it a powerful tool in the synthetic chemist's toolbox. Unsurprisingly, researchers still strive to develop new hydroboration methods, e.g., by expanding the scope or reducing the environmental impact. In contrast, haloboration - although discovered at roughly the same time has remained a niche technique that has gained only a small fraction of the attention even though it adds an additional highly valuable group (a halide) in the same step. The concomitant installation of a boron unit and a halide generates functionality rich molecules, containing a nucleophilic C-B and an electrophilic C-X unit. Thus, it is surprising that this reaction is so under-utilised, despite being potentially useful to many.

In this review, we give a brief discourse of the historic development of the haloboration reaction, from the curiosity driven fundamental research mainly by the group of Lappert, to

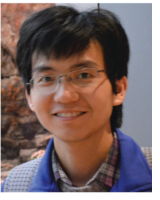
EaStCHEM School of Chemistry, University of Edinburgh, Edinburgh, EH9 3FJ, UK. E-mail: michael.ingleson@ed.ac.uk

† These authors contributed equally to this work.



Sven Kirschner

Sven Kirschner studied chemistry at the Goethe-University Frankfurt/ Main (Germany), where he received his PhD in 2019 for his contributions to the field of boron-doped polycyclic aromatic hydrocarbons. He joined the group of Michael Ingleson in 2019 as a postdoctoral research associate and is currently working on the development of main-group element mediated C-C coupling reactions.



Kang Yuan

Kang Yuan received his BSc degree in 2013 from Sun Yat-sen University under the guidance of Prof. Ming-Liang Tong. Then, he moved to Canada to explore main group chemistry and obtained his PhD degree under the supervision of Prof. Suning Wang at Queen's University. In 2018, he joined the Ingleson group as a postdoctoral research associate. His current research focuses on developing new methods for constructing novel boron containing molecules.

the usage of Pd catalysed cross-coupling reactions to demonstrate the full potential of haloboration from the group of Suzuki who broadened the scope and deepened the understanding of this reaction.8 We also provide an in-depth discussion of the underlying mechanism and select applications of the products from the haloboration reaction in the field of synthesis. Recently, haloboration has started to gain wider interest through its use in natural-product synthesis to introduce C=C double bonds stereoselectively, and in the synthesis of boron-doped polycyclic aromatic hydrocarbons (B-PAHs), thus these are the main applications focused on herein. B-PAHs are a relatively new class of organic materials with interesting optoelectronic properties and haloboration is a fast and convenient way to incorporate a borane unit and a halogen functionality into a PAH at the same time. Since the publication of the last reviews on haloboration, which were in the 1980s to the best of our knowledge, 8d the scope and utility of haloboration in synthetic chemistry has increased significantly. We hope that this review, focused on the use of boron electrophiles to haloborate C = Y and C = Y nucleophiles (Y = C, N) or O based substituents) facilitates the wider application of this useful, yet often overlooked, reaction.

2. Haloboration of simple alkynes using Y_2B-X (Y = X or R)

2.1 The early work

Chloroboration of alkynes was first explored by H. R. Arnold in 1946.9 With mercury(1) chloride on activated carbon as the catalyst, chloroboration of acetylene with boron trichloride (BCl₃) was achieved to afford 2-chlorovinyldichloroborane at 150-300 °C (Scheme 1a). In this patent, the stereoselectivity of chloroboration was not determined. Subsequently, Jensen et al. repeated the synthesis under similar conditions and measured the dipole moment of the obtained product.¹⁰ The experimentally



Michael J. Ingleson

Mike Ingleson received his PhD. in organometallic chemistry in 2004 working with Prof. Andrew S. Weller. He then pursued postdoctoral studies with Prof. Kenneth G. Caulton at Indiana University and with Professor Matthew J. Rosseinsky at the University of Liverpool. started his independent career at the University of Manchester in 2008 with a Royal Society University Research Fellowship. At Manchester, he was promoted

to Reader in 2012 and Full Professor in 2018. He joined the University of Edinburgh in 2019. His research focuses on main group chemistry, and he has a particular fondness for all things boron.

a)
$$H = H$$

$$= H$$

Scheme 1 Chloroboration of acetylene/phenylacetylene under various conditions

determined dipole moment (1.06(0.05) D) was found to be very close to the predicted value of trans-product 1 (1.05 D). For comparison, the dipole moment of the cis-isomer was predicted to be 3.23 D. Thus, trans-product 1 was believed to be formed. In addition, the trans-chloroboration product 1 was found to be significantly more stable than the cis-isomer by 145 kJ mol⁻¹ in electronic energy based on ab initio calculations.

Subsequent to Arnold's work, Gipstein et al. found that when EtBCl2 was used, in spite of the reduced Lewis acidity compared to BCl₃, the chloroboration of acetylene in the presence of activated carbon could be realised at 70 °C, affording product 2 (configuration not determined) in 90% vield (Scheme 1b).11 Later, Lappert and co-workers studied the chloroboration reaction with a variety of alkynes and boranes.12 For instance, phenylacetylene was reported to undergo chloroboration with one equivalent of BCl₃ readily even at -78 °C to afford the syn-addition product 3. The obtained product 3 was shown to react with another equivalent of phenylacetylene in a syn-manner to give compound 4 (Scheme 1c). In the initial report by Lappert and co-workers, the configurations of 3 and 4 were assigned with incomplete evidence. Subsequent studies confirmed that chloroboration of terminal alkynes with BCl₃ proceeds in a syn-manner. ¹³ Although the reaction between BCl₃ and terminal alkynes such as phenylacetylene occurs promptly, no reactivity was observed when internal alkynes such as diphenylacetylene and BCl₃ were mixed at 15 °C. In accordance with the reactivity of BCl₃, PhBCl₂ readily reacted with two equivalents of phenylacetylene and compound 5 was obtained (Scheme 2a).12 In contrast, when 1-hexyne was treated with half an equivalent of PhBCl2, both chloroboration and carboboration occurred to give the product 6 (Scheme 2b). In a controlled reaction of 1-hexyne with Ph2BCl, carboboration occurred exclusively yielding compound 7 (Scheme 2c).

Lappert et al. also investigated the haloboration of acetylene using BBr₃. The increased Lewis acidity of BBr₃ (relative to BCl₃) drastically facilitated the transformation which proceeded at room temperature and the addition product 8 was obtained (Scheme 3a). Alcoholysis of compound 8 with n-butanol afforded the known compound E-9Bu with specified configuration. The configuration of E-9^{Bu} led the authors to assign a trans-configuration to 8. However, a recent study suggested the

a) PhBCl₂
$$0 \,^{\circ}\text{C}$$
 Ph Cl

b) $n\text{-Bu} = \frac{PhBCl_2}{130\text{-}140 \,^{\circ}\text{C}}$ $n\text{-Bu} = \frac{PhBCl_2}{130\text{-}140 \,^{\circ}\text{C}}$ $n\text{-Bu} = \frac{Ph_2BCl}{130\text{-}140 \,^{\circ}\text{C}}$ $n\text{-Bu} = \frac{Ph_2BCl}{130\text{-}140 \,^{\circ}\text{C}}$ $n\text{-Bu} = \frac{Ph_2BCl}{Ph} =$

Chloroboration and carboboration of terminal alkynes

Scheme 3 Bromoboration reactions to gain insight into the selectivity.

bromoboration of acetylene typically gives a mixture of E/Zisomers at around 0 °C (for further details and mechanistic discussion, see Sections 2.2 and 2.3). Lappert et al. also found that the addition of pyridine (Py) to 8 resulted in the elimination of Py-BBr₃ and regeneration of acetylene, which indicated that the bromoboration of acetylene might be reversible (Scheme 3a). The bromoboration of acetylene with BBr₃ followed by esterification with alcohols provides a convenient route to halo-alkenylboronates that serve as versatile building blocks (vide infra).

Although BCl₃ does not react with internal alkynes, Lappert and co-workers found that bromoboration of diphenylacetylene occurs readily in neat BBr3 at room temperature within one hour. In this case, syn-addition product 10 was formed as confirmed by subsequent protodeboronation reactions with acetic acid (AcOH) to give 11 (Scheme 3b), with protodeboronation known to proceed with retention. 12,14a Blackborow performed detailed studies on the bromoboration of 1-hexyne under various conditions (Scheme 3c).¹⁴ Generally, the bromoboration of the terminal alkyne proceeds in a Markovnikov fashion. The stereoselectivity, however, was found to be highly dependent on the reaction conditions. For example, when the reaction was performed at -80 °C in petroleum or dichloromethane, syn-addition product 12 was found to be dominant (98%) as determined by analysis post protodeboronation with AcOD. In contrast, when the reaction was performed at -40 °C, the stereoselectivity decreased with the major product being the anti-bromoboration product 13 (64%) while only 30% synaddition product 12 was observed under these conditions. In addition to E/Z isomers, multiple haloborations to form the respective divinylbromoborane and trivinylborane also were observed. However, as the formation of these multiple borylation products is relatively slow under the reaction conditions and the product distribution was not well defined, these details are not discussed further here.

Subsequent to Lappert's work with BBr3, Eisch and co-workers found that the less Lewis acidic borane PhBBr2 reacts with diphenylacetylene in a reversible manner (Scheme 4a).15 Upon mixing PhBBr₂ and diphenylacetylene, cis-bromoboration product 14 was formed rapidly, which was confirmed by protodeboronation with acetic acid. However, prolonged storage of compound 14 in hydrocarbon solvents led to the irreversible formation of carboboration product 15. This suggests that the bromoboration of diphenylacetylene with PhBBr2 is a kinetically favoured but reversible process while the carboboration is a slower and irreversible competing process. Wrackmeyer studied the reaction of 3-hexyne with BBr₃ at -78 °C (Scheme 4b). ¹⁶ When the sample was kept at room temperature for one hour, the syn-addition product 16 was found to be the major product in the reaction mixture (16:17 = 15:1). However, after several days, the *E*-isomer 17 became dominant with the ratio of 16:17 switching to 1:7.

Eisch also studied the reactivity of MeBI₂ with diphenylacetylene. In this case, iodoboration occurred rapidly and compound 18 was obtained (Scheme 5a).17 Siebert found that iodoboration of 3-hexyne with BI3 furnished the syn-addition product 19 rapidly, which, in line with Wrackmeyer's observations, underwent slow isomerisation to form the anti-addition product 20 at room temperature (Scheme 5b). 18,19

As discussed above, bromoboration of terminal alkynes with BBr3 does not occur exclusively in a 1:1 stoichiometry due to further reactions of the vinylBBr2 species with additional alkyne. One solution to this problem is to use 9-halo-9borabicyclo[3.3.1]nonane (9-X-BBN; X = Br, I). Suzuki and co-workers found that 9-Br-BBN could react with one equivalent of 1-octyne to afford 21 in high regio- and stereoselectivity (Scheme 5c).20 They also found that 9-Br-BBN is inert to internal alkynes likely due to its lower Lewis acidity. These early studies clearly demonstrated the viability of alkyne haloboration, albeit complicated in many cases by formation of

Scheme 4 Isomerisation in reactions of internal alkynes with haloboranes

Scheme 5 Haloboration reactions on various alkynes with different haloboranes

different haloboration isomers. The origin of Z- and E-configured haloboration products was at the time unclear and required subsequent DFT calculations to provide mechanistic insight.

2.2 Mechanistic studies

The haloboration of alkynes was investigated computationally initially by Uchiyama and co-workers. 21 Ab initio calculations with second-order Møller-Plesset perturbation theory (MP2) were performed on the haloboration of acetylene and propyne using a dichloromethane continuum solvent model (Scheme 6). The two alkynes first form loose van der Waals complexes A with BX_3 (X = Cl, Br, I), which then may transform into π -bonded complexes **B** if X = Br or I but not Cl. Intermediates A (X = Cl) or B (X = Br, I) then convert into the syn-addition products cis-P via four-centred transition state TS. Consistent with the observed reactivity of BX_3 with alkynes ($BBr_3 > BCl_3$), the reaction energy barriers (energy of TS) decrease in order of $BCl_3 > BBr_3 > BI_3$ for the syn-addition pathway.

An energy barrier of 33.8 kcal mol⁻¹ was determined computationally for the cis-chloroboration of acetylene, which agrees with the experimental observations that chloroboration of acetylene required a high reaction temperature and a catalyst. In addition, the haloboration of propyne was found to have a lower energy barrier than acetylene, which is ascribed to hyperconjugation stabilising the developing positive charge

trans-P

Scheme 6 Reaction pathway for cis-haloboration of alkynes

at carbon in TS (when R = Me). Furthermore, the anti-Markovnikov pathway for the chloroboration of propyne was also explored computationally. The transition state of anti-Markovnikov pathway **TS**-a (X = Cl, ΔG^{\ddagger} = 35.6 kcal mol⁻¹) was found to be much higher than the Markovnikov pathway $(X = Cl, \Delta G^{\ddagger} = 15.5 \text{ kcal mol}^{-1})$. The results fit well with the high regioselectivity of terminal alkyne haloboration reactions. For all transformations, the trans-haloboration products trans-P also were computed. For acetylene, they were found to be thermodynamically more stable than the cis-products, which is in agreement with the observation that the trans-chloroboration product was formed exclusively at high temperature in the aforementioned reports (cf. Section 2.1). In contrast, the syn-addition products of propyne haloboration are very close in energy to their anti-addition isomers. Again, this result is consistent with the observation that bromoboration of 1-hexyne at temperatures above −40 °C gave a mixture of both isomers. After the exploration of several different potential reaction pathways, Uchiyama and co-workers proposed that the stereoconversion proceeded via a haloboration/retro-haloboration mechanism of cis-P with BX₃ (Scheme 7).

The chloroboration of the internal alkyne 2-butyne, and propene was also investigated. For both substrates, the chloroboration was found to be endergonic at 293 K ($\Delta G = 2.8 \text{ kcal mol}^{-1}$ for 2-butyne; $\Delta G = 9.4 \text{ kcal mol}^{-1}$ for propene) consistent with that lack of reactivity between BCl₃ and internal alkynes and olefins.

2.3 Recent studies into the bromoboration of simple alkynes with BBr₃

In Lappert's initial studies, the stereoselectivity for the bromoboration of acetylene was determined by analysis of the postesterification (conversion of C-BBr₂ to C-B(OR)₂) products. Recently, Mazal and co-workers carried out similar haloboration experiments at 0 $^{\circ}$ C and monitored the formation of E/Zvinylbromoborane 8 by NMR spectroscopy (Scheme 8a). With BBr₃ distilled from Mg turnings, a mixture of E/Z isomers was obtained (E:Z = 15:85). Interestingly, addition of small amounts of water, NEt3 or [n-Bu4N]Br to the bromoboration reaction facilitated the formation of anti-addition product E-8. It is even more notable that the authors exclusively found the

Scheme 7 Possible mechanism for stereoconversion mediated by BX₃.

Scheme 8 (a) Reaction of acetylene with BBr₃. (b) Brønsted-acid-free workup conditions allow for the selective isolation of Z- $\mathbf{9}^{Pin}$. (c) Possible reaction pathways and energies (kcal mol⁻¹) of acetylene bromoboration.

trans-vinylboronate E-9^{Pin} post esterification workup. They proposed radical or polar addition mechanisms involving adventitious HBr to rationalise the observed reactivity.²² By modifying the workup procedures to avoid the formation of HBr, a mixture of E/Z-bromovinylboronate E/Z-9^{Et} could be obtained, which was subjected to transesterification with pinacol, yielding E/Z- 9^{Pin} . Selective decomposition of the E-isomer (AcOK, MeOH) enabled the isolation of Z-bromovinylboronate Z- 9^{Pin} in useful yields (Scheme 8b).

In their proposed radical mechanism, a cis-bromoboration of acetylene occurs first to give Z-8 with an energy barrier of 24.1 kcal mol⁻¹ (MP2/6-31+(d)/PCM(DCM)). Attack of a bromine radical (proposed to be generated from adventitious HBr) on Z-8 proceeds with almost no barrier (1.8 kcal mol⁻¹). Bond rotation followed by bromine radical elimination was then assessed to afford the trans-bromoboration product E-8. An anti-addition of BBr3 and Br to acetylene was also suggested for the formation of E-8. Acetylene was calculated to first form a π -bonded complex with BBr₃, which was then attacked by Br⁻ to form a borate intermediate. Abstraction of Br with BBr3 from the intermediate gave E-8. The energy barrier for this process was determined to be 12.6 kcal mol⁻¹.

2.4 1,1-Bromoboration of internal alkynes

Recently, the group of Ingleson have revisited the reaction of internal alkynes with BBr₃. Consistent with Lappert's observations, they found that in non-polar solvents such as heptane,

Scheme 9 1,1-Bromoboration of diphenylacetylene yields mixtures of the syn-1,2-addition product 10 and the 1,1-addition product 22.

the reaction of diphenylacetylene with BBr₃ at room temperature afforded the cis-1,2-addition product 10 within a few minutes.²³ However, upon heating to 60 °C or prolonged standing at room temperature, the 1.1-addition product 22 was observed (Scheme 9a). When a polar solvent such as dichloromethane was used in the haloboration reactions, both 10 and 22 were formed within a few minutes with 22 being the major product (Scheme 9b). No trans-1,2-bromoboration product was observed by in situ NMR spectroscopy at any stage of the reaction. These observations suggest that 1,1-bromoboration proceeds through a polar transition state, likely a vinyl cation type intermediate(s) which is more stabilised by the more polar solvent dichloromethane. Upon reaction with pinacol and NEt3, compound 22 was converted into the corresponding pinacol ester 23 while 10 was converted back into diphenylacetylene. The 1,1-bromoboration could be extended to diarylalkynes and arylalkylalkynes, providing a convenient way to 1-bromo-2,2-diaryl substituted vinylboronate esters.

This is a further example that the reactions of alkynes and BX₃ can give conditions dependent outcomes. Nevertheless, the use of the appropriate conditions and work-up can lead to isolation of a single haloboration product in useful yield. This diversity in outcome (1,1-, syn-1,2- or anti-1,2-haloboration products being accessible) arguably increases the power of the haloboration transformation.

Other alkyne haloboration reactions

This section covers recent synthetic reports that use other (i.e., not Y_2B-X , Y = X or R) boron electrophiles or provides alkyne haloboration products that are distinct to those reported in the work discussed in Section 2.

3.1 Borocation mediated chloroboration of alkynes

As discussed in Section 2.2, Uchiyama and co-workers found the chloroboration of internal alkynes with BCl3 to be thermodynamically uphill. However, more electrophilic boranes such as BBr₃ and BI₃ react with internal alkynes readily. Therefore, Ingleson et al. envisioned the use of more electrophilic chloroborane species, such as borocations, to enable internal alkyne chloroboration.24 They prepared the boronium (tetracoordinate at B mono-cation) salt $[Cl_2B(2-DMAP)][AlCl_4]$ (2-DMAP = 2-dimethylaminopyridine) by sequential addition of 2-DMAP and AlCl₃ to BCl₃ (Scheme 10a). Due to the strain within the four-membered boracycle, this complex showed a low energy

Scheme 10 Reaction route and calculated energies (kcal mol⁻¹) for chloroboration of phenylacetylene with boronium/borenium complexes.

barrier to ring opening and reacted as a masked borenium (= tricoordinate B monocation) ion. [Cl₂B(2-DMAP)][AlCl₄] reacted with one equivalent of a terminal alkyne at room temperature with high regio- and stereoselectivity. In contrast to the haloboration of alkynes with neutral BX3 which proceeded via a four-membered transition state with concerted formation of B-C and Cl-B bonds, chloroboration of phenylacetylene with [Cl₂B(2-DMAP)][AlCl₄] was calculated to proceed via a vinyl-cation intermediate, 24a. Intramolecular chloride transfer in 24a occurs with an energy barrier of 24.9 kcal mol⁻¹ to give 24b (at the M06-2X/6-311G(d,p)/PCM(DCM) level of theory), which then rearranges to afford the chelated compound 24. Although the reaction between the boronium complex and terminal alkynes was facile, no reactivity was observed between [Cl₂B(2-DMAP)][AlCl₄] and internal alkynes. This is presumably due to the requirement to open the four-membered boracycle within [Cl₂B(2-DMAP)]⁺ prior to the haloboration reaction, coupled with the significant N \rightarrow B π donation (as shown in 24b) in the borenium isomer of [Cl₂B(2-DMAP)]⁺ (reducing the Lewis acidity at B) leading to unfavourable energetics.

Ingleson and co-workers also reported that when the borenium salt $[Cl_2B(Lut)][AlCl_4]$ (Lut = 2,6-lutidine) was used, syn-addition of the B-Cl bond across both terminal and internal alkynes occurred (Scheme 10b), presumably due to the enhanced electrophilicity at B in this borocation relative to the 2-DMAP analogue. Good chloroboration stereoselectivity was achieved for dialkyl, diaryl and arylalkylalkynes. Remarkably, for arylalkylalkynes, regioselective chloroboration could also be readily realised. All the products could be converted into the corresponding pinacol boronate esters by subsequent esterification with no loss in stereo-/regioisomeric purity.

3.2 1,2-trans-Chloroboration of alkynes

Chloroboration of alkynes typically proceeds in a syn-manner. The group of Ingleson found that treatment of 2-dimethylaminotolan

directed trans-chloroboration Scheme 11 Amino group

with BCl₃ gave the unusual anti-addition product 25 (Scheme 11a).²⁵ Subsequently, Pei and co-workers reported that 2-aminotolan also reacted with PhBCl₂ to give the trans-chloroboration product 26 (Scheme 11b).²⁶ The reaction was proposed to be initiated through the activation of the triple bond by the boron moiety, which was then followed by a nucleophilic attack of the alkyne with chloride. This method serves as a convenient way to prepare B,N-fused polycyclic aromatic hydrocarbons. The disparity between the formation of five membered 25 and six membered 26 is notable. This can be attributed to the all sp²-containing 26 having a strong preference for forming six membered boracycles via electrophilic borylation, while the incorporation of a single tetrahedral centre (as in 25) leads to five membered boracycles from electrophilic borylation being the favoured products.27

3.3 1,3-Chloroboration of alkynes

In 2018, Melen et al. reported a unique 1,3-chloroboration of propargyl esters.²⁸ By treating propargyl benzoate with 1 equivalent of PhBCl2, an intermediate dioxolanonium ion was proposed to be formed via boron promoted cyclisation. The intermediate was assumed to subsequently undergo ringopening and chloride migration to furnish the corresponding product 27 in high yield at room temperature (Scheme 12a). Interestingly, for a related propargyl ester with two methyl

Scheme 12 1,3-Chloroboration/1,1-carboboration of propargyl esters.

groups at the propargylic position, 1,1-carboboration occurred when PhBCl₂ was added to give 27a (Scheme 12b), presumably

due to dimethyl substituents leading to a higher barrier for the 1,4-chloride transfer.

Chloroboration of diynes 3.4

NJC

Ingleson and co-workers found that 1,6-heptadiyne reacts rapidly with BCl₃ at room temperature to afford a chlorinated cyclohexene featuring an exocyclic vinylBCl₂ moiety.²⁹ The corresponding pinacol protected compound 28 could be isolated in high yield post esterification with pinacol/NEt₃ (Scheme 13a). In this case terminal alkyne 1,2-haloboration must have a higher barrier than intramolecular reaction of the alkyne-BCl₃ adduct with the second alkyne. Furthermore, the reaction outcome was highly solvent dependent, with dichloromethane and dichloroethane affording 28 in good yield, whereas the use of chloroarenes resulted in very low yields of 28, suggesting the vinyl chloride is made via a carbocationic intermediate, which can engage in side-reactions with aromatic solvents.

Although BCl₃ shows no reactivity towards internal alkynes, facile transformations between BCl₃ and 1,2-dialkynyl benzenes have been reported by the groups of Erker, Yamaguchi and subsequently Ingleson. 30,31 Presumably these proceed from the adduct between the internal alkyne and BCl₃. In these reports, dibenzopentalenes or analogues were the major products in most cases (Scheme 13b). However, Ingleson et al. found that by introduction of electron donating groups such as p-methoxyphenyl (p-An) to 1,2-dialkynyl benzene substrates, the major product is the benzofulvene 29 via a formal 1,4-chloroboration reaction. In this case the dibenzopentalene 30 is only observed as the minor product (Scheme 13c).³¹

a)
$$= 2) \operatorname{pinacol/NEt_3}$$

$$= 2) \operatorname{pinacol/NEt_3}$$

$$= 2) \operatorname{pinacol/NEt_3}$$

$$= 20 °C$$

$$= 28$$
b)
$$Ph$$

$$Ph$$

$$BCl_3$$

$$rt$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$Ph$$

$$BCl_3$$

$$Ph$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$Ph$$

$$BCl_3$$

$$Ph$$

$$Ph$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$Ph$$

$$Ph$$

$$Ph$$

$$BCl_2$$

$$Ph$$

$$P$$

Scheme 13 Borylative cyclisation/chloroboration of diynes

Scheme 14 Change in product ratio on altering the boron electrophile or by the addition of an exogenous chloride source

Interestingly, the formation of 1,4-chloroboration products becomes favoured in other cases by the addition of an exogenous chloride donor such as [BCl₄] (Scheme 14). For example, 1,2-bis(p-tolylethynyl)benzene reacts with BCl₃ affording the corresponding benzofulvene 31 and dibenzopentalene 32 in a 1:2 ratio. However, in the presence of three extra equivalents of a [BCl₄] salt, the ratio of 31 and 32 switched to 1.5:1, indicating the role of [BCl₄] in promoting the formal chloroboration reaction potentially by transferring chloride to the vinyl cation zwitterionic intermediate. Furthermore, precluding the presence of [BCl₄] or [RBCl₃] species in the reaction mixture by using boronium salt [Cl2B(2-DMAP)][AlCl4] significantly reduced the amount of the 1,4-chloroboration product observed. These studies therefore reveal that in the haloboration of divnes the halide source must be considered carefully to ensure a successful reaction outcome.

Directed trans-bromoboration of alkynes

In addition to the aforementioned trans-bromoboration of acetylene and terminal alkynes, there are a limited number of other examples of internal alkyne trans-bromoboration reactions. Yamato and co-workers reported a BBr₃ induced transformation of a o,o'-dimethoxy-substituted tolan derivative (Scheme 15).³² The highly Lewis acidic BBr₃ induced a twofold ether cleavage and intramolecular trans-bromoboration, yielding benzofurochromene derivative 33.

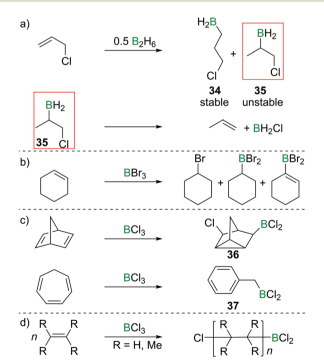
Pei and co-workers also reported a directed trans-bromoboration similar to their trans-chloroboration reaction discussed above (cf. Scheme 11).26 Due to the similarities the details are not discussed again herein.

Scheme 15 Directed trans-bromoboration of internal alkynes.

Sections 2 and 3 show the utility of alkyne haloboration, and this is by far the most developed reaction. The application of the haloboration reaction to other π systems is much less developed and the limited examples reported currently to the best of our knowledge are discussed in Section 4.

4. Haloboration of C=C double bonds

In their attempted hydroboration of vinyl chloride with B₂H₆ at -80 °C, DuPont et al. observed decomposition of the putative tris(2-chlorovinyl)borane upon warming to room temperature, whereas allyl chloride underwent hydroboration smoothly even at ambient conditions.³³ Those findings were confirmed by Brown and Köster, who investigated the selectivity of allyl chloride hydroboration. Hydroboration resulted in a formal 6:4 anti-Markovnikov (34)/Markovnikov (35) selectivity. However, the Markovnikov product underwent rapid elimination of the vicinal BH₂ and Cl groups to form H₂BCl and propene (Scheme 16a).^{34,35} These observations foreshadowed the subsequent reports that found formation of olefin 1,2-haloboration products to be an energetically unfavourable process. Haloboration experiments on cyclohexene by Lappert et al. supported olefin haloboration being energetically uphill: only a mixture of products could be identified, with no α-haloalkyl borane (the primary product from haloboration) observed, indicating that additional reactivity has to take place to lead to an overall exergonic process (Scheme 16b).36



Scheme 16 (a) Hydroboration of allyl chloride gives propylborane (34) and 1-chloro-2-propylborane (35); the latter decomposes under elimination of BH₂Cl to give propene. (b) Reaction of cyclohexene with BBr₃ yields a mixture of products. (c) and (d) Select examples of BCl3 initiated cationic rearrangements or polymerisation of alkenes

Scheme 17 Select examples of selectivity of alkyne haloboration over alkene haloboration and of haloboration reactions on C=C double bonds.

Rearrangements also can be used to trap initial olefin haloboration products, for example the reaction of BCl₃ with suitable olefins such as norbornadiene or cycloheptatriene, furnishes tricyclene 36 or BnBCl₂ (37, Scheme 16c). 36,37

These observation are consistent with the ability of boron trihalides to function as initiators in alkene polymerisation (Scheme 16d), instead of resulting in simple olefin haloboration.³⁸ These experimental findings were corroborated through the earlier discussed calculations by Uchiyama et al., who showed that haloboration of alkenes is thermodynamically unfavourable.²¹ This is further supported by the experimental finding that alkynyl substituted alkene 38 reacts selectively with BBr3 via the alkyne moiety to yield 39 (Scheme 17a).³⁹ Thus haloboration of olefins is an uncommon route to β-haloalkylboranes with only one inter- and one intramolecular synthesis reported to our knowledge. 40,41 Other methods are preferred to prepare this versatile structural motif.⁴²

While isolated olefins do not undergo haloboration, Lappert et al. showed that allenes reacted smoothly at -20 °C with BBr₃ to give the respective 1,2 adduct. Isolation of the bromoborane, however, was problematic as polymerisation of the remaining alkene function occurred at elevated temperatures during attempted distillation.³⁶ By esterification of the BBr₂ group through reaction with anisole, polymerisation was prevented and the adduct could be isolated (as 40), thus providing stereoselective access to allylboronic esters (Scheme 17b). 43 Iodoboration of terminal allenes with 9-I-BBN also can be a useful route to 2-iodoalkenes, if post haloboration the 9-BBN moiety is removed via acetolysis.44 Notably, if acetolysis is omitted, this reaction can furnish a sought after allylboronate without the need for allyl-metal species which are prone to 1,3 metallotropic shifts. 45 If silylated allenes such as 41 are employed, the outcome of the reaction changes. Instead of the 1,2 haloboration product, the formal 1,3 haloboration product 42 is obtained (Scheme 17c). Quantum chemical calculations suggest silyl migration followed by sterically driven 1,3 boryl shifts to be responsible for this special stereoselectivity. 46

Haloboration of E=C bonds (E=O, NR)

Lappert et al. investigated the reactivity of aldehydes towards BX₃ and suggested haloboration across C=O to be a first step

a)

R—CI +
$$\frac{1}{3}$$
BCI₃ + $\frac{1}{6}$ B2O₃ +

Scheme 18 Reaction of aldehydes with BCl₃ leading to complex mixtures, depending on the nature of the aldehyde. With iso(thio)cyanates, well defined products are obtained.

in a cascade of reactions that, depending on the nature of the aldehyde, eventually lead to borate esterification, formation of alkylhalides, haloalkylethers, or enolisation with concomitant polymerisation of the resulting vinylboric ester or alkenylether (Scheme 18a).⁴⁷ The reaction of acetone with BCl₃, however, only yielded an ill-defined mixture of products upon release of HCl, which precluded characterisation.⁴⁸ Remarkably, when perhalogenated ketones (Hal = F, Cl, Br) were employed, usually haloboration of the C=O bond occurred,49 but in some instances, simple halide exchange is favoured, giving a different boron halide and the respective perhaloketone.⁵⁰ In the case of isocyanates and isothiocyanates (43), multiple C=E double bonds exist. In these cases, haloboration proceeds stepwise via an R-N=C=E \rightarrow BX₃ (E = O, S) adduct 44, followed by 1,3 migration of X to C. The higher stability of a C=O bond compared to a C=N bond then leads to a B shift, giving aminoborane derivatives 45 (Scheme 18b).

Nitriles, which are isostructural and isoelectronic to alkynes can react with BX₃ in a fashion related to their C≡C analogues. However, due to the lone pair at the N atom, the resulting aminoboranes 46 exist in an equilibrium with their cyclic dimers (47, Scheme 19a). 51,52 In aminobenzonitriles, reactivity depends on the position of the two groups relative to each other.

a)
$$\begin{array}{c} N \\ R \end{array}$$
 $\begin{array}{c} BX_3 \\ R \end{array}$
 $\begin{array}{c} R \\ X \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_3 \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_3 \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_3 \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_3 \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_2 \end{array}$
 $\begin{array}{c} X_2 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$
 $\begin{array}{c} X_1 \\ X_2 \end{array}$
 $\begin{array}{c} X_1 \\ X_3 \end{array}$

Scheme 19 Select examples of E=C haloboration reactions.

For para- and meta-substituted aminobenzonitrile, simple adducts of the type $R-NH_2 \rightarrow BX_3$ form. In the case of ortho-aminobenzonitrile and PhBCl₂, haloboration of the C≡N triple bond occurs, furnishing aminoborane 48 which at elevated temperatures cyclises to give the heterocyclic product 49 (Scheme 19b).⁵³ In contrast to the meta- and para-substituted benzonitriles, ambidentate cyanamide reacts exclusively via haloboration of the C≡N triple bond giving dimeric iminoboranes, due to the strong and immediate mesomeric effect of the NH₂ group on the $C \equiv N$ triple bond and the thereby increased basicity of the cyano N-atom.⁵⁴ Isonitriles form stable adducts **50** with the higher haloboranes BX_3 (X = Cl, Br, I; for X = F, polymerisation occurs). If heated, these compounds undergo 1,1-haloboration and dimerisation to form 2,5-dihydrodiborapyrazines (51, Scheme 19c).⁵⁵

Finally, reports on the reaction of BX_3 with C=NR double bonds are, just like in the case of C=C double bonds, scarce, and limited to either highly electron deficient perfluorinated imines^{56,57} or chelating 1,4 diazabutadienes such as 52.⁵⁸ The latter is an early example of a simple route to 1,3,2-diazaborolidines (53, Scheme 20), which play a crucial role in accessing nucleophilic boranes.59

6. Application of alkyne haloboration products in organic synthesis

As described in Sections 2 and 3, haloboration of terminal alkynes is a versatile tool to access selectively 1,1- or 1,2difunctionalised alkenes. The stereoselectivity can be controlled and the resulting vinylboranes show ambiphilic reactivity: the halogenated C atom is a potential electrophile, whereas the boron-bonded C atom behaves as a nucleophile. In this section we focus on applications that take advantage of both these groups or use the electrophilic vinylBX₂ intermediate (particularly in reactions other than esterification). The applications highlighted are distinct to the plethora of reports on forming and utilising vinyl-boronate esters, and the reader is directed to the excellent recent reviews on these topics.60,61

Haloboration initially was used to stereoselectively produce singly or doubly halogenated terminal alkenes by reacting either terminal alkynes or haloalkynes with e.g., 9-Br-BBN^{13,14,62} to give the intermediate alkenylborane 54 followed by acetolysis (55, Scheme 21a). 63-65 1,2-Dihaloalkenes can also be accessed from terminal alkynes if post haloboration the boron moiety is transformed into an R-BF₃K salt and then treated with an electrophilic halogenating agent.66 Suzuki et al. first employed vinylBBr2 derivative 56 in a one-pot two-steps Negishi/Suzuki-Miyaura cross-coupling sequence to access 1,2-disubstituted alkenes (57)

DIPP-N N-DIPP

52

DIPP =
$$2,6-i$$
-Pr₂C₆H₃

CI CI

DIPP = $5,6-i$ -Pr₂C₆H₃

53

Scheme 20 Chloroboration of 1,4-diazabutadienes to access 1,3,2diazaborolidines.

Scheme 21 Early examples of functionalisation via B terminus

selectively (Scheme 21b).67,68 Although yields were good, the β-bromoalkenyl dibromoboranes 56 were found to be prone to retro-haloboration in the presence of Pd complexes, making further derivatisation to boronic esters expedient to lower the Lewis acidity of the B atom. 69-71 During their fundamental studies, Lappert et al. used esterification of R-BX2 with catechol or alkanols to transform their products into stable and conveniently analysed derivatives. 12,37 Esterification by ether cleavage 72 or by reaction with the respective alcohol⁷³ gave boronic esters with prolonged shelf-life.²⁵ Other functionalisation includes formation of R-B(dan) (dan = 1,8-diaminonaphthalen-N,N'diyl), 74 R-B(MIDA) complexes (MIDA = N-Methyliminodiacetate), 75 or R-BF₃K salts. 66,76 As expected, these classes of compounds are versatile reagents in transition-metal catalysed transformations such as Suzuki-Miyaura cross-coupling or Rh-catalysed [2 + 2 + 2] cycloadditions owing to their ambiphilic nature (electrophilic C-X/nucleophilic C-B). 39,77-80

The high B-centred Lewis acidity confers the alkenyl dihaloboranes a unique reactivity. The electrophilicity of the B atom paired with the nucleophilicity of the adjacent C atom enables the carboboration of alkynes. For example, the borocation $[LutBCl_2]^+$ (Lut = 2,6-lutidine) first reacts with R-C \equiv CH (R = alkyl, alkenyl, aryl) under 1,2-syn-haloboration to form compounds of general formula 58. Upon addition of Me-C≡C-SiMe₃, 1,2-syn-carboboration takes place, furnishing a borylated butadiene 59 which can be transformed into the corresponding pinacol ester by reaction with pinacol/NEt₃ (60, Scheme 22a). 81 Similar reactivity was observed for (F₅C₆)₂BCl. 82 If, however, $(F_5C_6)_2BX$ (X = Cl, Br) is reacted with enynes or cyclopropylacetylene, oligomerisation of the alkyne moiety to give 61 without interference of the alkene part of the molecule occurs. In this process, the newly formed C-B bond is added across the C≡C triple bond in a 1,2 carboboration reaction (Scheme 22b).83 Experiments by Eisch et al. suggest that haloboration is kinetically favoured over carboboration, but due to its irreversibility the latter is often the observed reaction outcome. 15 Taking advantage of boron's intrinsically high oxophilicity, haloboration can be exploited to form 1,4-dienes from alkynes, BX₃ and aldehydes under deoxygenative conditions in a one-pot process without isolation of the haloboration product 62 that is the intermediate in this process. Remarkably, the stereoselectivity is dependent on the halide employed with BCl_3 furnishing exclusively the (E,Z)diastereomer and BBr₃ the respective (Z,Z) isomer of 63 (Scheme 22c).84,85

a) [LutBCl₂][AlCl₄] R-C=CH CI
$$\oplus$$
 B(Lut)CI R 58 H

CI \longrightarrow Bpin Pinacol NEt₃ Pinacol NEt₃ SiMe₃

b) \longrightarrow BAr₂F R CI \longrightarrow BAR₂F R C

Scheme 22 Select examples of functionalisation of haloborated alkynes via the B terminus.

In a similar fashion, propargyl, 86 benzyl, 87 or allyl 88 alcohols can be used to access propargyl-, benzyl-, or allyl-substituted styrenes 64 (Scheme 22d). If instead of BX₃ 9-I-BBN is used, haloboration of ethoxyethyne yields a haloboration product which reacts with aldehydes via a formal C=O carboboration. The resulting secondary alkylborinic esters are readily hydrolysed to α,β unsaturated carbonic esters.89

The electron withdrawing nature of a BX2 group also renders alkenes like E-8 electron poor, making them good substrates in Diels–Alder reactions. Due to the instability of α-halogenated alkylboranes (cf. Section 4), BBr₃ elimination transforms the intermediate cyclohexene 65 into a 1,4-cyclohexadiene 66 (Scheme 23).90

6.1 Haloboration as a tool in natural product synthesis

Polyenes are common substructures in natural products (Scheme 24). 91,92 Usually, the double bonds are formed via Wittig-type olefinations, 93 and thus can suffer from forming mixtures of both E and Z isomers. Using the appropriate conditions the haloboration of terminal alkynes, as discussed above, proceeds stereoselectively (usually >98%^{70,94,95}) and furnishes ambiphilic halogenated vinyl boronic acid derivatives post workup.

Scheme 23 Diels-Alder reactions of haloboration products yield 1,4cvclohexadienes

Negishi, 2009

 $\dot{N}Me_2$ N-acyl spermidine Roulland, **2009** epothilone D (3Z.6E) α-farnesene

Scheme 24 Select natural products synthesised via haloboration/crosscoupling sequences. The fragment incorporated in such manner is coloured in red.

Whiting, 2018 HO

Br₂B
$$ii)$$
, iii)

OAc

 $O[Si]$
 $Si] = SiPh_2t$ -Bu

aonidiella citrina sex pheromone

Negishi, **2008**: 34% in 6 steps, >98% isomerically pure

Scheme 25 Example of a natural product synthesis sequence based on haloboration. Reagents and conditions: (i) BBr₃, DCM, -78 °C to 23 °C, 1 h; (ii) Me₂C=CH-CH₂ZnBr, Pd(PPh₃)₂Cl₂ (1 mol%), THF, 23 °C, 2 h; (iii) l₂, NaOAc, THF/H₂O, 23 °C, 1 h; (iv) zinc organyl, Pd(PPh₃)₄ (1 mol%), THF, 23 °C, 5 h; (v) [n-Bu₄N]F, THF, 23 °C, 1 h; (vi) Ac₂O, pyridine, DCM, 23 °C, 12 h.

Under the mild conditions of Pd catalysed cross-coupling reactions, the stereo information of the double bond is conserved. 94,95 Thus, haloboration/cross-coupling sequences can be applied in the synthesis of olefinic natural products (Scheme 25). 96 The inherent instability of polyene boronic acids can be circumvented by either using them directly without further purification or using the MIDA boronate derivatives. Indeed, only 12 different MIDA boronates derived inter alia from the haloboration product of acetylene are necessary to build most polyene natural products via consecutive Suzuki-Miyaura reactions. 75,92

6.2 Haloboration in the synthesis of B-doped PAHs

Besides providing a useful route to form β-functionalised boronic-acid derivatives, haloboration is an expedient tool to form boron-doped polycyclic aromatic hydrocarbons (B-PAHs), which are promising candidate materials in the field of organic electronics as they are electron deficient aromatics with a comparatively low LUMO energy. 23,97 Haloboration of 1-alkynylnaphthalenes 67 can lead selectively (under appropriate conditions) to either trans-1,2-haloboration products (R = H, alkyl) or 1,1-haloboration products 68 (R = aryl,

Scheme 26 (a) Synthesis of 1-boraphenalene 69 via haloboration/bora Friedel-Crafts reaction sequence. (b) Extension to pyrene derivatives.

Scheme 26), which then undergo bora Friedel-Crafts reactions furnishing 1-boraphenalenes (69, Scheme 26a). 97 If pyrene instead of naphthalene is used as the hydrocarbon core (70), singly or doubly (71) B-doped PAHs with very low LUMO energies are accessible.²³ The utility of bromoboration is then demonstrated by usage of the vinyl-Br unit in a subsequent Negishi coupling enabling donor-acceptor-donor complex 72 to be formed readily (Scheme 26b). In other materials synthesis applications the high Lewis acidity of BBr₃ allows for a one-shot double cyclisation of o,o'-dimethoxy-substituted tolan derivatives by ether cleavage and concomitant trans-haloboration, yielding benzofurochromene derivatives (cf. Scheme 15, Section 3).32 In a similar fashion, N-protected propargylamines underwent intramolecular N-B bond formation post haloboration.⁹⁸

Scheme 27 Haloboration of diynes can be used to construct B incorporating PAHs or to access poly(vinylboranes).

Perspective NJC

A related reaction sequence was used to construct even larger B,N doped PAHs. 99 Besides being useful to incorporate boron into PAHs, haloboration has also been demonstrated to be a valuable tool to construct B-incorporated polymers. This was achieved by subjecting diynes such as 73 to multiple haloboration reactions using either BBr3 or R2BBr as the B source to yield polymers such as 74 (Scheme 27). 100

7. Conclusion

Since its discovery almost 80 years ago, the haloboration of alkynes has developed into a reliable way to form bifunctional alkenes with excellent control of stereoselectivity possible. The haloboration of other substrates is more limited and further work is required to develop these into broadly useful transformations. Regarding alkyne haloboration, although some mechanistic details are still subject to debate, experimental evidence clearly demonstrates that the stereoselectivity of haloboration can be controlled. The addition of BX_3 (X = Cl, Br, I) to a terminal alkyne proceeds via syn-addition of a B-X bond across the C≡C triple bond and is usually very fast, even at low temperatures. Thus, low-temperature quenching furnishes the Z-adduct selectively. At higher temperatures, isomerisation to the E-adduct can occur. The initial addition as well as the subsequent isomerisation are highly efficient, allowing access to the respective adduct in high yields with stereoselectivities often > 98%. Haloboranes such as PhBBr₂ or 9-Br-BBN can be employed in haloboration as well. For internal alkynes, stronger electrophiles than BCl3 are needed to effect haloboration, and most simply BBr₃ can be used. The lower reactivity of internal C≡C triple bonds allows for the isolation of the synaddition product even at room temperature within minutes. Yet, prolonged storage in solution or elevated temperatures (or more polar solvents) can lead to rearrangement of the 1,2-syn addition product to the 1,1-haloboration product in a reversible process. Thus, the addition of BBr₃ to internal alkynes yields mixtures of both 1,1- and 1,2-adducts. Notably both the 1,2-adduct and the 1,1 adduct can be isolated selectively using the appropriate conditions.

The combination of halide and boron moieties in the addition product allows for broad diversification of the obtained olefin. For example, haloboration outperforms most other common C=C bond formation reactions in terms of selectivity and it has proved its worth in the field of polyene natural product synthesis. Notably, the primary products from haloboration contain vinylBX2 units which often have distinct reactivity compared to vinylB(OR)₂ analogues due to the stronger electron withdrawing nature of the BX₂ unit. An emerging application of the haloboration reaction is in the incorporation of B atoms into large delocalised π systems. This may either be achieved by using strategically positioned heteroatoms (e.g., O, N) to direct alkyne haloboration, or by alkyne haloboration followed by a bora Friedel-Crafts reaction on alkyne-substituted PAHs. These are attractive as the formed B-doped PAHs are halogenated enabling facile subsequent diversification. We hope

that further applications are forthcoming that help bring haloboration out of the shadow of the ubiquitous hydroboration

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was made possible by financial support from the Horizon 2020 Research and Innovation Programme (Grant No. 769599). Dr Jürgen Pahl and Saqib Iqbal (both University of Edinburgh) are thanked for useful comments.

Notes and references

- 1 B. M. Trost, Science, 1991, 254, 1471-1477.
- 2 H. C. Brown and B. C. Subba Rao, J. Am. Chem. Soc., 1956, 78, 5694-5695.
- 3 D. S. Matteson, K. M. Sadhu and M. L. Peterson, J. Am. Chem. Soc., 1986, 108, 810-819.
- 4 N. A. Petasis and I. Akritopoulou, Tetrahedron Lett., 1992, 34, 583-586.
- 5 (a) D. M. T. Chan, K. L. Monaco, R. Li, D. Bonne, C. G. Clark and P. Y. S. Lam, Tetrahedron Lett., 2003, 44, 3863-3865; (b) P. Y. S. Lam, G. Vincent, D. Bonne and C. G. Clark, Tetrahedron Lett., 2003, 44, 4927-4931.
- 6 N. Miyaura and A. Suzuki, J. Chem. Soc., Chem. Commun., 1979, 866-867.
- 7 For recent reviews on hydroboration of C=E bonds, cf. (a) C. M. Crudden and D. Edwards, Eur. J. Org. Chem., 2003, 4695-4712; (b) A. M. Carroll, T. P. O'Sullivan and P. J. Guiry, Adv. Synth. Catal., 2005, 347, 609-631; (c) C. C. Chong and R. Kinjo, ACS Catal., 2015, 5, 3238-3259; (d) J. V. Obligacion and P. J. Chirik, Nat. Rev. Chem., 2018, 2, 15-34; (e) M. L. Shegavi and S. K. Bose, Catal. Sci. Technol., 2019, 9, 3307-3336.
- 8 Reviews on the haloboration reaction were published by Suzuki and Hara, but the articles were either in Japanese or the journal they were published in ceased to exist: (a) S. Hara, J. Synth. Org. Chem., Jpn., 1990, 48, 1125–1137; (b) A. Suzuki and S. Hara, J. Synth. Org. Chem., Jpn., 1985, 43, 100-111; (c) A. Suzuki, Rev. Heteroat. Chem., 1997, 17, 271-314. To the best of our knowledge, the last broadly accessible review is: (d) A. Suzuki, Pure Appl. Chem., 1986, 58, 629-638.
- 9 H. R. Arnold, New Organoboron Compounds, US Pat., 2402589, E. I. DuPont de Nemours & Company, 1946.
- 10 R. J. Jensen, J. K. Hayes, C. L. Cluff and J. M. Thorne, IEEE J. Quantum Electron., 1980, 16, 1352-1356.
- 11 E. Gipstein, P. R. Kippur, M. A. Higgins and B. F. Clark, J. Org. Chem., 1961, 26, 943.
- 12 M. F. Lappert and B. Prokai, J. Organomet. Chem., 1964, 1, 384-400.

- 13 N. Zhou, Q. Wang, A. J. Lough and H. Yan, Can. J. Chem., 2012, 90, 625-630.
- 14 (a) J. R. Blackborow, J. Chem. Soc., Perkin Trans. 2, 1989, 1973; (b) J. R. Blackborow, J. Organomet. Chem., 1977, 128,
- 15 J. J. Eisch and L. J. Gonsior, J. Organomet. Chem., 1967, 8, 53-64
- 16 B. Wrackmeyer, Polyhedron, 1986, 5, 1709-1721.
- 17 J. J. Eisch and H. P. Becker, J. Organomet. Chem., 1979, 171, 141-153.
- 18 W. Siebert, R. Full, J. Edwin and K. Kinberger, Chem. Ber., 1978, 111, 823-831.
- 19 Y. Nie, S. Schwiegk, H. Pritzkow and W. Siebert, Eur. J. Inorg. Chem., 2004, 1630-1638.
- 20 S. Hara, Y. Satoh, H. Ishiguro and A. Suzuki, Tetrahedron Lett., 1983, 24, 735-738.
- 21 C. Wang and M. Uchiyama, Eur. J. Org. Chem., 2012, 6548-6554.
- 22 J. Polášek, J. Paciorek, J. Stošek, H. Semrád, M. Munzarová and C. Mazal, J. Org. Chem., 2020, 85, 6992-7000.
- 23 K. Yuan, R. J. Kahan, C. Si, A. Williams, S. Kirschner, M. Uzelac, E. Zysman-Colman and M. J. Ingleson, Chem. Sci., 2020, 11, 3258-3267.
- 24 J. R. Lawson, E. R. Clark, I. A. Cade, S. A. Solomon and M. J. Ingleson, Angew. Chem., Int. Ed., 2013, 52, 7518-7522.
- 25 A. J. Warner, A. Churn, J. S. McGough and M. J. Ingleson, Angew. Chem., Int. Ed., 2017, 56, 354-358.
- 26 F. D. Zhuang, J. M. Han, S. Tang, J. H. Yang, O. R. Chen, J. Y. Wang and J. Pei, Organometallics, 2017, 36, 2479-2482.
- 27 S. A. Iqbal, J. Pahl, K. Yuan and M. J. Ingleson, Chem. Soc. Rev., 2020, 49, 4564-4591.
- 28 L. C. Wilkins, Y. Soltani, J. R. Lawson, B. Slater and R. L. Melen, Chem. - Eur. J., 2018, 24, 7364-7368.
- 29 A. J. Warner, J. R. Lawson, V. Fasano and M. J. Ingleson, Angew. Chem., Int. Ed., 2015, 54, 11245-11249.
- 30 C. Chen, M. Harhausen, R. Liedtke, K. Bussmann, A. Fukazawa, S. Yamaguchi, J. L. Petersen, C. G. Daniliuc, R. Fröhlich, G. Kehr and G. Erker, Angew. Chem., Int. Ed., 2013, 52, 5992-5996.
- 31 A. J. Warner, K. M. Enright, J. M. Cole, K. Yuan, J. S. McGough and M. J. Ingleson, Org. Biomol. Chem., 2019, 17, 5520-5525.
- 32 Y. Uchikawa, K. Tazoe, S. Tanaka, X. Feng, T. Matsumoto, J. Tanaka and T. Yamato, Can. J. Chem., 2012, 90, 441-449.
- 33 M. F. Hawthorne and J. A. DuPont, J. Am. Chem. Soc., 1958, 80, 5830-5832.
- 34 H. C. Brown and K. A. Keblys, J. Am. Chem. Soc., 1964, 86, 1791-1795.
- 35 R. Köster, G. Griasnow, W. Larbig and P. Binger, Justus Liebigs Ann. Chem., 1964, 672, 1-34.
- 36 F. Joy, M. F. Lappert and B. Prokai, J. Organomet. Chem., 1966, 5, 506-519.
- 37 F. Joy and M. F. Lappert, Proc. Chem. Soc., 1960, 1, 353-354.
- 38 For examples of BX3 used as a catalyst in cationic polymerisation, cf. (a) E. L. Muetterties, J. Am. Chem. Soc., 1960, 82, 4163-4166; (b) L. Balogh, L. Wang and R. Faust,

- Macromolecules, 1994, 27, 3453-3458; (c) L. Balogh, Z. Fodor, T. Kelen and R. Faust, Macromolecules, 1994, 27, 4648-4651.
- 39 K. K. Wang, Z. Wang, A. Tarli and P. Gannett, J. Am. Chem. Soc., 1996, 118, 10783-10791.
- 40 B. M. Mikhailov and M. E. Nikolaeva, Russ. Chem. Bull., 1963, 12, 1246-1250.
- 41 Y. N. Bubnov, N. Y. Kuznetsov, F. V. Pastukhov and V. V. Kublitsky, Eur. J. Org. Chem., 2005, 4633-4639.
- 42 H. Abu Ali, V. M. Dembitsky and M. Srebnik, in Science of Synthesis 6: Category 1, Organometallics, ed. D. E. Kaufmann, D. S. Matteson, E. Schaumann and M. Regitz, Georg Thieme Verlag, Stuttgart, 1st edn, 2005, pp. 867–879.
- 43 S. Hara and A. Suzuki, Tetrahedron Lett., 1991, 32, 6749-6752.
- 44 S. Hara, S. Takinami, S. Hyuga and A. Suzuki, Chem. Lett., 1984. 345-348.
- 45 C. Diner and K. J. Szabó, J. Am. Chem. Soc., 2017, 139, 2-14.
- 46 Z. Yang, T. Liu, X. Chen, R. Wan, Y. Li, X. Wang, C. H. Yang and J. Chang, Org. Lett., 2019, 21, 9541-9544.
- 47 M. J. Frazer, W. Gerrard and M. F. Lappert, J. Chem. Soc., 1957, 739.
- 48 A. G. Massey, J. Chem. Soc., 1961, 1103-1105.
- 49 (a) G. W. Parshall, *Inorg. Chem.*, 1965, 4, 52-54; (b) E. W. Abel, D. J. Walker and J. N. Wingfield, Inorg. Nucl. Chem. Lett., 1969, 5, 139–140; (c) E. W. Abel, N. Giles, D. J. Walker and J. N. Wingfield, J. Chem. Soc. A, 1971, 1991-1993.
- 50 E. W. Abel, D. J. Walker and J. N. Wingfield, Inorg. Nucl. Chem. Lett., 1969, 5, 139-140.
- 51 A. Meller and W. Maringgele, Monatsh. Chem., 1968, 99, 2504-2513.
- 52 A. Meller and A. Ossko, Monatsh. Chem., 1969, 100, 1187-1194.
- 53 A. Meller, W. Maringgele and G. Maresch, Monatsh. Chem., 1974, 105, 637-647.
- 54 B. Bonnetot, B. Frange, F. Guilhon, H. Mongeot and W. Einholz, Polyhedron, 1994, 13, 2211-2216.
- 55 A. Meller and H. Batka, Monatsh. Chem., 1969, 100, 1823-1828.
- 56 K. Niedenzu, K. E. Blick and C. D. Miller, Inorg. Chem., 1970, **9**, 975-977.
- 57 C. D. Miller and K. Medenzu, Synth. React. Inorg. Met.-Org. Chem., 1972, 2, 217-222.
- 58 F. S. Mair, R. Manning, R. G. Pritchard and J. E. Warren, Chem. Commun., 2001, 1136-1137.
- 59 (a) M. Yamashita, Y. Suzuki, Y. Segawa and K. Nozaki, J. Am. Chem. Soc., 2007, 129, 9570-9571; (b) Y. Segawa, Y. Suzuki, M. Yamashita and K. Nozaki, J. Am. Chem. Soc., 2008, 130, 16069-16079; (c) A. Yagi, H. Kisu and M. Yamashita, Dalton Trans., 2019, 48, 5496-5499.
- 60 A. J. J. Lennox and G. C. Lloyd-Jones, Chem. Soc. Rev., 2014, 43, 412-443.
- 61 J. Carreras, A. Caballero and P. J. Pérez, Chem. Asian J., 2019, 14, 329-343.
- 62 M. J. Bayer, H. Pritzkow and W. Siebert, Z. Naturforsch. B, 2002, 57, 295-300.

- 63 S. Hara, H. Dojo, S. Takinami and A. Suzuki, Tetrahedron Lett., 1983, 24, 731-734.
- 64 S. Hara, T. Kato and A. Suzuki, Synthesis, 1983, 1005-1006.
- 65 R. V. Edwankar, C. R. Edwankar, J. R. Deschamps and J. M. Cook, J. Org. Chem., 2014, 79, 10030-10048.
- 66 (a) N. Zhou, O. Wang, A. J. Lough and H. Yan, Can. J. Chem., 2012, 90, 625-630; (b) M. L. Yao, G. W. Kabalka, D. W. Blevins, M. S. Reddy and L. Yong, Tetrahedron, 2012, **68**, 3738-3743.
- 67 S. Hyuga, Y. Chiba, N. Yamashina, S. Hara and A. Suzuki, Chem. Lett., 1987, 1757-1760.
- 68 Y. Satoh, H. Serizawa, N. Miyaura, S. Hara and A. Suzuki, Tetrahedron Lett., 1988, 29, 1811-1814.
- 69 M. L. Yao, M. S. Reddy, W. Zeng, K. Hall, I. Walflsh and G. W. Kabalka, J. Org. Chem., 2009, 74, 1385-1387.
- 70 C. Wang, Z. Xu, T. Tobrman and E. I. Negishi, Adv. Synth. Catal., 2010, 352, 627-631.
- 71 C. Mazal and M. Vaultier, Tetrahedron Lett., 1994, 35, 3089-3090.
- 72 N. Yamashina, S. Hyuga, S. Hara and A. Suzuki, Tetrahedron Lett., 1989, 30, 6555-6558.
- 73 S. Hyuga, N. Yamashina, S. Hara and A. Suzuki, Chem. Lett., 1988, 809-812.
- 74 Y. Sumida, R. Harada, T. Kato-Sumida, K. Johmoto, H. Uekusa and T. Hosoya, Org. Lett., 2014, 16, 6240-6243.
- 75 S. J. Lee, K. C. Gray, J. S. Paek and M. D. Burke, J. Am. Chem. Soc., 2008, 130, 466-468.
- 76 X. Fang, J. Sun and X. Tong, Chem. Commun., 2010, 46, 3800-3802.
- 77 K. K. Wang and Z. Wang, Tetrahedron Lett., 1994, 35, 1829-1832.
- 78 A. Tarli and K. K. Wang, J. Org. Chem., 1997, 62, 8841-8847.
- 79 B. M. Trost and F. D. Toste, J. Am. Chem. Soc., 2002, 124, 5025-5036.
- 80 M. Sato, Y. Yamamoto, S. Hara and A. Suzuki, Tetrahedron Lett., 1993, 34, 7071-7074.
- 81 J. R. Lawson, V. Fasano, J. Cid, I. Vitorica-Yrezabal and M. J. Ingleson, Dalton Trans., 2016, 45, 6060-6070.
- 82 K. Škoch, C. Pauly, C. G. Daniliuc, K. Bergander, G. Kehr and G. Erker, Dalton Trans., 2019, 48, 4837-4845.
- 83 A. Ueno, J. Li, C. G. Daniliuc, G. Kehr and G. Erker, *Chem.* Eur. J., 2018, 24, 10044-10048.
- 84 G. W. Kabalka, Z. Wu and Y. Ju, Org. Lett., 2002, 4, 1491-1493.
- 85 G. W. Kabalka, M. L. Yao, S. Borella, Z. Wu, Y. H. Ju and T. Quick, J. Org. Chem., 2008, 73, 2668-2673.
- 86 G. W. Kabalka, Z. Wu and Y. Ju, Org. Lett., 2004, 6, 3929-3931.

- 87 G. W. Kabalka, M. L. Yao, S. Borella and Z. Z. Wu, Org. Lett., 2005, 7, 2865-2867.
- 88 G. W. Kabalka, M. L. Yao, S. Borella and Z. Wu, Chem. Commun., 2005, 2492-2494.
- 89 Y. Satoh, T. Tayano, S. Hara and A. Suzuki, Tetrahedron Lett., 1989, 30, 5153-5156.
- 90 D. A. Singleton and S. W. Leung, J. Organomet. Chem., 1997, **544**, 157-161.
- 91 C. Thirsk and A. Whiting, J. Chem. Soc., Perkin Trans. 1, 2002, 999-1023.
- 92 E. M. Woerly, J. Roy and M. D. Burke, Nat. Chem., 2014, 6, 484-491.
- 93 B. E. Maryanoff and A. B. Reitz, Chem. Rev., 1989, 89, 863-927.
- 94 S. Xu, C. T. Lee, H. Rao and E. I. Negishi, Adv. Synth. Catal., 2011, 353, 2981-2987.
- 95 E. I. Negishi, T. Tobrman, H. Rao, S. Xu and C. T. Lee, Isr. J. Chem., 2010, 50, 696-701.
- 96 (a) N. Miyaura, Y. Satoh, S. Hara and A. Suzuki, Bull. Chem. Soc. Jpn., 1986, 59, 2029-2031; (b) A. B. Smith, G. K. Friestad, J. J. W. Duan, J. Barbosa, K. G. Hull, M. Iwashima, Y. Qiu, P. Grant Spoors, E. Bertounesque and B. A. Salvatore, J. Org. Chem., 1998, 63, 7596-7597; (c) M. D. Chappell, S. J. Stachel, C. B. Lee and S. J. Danishefsky, Org. Lett., 2000, 2, 1633-1636; (d) Z. Xu and E. I. Negishi, Org. Lett., 2008, 10, 4311-4314; (e) X. Guinchard, X. Bugaut, C. Cook and E. Roulland, Chem. - Eur. J., 2009, 15, 5793-5798; (f) C. Wang, T. Tobrman, Z. Xu and E. I. Negishi, Org. Lett., 2009, 11, 4092-4095; (g) A. M. Haydl and B. Breit, Chem. - Eur. J., 2017, 23, 541-545; (h) K. S. Madden, B. Laroche, S. David, A. S. Batsanov, D. Thompson, J. P. Knowles and A. Whiting, Eur. J. Org. Chem., 2018, 5312-5322.
- 97 R. J. Kahan, D. L. Crossley, J. Cid, J. E. Radcliffe and M. J. Ingleson, Angew. Chem., Int. Ed., 2018, 57, 8084–8088.
- 98 A. Meller, F. J. Hirninger, M. N. W. Maringgele, A. Meller, F. J. Hirninger and M. Noltemeyer, Chem. Ber., 1981, 114, 2519-2535.
- 99 F. D. Zhuang, Z. H. Sun, Z. F. Yao, Q. R. Chen, Z. Huang, J. H. Yang, J. Y. Wang and J. Pei, Angew. Chem., Int. Ed., 2019, 58, 10708-10712.
- 100 (a) Y. Chujo, I. Tomita and T. Saegusa, Macromolecules, 1990, 3, 687-689; (b) Y. Chujo, I. Tomita and T. Saegusa, Makromol. Chem., Macromol. Symp., 1993, 70-71, 47-56; (c) N. Matsumi and Y. Chujo, Polym. Bull., 1997, 39, 295-302; (d) M. Miyata, N. Matsumi and Y. Chujo, Polym. Bull., 1999, 42, 505-510.