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Relation	





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Borylstannylation of alkynes with inverse regioselectivity: copper-catalyzed three-component coupling using a masked diboron

Borylstannylation of terminal alkynes proceeds with inverse regioselectivity to those of the previous borylstannylations by utilizing a masked diboron as a boron source under copper catalysis.

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Borylstannylation of alkynes with inverse regioselectivity: copper-catalyzed threecomponent coupling using a masked diboron[†]

Pd-catalyzed borylstannylatior

H. Yoshida,*^{ab} Y. Takemoto^a and K. Takaki^a

A variety of terminal alkynes are facilely convertible into cisboryl(stannyl)alkenes with inverse regioselectivity to those of the previous borylstannylation by the copper-catalyzed three-component reaction using a masked diboron. The synthetic utility of the resulting boryl(stannyl)alkenes has been demonstrated by chemoselective coupling reactions.

Transition metal-catalyzed dimetallation of alkynes has commanded considerable attention¹ because it provides a convenient and direct method for constructing regio- and stereo-defined dimetallated alkenes, whose carbon-metal bonds are utilizable for carbon-carbon bond-forming processes² to give multisubstituted alkenes, which constitute an important class of biologically and pharmaceutically active molecules. One of the most valuable dimetallation reactions would be borylstannylation, in which the resulting hetero-dimetallic moieties can undergo chemoselective cross-coupling reactions (Suzuki-Miyaura³ and Migita-Kosugi-Stille coupling⁴) in tandem with high functional group compatibility under controlled reaction conditions. Since the pioneering work reported by Tanaka,^{5a} borylstannylation has hitherto been achieved by direct insertion of alkynes into a B-Sn bond of borylstannanes under palladium catalysis.⁵ On the other hand, we have recently disclosed a different mode of the borylstannylation by a copper-catalyzed three-component coupling using a diboron and a tin alkoxide.⁶⁻⁸ Irrespective of the catalytic systems and the reaction modes, terminal alkynes exclusively accept the regioselective addition of the boryl group at the terminal carbon and the stannyl group at the internal carbon in a *cis* fashion to give (Z)-1-boryl-2-stannyl-1-alkenes (Scheme 1), and thus we have focused our attention on the reversal of regioselectivity, which increases structural diversity of vic-boryl(stannyl)alkenes and

Pd(PPh₃)₄



thereby broadens the synthetic utility of the borylstannylation. Herein we report that the use of a masked diboron⁹ in the coppercatalyzed three-component borylstannylation of terminal alkynes completely inverts the regioselectivity, and that this method provides a convenient and direct access to unprecedented hetero-dimetallated alkenes having masked boryl and stannyl moieties.10

First we conducted the reaction of 1-octyne (1a) with a masked diboron ((pin)B-B(dan), pin: pinacolato, dan: naphthalene-1,8diaminato¹¹) and tributyltin methoxide in THF at room temperature in the presence of an N-heterocyclic carbene (NHC)-coordinated copper complex ((SIPr)CuCl), and found that the cis-borylstannylation took place with regioselectivity inverse to those of the previous borylstannylation (74% yield, 2a:2'a = 96:4), leading to the introduction of the boryl group at the internal carbon and the stannyl group at the terminal carbon (Table 1, entry 1). It is noteworthy that the B(dan) moiety was solely installed in the product, and a borylstannylation product having the B(pin) moiety was not formed at all. The regioselectivity for the formation of 2a was generally high with bulky ligands





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R -----(pin)B-B(pin) Bu₃SnOMe Cu(OAc)₂ PCy₃ B(pin) = Cu-catalvzed three-component borvlstannvlation

Table 1 Ligand effect on Cu-catalyzed borylstannylation of 1-octyne⁴



(bii i jouoi	,	74	50.4
(SIMes)CuCl	2	81	90:10
(tBu-SIPr)CuCl	20	69	94:6
(IPr)CuCl	5	75	96:4
(IPr*)CuCl	11	80	96:4
$P(tBu)_3$, CuCl	2	81	93:7
(PPh ₃) ₃ CuCl	48	75	44:56
PCy_3 , CuCl	14	60	84:16
(SIPr)CuCl	10	86	96:4

^a General procedure: 1a (0.30 mmol, 1 equiv.), (pin)B-B(dan) (0.36 mmol, 1.2 equiv.), Bu₃SnOMe (0.36 mmol, 1.2 equiv.), Cu catalyst (6.0 µmol, 2 mol%), THF (1 mL). ^b Isolated yield. ^c Determined using ¹H NMR. ^{*d*} Ligand = 4 mol%. ^{*e*} Bu₃SnOMe = 2 equiv.

(SIMes, *t*Bu-SIPr, IPr, IPr *12 and P(*t*Bu)₃) (entries 2–6), whereas the use of triphenylphosphine ((PPh₃)₃CuCl) led to the formation of regioisomeric mixtures (2a: 2'a = 44: 56, entry 7). In addition, the reaction with PCy₃, used for the previous borylstannylation with bis(pinacolato)diboron,^{6a} also afforded 2a preferentially (2a: 2'a = 84: 16, entry 8), which reveals that the choice of a diboron as well as a ligand is the key to achieving the present regioselectivity.13 Since an increase in the amount of tributyltin methoxide resulted in an increase in the yield with the highest regioselectivity (86% yield, entry 9), we selected the conditions for further studies.¹⁴

With the optimum conditions in hand (Table 1, entry 9), the substrate scope of alkynes was next investigated (Table 2). Such aliphatic terminal alkynes as 1-hexyne (1b), 4-methyl-1-pentyne (1c) and 4-phenyl-1-butyne (1d) also underwent the borylstannylation with high degrees of regioselectivity to give 2b, 2c and 2d in 78, 81 and 74% yield (entries 1–3). The functional group compatibility of the reaction was sufficiently high, and thus a C-Br bond¹⁵ in **1e** and a cyano group in **1f** remained intact throughout the reaction (entries 4 and 5). The present regioselectivity was also observed by using envne (1g) and phenylacetylene (1h) (entries 6 and 7), and furthermore the reaction of propargyl ethers (1i and 1j) or a THP-protected propargyl alcohol (1k) resulted in the exclusive formation of 2i-2k (entries 8-10). In addition, propargyl amine (11) and trimethylsilylacetylene (1m) accepted the addition of the B(dan) moiety at their internal carbon with perfect regioselectivity (entries 11 and 12).¹⁶ The versatility of

Table 2 NHC-Cu-catalyzed borylstannylation of terminal alkynes⁴

			(dan)B	SnBu₃ —∕
		Bu₃SnOMe (2 equiv) (SIPr)CuCl (2 mol%)	R	2
R— <u>—</u> 1 1	: + (pin)B—B(dan) : 1.2	THF, rt, 1 h	Bu₃Sn ≻= R	B(dan) —∕
Entry	R	Yield ^{b} (%)	2:2' ^c	Products
1	<i>n</i> Bu (1b)	78	94:6	2b, 2'b
2	iBu (1c)	81	99:1	2c, 2'c
3	$Ph(CH_2)_2$ (1d)	74	94:6	2d, 2'd
4	$Br(CH_2)_2$ (1e)	87	99:1	2e, 2'e
5	$NC(CH_2)_3$ (1f)	79	95:5	2f, 2'f
6	1-Cyclohexenyl (1g)	81	99:1	2g, 2'g
7	Ph (1h)	73	99:1	2h, 2'h
8	$MeOCH_2$ (1i)	66	>99:1	2i
9	$BnOCH_2$ (1j)	66	>99:1	2j
10	THPOCH ₂ $(1k)$	66	>99:1	2k
11	Et_2NCH_2 (11)	69	>99:1	21
	`. `			-

General procedure: 1 (0.30 mmol, 1 equiv.), (pin)B-B(dan) (0.36 mmol, 1.2 equiv.), Bu₃SnOMe (0.60 mmol, 2 equiv.), (SIPr)CuCl (6.0 µmol, 2 mol%), THF (1 mL). ^b Isolated yield. ^c Determined using ¹H NMR.

the borylstannylation was further expanded by application to 1,7-octadiyne¹⁷ (1n) and allenes¹⁸ (3a and 3b): both of the triple bonds were convertible regioselectively into the borylstannylalkenes in the former case, and the regio- and stereoselective reaction proceeded to provide (Z)-1-stannyl-2-boryl-2-alkenes (4a and 4b) as the single product, although the regioselectivity is similar to that of the previous borylstannylation with bis(pinacolato)diboron^{6b} in the latter case (Scheme 2).

Similarly to the previous copper-catalyzed borylstannylation with bis(pinacolato)diboron,⁶ generation of a borylcopper species, Cu-B(dan), from Cu-OMe and a masked diboron commences the reaction (Scheme 3, step A). Subsequent insertion of an alkyne



Scheme 2 Borylstannylation of a diyne and allenes.

3

4

5

 6^d 7 8^d

٩é



Scheme 3 A plausible catalytic cycle for borylstannylation.

into the Cu-B(dan) bond, which produces a β-borylalkenylcopper species (borylcupration, step B),¹⁹ followed by capturing with tin methoxide furnishes the product (step C).²⁰ The formation of Cu-B(dan) (vs. Cu-B(pin)) can be rationally explained by selective interaction between the Lewis acidic B(pin) moiety of (pin)B-B(dan) and the methoxy moiety of Cu-OMe in step A, leading to the exclusive introduction of the masked boryl moiety across the triple bond of alkynes. The orientation of a borylcopper species in the borylcupration step entirely governs the regiochemical outcome of the borylstannylation (Scheme 4), and the mode of the borylcupration with Cu-B(dan) would simply be controlled by steric repulsion between a substituent on alkynes and a bulkier copper moiety as was the case with the hydroboration.^{10a} Hence, the B(dan) moiety is solely installed into the internal carbon of terminal alkynes,^{21,22} which results in the inverse regioselectivity in the present borylstannylation.

Synthetic utility of the boryl(stannyl)alkenes was demonstrated by the chemoselective cross-coupling: a C–Sn bond of **2i** was solely convertible into a C–C bond by the palladium-catalyzed Migita– Kosugi–Stille reaction to provide an 82% yield of **5** with a masked boryl moiety remaining intact (Scheme 5). Furthermore, the masking enabled the copper-mediated oxidative homocoupling to take place at the C–Sn bond selectively, affording 1,4-diboryl-1,3-butadienes (**6–8**) stereoretentively in high yield. Unmasking of the resulting 1,4-diboryl-1,3-butadiene, followed by the Suzuki–Miyaura reaction with 4-iodotoluene furnished 1,1,4,4-tetraarylbutadiene **9**.

In conclusion, we have disclosed that the borylstannylation of terminal alkynes proceeds with inverse regioselectivity by the



Scheme 4 Regioselectivity in the borylcupration



Scheme 5 Transformation of borylstannylation products.

copper-catalyzed three-component reaction using a masked diboron, which gives us a convenient and potent approach to diverse *cis*-boryl(stannyl)alkenes bearing the masked boryl moiety at the internal carbons. Moreover, the synthetic versatility of the resulting boryl(stannyl)alkenes has been shown by the chemoselective coupling reactions depending on the difference in the reactivity between the masked boryl and the stannyl moieties. Further studies on copper-catalyzed borylation reactions using a masked diboron as well as on the details of the mechanism are in progress.

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