

Two-Step Regio- and Stereoselective Synthesis of *cis*-Vinylstannanes

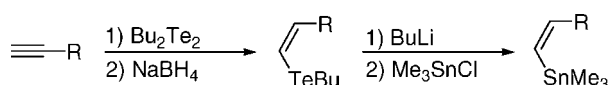
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ABSTRACT



Herein we describe a novel stereoselective synthesis of *cis*-vinylstannanes employing the widely established Li/Te exchange pathway. In contrast to previously reported methods of *cis*-selective hydrostannylation (i.e., ZrCl_4), this method demonstrates compatibility toward oxygenated substrates.

The Stille coupling¹ of vinyltin reagents with vinyl² and aryl halides² and triflates² is a synthetically useful means for obtaining conjugated double bonds with defined geometry.¹ Unfortunately, methods to access *cis*-vinylstannanes are somewhat limited and often exhibit poor stereoselectivity. To date, *cis*-vinylstannanes have been most commonly synthesized by radical,³ Lewis acid,⁴ and transition metal⁵ mediated hydrostannylation of alkynes, hydrozirconation⁶ of stannyl alkynes, and the transmetalation of vinyl metallic substrates.⁷ The use of ZrCl_4 as a Lewis acid catalyst in the hydrostannylation of alkynes has proven especially valuable,^{4b} mainly because the reaction proceeds in an entirely *cis*-selective manner. However, the reaction is not compatible with oxygen-containing functionality, possibly due to coordination to the Lewis acid, as it requires bulky substituents (e.g., TBDMS) on oxygen to facilitate reaction.^{4b} The use of bulky substituents on oxygen, however, has not proven

to be a general method in order to circumvent this problem.^{8,9} Furthermore, in the case of di- and tristannylation the high control over stereochemistry is lost.^{4c}

The hydrometalation of alkynes tends to occur in a *syn* fashion to give *trans*-organometallics. However, isomerization often leads to mixtures of *cis* and *trans* isomers. An exception is found in the hydrotelluration of alkynes, which occurs in a highly selective *anti* fashion to give *cis*-vinyltellurides, which are not susceptible to isomerization.¹⁰ Vinyl tellurides are also known to undergo transmetalation to give, for example, vinyl lithium, copper, zinc, magnesium, aluminum, calcium, and sodium reagents with retention of configuration.¹¹

With a view to finding a method to selectively synthesize *cis*-vinylstannanes that could also accommodate oxygen functionality, a two-step hydrotelluration lithium–tin exchange protocol was investigated, the results of which are presented herein.

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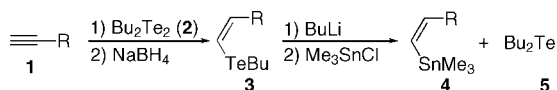
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Table 1. Hydrotelluration of Alkynes **1** Affording *cis*-Vinyltellurides **3**

entry	substrate	vinyltelluride	time (h)	yield (%) ^a
1			20	67%
2			18	71%
3			22	79%
4			2.5	61%
5			5	78%
6			3	90%
7			7	78%
8			2.5	75%
9			15	48%
10 ^b			21	42%

^a Isolated yields. ^b Inseparable mixture (ratio 2:1).

Alkynes (i.e., **1**) (Table 1, entries 1–10) were treated with dibutyl ditelluride **2** and sodium borohydride under standard conditions¹¹ to give *cis*-vinyltellurides (i.e., **3**) in 48–90% yield (Table 1) (Scheme 1). In all conjugated alkyne cases,

Scheme 1. Synthesis of *cis*-Vinyl Tellurides **3** and *cis*-Vinylstannanes **4**

regio- and stereoselectivity were maintained. Isolated alkynes (Table 1, entry 10), however, posed regioselectivity problems yielding mixtures of 1,2- and 2,2-disubstituted vinylic tellurides, in favor of the 1,2-disubstituted vinylic tellurides, as previously documented.^{12,13}

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The *cis*-vinyl tellurides were uneventfully transmetalated with *n*-butyllithium to the corresponding vinyl lithium, which were subsequently quenched with trimethyltin chloride affording the *cis*-vinylstannanes in 40–68% yield without any detectable *trans*-vinylstannane (Table 2, entries 1–9) (Scheme 1).

Table 2. Transmetalation of *cis*-Vinyl Telluride **3** Affording *cis*-Vinylstannanes **4**

entry	vinyltelluride	vinylstannane	yield (%) ^a
1			46%
2			68%
3			60%
4			50%
5			68%
6			61%
7			40%
8			64%
9			50%

^a Isolated yields.

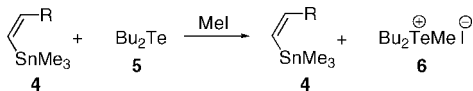
Vinylstannanes **4** were accompanied by small amounts of the corresponding protonated alkenes, which were readily removed under vacuum. The removal of dibutyltelluride **5**, however, proved very difficult by distillation or by column chromatography of the nonpolar and/or low molecular weight compounds (i.e., Table 2, entries 4–7, 9¹⁴). Hence, an efficient method for separation of nonpolar *cis*-vinylstannanes **4** from dibutyl telluride **5** was required. Capitalizing on the fact that dibutyl telluride **5** reacts with methyl iodide to give methyl di-*n*-butyltelluronium iodide **6** under catalyzed (AgNO₃, AgBF₄)¹⁵ and noncatalyzed conditions,¹⁶ excess methyl iodide (6 equiv) was added to the reaction mixture after

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(14) Purification by silica column chromatography was achieved; however, the general procedure should be followed for ease.

trimethyltin chloride. This workup additive reacted selectively with the dibutyl telluride **5** affording telluronium salt **6**, which could be readily separated from the *cis*-vinylstannane **4** by trituration followed by column chromatography (Scheme 2).

Scheme 2. Removal of Dibutyl Telluride From **4**



In conclusion, developing new methods that access *cis* double bonds selectively for application to the construction

of conjugated *cis* double bond systems is of considerable importance, for example, construction of *cis*-configured 1,3,5-trienes for 6π -electrocyclization.^{17,18} This body of work demonstrates that access to *cis*-vinylstannanes containing oxygen functionality is now readily achievable allowing further utilization of the Stille reaction to incorporate *cis* double-bond conjugation.

Acknowledgment. We thank the University of Queensland for financial support.

Supporting Information Available: Copies of ^1H and ^{13}C NMR spectra for all compounds, including experimental conditions and characterization data. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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