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Palladium-Catalyzed 1,4-Difunctionalization of Butadiene to Form Skipped Polyenes

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Supporting Information Placeholder

ABSTRACT: A palladium-catalyzed 1,4-addition across the commodity chemical 1,3-butadiene to afford skipped polyene products is reported. Through a palladium $\sigma \rightarrow \pi \rightarrow \sigma$ allyl isomerization, two new carbon-carbon bonds are formed with high regio- and trans stereochemical selectivity of the newly formed alkene. The utility of this method is highlighted by the successful synthesis of the ripostatin A skipped triene core.

The four-carbon feedstock, 1,3-butadiene, is a product of petroleum cracking on a >10 million ton scale annually, which makes it an especially attractive candidate for converting into more complex building blocks for synthesis. In this regard, we envisioned using this simple and symmetric diene² in a similar approach to that of the venerable Diels-Alder reaction by forming carbon-carbon bonds at both termini with the preservation of a single alkene in the product. In contrast to cycloadditions of 1,3-butadiene with dienophiles, we wanted to controllably form the carbon-carbon bonds using two distinct reaction partners through a 1,4difunctionalization process (Figure 1a). This would efficiently generate significant molecular complexity from a simple feedstock. In considering possible coupling partners, a specific goal was to ultimately form skipped diene and triene products as these are common motifs in various natural products³⁻¹⁵ and often challenging to prepare.¹⁶⁻¹⁹ Herein we present the development of a Pd-catalyzed 1,4-addition of vinyl electrophiles and boronic acid derivatives across 1,3butadiene to access these important structural motifs highlighted by the synthesis of the skipped triene core of ripostatin A.3

To accomplish a 1,4-difunctionalization of butadiene using a three-component coupling strategy, several important selectivity issues needed to be considered.²⁰⁻²¹ We recently reported a three-component coupling of substituted 1,3dienes, vinyl triflates, and boronic acids wherein selective formation of the 1,2-addition product is observed (Figure ib).^{21d} The use of vinyl triflates is proposed to account for the high selectivity of the three-component coupling product rather than either the Heck or Suzuki products presumably due to the cationic nature of intermediates along the reaction path. The use of these and related reagents is an obvious starting point for reaction development. Achieving a 1,4addition rather than a 1,2-addition across butadiene after oxidative addition to form **A** requires a $\sigma \rightarrow \pi \rightarrow \sigma$ isomerization of the allyl complex²² (Figure 1c, $\mathbf{A}\rightarrow\mathbf{B}\rightarrow\mathbf{C}$) with subsequent cross-coupling of **C** with a boronic acid derivative. A key previous report suggests that this isomerization is facile (Figure 1d). Specifically, a systematic study of substituent effects in the hydroarylation of terminal 1,3-dienes, which proceeds through a similar π -allyl species, reveals a linear free-energy relationship between Charton steric parameters and the logarithm of the ratio of 1,2 and 1,4-regioisomers.^{21e} If 1,3-butadiene was employed, extrapolation of the Charton relationship would achieve >20:1 selectivity for the 1,4-addition reaction. However, the use of a Pd-Ar rather than a Pd-H species to initiate the alkene functionalization, as well as the likely requirement of different reaction conditions than those used in the hydroarylation reaction, makes the outcome of this reaction less predictable.





Figure 1. Proposed 1,4-difunctionalization of 1,3-butadiene with vinyl triflates and boronic acids

To explore the possibility of a 1,4-difunctionalization of 1,3butadiene, vinyl triflate 1 and phenyl boronic acid 3 were chosen for reaction optimization (Table 1). The initial reaction conditions tested were those found optimal for the

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Table 2. Scope of 1,4-vinylarylation of 1,3-butadiene



All yields and ratios represent the average of two experiments on 0.5 mmol scale. E:Z ratios determined by ¹H NMR. Yields represent a mixture of stereo and regioisomers. ^a R⁴ = Tf. ^b R¹ = Nf. ^c Ratio of **7:8** as determined by ¹H NMR.

Table 1. Optimization of three-component coupling with 1,3-butadiene

1 x equiv	OTf + 2 y equiv	+ PhB(C 3 1.5 ec	3 mol% DH) ₂ <u>1.7 e</u> DMA 55 °	Pd₂(dba)₃ equiv KF (0.05 M) C, 16 h		4 product
_					5 1,2-pr	oduct
Entry	Conc. (M)	х	у	% Yield ^a	4 : 5 ^b	$E: Z^b$
1	0.05	1.0	balloon	81	2.4	88:12
2	0.05	1.0	1.0 ^c	50	12.6	91:9
3	0.2	1.0	1.0 ^c	79	15.0	91:9
4	0.05	1.5	1.0 ^c	71	16.7	90:10
5	0.2	1.5	1.0 ^c	75	>20	92:8
6 ^d	0.2	1.5	1.0 ^c	73	14.3	92:8

^aIsolated yield. ^bDetermined by ¹H NMR. ^c15 wt% solution **2** in hexanes used. ^dNonaflate used in place of triflate.

previously reported three-component coupling.^{21d} While the three-component coupling products were formed in an excellent yield, the product distribution was suboptimal with a 3:1 ratio of the desired 1,4-addtion product 4 to the 1,2addition product 5 as an inseparable mixture (entry 1). Initially, butadiene was introduced in the gaseous phase via a balloon, where the stoichiometry cannot be easily controlled. Therefore, we moved to a standard, commercial solution as the source of 1,3-butadiene (15 wt% in *n*-hexane). Upon this change, a significant improvement in selectivity for the 1,4addition product is observed albeit at the cost of yield. An origin for this improvement may be inhibition of $\sigma \rightarrow \pi \rightarrow \sigma$ isomerization when 1,3-butadiene is in substantial molar excess as it may act as a ligand on Pd. Modest concentration deviations of the diene had negligible effects.²³ Further improvement of both selectivity and yield is observed upon either performing the reaction at higher overall concentration (entry 3) or adding an excess of the vinyl triflate (entry 4). Combining these changes provides an optimized procedure where a >20:1 selectivity for the 1,4-addition product and a 92:8 ratio for the desired trans alkene is observed (entry 5). It should be noted that some of the enhancement of the observed selectivity is from consumption of the undesired terminal alkene product 5 with the excess vinyl triflate. Therefore, an improvement of selectivity and ease of purification comes with a cost of overall yield. Finally, the vinyl triflate can be replaced with the more economical vinyl nonaflate,²⁴ which produces nearly identical results (entry 6).

Using the optimal conditions, the scope of this 1,4vinylarylation of 1,3-butadiene was assessed, which was initially focused on a variety of substituted arenes and heterocyclic vinyl triflates/nonaflates resulting in excellent selectivity for the 1,4-addition product. Electronic effects of the aryl boronic acid had little influence on product yields (7a-f), although a lower regioselectivity was observed for the more electron rich aryl group installed in **7c**. Examining the steric influence of o-tolyl boronic acid, 7g, resulted in good selectivity and good yield. When larger arenes, including several heteroaromatic groups, were examined, the yield and selectivity of the process were not significantly influenced (7h-k). Unfortunately, boronic acids incorporating more Lewis basic heteroatoms (e.g., pyridine) fail to yield the threecomponent coupling product in an appreciable amount. Acyclic Z-vinyl triflates derived from β-dicarbonyls were successfully used, yielding 7l and 7m without isomerization of alkene geometry suggesting that a wide variety of skipped dienes can be accessed using this method. Finally, the use of a simple *E*-vinyl triflate (6n) resulted in the desired product (7n), again, without loss of stereochemical integrity originating from the vinyl triflate (Eq. 1). It should be noted that the internal alkene derived from butadiene is formed in a >8:1 ratio favoring the trans configuration in all cases, which is consistent with other difunctionalization reactions of this substrate. 20k,22c,h-j

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Encouraged by the successful vinylarylation threecomponent coupling reactions, we sought to explore a threecomponent vinylvinylation of 1,3-butadiene to form skipped trienes (Table 3). A styrenyl boronic acid can be coupled with a cyclic and an acyclic boronic acid with 1,3-butadiene to effectively form skipped trienes 9a and 9b in modest to good yields. A simple acyclic vinyl boronic is also an effective substrate in the synthesis of the skipped triene **9c**. The use of a considerably more complex organometallic coupling partner requires the use a higher catalyst loading to prepare od. In this case, the pinacol boronic ester was used as it was more easily accessed (9d-9e). In these cases, the alkene geometry in both the vinyl triflate and the boronic ester remain intact. Additionally, the alkene derived from 1,3-butadiene is again formed in the trans configuration suggesting a trans σallyl intermediate as depicted in C in Figure 1. The selectivity for 1,4-addition relative to 1,2-addition is diminished relative to the vinylarylation process but the 1,4-addition product of the skipped dienes can be accessed by re-exposing the mixture to the reaction conditions, which selectively consumes the terminal alkene.

Table 3. Scope of 1,4-vinylvinylation of 1,3-butadiene



All yields and ratios represent the average of two experiments on 0.5 mmol scale. Yields represent a mixture of stereo and regioisomers. ^a ratio of **9:10** and *E:Z* ratio determined by ¹H NMR of the crude reaction mixture. ^b Using the pinacol boronic ester and 6 mol% Pd₂(dba)₃

To showcase the utility of this method, the skipped diene core of ripostatin A, a bacterial RNA-polymerase inhibitor,³ was targeted. To our delight, the desired product (**9e**) was formed with the appropriate display of the skipped triene core and necessary functional groups for further manipulation. This example effectively illustrates the synthetic potential of this method by incorporating a simple feedstock chemical between two relatively complex fragments.

In summary, we have developed a new method for the efficient access to skipped polyenes. This process takes advantage of the simplicity of 1,3-butadiene and the ability to selectively and independently functionalize both terminal alkenes. Additionally, the reaction proceeds with high stereochemical integrity of the alkenes formed wherein the geometry of the starting materials is relayed into the product and the alkene fashioned from 1,3-butadiene is mainly of the trans geometry. The scope of the method is highlighted by the synthesis of the skipped triene core structure of ripostatin A in only three linear steps from simple starting materials. Many mechanistic questions arise when considering substituted terminal 1,3-dienes where more highly functionalized skipped polyene products can be envisioned. Future work is focused on examining these effects and expanding the scope of coupling partners.

ASSOCIATED CONTENT

Supporting Information. Experimental procedures and characterization data for new substances. This material is available free of charge via the Internet at http://pubs.acs.org.

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	1,4-difunctionalization of b	utadiene				
HO + Pd(0), KF, DMA 55 °C, 16 h FtO_{PC} OEt						
		skipped triene fragment of ripostatin A				
	- V VOEt					