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Operationally Simple and Highly (*E*)-Styrenyl-Selective Heck Reactions of Electronically Non-Biased Olefins

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Abstract

Simple, mild, and efficient conditions are reported for a Pd^0 -catalyzed Heck reaction that delivers high yields and selectivity for (*E*)-styrenyl products using electronically non-biased olefin substrates bearing a range of useful functionality. Preliminary mechanistic studies demonstrate that the σ -donating DMA solvent is crucial for high selectivity. Further studies suggest that the catalyst distinguishes between β -hydrogens based on their relative hydridic character, which is in contrast to previously reported Pd^{II} -catalyzed oxidative reaction conditions.

The Heck reaction, or palladium-catalyzed substitution of a vinylic C-H bond for an arene, is a transformation utilized extensively in synthesis despite significant limitations.¹ Arguably, the most restrictive limitation is the required use of an electronically biased olefin, such as a styrene or α,β -unsaturated carbonyl, in order to deliver a single arylated alkene isomer. In the absence of substrate bias, a mixture of olefin products arises under both typical Pd⁰-catalyzed, and most oxidative Pd^{II}-catalyzed,² Heck reaction conditions. This mixture of products likely results from Pd^{II}- σ -alkyl intermediate **A** undergoing indiscriminate β -hydride elimination with either H_S or H_A (eq 1).³ Additionally, there exists no generally applicable set of Heck conditions; parameters typically must be optimized for a specific desired reaction.^{1b}



(1)

Recently, we reported a Pd^{II}-catalyzed oxidative Heck reaction capable of delivering a variety of (*E*)-styrenyl products from electronically non-biased terminal alkene substrates, where the high selectivity observed was attributed to catalyst control (eq 2).⁴ Specifically, preliminary mechanistic studies suggested that the unusually high selectivity resulted from a cationic palladium catalyst stabilized by a strongly σ -donating ligand. It appears that, when undergoing β -hydride elimination from **A**, this metal center is capable of distinguishing between H_A and H_S based on relative C-H bond strength. This surprising result, which suggests that the hydrogen leaving in the selectivity-determining step may have partial radical character, has engaged our interest in this fundamental organometallic process.

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Supporting Information Available: Optimization data, experimental procedures, and characterization data. This material is available free of charge via the Internet at http://pubs.acs.org

While the aforementioned report significantly improved the substrate scope of the oxidative Heck reaction, it requires catalyst synthesis, a three-fold excess of the arene organometallic, and a catalytic amount of copper salts to proceed with high yield and excellent selectivity. Additionally, these conditions are incompatible with some common functional groups, such as carboxylic acids and nitriles. Guided by our previous mechanistic studies, we report herein the development of a complementary and operationally simple Pd^0 -catalyzed Heck reaction using electronically unbiased terminal alkenes, which allows for the introduction of diverse functional groups and is highly selective for (*E*)-styrenyl products.

(2)

In order to achieve high selectivity, it was hypothesized that mimicking the electrophilic nature of the proposed Pd^{II} - σ -alkyl intermediate **A** capable of distinguishing between β -hydrogens would be essential, and that the elevated temperatures typically required for oxidative addition should be avoided. As such, it was envisioned that aryl diazonium tetrafluoroborates, pioneered by Matsuda, would be well-suited as the oxidant in light of both their tendency to undergo facile oxidative addition⁵ and the non-coordinating nature of the BF₄ counterion of the resultant aryl-Pd^{II} complex.⁶ An initial experiment utilizing this oxidant resulted in a modest yield and relatively low selectivity (Table 1, entry 1).⁷

A control experiment (entry 2) excluding the *N*-heterocyclic carbene ligand delivered higher selectivity, prompting the elimination of this additive in future experiments. A significant disparity between consumption of substrate and product yield dictated the careful monitoring of this ratio over time, revealing reaction completion after only 15 minutes and further consumption of the product if the reaction mixture is not quenched (entries 2 vs 3).⁸ Decreased catalyst and arene loadings resulted in improved yield and selectivity (entries 4 and 5), while entry 6 confirms that palladium is required. Of particular note, the use of MeOH or MeCN, solvents commonly employed in Matsuda-Heck reactions,⁵ results in very poor selectivity (entries 7 and 8).

The optimized conditions were evaluated for compatibility with a variety of substrates that performed well under oxidative conditions.⁴ For example, substrates bearing an ester (Table 2, entry 1), a ketone (entry 2) and a silyl ether (entry 3) all proceed with excellent yield and selectivity. A free homoallylic alcohol (entry 4) was compatible and, more surprisingly, an allylic acetate is an excellent substrate that does not undergo oxidative addition under these conditions (entry 5).⁹ A doubly protected allylic amine is a good substrate,¹⁰ as is a distal free alcohol (entries 6 and 7). The Pd^{II}-catalyzed conditions are incompatible with carboxylic acids,⁴ but gratifyingly, **3h** and **3i** were prepared in good to excellent yield using the present system. Of note, the heteroatom-free substrate dodecene gives excellent results, providing strong evidence that the selectivity observed under these conditions is catalystcontrolled rather than dependent on substrate chelation (entry 10).¹¹ A substrate bearing a nitrile (also incompatible with oxidative conditions) is arylated with either a phenyl group or an arene bearing a methyl ester (entries 11 and 12). A free 1,2-diol reacts cleanly when installing a phenyl group, but the yield suffers when installing an arene with more steric bulk (cf. entries 13 and 14). Free phenols and aryl chlorides are compatible with these conditions (entries 15 and 16), and submission of a free allylic alcohol gives the desired product, but the reaction proceeds more slowly and the yield is diminished due to the formation of a ketone by-product (entry 17).¹² Challenging nitro- and iodo-substituted arenes are also compatible under the conditions described, providing valuable handles for further functionalization (entries 18 and 19).¹³ The reaction proceeds rapidly and in high

yield using α -methylstyrene as the substrate, but unfortunately a mixture of olefin isomers is observed (entries 20 and 21).¹⁴ Finally, a highly enantiomerically enriched substrate, that may be susceptible to racemization, suffers no erosion of enantiomeric excess (entry 22).

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While evaluating the scope of this transformation, it was observed that the reaction is highly exothermic raising concerns that the selectivity of a reaction performed on larger scale may suffer. Therefore, on 5 mmol scale, **1a** was subjected to the conditions described to prepare **3a**, except that the catalyst loading was decreased to 2 mol % and the reaction was performed at -15 °C (eq 3). Under these conditions, **3a** was obtained in comparable yield and improved selectivity for the (*E*)-styrenyl product after 5.5 h.

Having established that the active catalyst described in this report indeed exhibits a unique preference for (*E*)-styrenyl products, we wished to gain an understanding of how, exactly, the catalyst imparts selectivity. The mechanistic origin of selectivity under oxidative conditions was previously determined by submitting β , γ -unsaturated ester **1t** to Pd^{II}- catalyzed conditions using a variety of electronically disparate arene sources; an experiment designed to probe the selectivity-determining β -hydride elimination step where a significant substituent effect was observed.⁴ Interestingly, the product distribution measured under Pd⁰- catalyzed conditions reveals no clear trend, with styrene products favored in similar ratios regardless of the electronic nature of the arene (Table 3).¹⁵

A more sensitive mechanistic experiment was designed, whereby allyl benzene was submitted to the optimal conditions with a variety of electronically disparate aryl diazonium salts. In this product-partitioning experiment, the catalyst must distinguish between two benzylic hydrogens, which presumably differ only by virtue of the arenes immediately adjacent (Figure 1). The linear free energy relationship observed suggests that the origin of selectivity under these conditions is related to the ability of the metal center in intermediate **X** to distinguish between β -hydrogens as a function of their relative *hydridic* nature.^{16,17} Therefore, submission of electron-rich diazonium salts results in relatively more **5**_{Ar}, due to the greater ability of the newly installed arene to stabilize partial positive charge developing during β -hydride elimination. Consistent with this proposal, submission of partial positive charge by the newly installed arene. These results contrast markedly with those observed under oxidative Pd^{II} catalysis,⁴ suggesting an interesting mechanistic complementarity along with the more intuitive counterpart arising from the differing oxidation states of the two precatalysts.

In conclusion, we have developed simple and efficient conditions for a Pd⁰-catalyzed Heck reaction that delivers high selectivity for (*E*)-styrenyl products in the absence of substrate bias. This reaction is compatible with a greater range of functional groups than the related Pd^{II} system and utilizes a commercially-available catalyst. Additionally, the reaction requires no base, elevated temperatures, nor additional oxidant. For most substrates evaluated, the reaction is completed rapidly, but the rate is retarded when using substrates with allylic coordinating groups. Some functional groups are incompatible,¹⁰ but it is reasonably easy to predict these, providing another advantage over other Heck protocols, which can require optimization for each substrate. Initial mechanistic experiments suggest that the selectivity observed is highly sensitive to the identity of the solvent. A linear free energy relationship probing product distribution as a function of the electronic nature of the

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(3)

introduced arene suggests that this catalyst selects between β -hydrogens based on their hydridic nature, which is in contrast with the Pd^{II} system previously reported. This reactivity manifold has inspired further experiments designed to probe the factors that dictate selectivity in β -hydride elimination from related organometallic intermediates. Investigations in this pursuit are currently underway in our laboratory.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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- 3. Additional byproducts can arise from Heck insertion in the opposite orientation resulting in terminal olefin products, and internal olefin isomeric products arising from Pd-H migration. Z-olefin isomers may also be observed.
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- 6. The pK_A of the conjugate acid provides a commonly used estimation of the coordinative abilities of counterions in catalysis; the pK_A of HBF₄ is -5.
- 7. Throughout the paper, when isomers are detected they arise either from migratory insertion in the opposite direction to give terminal olefin products, or from olefin migration via non-selective β-hydride elimination. (*Z*)-Olefin isomers were not observed.
- 8. A minor byproduct arises from a subsequent Heck reaction of the resultant styrene utilizing the remainder of the aryldiazonium salt. The desired product is also believed to be oligomerized in the

reaction mixture overtime, likely due to the equivalent of HBF₄ generated in the reaction. See supporting information for details.

- 9. Allylic acetates are known to be displaced by Pd^0 to form π -allyl complexes. For a review, see: Trost BM. Angew. Chem. Int. Ed. 1989; 28:1173.
- 10. More basic nitrogen functionality, such as a trialkyl amine-containing substrate, was incompatible with these conditions. Other incompatible functionality included quinoline and indole arenes. See supporting information for further details.
- For examples of chelation-controlled Heck reactions see: (a) Filippini L, Gusmeroli M, Riva R. Tetrahedron Lett. 1993; 34:1643–1646. (b) Kang SK, Lee HW, Jang SB, Kim TH, Pyun SJ. J. Org. Chem. 1996; 61:2604–2605. [PubMed: 11667088] (c) Olofsson K, Sahlin H, Larhead M, Hallberg A. J. Org. Chem. 2001; 66:544–549. [PubMed: 11429827] (d) Buezo ND, Rosa JC, Priego J, Alonso I, Carretoero JC. Chem. Eur. J. 2001; 7:3890–3900. (e) Delcamp JH, Brucks AP, White MC. J. Am. Chem. Soc. 2008; 130:11270–11271. [PubMed: 18671350]
- 12. Ketones like **6** are usually the major product observed using free allylic alcohols under Heck conditions; see above references.

- 13. It is important to note that while most aryl diazonium salts submitted to this reaction performed well, there are several examples of synthetically inaccessible aryl diazonium salts. See supporting information for details.
- 14. See supporting information for details.
- 15. Interestingly, allylic products 5_{Allyl} were typically favored under oxidative Heck conditions, while they were the minor products under these conditions regardless of arene. See supporting information for details.
- For examples of Hammett analyses of other Heck reactions see: (a) Benhaddou R, Czernecki S, Ville G, Zegar A. Organometallics. 1988; 7:2435–2439. (b) Fristrup P, Le Quement S, Tanner D, Norrby PO. Organometallics. 2004; 23:6160–6165.
- 17. The modest slope of the Hammett plot may explain why no trend was apparent when attempting to construct a similar Hammett plot using a β , γ -unsaturated ester as the substrate. It is likely that the hydridic nature of benzylic hydrogens and those α to an ester are significantly different.



Figure 1. Hammett plot.

Table 1

$\hat{ ho}$			+ PhN ₂ BF	-4 X mol % Pd ₂ dt		^H
	1 a	~	4a (y edu	iv)	5	3a 3a
entry	×	×	time	% conversion ^a	% yield ^a	selectivity ^{a,b}
1^c	S	1.5	16 h	66<	68.4	3.8
2	5	1.5	16 h	66<	43.3	6.8
3	5	1.5	15 min	66<	62.6	7.1
4	ю	1.5	15 min	66<	86.2	7.5
5	ю	1.1	15 min	>99	66<	10.7
9	0	1.1	15 min	3.6	0	
pL	ю	1.1	15 min	66<	20	0.2:1
86	ю	1.1	15 min	98.1	15	0.3:1

⁴Conversion and yield calculated by comparing starting material and product peak integration to integration of internal standard using corrected GC analysis. Yields refers to the sum of all product isomers.

 b Selectivity refers to the ratio of (*E*)-styrene to all other isomers.

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 c 12.5 mol % l $^{i}\mathrm{Pr}$ carbine added.

 d MeOH used as solvent.

 e MeCN used as solvent.

Scope of the Heck reaction.

	$\sum_{i} \sum_{i \in I \text{ sum}} \frac{\lim_{i \in I \text{ sum}} \sum_{i \in I \text{ sum}} \sum_$		
entry	product	time	% yield ^a
1	ي 3a ب	20 min	97 ^b
2	3h Ph	20 min	89
	ox 30		
3	$\begin{array}{c} Ph^{\prime} & \checkmark & \checkmark \\ X = TBS \ \mathbf{3c} \end{array}$	16 h	87
4	X = H 3d	1.5 h	72
5	OAc	16 h	87 ^{<i>c</i>}
6	3e Ph Cbz N Ph Boc 3f	16 h	96
	R M R'		
7	$\mathbf{R} = \mathbf{CH}_2\mathbf{OH}, \mathbf{R'} = \mathbf{H} \mathbf{3g}$	40 min	77 ^b
8	$\mathbf{R} = \mathbf{CO}_2\mathbf{H}, \mathbf{R}' = \mathbf{H} \mathbf{3h}$	2 h	95 ^b
9	$R = CO_2H$, $R' = 3,5$ -dimethoxy 3i	2 h	69 ^b
10	R = CH ₂ CH ₃ , R' = H 3j	20 min	77b
	NC R'		
11	$R' = H \mathbf{3k}$	3 h	96
12	R' = CO ₂ Me 3I OH R'	3 h	98
	но		
13	$\mathbf{R}' = \mathbf{H} \mathbf{3m}$	2 h	83
14	R' = Me 3n R'	2 h	66 ^C
15	R = OH, R' = H 30	1.5 h	98
16	$R = CI, R' = OMe \mathbf{3p}$	1.5 h	98

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^{*a*}Yields are averages of two experiments performed on a 0.5 mmol scale, and are the sum of all product isomers. The selectivity for (*E*)-styrene >20:1 unless otherwise noted.

^bSelectivity for (*E*)-styrene was 10:1.

^cUsing 5 mol % Pd2dba3 and 1.5 equiv ArN2BF4.

^dIsolated 44% keton product.

^eObtained ~1:1 mixture of olefin isomers.

Table 3

Product distribution using a β , γ -unsaturated ester.

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Ar	$5_{\rm Sty}/5_{\rm Allyl}$	Ar	5 _{Sty} /5 _{Allyl}				
4-H	6.03	4-CO ₂ Me	7.69				
4-NO ₂	6.11	4-Br	6.47				
4-OMe	7.87	4-F	6.12				