Rhodium-Catalyzed Enantioselective Hydroamination of Alkynes with Indolines

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Supporting Information

ABSTRACT: The hydroamination of internal alkynes via tandem rhodium catalysis gives branched N-allylic indolines with high regio- and enantioselectivity. An acid switch provides access to the linear isomer in preference to the branched isomer by an isomerization mechanism. Mechanistic studies suggest formation of an allene intermediate, which undergoes hydroamination to generate allylic amines instead of the enamine or imine products typically observed in alkyne hydroaminations. Constructing C−N bonds with high regio- and enantiocontrol represents an important challenge given the occurrence of amines in agrochemicals, fine chemicals, and pharmaceuticals. Transition-metal catalysis has enabled the coupling of amines with allylic electrophiles to provide either the branched or linear isomers (Figure 1a).1 The hydroamination of alkynes has emerged as an attractive approach for C−N bond formation because of its high atom economy.2 The majority of metal catalysts investigated provide enamine or imine products (Figure 1b). In contrast, Yamamoto demonstrated a Pd-catalyzed hydroamination3 to yield allylic amines, with preference for the linear isomers (Figure 1c).3 While a promising approach, no intermolecular variants have been shown to access the corresponding branched isomers. Herein, we demonstrate a Rh-catalyzed alkyne hydroamination4 that allows access to either the branched or linear isomers, with high regiocontrol by the choice of carboxylic acid additive used (Figure 1d). This communication showcases the first enantioselective intermolecular hydroamination of alkynes.

Late transition-metal hydrides (e.g., Ru, Rh, Ir, Pd) are known to convert alkynes to π-allyl metal intermediates. Since Yamamoto’s report,3 Ishii,5 Krische,6 and our group7 have used metal-hydride catalysis to couple internal alkynes with alcohols or aldehydes to generate C=C bonds. The Breit group pioneered the use of terminal alkynes to form allylrhodium intermediates that undergo C=O and C=S bond formations.8 Encouraged by this emerging concept, we focused on the functionalization of alkynes to generate C−N bonds with both regio- and enantiocontrol.

To test our hypothesis, we chose the coupling of indoline 1a and 1-phenyl-1-propyne 2a as model substrates (Table 1). Commercially available metal hydride complexes (e.g., RuHCl(CO)(PPh3)3, RhH(CO)(PPh3)3, or IrH(CO)(PPh3)3) did not promote hydroamination. In contrast, a mix of [Rh(COE)2Cl]2/L1 and benzoic acid8 afforded the desired branched indoline 3aa in trace amounts (4% yield, entry 1).9 More electron-rich and bulky ligands L2 and L3 improved the reactivity and gave high regioselectivities (3aa/4aa) and moderate enantioselectivities (47−55% ee, entries 2−3). Among the bidentate phosphines bearing point chirality that we studied, (S,S)-BDPP L6 gave the most promising regio- and enantioselectivity (87% ee, entry 6). We assume this ligand shows enhanced reactivity and selectivity due to its ability to promote isomerization and hydroamination in parallel. In the absence of acid, no hydroamination was observed. By investigating other acid additives, we found higher enantioselectivity using m-xylic acid (90% ee, entry 7).

With this protocol in hand, we explored the hydroamination of alkyn 2a with various amines 1 (Table 2). Good yields (72% and 81%) and high enantioselectivities (90% ee) were obtained using indolines with varying substituents at the 4-position (3ba and 3ca). Substrates bearing fluoro, chloro, and bromo at the 5-position were transformed into allylic amines with similar efficiency (3da−3fa). A slight decrease in enantioselectivity was observed with indolines containing electron-donating substituents at the 5- or 6-position (3ga vs 3ha, 3ia). In contrast, an aliphatic amine, such as morpholine, showed poor reactivity.
under this protocol (8% yield, 3a). An improved yield (47%, 3a) could be obtained by using more m-xyllic acid (50 mol %). Aniline 1k was coupled with alkyne 2a in good selectivity (>20:1 b/l, 76% ee, 3ka) without bisallylation. In all cases, we observed excellent regioselectivity for the branched isomers in preference to the linear isomers.

Next, we examined the scope of alkynes (Table 3). The use of electron-rich 1-aryl-1-propynes 2b–c gave branched allicic amines in >20:1 regioselectivities, 70–73% yields, and 82–91% ee (entries 1–2). Although a decrease in regioselectivity (from 20:1 to 3:1) was observed with electron-deficient alkynes 2d–g, high enantioselectivities (86–92%) were still achieved (entries 3–6). The coupling of indole with meta-substituted 1-phenyl-1-propynes 2h–j gave a slight increase in enantioselectivity (90–94%, entries 7–9). This protocol tolerated a heterocycle-substituted alkyne 2k (80% yield, entry 10). However, low reactivity (15% yield) and selectivity (27% ee) were obtained using the internal alkyne 2l bearing an aliphatic group (entry 11).

The hydroamination of terminal alkyne 2m gave the same product as that of 2a, with similar regio- and enantioselectivity but lower reactivity (entry 12). Poor enantioselectivity (7%) was observed in the hydroamination of 1-octyne 2n (entry 13).10 These results suggest that an aromatic substituent on alkyne 2 is critical for high yield and selectivity.

A one-pot protocol was developed, using Rh-catalyzed asymmetric hydroamination and subsequent oxidative dehy-droromatization of indolines 3, to provide facile access to N-allylic indoles11 5aa (90% ee) and 5ba (85% ee) (eq 1). The absolute configuration of 5aa was determined to be (S) by comparison of its optical rotation with literature data.11c,d,12
During our optimization studies, we found that using phthalic acid (benzene-1,2-dicarboxylic acid) instead of \( m \)-xylylic acid gave linear allylic amines with high regioselectivity (>20:1). A number of amines are suitable coupling partners (4aa, \( 4a \)−\( 4o \), Table 4). By tuning the stoichiometry of aniline \( 1k \) (PhNH\(_2\)) and \( 2 \)a, either monoallylated product \( 4ka \) or bisallylated product \( 4ka' \) could be formed exclusively. This protocol can be applied to 1-phenyl-1-butyne \( 2n \), which bears an ethyl group, to generate allylic amine \( 4ao \).

On the basis of literature\(^{1,8} \) and our observations, we propose a pathway involving tandem Rh-catalysis (Figure 2). The oxidative addition of carboxylic acid with a Rh(I) precursor generates a rhodium(III)-hydride species. The insertion of alkyne \( 2a \) into the Rh(III)−H gives Rh-vinyl intermediate \( A \), which subsequently undergoes \( \beta \)-hydride elimination to form intermediate allene \( 6 \) and regenerate the Rh(III)−H species. Reinsertion of the terminal allene \( 6 \) into Rh(III)−H yields chiral \( \pi \)-allyl rhodium complex \( B \). Two competing pathways can be considered for formation of product \( 3 \).\(^{1,4} \) For path \( A \), a ligand exchange of amine \( 1 \) with the benzoate \( X \) on complex \( B \) will give \( \pi \)-allyl rhodium species \( C \), which can undergo reductive elimination to yield allylic amine \( 3 \). In path \( B \), amine \( 1 \) undergoes nucleophilic attack at the carbon center to deliver allylic amine \( 3 \).

To support the proposed allene intermediate, phenyllallene \( 6 \) was prepared and then subjected to coupling with indoline \( 1a \) under otherwise standard conditions (eq 2).\(^{15} \) Only a trace amount of the expected allylic amine was generated using a high concentration (~1.20 M) of allene \( 6 \). However, the allene was transformed into the expected allylic amine when using a lower concentration (~0.16 M). Our results support the feasibility of allene \( 6 \) as the intermediate and suggest that high concentrations of allene inhibit catalysis. Using alkyens as allene surrogates can enable transformations that would be sluggish with allene precursors.\(^{6a,7} \)

When the hydroamination was performed with deuterated alkyne \( 2a-d_3 \), the deuterium label was scrambled into the \( \alpha \)-, \( \beta \)-, and \( \gamma \)-positions of amine \( 3aa-d_n \) (eq 3). The observed incorporation of hydrogen at the \( \gamma \)-position of amine \( 3aa-d_n \) indicates the reversibility of \( \beta \)-hydride elimination during allene formation.

To better understand the regioselective switch (Table 2 vs 4), we isolated and subjected racemic \( 3aa \) to various conditions (Figure 3). No isomerization was observed in the absence of

![Figure 2. Tandem Rh-catalysis: proposed mechanism for hydroamination of alkyens.](image-url)
either Rh or acid. However, the branched isomer 3aa underwent isomerization to the linear product 4aa faster in the presence of a more acidic additive (e.g., m-xylene vs phthalic acid). Thus, the branched isomer is the kinetic product, which can be generated in high yield using m-xylene acid. In contrast, our phthalic acid protocol generates linear products by a thermodynamically controlled isomerization. Indeed, when we monitored the reaction profile for our linear-selective hydrosamination, we observed formation of the branched isomer at early time points followed by its conversion to the linear. Yudin observed a related isomerization of allylic amines under Pd-catalysis, whereby the kinetic (branched) product was favored by addition of DBU.16

Our Rh-catalyzed hydrosamination provides an atom economical synthesis of allylic amines that complements traditional allylic aminations, which require the use of leaving groups. Mechanistic studies support the in situ formation of an allene which undergoes hydrosamination to provide allylic amines, rather than the typically observed products of alkyne hydrosamination (e.g., imines and enamines). Phthalic acid promotes isomerization of the kinetically favored isomers to yield the more stable linear isomers. Future studies will focus on catalyst design to extend substrate scope and develop other variants.

**REFERENCES**


(9) For the absolute configuration of all other amines 3 were assigned by analogy.


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