

Rh₂(II)-Catalyzed Intramolecular Aliphatic C–H Bond Amination Reactions Using Aryl Azides as the N-Atom Source

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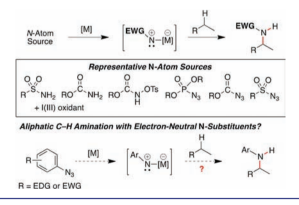
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Supporting Information

ABSTRACT: Rhodium(II) dicarboxylate complexes were discovered to catalyze the intramolecular amination of unactivated primary, secondary, or tertiary aliphatic C–H bonds using aryl azides as the N-atom precursor. While a strong electron-withdrawing group on the nitrogen atom is typically required to achieve this reaction, we found that both electron-rich and electron-poor aryl azides are efficient sources for the metal nitrene reactive intermediate.

The development of transition metal-catalyzed aliphatic C– H bond amination reactions that are stereoselective, use readily accessible starting materials and catalysts, and are environmentally benign continues to inspire the efforts of research groups around the world.^{1,2} While considerable progress has been made, the current methods are still limited by the oxidative conditions to form the nitrene and the requirement for strong electron-withdrawing groups on the nitrene (Scheme 1). The use of azides as the N-atom source

Scheme 1. Nitrogen Substituent Requirements for Aliphatic C–H Bond Amination



would address these limitations because no oxidant would be required and the only byproduct of the reaction would the environmentally benign N_2 gas.³ While azides have been used for a variety of N-atom-transfer reactions,^{2b,4} these transformations also require electron-withdrawing groups on the azide. We anticipated that a more general, complementary solution for aliphatic C–H bond amination might emerge if conditions were found to use electron-neutral aryl azides as the N-atom source. While we have reported a number of sp² C–H bond amination reactions using aryl azides,⁵ our mechanism studies suggest that C–N bond formation occurs via a 4π electron-5-atom electrocyclization.^{5b} In contrast, aliphatic C–H bond amination requires metal-catalyzed C–H insertion or Hatom abstraction mechanisms—processes that have remained elusive to control using aryl azides as the N-atom source.⁶

In search of the optimal conditions to achieve intramolecular aliphatic C–H bond amination using aryl azides, the reactivity of *o-tert*-butylaryl azide 1a toward transition metal complexes was examined (Table 1).⁷ This aryl azide is relatively thermally

Table 1. Development of Optimal Conditions

	Me Me MX _n CH ₂ (5 mol %) N ₃ PhMe, 120 °C additive			Me Me N a P.G.		
entry	catalyst	additive	conv, % ^a	yield, % ^b		
1	none	n.a.	0	0		
2	FeBr ₂	n.a.	0	dec ^c		
3	CuBr	n.a.	0	0		
4	CoTPP	n.a.	0	0		
5	$RuCl_3 \cdot nOH_2$	n.a.	0	0		
6	$[Ir(cod)OMe]_2$	n.a.	0	0		
7	$[Rh(cod)OMe]_2$	n.a.	10	0^{c}		
8	$Rh_2(O_2CC_7H_{15})_4$	n.a.	35	35		
9	$Rh_2(esp)_2$	n.a.	99	75		
10	$Rh_2(esp)_2$	Boc ₂ O	99	90		
11	$Rh_2(esp)_2$	Ac ₂ O	99	83		
12	$Rh_2(esp)_2$	Bz ₂ O	99	aniline		
13	$Rh_2(esp)_2$	Tf_2O	99	aniline		
and the second second by NMD and the second by the second se						

"As determined using ¹H NMR spectroscopy. ^bIsolated after silica gel chromatography. ^cAniline formed.

robust, with no reaction observed at 120 °C (entry 1).⁸ Exposure of **1a** to commercially available transition metal complexes known to catalyze N-atom-transfer reactions was met with limited success. Indoline **3a** was not observed in the presence of iron,⁹ copper,¹⁰ cobalt,^{2b,4a-c} ruthenium,^{2a,6,11} or iridium¹² complexes (entries 2–6). Partial conversion to indoline **2** was observed when rhodium octanoate was used (entry 8). We anticipated that the partial conversions resulted from catalyst decomposition, and we found that using more thermally robust $Rh_2(esp)_2^{13}$ improved both the conversion and yield of the process (entry 9). Examination of alternative solvents, concentration, and temperatures, however, did not

Received: February 15, 2012 Published: April 21, 2012 increase the yield, and control experiments revealed that oxidative decomposition of the indoline occurred during purification.

To improve the isolated yield of the amination reaction, in situ protection of the nitrogen atom was attempted. In line with our assumption, we found that the reaction yield was improved when the indoline was protected with either a Boc or Ac group (entries 10 and 11). The reaction outcome appeared to correlate with the pK_a of the acid byproduct of the protection reaction: aniline was produced when the stronger benzoic and triflic acids were produced (entries 12 and 13).¹⁴

Using these optimal conditions, the electronic and steric constraints of the aliphatic C–H bond amination reaction were investigated (Table 2). In contrast to existing amination

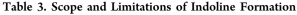
Table 2. Scope and Limitations of Indoline Formation

Me Me CH N ₃ H	Rh ₂ (esp (5 mol 9 PhMe, 12 Boc ₂ C		R Me Me				
entry	1	R	yield, % ^a				
1	1a	Н	84				
2	1b	OMe	63				
3	1c	Me	54				
4	1d	CH_2CH_2Ph	64				
5	1e	CHCHPh	70				
6	1f	Ph	54				
7	1g	3,5-OMe ₂ C ₆ H ₃	58				
8	1h	Br	73				
^{<i>a</i>} Isolated after silica gel chromatography.							

processes,^{2,4} our method does not require an electronwithdrawing group on the nitrogen: aryl azides bearing *para*electron-releasing, -neutral, or -withdrawing groups were converted to indolines. Illustrating the chemoselectivity of our process, olefins were tolerated as substituents (entry 5).

To further examine the scope of the transformation, the identity of the ortho-alkyl substituent was varied (Table 3). While replacing one of the methyl groups with an ester group did not diminish the yield (entry 1), substitution with hydrogen reduced the reaction efficiency (entry 2). Amination can be achieved at tertiary and secondary C-H bonds, although dehydrogenation of 5d occurred to afford indole. Dehydrogenation could be circumvented if an additional substituent was introduced at the benzylic position on the aryl azide. Submission of 4e-4k to reaction conditions produced indolines as single diastereomers (entries 5-11). In contrast, pyrolysis of aryl azide 4g was reported by Smolinsky to produce a 1:1 mixture of diastereomers.¹⁵ Although reaction with the methyl C-H bond in azide 4l could produce a six-membered ring, only amination of the methyl C-H bond was observed to afford indoline 51 as the sole product. While single diastereomers were obtained from substrates bearing ocyclopentyl or cyclohexyl groups, diminished stereoselectivity was observed with o-cycloheptyl-substituted aryl azide 4m (entry 13).

While indoline formation could occur through several different mechanisms,^{10a,16} our reactivity trends suggest that C–N bond formation occurs through N-atom transfer (Scheme 2). Coordination of the rhodium(II) carboxylate to either the α - or γ -nitrogen of the aryl azide produces 7.¹⁷ Extrusion of N₂ then forms the rhodium nitrene 8.¹⁸ While rhodium could

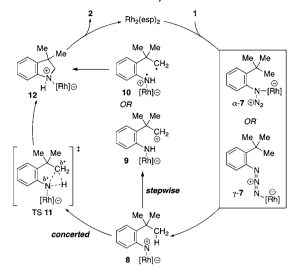


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R	γ	$\begin{array}{c} R & Rh_2(esp)_2 \\ (5 \text{ mol }\%) \\ H & PhMe, 120 \ °C \\ Boc_2O \end{array} R^-$	$\begin{bmatrix} & & \\ & $	6 Boc
entry	4	aryl azide	indoline	yield, % ^a
1	a	EtO ₂ C Me Me N ₃	EtO ₂ C Me	70
2	b	Me Me N ₃	H Me N Boc	20
3	с	Me Ph N ₃ H	Ne H	55 ^b
4	d	Me H N ₃	N H H	30 ^c
5	e	MeO	MeO H H H Boc	80
6	f	Me N3	Me H N H N H Boc	73
7	g	H	H H H H H H H H H H H H H H H H H H H	70
8	h	F ₃ C N ₃	F ₃ C H H H Boc	63
9	i		H N H Boc	85
10	j	Me N ₃	Me H H Boc	73
11	k	F ₃ C N ₃	F ₃ C N H Boc	82
12	1	Me Me N ₃	Me H N H Boc	86
13	m	CUC _{N3}	H N H Boc	63 d.r. 82:18

 a Isolated after silica gel chromatography. $^b20\%$ aniline observed. $^c30\%$ aniline observed.

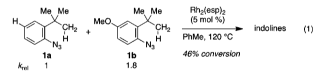
mediate a reversible one-electron oxidation to generate free nitrene,¹⁹ our current mechanistic hypothesis is that the nitrene remains metal bound for the C–H bond amination step since pyrolysis involving free nitrene is not diastereoselective.¹⁵ The mechanism of the amination step could be stepwise or concerted: hydride²⁰ or H-atom abstraction^{16a,21} (to form **9** or **10**) followed by recombination produces the C–N bond; alternatively this bond could be formed through the concerted

Scheme 2. Possible Mechanisms for Intramolecular Aliphatic C–H Bond Amination from Aryl Azides



insertion^{16c,22} of the metal nitrene into the proximal C–H bond via transition state **11**. Finally, the indoline is produced upon dissociation of the rhodium complex from **12**.

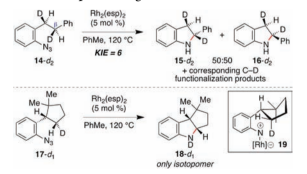
Insight into the initial steps of the catalytic cycle was provided by the reactivity of 4-substituted aryl azides toward reaction conditions (eq 1). Underscoring the difference



between our method for aliphatic C–H bond amination and others, we found that more electron-rich aryl azides (e.g., **1b**) were more reactive to the reaction conditions. The increased reactivity of **1b** relative to **1a** could be due to either preferred coordination of **1b** to $Rh_2(esp)_2$ or an accelerated N_2 extrusion from the resulting azide–metal complex 7.

To examine the nature of the C-H bond cleavage step, twolabeled aryl azides were examined (Scheme 3). We anticipated

Scheme 3. Isotope Labeling Studies



that the number of indoline diastereomers from $14-d_2$ would reveal if the mechanism of C–H bond amination was stepwise or concerted. If the reaction was concerted, insertion into either the β -C–H or β -C–D bond would produce only two products. In contrast, if a radical (or cation) was formed at the β -carbon, then scrambling of the C2-stereocenter could occur before recombination to form both 15- d_2 and 16- d_2 . In support of a stepwise C–H bond amination, two diastereomers of 2phenylindoline (dr 50:50) and an intramolecular kinetic isotope effect (KIE) of 6.7 were observed.²³ The magnitude of this isotope effect is significantly smaller than for reactions involving an H-atom abstraction by an aryl nitrene²¹ or an aryl metal nitrene^{6c} ($k_{\rm H}/k_{\rm D} = 12-14$), but larger than hydride shift reactions ($k_{\rm H}/k_{\rm D} \approx 2$ for Cannizzaro reaction and Meerwein– Ponndorf–Verley reduction).^{24–26} Smaller KIEs were observed at lower reaction temperatures, revealing that our amination reaction occurs above the isokinetic temperature and, as a consequence, is under entropic control.^{27,28} We found, however, that the spatial constraints of this reaction override these isotope effects: cyclopentanone-derived aryl azide 17- d_1 reacted preferentially with the *syn*-C–D bond to form 18 exclusively.²⁹

In conclusion, we have developed an efficient and diastereoselective rhodium(II)-catalyzed aliphatic C–H bond amination reaction that uses an aryl azide as the N-atom source. Our method distinguishes itself from previously reported aliphatic C–H bond amination reactions by not requiring a strong electron-withdrawing group on the nitrogen atom. The reactivity of stereospecific labeled aryl azides revealed that the amination reaction occurred stepwise with the *syn*-C–H bond. Our current aims are to more deeply examine the nature of the catalytic intermediates in this C–H bond amination reaction and to extend this newfound reactivity of aryl azides to the stereoselective synthesis of complex, functionalized *N*-heterocycles.

ASSOCIATED CONTENT

S Supporting Information

Experimental procedures and spectroscopic and analytical data for the products. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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