Rhodium-Catalyzed Oxidative C–H Allylation of Benzamides with 1,3-Dienes by Allyl-to-Allyl 1,4-Rh(III) Migration

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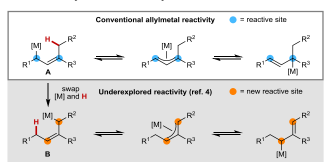
ABSTRACT: The Rh(III)-catalyzed oxidative C–H allylation of *N*-acetylbenzamides with 1,3-dienes is described. The presence of allylic hydrogens *cis*- to the less substituted alkene of the 1,3-diene is important for the success of these reactions. With the assistance of reactions using deuterated 1,3-dienes, a proposed mechanism is provided. The key step is postulated to be the first reported examples of allyl-to-allyl 1,4-Rh(III) migration.

INTRODUCTION

Allylmetal species are important intermediates in organic synthesis. 1,2 For example, π -allylmetal species are usually electrophilic, and can be intercepted by diverse nucleophiles in allylic substitutions. 1 On the other hand, σ -allylmetal species are usually nucleophilic, and can be employed in a huge range of allylations of π -electrophiles. 2 Numerous catalytic, diastereoselective, and/or enantioselective variants of these processes have also been reported. 1,2

A well-recognized feature of allylmetal reactivity is the often facile 1,3-transposition of the metal from one end of the allylic fragment to the other, which potentially enables new bond-forming reactions at either side (Scheme 1, top). Isomerizations of allylmetal species that open up reactions at sites beyond those resulting from conventional 1,3-allylic transposition would be highly enabling for reaction discovery.^{3,4,5} As part of a program in enantioselective rhodium-catalyzed additions of allylboron reagents to imines, 3,4,6 we have described the allyl-to-allyl 1,4-Rh(I) migration of allylrhodium(I) species (as in A to B, Scheme 1).^{4,7} This isomerization allows subsequent carbon-carbon bond formation at sites not immediately expected from the structure of the allylboron reagents (Scheme 1, bottom). Given the synthetic potential of this underexplored mode of reactivity, its investigation in other classes of reactions is warranted. In particular, demonstration of metals other than Rh(I) to engage in allyl-to-allyl 1,4migration would be highly valuable.

Scheme 1. Allylmetal Reactivity



Scheme 2. System to Test Allyl-to-Allyl 1,4-Rh(III) Migration

In connection with our work on Rh(III)-catalyzed C-H functionalization^{8,9} in combination with alkenyl-to-allyl 1,4-Rh(III) migration to prepare heterocyclic 10 and carbocyclic 11 products, we became interested in whether allyl-to-allyl 1,4-Rh(III) migrations would be possible. 12 Our design for investigating the feasibility of this migration is shown in Scheme 2. The directing-group-assisted cyclorhodation of substrate C with a Rh(III) complex to give rhodacycle **D** is well-known.⁸ Migratory insertion of **D** with a 1,3-diene **E**, which contains allylic hydrogens cis- to the less-substituted alkene, would give allylrhodium species F, which is likely to be in equilibrium with the π -haptomer G.¹³ If allyl-to-allyl 1,4-Rh(III) migration of F were then to occur, a new allylrhodium species H would form. Although the final fate of H could not be predicted, this process could serve as a valuable addition to the currently limited number of catalytic C-H functionalizations involving 1,3-dienes,14 provided that high overall chemo-, regio-, and stereoselectivity is exhibited. Herein, we describe the successful use of allyl-to-allyl 1,4-Rh(III) migration in the oxidative C-H allylation of benzamides with 1,3-dienes. These reactions are distinct from other metal-catalyzed C-H allylations of arenes, which employ allylic electrophiles,15 allenes, 16 or terminal alkenes 17 as the reaction partners.

RESULTS AND DISCUSSION

After attempting Rh(III)-catalyzed reactions of various aromatic substrates of type \mathbf{C} with 1,3-dienes of type \mathbf{E} (see Scheme 2), ¹⁸ we found that N-acetylbenzamides $\mathbf{1}$ gave productive reactions under oxidative conditions to form allylation products $\mathbf{3}$. For example, the reaction of N-acetylbenzamide $\mathbf{1a}$

Table 1. Scope of the 1,3-Diene^a

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entry	1,3-diene	product		R	yield (%)		
1 ^c 2	2a 2b	O NHAC R	3aa 3ab	CH ₂ OBn Ph	50 69 (27) ^d		
3 ^c	2c	NHAc 9:1 E/Z	3ac		61		
4 5 6 ^c	2d 2e 2f	NHAc R	3ad 3ae 3af	Me CH ₂ OBn Ph	77 77 63		
7	2g	NHAC N	3ag		60		
8	2h	NHAc Me N-Bu 1.4:1 E/Z	3ah		31		
9° 10°	2i 2j	NHAC Ph	3ai 3aj	(CH ₂) ₄ Cl CH ₂ OMe	62 76		
11	2k	NHAc NHAc	3ak		82		

^aUnless stated otherwise, reactions were conducted using 0.30 mmol of **1a** and 0.60 mmol of **2**. ^bYield of isolated product. ^cConducted using 0.60 mmol of **1a** and 0.30 mmol of **2**. ^dValues in parentheses refer to the yield of the product **4ab** resulting from reaction at both *ortho*-positions of **1a**.

with diene **2a** in the presence of [Cp*RhCl₂]₂ (2.5 mol %) and Cu(OAc)₂ (2.1 equiv) in DMA at 70 °C for 15 h gave product **3aa** in 50% yield (Table 1, entry 1). Other dienes **2b–2k**, containing either a methyl or a methylene group *cis*- to the less-substituted alkene, are also effective and gave products **3ab–3ak** in 31–82% yield (Table 1, entries 2–11). The mass balance in these reactions was mainly composed of unreacted starting materials. In some cases, a 1:2 ratio of benzamide and diene, respectively, was optimal to maximize the yield of the products **3** (entries 2, 4, 5, 7, 8, and 11). However, in other cases a 2:1 ratio of **1a:2** was chosen to minimize the formation

of products 4, which result from C–H functionalization at both ortho-positions of **1a** (entries 1, 3, 6, 9, and 10). In one reaction, the diallylated product **4ab** was isolated (entry 2). Dienes containing a terminal alkene are effective (entries 1-3), and hydrogen, phenyl, and various alkyl groups at the alkenes are well-tolerated. With 2a, the product 3aa is derived from loss of a hydrogen atom from the methyl substituent cis- to the vinyl group, rather than from the benzyloxymethyl substituent trans- to the vinyl group (entry 1). Diene 2c, which contains a 1,2-disubstituted Z-alkene, reacted to give dienol benzyl ether 3ac as a 9:1 mixture of E/Z isomers, along with traces of unidentified decomposition products (entry 3). Dienes 2d-2k, which contain a 1,2-disubstituted alkene and a trisubstituted alkene, were also effective (entries 4-11). Here, carboncarbon bond formation occurs exclusively at the 1,2disubstituted alkene, at the carbon distal to the trisubstituted alkene. As with diene 2a (entry 1), when there are different geminal alkyl groups at the trisubstituted alkene, the products are derived from loss of a hydrogen atom at the alkyl group cis- to the disubstituted alkene (entries 4, 5, 7, and 8). This point is further exemplified by the outcomes with dienes 2e and 2h, which are geometric isomers of each other (entries 5 and 8). The reaction with 2h did not go to completion but dienol benzyl ether 3ah was obtained in 31% yield as a 1.4:1 mixture of E:Z isomers (entry 8). No evidence of 3aa was detected in this reaction.

Attention was then turned to the scope of the reaction with respect to the *N*-acetylbenzamide (Table 2). Benzamides containing a methyl group at the *para*-, *meta*-, or *ortho*-positions (entries 1, 2, 5, and 6) are tolerated, as are those bearing *para*-methoxy (entry 3) or *para*-nitro substituents (entry 4). Electron-withdrawing substituents on the aromatic ring of the benzamide appear to be beneficial, as shown by the formation of **3dj** in 75% yield compared with a 46% yield for **3cj** (compare

Table 2. Scope of the N-Acetylbenzamide^a

entry	1,3-diene	product		R	yield (%) ^b
1 2	2d 2e	NHAc R	3bd 3be	Me CH ₂ OBn	53 53
3° 4°	2j 2j	NHAc Ph NHAc Ph	3cj 3dj	OMe NO ₂	46 75
5	2d	Me NHAc Me	3ed		64
6	2b	Me O NHAc Ph	3fb		98

^aUnless stated otherwise, reactions were conducted using 0.30 mmol of 1 and 0.60 mmol of 2. ^bYield of isolated product. ^cConducted using 0.60 mmol of 1 and 0.30 mmol of 2.

entries 3 and 4). C–H functionalization of a furan-containing substrate **5** is also possible, although the yield of the product **6** was modest (eq 1).

A possible catalytic cycle for these reactions begins with formation of Cp*Rh(OAc)₂ from [Cp*RhCl₂]₂ and Cu(OAc)₂ (Scheme 3), which reacts with *N*-acetylbenzamide **1a** to give rhodacycle **7** and AcOH. Coordination and migratory insertion of 1,3-diene **2d** at the less-substituted alkene gives rhodacycle **8**, in which there is also an allylrhodium(III) moiety. Acetolysis of **8** gives allylrhodium(III) species **9**, which can undergo 1,4-Rh(III) migration^{10,11,12} to the *cis*-allylic carbon to give a

Scheme 3. Postulated Catalytic Cycle

new σ -allylrhodium intermediate 10. A σ - π - σ isomerization of 10 provides σ -allylrhodium species 11, which undergoes β -hydride elimination to give product 3ad and $Cp^*Rh(OAc)H$. Reaction of $Cp^*Rh(OAc)H$ with $Cu(OAc)_2$ (2.0 equiv) leads to reductive elimination to give AcOH and $Cp^*Rh(I)$, which is oxidized to regenerate $Cp^*Rh(OAc)_2$. Although we have proposed the acetolysis of 8 into 9, we cannot discount the possibility that the directing group remains coordinated to rhodium in one or more of the subsequent intermediates.

An alternative mechanism involves the isomerization of 9

Scheme 4. Alternative Mechanistic Pathway

A. Reaction of 1a with a hexadeuterated 1,3-diene

B. Deuterium depletion by β-deuteride elimination of [D]₆-11a

C. Deuterium depletion at the alkenyl methylene

Figure 1. Investigation of deuterium transfer with 1,3-diene [D]6-**3ad** and mechanistic rationale.

into σ -allylrhodium species 12, which undergoes β -hydride elimination to give 3ad (Scheme 4). If this mechanism was operative, it would be expected that dienes 2e and 2h, which differ only in the geometry of the trisubstituted alkene, would react to provide similar outcomes. The fact that different products are obtained in their reactions with 1a (Table 1, entries 5 and 8) suggests this pathway is less likely.

Further support for the mechanism proposed in Scheme 3 is provided by the reaction of **1a** with the hexadeuterated diene $[D]_6$ -**2d** (Figure 1A). This experiment gave $[D]_n$ -**3ad** in 67% yield, in which there was significant, but incomplete, deuterium transfer (78% D) from one of the CD_3 groups to the alkenyl carbon proximal to the benzene ring. This outcome may be rationalized by considering that σ - π - σ -isomerization of $[D]_6$ -**10** could provide $[D]_6$ -**11a** or $[D]_6$ -**11b** (Figure 1B). Deuterium depletion can then occur by β -deuteride elimination of $[D]_6$ -**11a** to give $[D]_5$ -**3ad**, whereas β -hydride elimination of $[D]_6$ -**11b** would give $[D]_6$ -**3ad**.

Another outcome of the experiment shown in Figure 1A is partial deuterium depletion (88% D) at the alkenyl methylene of $[D]_n$ -3ad. This result may be explained by reversible allylto-allyl 1,4-migration between $[D]_6$ -10, $[D]_6$ -9, and $[D]_6$ -10a, which leads to deuterium—hydrogen exchange between the two *cis*-allylic substituents (Figure 1C).⁴ σ - π - σ -Isomerization of $[D]_6$ -10a would provide $[D]_6$ -11c, from which β -deuteride elimination would give $[D]_5$ -3ad, in which there is deuterium depletion at the alkenyl methylene group.

Regarding the actual mechanism of allyl-to-allyl 1,4-Rh(III) migration, there are a number of possibilities (Scheme 5). First, in a manner similar to that proposed for the alkenyl-to-allyl 1,4-Rh(III) migrations we described previously, ^{10,11} an acetate-promoted, concerted metalation–deprotonation of [D]₆-9 would give rhodacycle 13, which could undergo acetolysis to give 10. Alternatively, 9 could undergo a C–H oxidative addition to give a Rh(V) hydride species 14, which can then form 10 by a C–H reductive elimination. The participation of Rh(V) intermediates has been suggested in various other Rh(III)-catalyzed C–H functionalization reactions¹⁹ and has gained some experimental and theoretical support.²⁰ Finally, 9 could undergo a σ-complex-assisted metathesis (σ-CAM)^{12a,21,22} *via* 15 to give 10.

Scheme 5. Possible Mechanisms for 1,4-Rh(III) Migration

To investigate the possibility of an acetate-assisted concerted metalation–deprotonation pathway to give 13, the reaction of *N*-acetylbenzamide 1a with 1,3-diene 2d was conducted in in a 9:1 mixture of DMA/D₂O. The presence of D₂O would be expected to provide some of deuterated 3ad as a result of deuteronolysis of 13, as we have observed previously in related alkenyl-to-allyl 1,4-Rh(III) migrations. ^{10,11} In the event, D₂O markedly decreased the efficiency of oxidative C–H allylation. Nevertheless, 3ad was isolated in 10% yield but no deuterium incorporation was detected. This result suggests that the intermediacy of 13 is less likely and that C–H oxidative

addition/reductive elimination or σ -CAM pathways may be more probable mechanisms for allyl-to-allyl 1,4-Rh(III) migration.

Thus far, all of the 1.3-dienes tested contain allylic hydrogens cis- to the less-substituted alkene, which enables facile allyl-to-allyl 1,4-Rh(III) migration. To test whether 1,3-dienes lacking this structural feature would also be effective substrates, the reaction of **1a** (2.0 equiv) with 1,3-diene **14**, the Eisomer of diene 2c (see Table 1, entry 3), was conducted (Figure 2A). This experiment did give allylation product 3ac as a 9:1 mixture of E/Z isomers, but in a much lower yield of 31% compared with the 61% yield obtained when the corresponding Z-diene 2c was used (Table 1, entry 3). In addition, alkenylation product 15 was isolated in 12% yield, which is notable as analogous alkenylation products were not formed in any of the reactions examined up till this point. The corresponding reaction conducted with dideuterated diene [D]₂-14 gave deuterated products [D]_n-15 and [D]_n-3ac, each in 16% yield, in which appreciable 1,4-deuterium transfer was observed (Figure 2B). This time, [D]_n-3ac was obtained as a 6:1 mixture of E/Z isomers.

A. Allylation and alkenylation of 1a with 1,3-diene 14

B. Reaction of 1a with a dideuterated 1,3-diene [D]₂-14

Figure 2. Reaction of a 1,3-diene lacking *cis*-allylic hydrogens.

The appreciable 1.4-deuterium transfer in both [D]_n-15 and [D]_n-3ac suggests a complex mechanism is operative, involving the interconversion between numerous allylrhodium(III) species by σ - π - σ isomerization (1,3-allylic transposition), E/Zisomerization, and allyl-to-allyl 1,4-Rh(III) migration pathways (Scheme 6). First, the reaction of 1a, [D]₂-14, and [Cp*RhCl₂]₂ following the initial steps of the catalytic cycle shown in Scheme 3 leads to the formation of (E)-16a, which can give a dideuterated isomer of alkenylation product [D]_n-15 by β -hydride elimination. Intermediate (E)-16a can also undergo σ - π - σ isomerization into (E)-17a, which, after β deuteride elimination, would give a monodeuterated allylation product $[D]_n$ -3ac. Alternatively, (E)-16a can undergo σ - π - σ isomerization with concomitant E/Z isomerization to give (Z)-16a, from which a series of reversible allyl-to-allyl 1,4-Rh(III) migrations involving either a 1,4-deuterium or a 1,4-hydrogen shift can give new allylrhodium(III) species (Z)-18a, (Z)-16b,

Scheme 6. Mechanistic Rationale to Explain the Outcome of the Reaction of 1a with [D]₂-14

and (*Z*)-18b. These latter three intermediates can undergo σ - π - σ isomerization to provide (*E*)-19a, (*E*)-17b, and (*E*)-19b, respectively, from which β -hydride or β -deuteride elimination would give various mono- and dideuterated isomers of [D]_n-3ac. Finally, σ - π - σ isomerization of (*E*)-17b into (*E*)-16b followed by β -hydride elimination would provide a dideuterated isomer of [D]_n-15.

To demonstrate the synthetic utility of the allylation products, 1,3-diene **3aa** was heated with *N*-phenylmaleimide in toluene at 80 °C to give Diels–Alder adduct **18** in 67% yield with >19:1 *endo:exo* selectivity (eq 3). Furthermore, allylation product **3a** reacted smoothly with 1,3-enyne **19** in a Rh(III)-catalyzed oxidative annulation to give isoindolinone **20** in 67% yield (eq 4). In this reaction, 1,3-enyne **19** functions as a one-carbon annulation partner as a result of an alkenyl-to-allyl 1,4-Rh(III) migration. ¹⁰

CONCLUSION

In summary, we have described the oxidative C-H allylation of N-acetylbenzamides with 1,3-dienes, which involve, to our knowledge, the first reported examples of allyl-to-allyl 1,4-Rh(III) migration. This new mode of Rh(III) reactivity enables reaction at sites not available from conventional 1,3allylic transposition. The results of reactions of deuterated 1,3dienes indicate that reversible interconversion of numerous allylrhodium(III) species by σ - π - σ isomerization, E/Z isomerization, and allyl-to-allyl 1,4-Rh(III) migration pathways occurs on timescales that are rapid compared to product-forming β -hydride (or β -deuteride) elimination steps. This work suggests that the possibility that these isomerization processes might occur should be taken into consideration in any future design of new reactions involving allylrhodium(III) species. Further investigation of the synthetic potential of allyl-to-allyl 1,4-metal migrations is ongoing in our group.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Experimental procedures and full spectroscopic data for all new compounds

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Notes

The authors declare no competing financial interest.

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