

1,2-Boryl Migration Empowers Regiodivergent Synthesis of Borylated Furans

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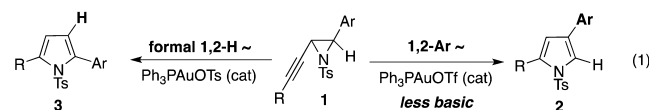
S Supporting Information

ABSTRACT: A regioselective transition metal-catalyzed cycloisomerization reaction of boron-containing alkynyl epoxides toward C2- and C3-borylated furans has been developed. It was found that the copper catalyst as well as the gold catalyst with more basic triflate counterion favor boryl migration toward C3-borylated furans, whereas employment of the cationic gold hexafluoroantimonate affords C2-borylated furan via a formal 1,2-hydrogen shift.

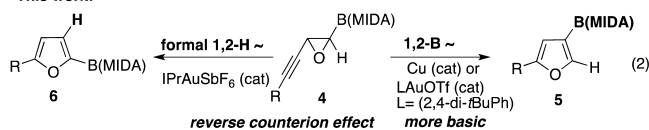
Migratory cycloisomerization is a powerful approach effectively used in synthesis of densely functionalized heterocycles.¹ Our group and others have developed a set of transition metal-catalyzed heterocyclization reactions proceeding with migration of various functional groups.² Recently, Davies' group efficiently employed alkynyl aziridine **1** in a gold-catalyzed cycloisomerization toward either 2,4- or 2,5-substituted pyrroles **2** and **3** (Scheme 1, eq 1).³ The

Scheme 1. Regiodivergent Synthesis of Furans and Pyrroles via Cycloisomerization of Alkynyl Epoxides and Aziridines

Previous work: Davies, 2009 (Ref. 3)



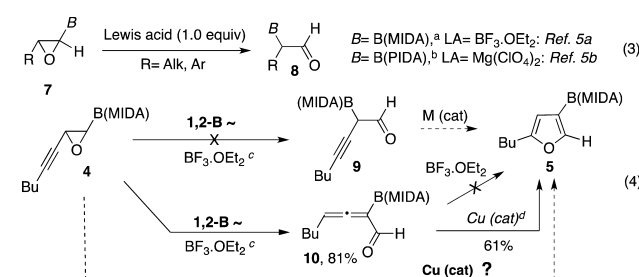
This work:



regioselectivity of this reaction is governed by the basicity of counterion, where the less basic triflate favors migration of an aryl group to afford **2**, while a more basic tosylate leads to pyrrole **3**, the product of a formal 1,2-hydrogen shift. Herein we report a regiodivergent copper- and gold-catalyzed cycloisomerization reactions of boron-containing alkynyl epoxide **4** toward C3- and C2-borylated furans **5** and **6** (Scheme 1, eq 2). To the best of our knowledge, this represents the first example of synthesis of heterocycles involving a 1,2-boryl group migration.⁴ Moreover, this gold-catalyzed cycloisomerization exhibits a reverse counterion effect, where the gold catalyst with *more basic* triflate counterion favors boryl migration to produce C3-borylated furan **5**, whereas employment of a cationic gold hexafluoroantimonate triggers a formal 1,2-hydrogen migration producing C2-borylated furan **6**.

Inspired by the recently reported Lewis acid-mediated 1,2-boryl migration in oxirane **7** into α -borylated aldehyde **8** (Scheme 2, eq 3),⁵ we attempted analogous transformation on

Scheme 2. Metal-Catalyzed 1,2-Boryl Migration in the Synthesis of 3-Borylated Furans

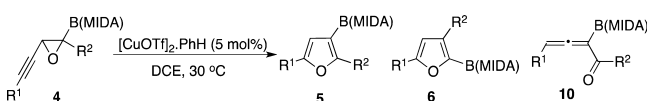


^aMIDA = *N*-methyliminodiacetic acid; PIDA = pinene-derived iminodiacetic acid; BF₃·OEt₂ (1.0 equiv), DCM, -30 °C.; [CuOTf]₂·PhH (5 mol %), DCE, 30 °C.

boron-containing alkynyl epoxide **4**. We expected formation of a skipped propargyl aldehyde **9** (analogue of **8**), which was thought to be a key precursor for the synthesis of very useful, yet otherwise challenging to access borylated furans such as **5**,^{6,7} via a subsequent cycloisomerization reaction (Scheme 2, eq 4).^{1,8} However, the reaction of **4** in the presence of BF₃·OEt₂^{5a} did not provide **9**; allenyl aldehyde **10** was efficiently formed instead (Scheme 2, eq 4).⁹ All attempts to cyclize allene **10** into furan **5** under forcing reaction conditions failed. Since allenyl aldehydes and ketones are capable substrates for the Cu-catalyzed cycloisomerization reaction toward furans,^{1,10} we tested transformation of allenyl boronate **10** into furan **5** in the presence of copper catalysts (Scheme 2, eq 4). Indeed, employment of [CuOTf]₂·PhH catalyst led to the formation of borylated furan **5** in a reasonable yield. Naturally, next, we explored the possibility of the direct transformation of alkynyl epoxide **4** into furan **5** in the presence of Cu catalyst (Table 1). To our delight, boron-containing alkynyl epoxide **4a** under the same copper-catalyzed conditions produced furan **5** in a good yield; although, a detectable amount of unexpected C2-borylated furan **6** was also formed (entry 1).¹¹ Likewise, phenyl-substituted alkynyl epoxide **4b** produced furan **5b** in good regioselectivity albeit in a low yield (entry 2). Cyclization of alkynyl epoxide **4c** possessing an electron-rich aryl group proceeded smoothly to produce furan **5c** in an excellent

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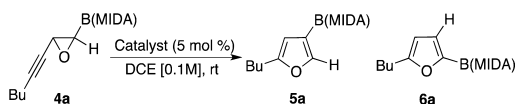
Table 1. Copper-Catalyzed Migratory Cycloisomerization Reaction toward Borylated Furans

entry	substrate	R ¹ , R ²	yield, % ^a (5:6) ^b
1	4a	<i>n</i> Bu, H	80 (97:3:0)
2	4b	Ph, H	36 (89:11:0) ^c
3	4c	2-OMe-Ph, H	51 (99:1:0)
4	4d	2-CF ₃ -Ph, H	96 (0:0:100) ^d
5	4e	<i>n</i> Bu, <i>n</i> Bu	0 ^e

^aIsolated yields. ^bNMR ratios. ^c*T* = 55 °C. ^dHeating of the reaction did not lead to furan. ^eDecomposition of 4.

regioselectivity (entry 3). However, attempts to cycloisomerize 4d possessing an electron-deficient aryl group failed (entry 4). Employing trisubstituted alkynyl epoxide 4e in this transformation resulted in complete decomposition of starting material (entry 5).

Seeking for more general and efficient conditions for regioselective synthesis of borylated furans, we next turned our attention to π -philic gold catalysts (Table 2).^{3,11,12} We

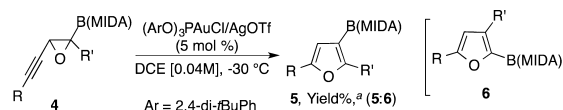
Table 2. Optimization of the Catalyst

entry	catalyst	NMR yield, %	5a:6a
1	Ph ₃ PAuCl/AgSbF ₆	>99	50:50
2	Ph ₃ PAuCl/AgOTf	85	82:18
3	(ArO) ₃ PAuCl ^a /AgOTf ^{b,c}	81 ^d	96:4 ^e
4	(ArO) ₃ PAuCl ^a /AgSbF ₆ ^b	76	22:78
5	IPrAuCl/AgOTf	79	33:67
6	IPrAuCl/AgSbF ₆	81	0:100
7	AgOTf ^b	39	87:13
8	AgSbF ₆	39	86:14

^aAr = 2,4-di-*t*BuPh. ^bSolution [0.04M]. ^c*T* = -30 °C. ^d76% NMR yield at rt. ^eRatio of 91:9 at rt.

found that in the presence of Ph₃PAuCl, the reaction was more regioselective for 5a with triflate counterion (entry 2) rather than with hexafluoroantimonate (entry 1). Gratifyingly, employment of a gold catalyst possessing electron-rich phosphite ligand with triflate led to furan 5a with an excellent regioselectivity and yield (entry 3). Conversely, switching counterion to hexafluoroantimonate favored formation of 6a (entry 4). Analogous results were obtained in the presence of IPrAuCl with triflate counterion (entry 5). Remarkably, employment of the same gold catalyst with hexafluoroantimonate led to furan 6a, exclusively (entry 6).¹³ Control gold-free experiments indicated that the reactions in the presence of silver triflate (entry 7) and silver hexafluoroantimonate (entry 8) were both less efficient and regioselective.

After finding conditions for regiodivergent synthesis of boryl furans 5 and 6, we first examined the scope of the migratory cycloisomerization reaction toward C3-borylated furan 5 (Table 3). Thus, C2-alkyl-substituted furans 5a, 5f, and 5g were produced in good yields and excellent regioselectivity. Phenyl-substituted alkynyl epoxide 4b reacted smoothly to afford 5b in 87% yield and good regioselectivity. Likewise,

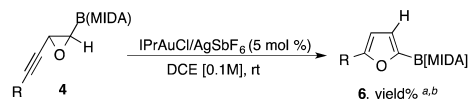
Table 3. Gold-Catalyzed Synthesis of C3-Borylated Furans

5a, 83, (92:8)	5f, 67, (97:3)	5g, 74, (96:4)
5b, 87, (90:10)	R = OMe: 5c, 81, (100:0) R = CF ₃ : 5d, 61, (100:0)	R = CF ₃ : 5h, 65, (91:9) R = Br: 5i, 79, (95:5)
5j, 67, (100:0)	5k, 83, (100:0)	5e, 75, (100:0)

^aIsolated yield. ^bReaction was performed at rt.

ortho-substituted aryl-containing furans 5c and 5d were exclusively isolated in good yields. Analogously, cyclization of alkynyl epoxides 4h and 4i provided furans 5h and 5i possessing *para*-substituted aryls. Diynyl epoxide 4j afforded alkynylated furan 5j exclusively in 67% yield. Remarkably, employment of trisubstituted oxirane 4k in this transformation led to the exclusive formation of furan 5k in 83% yield, thus, indicating overwhelming preference of boryl group versus aryl group^{1,2a-c,3} migration. Analogously, dialkyl-containing furan 5e was selectively obtained under these conditions.

Next, we investigated the gold-catalyzed cycloisomerization reaction of alkynyl epoxide 4 leading to C2-borylated furan 6 (Table 4). Thus, alkyl-containing furans 6a, 6f, and 6g were exclusively obtained in reasonable to good yields. Analogously, alkynyl epoxide 4b possessing phenyl substituent worked well under these reaction conditions. Likewise, electronically different *ortho*- and *para*-substituted aryl-containing furans 6c, 6d, 6h, and 6i were obtained in modest to good yields and

Table 4. Gold-Catalyzed Synthesis of C2-Borylated Furans

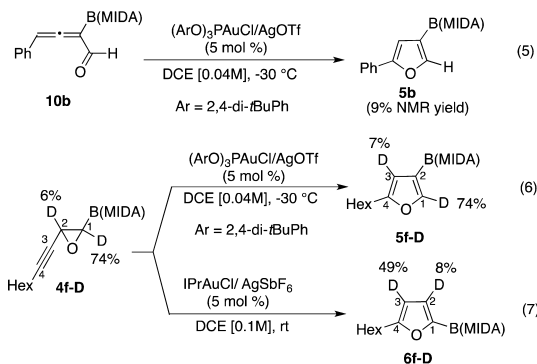
6a, 75	6f, 57	6g, 64
6b, 50	R = OMe: 6c, 49 ^c R = CF ₃ : 6d, 42 ^{c,d}	R = CF ₃ : 6h, 65 R = Br: 6i, 84
6j, 66	6l, 60 ^{e,f}	

^aIsolated yields. ^bExclusive formation of C2-borylated furans was observed in all examples. ^cSubstantial decomposition of substrate was observed. ^dAt 60 °C. ^eTHF as the solvent. ^fNMR yield.

excellent regioselectivity. In addition, diynyl epoxide **4j** was smoothly converted into alkynylated furan **6j** in good yield. Finally, cyclization of **4l** afforded monosubstituted boryl furan **6l** in 60% yield.¹⁴

We also performed some initial mechanistic experiments. First, we verified whether the Au-catalyzed reaction proceeds via intermediacy of allene **10** (Scheme 3, eq 5). However, a very

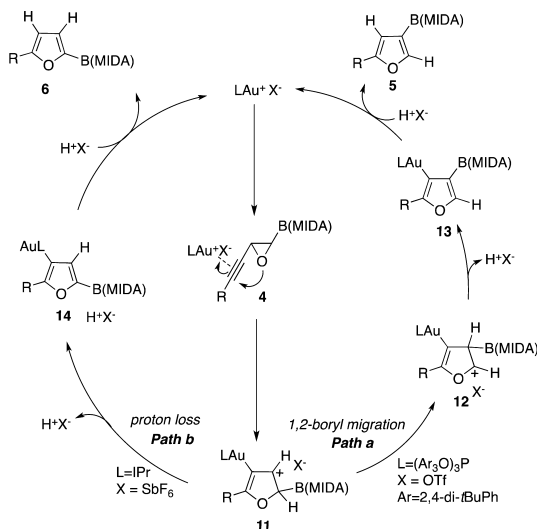
Scheme 3. Mechanistic Studies of the Gold-Catalyzed Cycloisomerization Reactions



low yield of furan **5b** was observed when **10b** was subjected to the reaction conditions. This indicates that in contrast to the Cu-catalyzed version (*vide supra*), generation of allene intermediate **10** in the gold-catalyzed transformation is less likely. Next, cycloisomerization of the deuterium-labeled alkyne epoxide **4f-D** was examined. It was found that under the gold-catalyzed migratory conditions, furan **5f-D** was formed in which deuterium label stayed intact at the C1 position (Scheme 3, eq 6). However, subjecting **4f-D** to the non-migratory cycloisomerization conditions produced furan **6f-D** with deuterium label at the C3 position substantially scrambled (Scheme 3, eq 7).¹⁵

Obviously, more detailed studies toward understanding the role of both ligand and counterion on the regioselectivity of this cycloisomerization reaction are required. At this point, based on the above observations, we propose the following working rationale for this regiodivergent transformation (Scheme 4).

Scheme 4. Mechanistic Proposal for the Regiodivergent Gold-Catalyzed Cycloisomerizations



First, activation of the π system of **4** by the gold catalyst triggers a nucleophilic attack of the oxygen at the distal position of alkyne moiety producing the heterocyclic cation **11**.¹² In the case of triflate counterion (*Path a*), a 1,2-boryl migration to the cationic center of **11** takes place to generate cation **12**, which, upon proton loss affords the furyl gold specie **13**. Protodemetalation of the latter produces the C3-borylated furan **5**. Alternatively, in the presence of hexafluoroantimonate counterion (*Path b*), a proton loss¹⁵ in intermediate **11** takes place to afford the furyl gold species **14**, which upon protodeauration produces C2-borylated furan **6**. It should be mentioned that the observed counterion effect on the regioselectivity of boryl group versus hydrogen migration is in a sharp contrast with the migration trend for carbon and silyl group versus hydrogen migration previously reported by our^{2d,k} and other¹⁶ groups. Further studies to elucidate the origins of the observed migratory trend are underway in our group.

In summary, complementary regioselective copper- and gold-catalyzed cycloisomerization reactions of boron-containing alkyne epoxides toward C2- and C3-borylated furans have been developed. It was found that the copper-catalyzed transformation triggers the initial 1,2-boryl group migration forming an allenyl derivative, which upon cycloisomerization produces C3-borylated furan. However, the Cu-catalyzed cycloisomerization appeared to be not general. Accordingly, we developed two efficient and general Au-catalyzed protocols, where depending on the choice of ligand and counterion, regioisomeric boryl furans can be selectively obtained. Thus, phosphite gold complex with triflate counterion strongly favors a 1,2-boryl migration during the cycloisomerization process to produce the C3-borylated furans. In contrast, employment of NHC gold hexafluoroantimonate affords C2-borylated furans, exclusively. MIDA boronate moiety proven to be a highly valuable group for a variety of useful transformations.¹⁷ Thus, it is believed that the obtained regioisomeric MIDA boronate-containing furans would become useful building blocks for organic synthesis.

■ ASSOCIATED CONTENT

Supporting Information

Experimental details, compound characterization, and spectra. 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.

■ ACKNOWLEDGMENTS

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