See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/6179835

Mechanistically Diverse Copper-, Silver-, and Gold-Catalyzed Acyloxy and Phosphatyloxy Migrations: Efficient Synthesis of Heterocycles via Cascade Migration/Cycloisomerization Appr...

ARTICLE in JOURNAL OF THE AMERICAN CHEMICAL SOCIETY · SEPTEMBER 2007

Impact Factor: 12.11 · DOI: 10.1021/ja072446m · Source: PubMed

CITATIONS READS

193 28

## **5 AUTHORS**, INCLUDING:



**Todd Schwier** 

Hospira

13 PUBLICATIONS 529 CITATIONS

SEE PROFILE



**Anna Sromek** 

McLean Hospital

22 PUBLICATIONS 1,434 CITATIONS

SEE PROFILE



Am Chem Soc. Author manuscript; available in PMC 2008 September 15.

Published in final edited form as:

J Am Chem Soc. 2007 August 15; 129(32): 9868–9878. doi:10.1021/ja072446m.

# Mechanistically Diverse Copper-, Silver-, and Gold- Catalyzed Acyloxy and Phosphatyloxy Migrations: Efficient Synthesis of Heterocycles via Cascade Migration/Cycloisomerization Approach

**Todd Schwier, Anna W. Sromek, Dahrika M. L. Yap, Dmitri Chernyak**, and **Vladimir Gevorgyan** *University of Illinois at Chicago, Department of Chemistry, 845 West Taylor Street, Chicago, Illinois* 60607–7061

#### Abstract

A set of cycloisomerizaton methodologies of alkynyl ketones and imines with concurrent acyloxy, phosphatyloxy, or sulfonyloxy group migration, which allow for the efficient synthesis of multisubstituted furans and N-fused heterocycles has been developed. Investigation of the reaction course by way of employing <sup>17</sup>O-labelled substrates allowed for elucidation of the mechanisms behind these diverse transformations. It was found that, while the phosphatyloxy migration in conjugated alkynyl imines in their cycloisomerization to N-fused pyrroles proceeded via a [3,3]sigmatropic rearrangement, the analogous cycloisomerization of skipped alkynyl ketones proceeds through two consecutive 1,2-migrations, resulting in an apparent 1,3-shift, followed by a subsequent 1,2-migration through competitive oxirenium and dioxolenylium pathways. Investigations of the 1,2acyloxy migration of conjugated alkynyl ketones en route to furans demonstrated the involvement of a dioxolenylium intermediate. The mechanism of cycloisomerization of skipped alkynyl ketones containing an acyloxy group was found to be catalyst dependent; Lewis and Brønsted acid catalysts caused an ionization/ $S_N$ 1' isomerization to the allene, followed by cycloisomerization to the furan, while transition metal-catalysts evoked a Rautenstrauch-type mechanistic pathway. Furthermore, control experiments in the cycloisomerization of skipped alkynyl ketones under transition metal catalysis revealed that, indeed, these reactions were catalyzed by transition metal complexes as opposed to Brønsted acids resulting from hydrolysis of these catalysts with eventual water. Further synthetic utility of the obtained phosphatyloxy-substituted heterocycles was demonstrated through their efficient employment in the Kumada cross-coupling reaction with various Grignard reagents.

## Introduction

Furans and pyrroles are frequently occurring heterocyclic units found in numerous naturally occurring and biologically active molecules,  $^1$  and find broad application as synthetic intermediates  $^2$  and in the material sciences.  $^3$  Thus, not surprisingly, numerous methods have been developed for the assembly of the furan and pyrrole ring,  $^4$ ,  $^5$  among which transition metal-catalyzed cycloisomerizations,  $^6$  perhaps, are the most general and powerful approaches toward these heterocycles.

Recently, we reported the Cu-catalyzed cycloisomerization of alkynyl imines and ketones to afford monocyclic and fused pyrroles and furans. <sup>6n,p</sup> It was proposed that the alkynyl imine or ketone 1 undergoes a base-assisted prototropic 1,3-H shift, leading to allene 2 which, upon a copper-catalyzed cycloisomerization, transforms into pyrrole or furan 3 (eq 1). This methodology, however, is limited to the synthesis of heterocycles lacking substituents at the C-3 and C-4 positions, as the hydrogen substituent at C-4 of heterocycle 3 arises from the prototropic 1,3-H migration and H at C-3 comes from the cycloisomerization of 2 to 3.

$$\begin{array}{c|ccccc}
H & R^2 & \text{prototropic} \\
H & R^1 & X & R^2 & R^2 \\
1 & 2 & X & R^2 & R^2 \\
X = O, NR & 3 & X & R^2 & R^2 \\
\end{array}$$
(1)

As a partial solution to this problem, we have recently developed a set of methods for the synthesis of pyrroles and furans **6** possessing a thio-<sup>6m</sup> or halogen<sup>6f</sup> substituent at C-3 of heterocycle. These novel transition metal-catalyzed cascade transformations involve isomerization of **4** into allene **5**, followed by cycloisomerization of the latter into **6**, during which, a 1,2-shift of the thio- or halogen group occurs (eq 2). Aiming at expanding the scope of the migrating group (MG), we explored the possibility of

engaging a 1,3-migration of different oxygen-based migrating groups, known to occur in propargyl/allenyl systems via a [3,3]-sigmatropic shift. It was found, however, that depending on the substitution pattern in 7 and/or reaction conditions, different regioisomeric products 9 or 11 were formed (eq 3). It was reasoned that, in the former case, the expected [3,3]-shift of the migrating group occurs to produce allene 8, followed by its cycloisomerization to 9. In the latter case, a prototropic rearrangement of 7 gives rise to allene 10, which cycloisomerizes with concomitant 1,2-migration into furan 11.

Herein, we describe a more detailed study of the scope and mechanisms of different modes of cycloisomerization<sup>8</sup> of alkynyl ketones and cyclic imines proceeding with concurrent migration of acyloxy and phosphatyloxy groups to produce multisubstituted furans and fused

pyrroles, together with the synthetic application of the obtained heterocyclic phosphates in Kumada cross-coupling reactions.

#### **Results and Discussion**

# Synthesis of Trisubstituted Heterocycles via [3,3]-Phosphatyloxy Migration in Conjugated Alkynyl Ketones and Imines

Our initial attempts explored the possibility of engaging a [3,3]-phosphatyloxy migration in alkynyl ketones 12 into 13 as an approach toward furans 14 (Scheme 1). To this end, cycloisomerization of several phosphatyloxy alkynyl ketones  $^9$  was examined in the presence of CuCl (5 mol%) in DMA. Since this transformation no longer involves a base-assisted prototropic H shift, we tested this reaction in the absence of amine. We were pleased to find that under these conditions the cycloisomerization proceeded smoothly, furnishing the corresponding furans 14a and 14b in good yield. While CuCl appeared to be a poor catalyst for the cycloisomerization of phenyl-substituted 12c, we found that the employment of AgBF<sub>4</sub> in dichloroethane at  $80^{\circ}$ C gave the desired furan 14c in good yield (Scheme 1).

Next, we investigated the applicability of this approach for the synthesis of phosphatyloxy-substituted indolizines **17** (Scheme 2). It was found that the cycloisomerization of alkynyl pyridines **15** under the same conditions proceeded uneventfully to furnish the desired indolizines **17a-e** in good to excellent yields. Of note, while the standard cycloisomerization of alkynyl imines to give monosubstituted indolizines required 50 mol% Cu salts, <sup>6p</sup> this novel cascade [3,3]-migration/cycloisomerization protocol proceeded efficiently in the presence of 5 mol% of CuCl only (Scheme 2).

It deserves mentioning that in both cases (Schemes 2 and 3) we did not observe the intermediate allenes (13 and 16), which supports their formation at the rate limiting event. In an effort to obtain a better understanding of the mechanism behind the propargyl-allenyl isomerization and their subsequent transformation into heterocycles, we investigated the cycloisomerization of phosphoryl oxygen [<sup>17</sup>O]-enriched alkynyl pyridine 18, the labeled analogue of 15c (Scheme 3), by means of <sup>17</sup>O NMR spectroscopy. <sup>10</sup> It was reasoned that if the isomerization indeed proceeds via a [3,3]-sigmatropic shift, the oxygen label should be located at the phosphate ester position in the intermediate allene 19, and thus in the indolizine, as shown in 20. Unsurprisingly, the NMR spectrum of the isolated indolizine product displayed a broad signal at 71 ppm corresponding to the bridging phosphate oxygen. <sup>11</sup> Only traces of the phosphoryl oxygen signal, resulting from formation of 21 were detected. Thus, the selective formation of 20 strongly supports the mechanistic rationale involving a propargyl-allenyl isomerization proceeding via a sigmatropic [3,3]-phosphatyloxy shift (Schemes 1-3).

## Synthesis of Trisubstituted Furans via 1,2-Migration of Acyloxy Group in Conjugated Alkynyl Ketones

Encouraged by the successful development of conditions for the efficient [3,3]-migration/cycloisomerization of phosphatyloxy-containing propargyl ketones, we next turned our attention to the elaboration of analogous transformations involving acyloxy systems. Our initial attempts at migration/cycloisomerization were disappointing. In the presence of catalytic CuCl, acetate **22a** gave a low yield of the expected cycloisomerization product **23a**<sup>7</sup> and trace amounts of an unexpected regioisomeric product **23b** (Scheme 4). The same reaction in the presence of Et<sub>3</sub>N additive however, produced the other regioisomer, **23b**, as a major product. Intrigued by this finding, we decided to investigate the latter transformation further, as it may provide access to heterocycles of a different substitution pattern. Switching to phenyl and tertbutyl alkynyl ketones resulted in increases in both the yields and the regioselectivity of the products (Table 1). Thus, a series of alkynyl ketones **22** with different acyloxy groups under

these reaction conditions, underwent smooth cycloisomerization to produce acyloxy furans **23** in good to excellent yields.

We propose the following plausible mechanisms to account for the observed unusual regiochemistry of the cycloisomerization products obtained (Scheme 5). The upper path **A** depicts the formation of allene **25** from **24** via a base-assisted prototropic shift which, upon metal activation of the carbonyl, forms dioxolenylium species **26**. Subsequent transformation of the latter to furan **27** occurs by an intramolecular Ad<sub>N</sub>-E process. <sup>12</sup> Alternatively (path **B**), **24** could undergo a Rautenstrauch-type <sup>13</sup> metal-assisted 1,2-acyloxy<sup>14</sup> migration <sup>15</sup> via dioxolenylium species <sup>16</sup> **28** to give metal carbenoid **29**. Intramolecular attack of the ketone to give **30**, followed by tautomerization, gives furan **27**. While, on the basis of our current studies, we cannot rule out either mechanistic pathway, the prerequisite of base for the selective formation of the observed regioisomer of furan **23b** over **23a** (Scheme 4) supports possible involvement of allene intermediate **25**. <sup>17</sup>

We hypothesized that the acyloxy group migration may also proceed via a three-membered oxirenium species (Scheme 6), analogously to sulfur and halogen group migrations reported earlier.  $^{6f,m}$  The involvement of either dioxolenylium (Scheme 5) or oxirenium (Scheme 6) motifs could be established through reaction of a labeled substrate 24. While dioxolenylium-involving pathway  $\bf A$  and  $\bf B$  both lead to the same isotopomer 27 (Scheme 5), a migration occurring via oxirenium intermediates (paths  $\bf A'$  and  $\bf B'$ ) would lead to a different isotopomer 29 (Scheme 6).

To test the above hypothesis, we synthesized <sup>17</sup>O enriched **33**<sup>9</sup> and subjected it to the cycloisomerization conditions (Scheme 7). The reaction was stopped at about 60% conversion. NMR analysis indicated that the reaction resulted in the formation of a 7:1 mixture of isotopomeric furans **34** and **35**. Additionally, it was found that the recovered starting material showed no additional signals in the <sup>17</sup>O spectrum, thus indicating no scrambling during the reaction. Furthermore, resubmission of the isolated furans to the reaction conditions did not result in a discernable change in the ratio of **34**:35, thus ruling out possible scrambling of the product. These observations, confirm that the major product, furan **34**, arises through either of the mechanistic pathways outlined in Scheme 5, with migration occurring via a dioxolenylium species. Formation of minor product **35**, however, is consistent with either of the two mechanisms depicted in Scheme 6 or an irreversible ionization/recombination to form the allene (see below, Scheme 11).

#### Synthesis of Tetrasubstituted Furans via a Cascade Migration in Skipped Alkynyl Ketones

Encouraged by successful cycloisomerization of various alkynyl ketones to furans (*vide supra*), we proposed that a propargyl-allenyl isomerization of isomeric propargyl ketone **36** via a [3,3]-migration might provide access to allene **37**, the fully substituted analogue of **10** (eq 3), which, upon cycloisomerization with subsequent 1,2-migration, would lead to fully substituted furan **38** (eq 4). Additionally, if successful, formation of allenyl intermediate **37** would provide additional support for our mechanistic proposal above (Scheme 5, path **A**).

$$Y = CR^4$$
,  $P(OR^4)_2$ ,  $S(O)R^4$ 

First, we examined the potential migration of acyloxy group in  $\bf 39~(Y=CR^4)$  *en route* to tetrasubstituted furan  $\bf 40$ . Our optimization studies indicated that several metal salts are able to efficiently catalyze this transformation (Table 2). While the silver salts AgBF<sub>4</sub> and AgOTf proved efficient for the cycloisomerization of *tert*-butyl substituted  $\bf 39b$  (entries 5 and 7), they were less desirable for *n*-butyl substituted  $\bf 39a$  (entries 4 and 6). However, it was found that  $Cu(OTf)_2$  was effective for the formation of *n*-butyl-substituted furan  $\bf 40a$  (entry 1). AuCl<sub>3</sub> worked reasonably well with both  $^tBu$ - and  $^nBu$  substituted substrates (entries 10 and 11). Interestingly, Lewis acids  $^{18}$  also were effective in promoting this transformation (entries 13 –16). Remarkably, even Brønsted acid (HOTf) efficiently catalyzed this transformation (entry 17).

Next, cycloisomerization of several differently-substituted propargyl acetates and pivaloates **39** under the optimized conditions was examined. It was found that the cascade cycloisomerization of **39** proceeded smoothly to produce fully-substituted furan **40** in good to excellent yields (Table 3). Noteworthy, lengthened reaction times were required for the reaction of pivaloates as compared to the corresponding acetate (entries 2 and 9 vs 1 and 8, respectively). As the TMS substituted alkyne function in **39i** is incompatible with silver salts, <sup>19</sup> employment of AuCl<sub>3</sub> gave the desired furan **40i** in good yield (entry 10).

Next, we examined possible cascade involving migration of phosphatyloxy group cycloisomerization (Scheme 8). It was found that 41, in the presence of  $AgBF_4$  in dichloromethane at room temperature, smoothly converted to allene 42, thus, providing evidence for the formation of allenyl intermediates in the reaction. Subsequent treatment of the allene 42 with  $AgBF_4$  in dichloroethane at  $60^{\circ}C$  afforded furan 43 in 77% yield. Additionally, the latter conditions were effective for the formation of furan 43 directly from the propargyl phosphate 41 (Scheme 8).

Aiming at expanding the scope of this transformation, we examined the analogous transformation of propargyl tosylates under similar reaction conditions. Unexpectedly, our attempts to synthesize **44** led to the direct formation of tosyl allene **45** (Scheme 9). When subjected to  $AgBF_4$  in dichloroethane at 60°C, allene **45** was smoothly converted to the tosyl-substituted furan **46** in 82% yield.

Understandably, we were interested in establishing the mechanism behind the cascade cycloisomerization in skipped systems. From the outset, we postulated two general mechanisms to explain the formation of furans from propargyl substituted ketones. The first operates via an allenyl intermediate **48** (or **37**, as depicted in eq 4), whereas the second goes through carbenoid species **50** (Scheme 10). If one of the oxygens in the migrating group is selectively labeled, as in **47**, it would be possible to distinguish between the two mechanistic paths. Thus, as depicted in the upper pathway, **47** first undergoes a [3,3]-shift, with "inversion" of the labeled oxygen to form allene **48**. A subsequent 1,2-shift via a dioxolenylium species would result in a second "inversion" to produce the Y=O oxygen-labeled furan **49**. Alternatively, if **47** undergoes a 1,2-shift to form metal carbenoid **50**, the label would be located at the bridged oxygen atom. The intramolecular nucleophilic attack and subsequent cycloisomerization proceeds with no migration, so the label would remain at the bridged oxygen position in the furan product **51**.

Thus, labeled acetate  $52^9$  was tested under several reaction conditions proved efficient in catalyzing this transformation (Table 4). When subjected to the CuCl/Et<sub>3</sub>N conditions, carbonyl oxygen-labeled 53 was formed as a *minor* product, with the major product being the ester oxygen-labeled 54 (entry 1). More remarkably, reactions using AgBF<sub>4</sub>, AuCl<sub>3</sub>, or Cu (OTf)<sub>2</sub> (entries 2, 3, and 4) gave 54 exclusively. Furthermore, the Lewis acids Sn(OTf)<sub>2</sub> and TMSOTf (entries 5 and 6) gave mixtures of nearly equal amounts of isotopomers 53 and 54.

Employment of the Brønsted acid (TfOH, entry 7) also resulted in the formation of a mixture of isotopomers (Table 4).  $^{21}$ 

In light of the recent observations that eventual Brønsted acids are the true catalysts in some transition metal-catalyzed transformations, <sup>22</sup> we investigated what role, if any, Brønsted acids may play in the described herein cycloisomerization reaction. <sup>23</sup> To this end, in the test experiment, we examined the cycloisomerization of **39b** in the presence of Brønsted base additive, which expected to quench eventual Brønsted acids which may be present in the reaction media. It was found that hindered base, 2,4,6-tri-tertbutylpyrimidine (TTBP), serves the purpose: addition of 15 mol% of TTBP to reaction mixture containing 10 mol% TfOH completely shuts down the reaction, which is otherwise complete in 6 hours (Table 5, entry 1). With this tool in hand, we then examined cycloisomerization of **39b** catalyzed by several transition metal catalysts in the presence and absence of TTBP (entries 2–6). It was found that in all cases, the reaction course was unaffected by the presence of the base: the reactions with TTBP additive proceeded with rates comparable to those lacking the base, thus ruling out possible involvement of Brønsted acids in these transition metal-catalyzed transformations. Moreover, it was found that the cycloisomerization of **52** in the presence of Au catalyst and TTBP (Table 4, entry 8) did not change the isotopomeric distribution over the base-free experiment (entry 3).

Based on the results of the labeling studies, we believe that the cycloisomerization of acetates follows one of two mechanistic pathways, depending on the catalyst. Transition metal-catalysts such as Ag(I), Au(III), and Cu(II) likely cause an initial 1,2-migration to form a metal carbenoid (Scheme 10, lower path). Subsequent cycloisomerization of this intermediate then provides the corresponding furan (Scheme 16). Alternatively, Lewis/Brønsted acid catalysts, such as TMSOTf,  $Sn(OTf)_2$ , and HOTf, or under high temperature conditions, a partial ionization of the propargyl acetate **36** to a propargylium/allenylium cation<sup>24</sup> occurs during which the <sup>17</sup>O label is scrambled: consistent with the observed mixture of isotopomers **53** and **54** (Scheme 10 and 11). To obtain additional support for the ionization mechanism, we tested cycloisomerization of acetate **39b** in the presence of an external nucleophile, methallyl TMS (Scheme 12). Unsurprisingly, the 1,5-enyne **57** was obtained in high yield, along with small amounts of the allene **58**. Further studies indicated that enyne **57** does not convert to allene **58** under the reaction conditions. As the allylation of propargylic acetates is known to follow an  $S_N$ 1 mechanism, <sup>18</sup> these results support the proposed ionization pathway (Scheme 11).

Since allenes were cleanly obtained in the cycloisomerization of propargyl phosphates and tosylates, it occurred to us that phosphates and acetates were following distinct mechanistic pathways. Thus, we decided to further investigate the mechanism of this intriguing transformation using labeled phosphates. Initially, we subjected phosphoryl oxygen-labeled 599 to the conditions under which the allene could be obtained (Scheme 13). Surprisingly, the obtained allene 60 maintained the phosphoryl oxygen-labeling pattern, with no traces of the other isotopomer, thus completely ruling out possible involvement of [3,3]-sigmatropic shift in this transformation! Moreover, conversion of the allene 60 to furan resulted in the formation of phosphoryl oxygen-labeled 61 as a major component over the bridged oxygen-labeled 62. Similar results were obtained when alkyne 59 was converted directly to the furan under elevated temperatures. In a control experiment, 59 rapidly (*ca* in 10 min) isomerized to the allene with 10 mol % triflic acid. Consistent with an ionization pathway, as experienced with acetates, allene 60 was formed accompanied with comparable amounts of its isotopomer.<sup>27</sup> When left for over 15 hours, the cycloisomerization of isotopomeric allenes proceeded to give furans 61 and 62 in 83% combined yield.

Apparently, the mechanism of migration of the phosphatyloxy group presented an interesting challenge. Involvement of the allene was demonstrated, as it was isolated and shown to convert

to the furan. Furthermore, the position of the oxygen label ruled out a [3,3]-sigmatropic shift (Scheme 10, upper path) as the product allene 60 from alkyne 59 possessed the oxygen label solely at the phosphoryl oxygen (Scheme 13). Such a migration would have resulted in the other isotopomer and ionization would result in the formation of both. It was clear that both of these proposed mechanisms were in disagreement with the obtained experimental results. To rationalize the selective formation of the observed isotopomer of allene, we propose the following potential mechanisms (Scheme 14). The first path (path A) presumes two consecutive 1,2-migrations via dioxolenylium species (64 and 66) and carbenoid intermediate 65. Net retention of the Y=O label is the result of two "inversions." Alternative path B occurs via two retentive 1,2-shifts proceeding through oxirenium entities 68 and 70 and carbenoid 69, which is isotopomeric to carbenoid 65 (Scheme 14). While path A is more consistent with previous mechanistic studies, based on our current results, we cannot rule out path B.

The formation of two furan isotopomers in the cycloisomerization of isotopomerically pure allene **60** (Scheme 13) can be explained via two competing mechanisms for the 1,2-phosphatyloxy migration (Scheme 15). Apparently, "retentive" 1,2-migration via oxirenium species **71** would provide the major isotopomer, phosphoryl oxygen-labeled furan **72**, whereas 1,2 migration via dioxolenylium species **73** produces furan **74**, with the bridged oxygen labeled.

Our mechanistic studies discussed above suggest that the mechanism for the migration/ cycloisomerization of propargyl substituted ketones depends on both the nature of the migrating group and catalyst employed. A generalized mechanism for these transformations is depicted in Scheme 16. Our initial postulate that allene 37 could be accessed via a [3,3]sigmatropic migration is in conflict with our current experimental data (Scheme 16). In the case of acyloxy migration in the presence of transition metal catalysts, the metal carbenoid 75, formed via a 1,2-migration, could be trapped by the carbonyl oxygen of the ketone to give furan 38 directly (Rautenstrauch-type). Alternatively, metal carbenoid 75 could be trapped by the carbonyl of the ester moiety to give the allene 37. A 1,2-migration via a dioxolenylium would result in the formation of furan 38 with inversion of <sup>17</sup>O label. As the isotopomeric outcome is the same in either of these two mechanisms, neither mechanistic possibility can be ruled out. Analogous transformation in the presence of Lewis acids proceeds with substantial <sup>17</sup>O scrambling in the product, which implies formation of scrambled allene 37 via an S<sub>N</sub>1' mechanism. In the cycloisomerization of phosphates, the first step is likely also a 1,2migration via a dioxolenylium species to metal carbenoid 75. However, since the allene 37 was isolated without formation of furan 38, a second 1,2-migration, also through a dioxolenylium intermediate via attack of the phosphoryl oxygen at the carbenoid center, produces the observed allene with exclusive retention of <sup>17</sup>O label. The 1,2-migration required to go from allene **37** to furan 38 likely follows two competing mechanistic pathways, among which a retentive 1,2migration via oxirenium species predominates, producing furan 38 with the <sup>17</sup>O label at the Y=O position as a major isotopomer (Scheme 16).

#### **Kumada Cross-Coupling of Hetaryl Phosphates**

To demonstrate the synthetic utility of the obtained phosphatyloxy-substituted heteroaromatics, we examined their reactions with Grignard reagents under Kumada cross-coupling conditions. It deserves mentioning that, while successful examples of Kumada cross-coupling of Grignard reagents with enol phosphates <sup>28</sup> and aryl phosphates <sup>29</sup> are well precedented, to the best of our knowledge, no examples of the analogous reaction involving heteroaromatic phosphates have ever been reported. We have found that Kumada cross-coupling of furyl- and indolizinyl phosphates is possible. Substantial optimization indicated that this transformation proceeded most efficiently in the presence of Pd<sub>2</sub>dba<sub>3</sub>/CyPF-<sup>t</sup>Bu<sup>30</sup> combination. The scope of cross-coupling was examined employing these optimized conditions (Table 6). Kumada cross-coupling of phosphatyloxy-containing furans **14c** and

indolizines **17b**, proceeded smoothly with various Grignard reagents to give C-3 substituted furans **76a-d** and C-1 substituted indolizines **77a-d** in good to excellent yields (Table 6).

## Conclusion

In summary, we have developed different modes of cascade cycloisomerizations of alkynyl ketones and imines proceeding via various types of migration of acyloxy, phosphatyloxy, and tosyloxy groups to give multisubstituted furans and indolizines in good to high yields. This set of methodologies allows for the efficient synthesis of tri- and tetrasubstituted furans and Nfused heterocycles. Mechanistic studies employing <sup>17</sup>O-labeled starting materials indicated that the reaction mechanism depends on the cyclization mode and nature of migrating group. It was shown that 1,3-phosphatyloxy migration in the cycloisomerization of conjugated alkynyl imines proceeds via a [3,3]-shift. The 1,2-migration of the acyloxy group in conjugated alkynyl ketones was found to proceed via a dioxolenylium intermediate. In contrast to acyloxy migration in conjugated alkynyl ketones, the mechanism of acyloxy migration in skipped systems depends on the reaction conditions. While Lewis and Brønsted acids and transition metal complexes catalyze the cycloisomerization of skipped alkynyl ketones, they follow different mechanistic pathways! When an acid (either Lewis or Brønsted) catalyst is used, formation of the postulated allenyl intermediate occurs through an  $S_N1$  pathway. However, when transition metal catalysts are used, the cycloisomerization follows either a Rautenstrauchtype 1,2-migration of the acyloxy group followed by the cycloisomerization to the furan directly or, alternatively, two 1,2 migrations of the acyloxy group to form the postulated allenyl intermediate, which cycloisomerizes into the furan via another 1,2-acyloxy shift. Furthermore, control experiments and <sup>17</sup>O-labeling studies demonstrated that the transition metal-catalyzed reactions were actually catalyzed by the metal and not by eventual protons. The analogous cycloisomerization of skipped phosphatyloxy alkynyl ketones was also shown to be induced by both Brønsted acid and transition metal catalysts. As in acetates, acids induce an ionization to form the allene via an  $S_N$ ! mechanistic pathway, whereas under transition metal catalysis, the allene forms via two 1,2-phosphatyloxy migrations. Furthermore, as an extension, we demonstrated the synthetic utility of phosphatyloxy furans and indolizines in the efficient Kumada cross-coupling reaction.

## **Experimental Section**

Concise representative procedures for the preparation of starting materials are provded here. Detailed procedures for the preparation of all starting materials, as well as for the cycloisomerization reactions, can be found in the Supporting Information.

## 12a

The preparation of **12a** is representative. To a microreactor were sequentially added DMAP (11 mg, 0.09 mmol), anhydrous dichloromethane (1 mL), triethylamine (91 µL, 0.65 mmol), 5-hydroxy-3-decyne-2-one (93 µL, 0.50 mmol), and diethylchlorophosphate (94 µL, 0.65 mmol). The reaction was stirred at room temperature for 2.5 hours until judged complete by TLC and GC analysis. The reaction was washed with saturated sodium chloride solution (10 mL), the aqueous layer was extracted twice with dichloromethane (3 mL), and the organic extracts were dried over sodium sulfate and then concentrated. The residue was purified by column chromatography (silica gel, 1:2 ethyl acetate/hexanes) to afford **12a** (111 mg, 73%). <sup>1</sup>H NMR (500.13 MHz, CDCl<sub>3</sub>)  $\delta$  5.09 (q, J=6.85 Hz, 1H), 4.18–4.10 (m, 4H), 2.35 (s, 3H), 1.87 (dt, J=15.04 Hz, J=7.52 Hz, 2H), 1.51–1.43 (m, 2H), 1.37–1.29 (m, 10H), 0.93–0.85 (m, 3H). <sup>13</sup>C NMR (125.77 MHz, CDCl<sub>3</sub>)  $\delta$  184.1, 88.4, 85.5, 67.6, 64.6, 36.2, 36.1, 33.0, 31.5, 24.7, 22.8, 16.6, 16.5, 14.4; LRMS m/z (M<sup>+</sup>+H, 275), 155 (100), 99 (100).

15a

The preparation of 15a is representative. To a solution of 2-ethynyl pyridine (300  $\mu$ L, 3 mmol) in THF (10 mL) at -78°C was added, dropwise, <sup>n</sup>BuLi (2.67M, 1.2 mL, 3.3 mmol). The reaction was stirred for a further 5 minutes before being brought to 0°C and stirred for 20 minutes. The reaction was returned to  $-78^{\circ}$ C and butyraldehyde (280  $\mu$ L, 3.3 mmol) was added slowly. The reaction was brought to and stirred at 0°C for 30 minutes and treated with chlorodiethylphosphate (564  $\mu$ L, 3.9 mmol). The reaction was brought to room temperature and stirred for an hour, until TLC analysis indicated the reaction was complete. The reaction was diluted with ether (90 mL) and poured into sat. NH<sub>4</sub>Cl<sub>(aq)</sub> (75 mL). The organic phase was washed (sat. NH<sub>4</sub>Cl<sub>(aq)</sub>, 75 mL), dried (MgSO<sub>4</sub>), filtered, and concentrated. The residue was purified by column chromatography (silica gel, EtOAc) to afford 15a as a viscous oil (880 mg, 2.8 mmol, 94%). <sup>1</sup>H NMR (500.13 MHz, CDCl<sub>3</sub>) δ 8.24 (d, *J*=4.0 Hz, 1H), 7.36 (tt, *J*=7.8 Hz, J=1.9 Hz, 1H), 7.11 (d, J=7.9 Hz, 1H), 6.92–6.97 (m, 1H), 4.87–4.93 (m, 1H), 3.78–3.90 (m, 4H), 1.52–1.67 (m, 2H), 1.21–1.30 (sext, J=7.3 Hz, 2H), 0.98–1.04 (m, 6H), 0.65 (t, J=7.4 Hz, 3H); <sup>13</sup>C NMR (125.76 MHz, CDCl<sub>3</sub>) δ 149.3, 141.6, 135.5, 126.5, 122.6, 85.4, 84.9, 67.2 (d, J=5.5 Hz), 63.2 (d, J=5.5 Hz), 63.1 (d, J=5.5 Hz), 37.7 (d, J=5.5 Hz), 17.4, 15.3 (d, J=5.5 Hz)Hz), 12.8.

39

To a round bottomed two-necked flask were successively added anhydrous THF (10 mL) and 3,3-dimethyl-1-butyne (0.62 mL, 5.0 mmol). The solution was cooled to  $-78^{\circ}$ C and  $^{n}$ BuLi (2.5M, 2.1 mL, 5.2 mmol) in hexanes was added dropwise. The resulting solution was allowed to warm to room temperature. To an additional 50 mL round bottomed two-necked flask the appropriate dione (4.8 mmol) was dissolved in anhydrous THF (10 mL). The dione solution was cooled to  $-78^{\circ}$ C and the acetylide solution from the first flask was added slowly via cannula to the dione solution at  $-78^{\circ}$ C. The resulting solution was allowed to warm to room temperature, then cooled down again to  $-78^{\circ}$ C and acetic anhydride (0.76 mL, 8.0 mmol) was added. The reaction was brought to room temperature and the solution was then poured into a separatory funnel containing saturated NH<sub>4</sub>Cl<sub>(aq)</sub> (200 mL) and diethyl ether (50 mL). After extraction, the organic layer was separated, dried (MgSO<sub>4</sub>), concentrated, and purified (silica gel) to afford pure acetoxy propargyl ketone **39**.

41

To a round bottomed, two-necked flask were successively added anhydrous THF (10 mL) and 3,3-dimethyl-1-butyne (0.62 mL, 5.0 mmol). The solution was cooled to -78 °C while stirring and <sup>n</sup>BuLi (2.5 M, 2.2 mL, 5.5 mmol) was added dropwise. The flask was removed from the cooling bath and allowed to warm to room temperature. Benzil (1.05 g, 5.0 mmol) was dissolved in anhydrous THF (10 mL) in a 50 mL round bottomed, two-necked flask. The solution was cooled to -78°C, and the acetylide solution from the first flask was transferred dropwise via cannula. The resulting purple-black solution was allowed to warm to room temperature while stirring and then cooled to -78 °C again. Diethylchlorophosphate (0.8 mL, 5.5 mmol) and anhydrous triethylamine <sup>31</sup> (0.8 mL, 5.5 mmol) were successively added. The solution was allowed to return to room temperature. The resulting amber solution was then poured into a separatory funnel containing 300 mL saturated NH<sub>4</sub>Cl solution and 50 mL diethyl ether. After thorough extraction, the organic layer was separated, dried (MgSO<sub>4</sub>), and concentrated. The resulting residue was purified by column chromatography (silica gel, 1:3 ethyl acetate/hexanes) to afford pure 41 (653 mg, 30%) and a second fraction of slightly contaminated **41** (878 mg, 41%). <sup>1</sup>H NMR (500.13 MHz, CDCl<sub>3</sub>) δ 7.89 (d, *J*=8.4 Hz, 2H), 7.70 (dd, J=7.8 Hz, J=0.9 Hz, 2H), 7.43-7.34 (m, 4H), 7.26 (t, J=7.7 Hz, 2H), 4.24-4.10 (m, 2H), 4.07-3.97 (m, 2H), 1.31 (t, J=7.1 Hz, 3H), 1.24 (t, J=8.1 Hz, 3H), 1.23 (s, 9H);  $^{13}$ C NMR  $(125.76 \text{ MHz}, \text{CDCl}_3) \delta 191.6, 137.8, 134.1, 133.0, 131.0, 129.7, 129.1, 128.0, 127.8, 103.8,$ 

82.6, 75.3, 64.1 (d, J=38.9 Hz), 64.1 (d, J=39.0 Hz), 30.6, 28.3, 16.5 (d, J=12.7 Hz), 16.4 (d, J=12.8 Hz); <sup>31</sup>P NMR (202.46 MHz)  $\delta$  -7.30; LRMS m/z 428 (M<sup>+</sup>, 15), 155 (66), 105 (PhCO, 100).

45

To a 25 mL round-bottomed, two-necked flask were successively added anhydrous THF (10 mL) and 3,3-dimethyl-1-butyne (5.0 mmol, 0.62 mL). The solution was allowed to cool to -78°C while stirring and <sup>n</sup>BuLi (2.5M, 2.2 mL, 5.5 mmol) in hexanes was added dropwiseThe reaction was allowed to warm to room temperature. Sepearately, a round bottomed, two-necked 50 mL flask was successively loaded with benzil (5.0 mmol, 1.05 g) and anhydrous THF (10 mL). The solution was stirred and cooled to  $-78^{\circ}$ C, and the acetylide solution was then transferred dropwise to the benzil solution via cannula. The resulting solution was allowed to reach room temperature before being returned to -78°C and treated sequentially with ptoluenesulfonyl chloride (1.05 g, 5.5 mmol) in anhydrous THF (1 mL) and anhydrous triethylamine (0.77 mL, 5.5 mmol) were successively added. The flask was then removed from the cooling bath and allowed to warm to room temperature. The solution was then poured into a separatory funnel containing saturated NH<sub>4</sub>Cl<sub>(aq)</sub> (300 mL) and diethyl ether (50 mL). After thorough extraction, the organic layer was separated, dried (MgSO<sub>4</sub>), and concentrated. The residue was purified by column chromatography(silica gel, 1:10 ethyl acetate/hexanes) to afford pure 45 (1.32 g, 60%) as a yellow oil which solidifies upon refrigeration. <sup>1</sup>H NMR  $(500.13 \text{ MHz}, \text{CDCl}_3) \delta 7.81 \text{ (dd, } J=8.3 \text{ Hz, } J=1.2 \text{ Hz, } 2\text{H}), 7.63 \text{ (d, } J=8.4 \text{ Hz, } 2\text{H}), 7.56 \text{ (t, } J=8.4 \text{ Hz, } 2\text{H}), 7.56 \text{ (t, } J=8.4 \text{ Hz, } 2\text{H}), 7.56 \text{ (t, } J=8.4 \text{ Hz, } 2\text{Hz, } 2\text{$ J=7.4 Hz, 1H), 7.41 (t, J=7.8 Hz, 2H), 7.32–7.38 (m, 5H), 7.10 (d, J=7.9 Hz, 2H), 2.35 (s, 3H), 1.01 (s, 9H); <sup>13</sup>C NMR (125.76 MHz, CDCl<sub>3</sub>) δ 196.6, 192.2, 145.1, 137.0, 136.2, 133.3, 132.8, 132.7, 129.6, 129.5, 128.8, 128.6, 128.3, 128.1, 120.5, 36.0, 27.4, 21.6.

## **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgment

The support of the National Institutes of Health (GM-64444) is gratefully acknowledged.

#### References

- a Bellina F, Rossi R. Tetrahedron 2006;62:7213.Hou, XL.; Yang, Z.; Wong, HNC. Progress in Heterocyclic Chemistry. Gribble, GW.; Gilcrist, TL., editors. 15. Pergamon; Oxford: 2003. p. 167Keay, BA.; Dibble, PW. Comprehensive Heterocyclic Chemistry II. Katritzky, AR.; Rees, CW.; Scriven, EFV., editors. 2. Elsevier; Oxford: 1997. p. 395Donnelly, D.; M. X.; Meegan, MJ. Comprehensive Heterocyclic Chemistry. Katritzky, AR.; Rees, CW., editors. 4. Pergamon; Oxford: 1984. p. 657Dean, FM. Naturally Occuring Oxygen Ring Compounds. Butterworths; London: 1963. p. 1Chapter 1
- a Lee H-K, Chan K-F, Hui C-W, Yim H-K, Wu X-W, Wong HNC. Pure Appl. Chem 2005;77:139. b Wong HNC, Yu P, Yick C-Y. Pure Appl. Chem 1999;71:1041. c Lipshutz BH. Chem. Rev 1986;86:795.
- 3. a Ferrero F, Napoli L, Tonin C, Varesano A. J. Appl. Polym. Sci 2006;102:4121. b Venkatatraman S, Kumar R, Sankar J, Chandrashekar TK, Senhil K, Vijayan C, Kelling A, Senge MO. Chem. Eur. J 2004;10:1423. c Facchetti A, Abbotto A, Beverina L, van der Boom ME, Dutta P, Evmenenko G, Pagani GA, Marks TJ. Chem. Mater 2003;15:1064. d Zhang L-Z, Chen C-W, Lee C-F, Wu C-C, Luh T-Y. Chem. Commun 2002:2336. e Lee C-F, Yang L-M, Hwu T-Y, Feng A-S, Tseng J-C, Luh T-Y. J. Am. Chem. Soc 2000;122:4992. f Novak P, Müller K, Santhanum KSV, Haas O. Chem. Rev 1997;97:207. [PubMed: 11848869]
- 4. For recent reviews, see: a Patil NT, Yamamoto Y. ARKIVOC 2007;10:121. b Kirsch SF. Org. Biomol. Chem 2006;4:2076. [PubMed: 16729118] c Brown RCD. Angew. Chem., Int. Ed 2005;44:850. d

Jeevanandam A, Ghule A, Ling Y-C. Curr. Org. Chem 2002;6:841. e Keay BA. Chem. Soc. Rev 1999;28:209. f Hou XL, Cheung HY, Hon TY, Kwan PL, Lo TH, Tong SY, Wong HNC. Tetrahedron 1998;54:1955.

- 5. For recent examples, see: a Galliford CV, Scheidt KA. J. Org. Chem 2007;72:1811. [PubMed: 17256992] b St. Cyr DJ, Martin N, Arndtsen BA. Org. Lett 2007;9:449. [PubMed: 17249784] c Zhang J, Schmalz H-G. Angew. Chem., Int. Ed 2006;45:6704. d Gorin DJ, Davis NR, Toste FD. J. Am. Chem. Soc 2005;127:11260. [PubMed: 16089452] e Kamijo S, Kanazawa C, Yamamoto Y. J. Am. Chem. Soc 2005;127:9260. [PubMed: 15969607] f Yao T, Zhang X, Larock RC. J. Org. Chem 2005;70:7679. [PubMed: 16149799] g Larionov OV, de Meijere A. Angew. Chem., Int. Ed 2005;44:5664. h Duan, X.-h.; Liu, X.-y.; Guo, L.-n.; Liao, M.-c.; Liu, W-M.; Liang, Y.-m. J. Org. Chem 2005;70:6980. [PubMed: 16095331] i Lee KY, Lee MJ, Kim JN. Tetrahedron 2005;61:8705. j Jung C-K, Wang J-C, Krische MJ. J. Am. Chem. Soc 2004;126:4118. [PubMed: 15053596] k Dhawan R, Arndtsen BA. J. Am. Chem. Soc 2004;126:468. [PubMed: 14719935] l Nishibayashi Y, Yoshikawa M, Inada Y, Milton MD, Hidai M, Uemura S. Angew. Chem., Int. Ed 2003;42:2681. m Braun RU, Zeitler K, Müller TJJ. Org. Lett 2001;3:3297. [PubMed: 11594818] n Brown CD, Chong JM, Shen L. Tetrahedron 1999;55:14233. o Shiraishi H, Nishitani T, Sakaguchi S, Ishii Y. J. Org. Chem 1998;63:6234. [PubMed: 11672254]
- 6. For recent examples, see: a Sniady A, Durham A, Morreale MS, Wheeler KA, Dembinski R. Org. Lett 2007;9:1175. [PubMed: 17328552] b Smith CR, Bunnelle EM, Rhodes AJ, Sarpong R. Org. Lett 2007;9:1169. [PubMed: 17309277] c Díaz-Álvarez AE, Crochet P, Zablocka M, Duhayon C, Cadierno V, Gimeno J, Majoral JP. Adv. Synth. Catal 2006;348:1671. d Zhou C-Y, Chan PWH, Che C-M. Org. Lett 2006;8:325. [PubMed: 16408906] e Kawai H, Oi S, Inoue Y. Heterocycles 2006;67:101. f Sromek AW, Rubina M, Gevorgyan V. J. Am. Chem. Soc 2005;127:10500. [PubMed: 16045332] g Suhre MH, Reif M, Kirsch SF. Org. Lett 2005;7:3925. [PubMed: 16119933] h Sromek AW, Kel'in AV, Gevorgyan V. Angew. Chem., Int. Ed 2004;43:2280. i Hashmi ASK, Sinha P. Adv. Synth. Catal 2004;346:432. j Ma S, Zhang J. J. Am. Chem. Soc 2003;125:12386. [PubMed: 14531663] k Gabriele B, Salerno G, Fazio A. J. Org. Chem 2003;68:7853. [PubMed: 14510566] I Ma S, Zhang J. Angew. Chem., Int. Ed 2003;42:183. m Kim JT, Kel'in AV, Gevorgyan V. Angew. Chem., Int. Ed 2003;42:98. n Kel'in AV, Gevorgyan V. J. Org. Chem 2002;67:95. [PubMed: 11777444] o Aurrecoechea JM, Pérez E, Solay M. J. Org. Chem 2001;66:564. [PubMed: 11429830] p Kel'in AV, Sromek AW, Gevorgyan V. J. Am. Chem. Soc 2001;123:2074. [PubMed: 11456838] q Gabriele B, Salerno G, Fazio A, Bossio MR. Tetrahedron Lett 2001;42:1339. r Hashmi ASK, Schwarz L, Choi J-H, Frost TM. Angew. Chem., Int. Ed 2000;39:2285. s Gabriele B, Salerno G, Lauria E. J. Org. Chem 1999;64:7687. t Hashmi ASK, Ruppert TL, Knöfel T, Bats JW. J. Org. Chem 1997;62:7295. [PubMed: 11671843] u Arcadi A, Marinelli F, Pini E, Rossi E. Tetrahedron Lett 1996;37:3387. v Vieser R, Eberbach W. Tetrahedron Lett 1995;35:4405. w Seiller B, Bruneau C, Dixneuf PH. 1995;51:13089. x Fukuda Y, Shiragami H, Utimoto K, Nozaki H. J. Org. Chem 1991;56:5816. y Marshall JA, Robinson ED. J. Org. Chem 1990;55:3450.
- a Zhao J, Hughes CO, Toste FD. J. Am. Chem. Soc 2006;128:7436. [PubMed: 16756286] b Zhang L, Wang S. J. Am. Chem. Soc 2006;128:1442. [PubMed: 16448102] c Wang S, Zhang L. Org. Lett 2006;8:4585. [PubMed: 16986956] d Wang S, Zhang L. J. Am. Chem. Soc 2006;128:8414. [PubMed: 16802803] e Buzas A, Gagosz F. J. Am. Chem. Soc 2006;128:12614. [PubMed: 17002332] f Zhang L. J. Am. Chem. Soc 2005;127:16804. [PubMed: 16316224] g Schlossarczyk H, Sieber W, Hesse M, Hansen H-J, Schmid H. Helv. Chim. Acta 1973;56:875.
- 8. Preliminary results of this work (Schemes 4, 8, and 9, Table 1 and portions of Table 3) have been previously communicated (see ref. <sup>6h</sup>). All other results, described herein (Schemes 5-7, 10-16, Tables 2, 4, 5, and part of Table 3) are new.
- 9. See Supporting Information for synthesis of starting materials.
- 10. For reviews on <sup>17</sup>O NMR, see: BoykinDW<sup>17</sup>O NMR Spectroscopy in Organic Chemistry1991CRC PressBoca Raton, FLbMcFarlaneWMcFarlaneHCEMasonJMultinuclear NMR1987403Plenum PressNew York, NYChapter 14cRodgerCSheppardNMcFarlaneCMcFarlaneWHarrisRKMannBENMR and the Periodic Table1978383Academic PressNew York, NYChapter 12dKintzingerJPLaszloPNMR of Newly Accessible Nuclei1983279Academic PressNew York, NYChapter 4 e Boykin DW, Baumstark AL. Tetrahedron 1989;45:3613.

11. For discussions on <sup>17</sup>O NMR of phosphates in cyclic and biological systems, see:aTsaiM-DBruzikKBerlinerLJReubenJBiological Magnetic Resonance19835129Plenum PressNew York, NYChapter 4 b Gordillo B, Eliel EL. J. Am. Chem. Soc 1991;113:2172. c Sopchik AE, Cairns SM, Bentrude WG. Tetrahedron Lett 1989;30:1221. d Eliel EL, Chandrasekaran S, Carpenter LE II, Verkade JG. J. Am. Chem. Soc 1986;108:6651. e Sammons RD, Frey PA, Bruzik K, Tsai M-D. J. Am. Chem. Soc 1983;105:5455.

- 12. See ref <sup>6m</sup> for details.
- For Rautenstrauch-type 1,2-acyloxy migrations, see:a Marion N, Nolan SP. Angew. Chem., Int. Ed 2007;46:2750. b Marion N, Díez-Gonzalález S, de Frémont P, Noble AR, Nolan SP. Angew. Chem., Int. Ed 2006;45:3647. c Fehr C, Galindo J. Angew. Chem., Int. Ed 2006;45:2901. d Pujanauski BG, Prasad BAB, Sarpong R. J. Am. Chem. Soc 2006;128:6786. [PubMed: 16719448] e Marco-Contelles J, Arroyo N, Anjum S, Mainetti E, Marion N, Cariou K, Lemière G, Mouriès V, Festerbank L, Malacria M. Eur. J. Org. Chem 2006:4618. f Faza ON, López CS, Álvarez R, de Lera AR. J. Am. Chem. Soc 2006;128:2434. [PubMed: 16478199] g Fürstner A, Hannen P. Chem. Eur. J 2006;12:3006. h Ohe K, Fujita M, Matsumoto H, Tai Y, Miki K. J. Am. Chem. Soc 2006;128:9270. [PubMed: 16848428] i Cho EJ, Kim M, Lee D. Eur. J. Org. Chem 2006:3074. j Kim M, Miller RL, Lee D. J. Am. Chem. Soc 2005;127:12818. [PubMed: 16159273] k Johansson MJ, Gorin DJ, Staben ST, Toste FD. J. Am. Chem. Soc 2005;127:18002. [PubMed: 16366541] l Shi X, Gorin DJ, Toste FD. J. Am. Chem. Soc 2005;127:5802. [PubMed: 15839674] m Prasad BAB, Yoshimoto FK, Sarpong R. J. Am. Chem. Soc 2005;127:12468. [PubMed: 16144376] n Mamane V, Gress T, Krause H, Fürstner A. J. Am. Chem. Soc 2004;126:8654. [PubMed: 15250709] o Miki K, Ohe K, Uemura S. J. Org. Chem 2003;68:8505. [PubMed: 14575478] p Rautenstrauch V. J. Org. Chem 1984;49:950.
- 14. For migrations in β-(acyloxy)alkyl radicals, see:a Crich D, Sartillo-Piscil F, Quintero-Cortes L, Wink DJ. J. Org. Chem 2002;67:3360. [PubMed: 12003547] b Zipse H, Bootz M. J. Chem. Soc., Perkin Trans. 2 2001:1566. c Crich D, Huang X, Beckwith ALJ. J. Org. Chem 1999;64:1762. [PubMed: 11674254] d Beckwith ALJ, Duggan PJ. J. Am. Chem. Soc 1996;118:12838.
- 15. For selected examples of metal-catalyzed 1,2-migrations of nonacyloxy groups, see refs <sup>6f,m</sup> anda Peng L, Zhang X, Zhang S, Wang J. J. Org. Chem 2007;72:1192. [PubMed: 17288372] b Seregin IV, Gevorgyan V. J. Am. Chem. Soc 2006;128:12050. [PubMed: 16967938] c Xu F, Shi W, Wang J. J. Org. Chem 2005;70:4191. [PubMed: 15876118] d Maiereanu C, Kanai A, Weibel J-M, Pale P. J. Carbohydr. Chem 2005;24:831.
- a Allen AD, Kitamura T, McClelland RA, Stang PJ, Tidwell TT. J. Am. Chem. Soc 1990;112:8873.
   b Lorenz W, Maas G. J. Org. Chem 1987;52:375.
- 17. For base-assisted propargyl-allenyl isomerization, see refs <sup>6h,n,p</sup> and Sonye JP, Koide K. J. Org. Chem 2006;71:6254. [PubMed: 16872214] b Huang J, Xiong H, Hsung RP, Rameshkumar G, Mulder JA, Grebe TP. Org. Lett 2002;4:2417. [PubMed: 12098261]
- 18. For Lewis acid-catalyzed substitution of acyloxy groups in propargyl systems, see: a Netz A, Polborn K, Nöth H, Müller TJJ. Eur. J. Org. Chem 2005:1823. b Schwier T, Rubin M, Gevorgyan V. Org. Lett 2004;6:1999. [PubMed: 15176803] c Müller TJJ. Eur. J. Org. Chem 2001:2021.
- 19. Orsini A, Viterisi A, Bodlenner A, Weibel J-M, Pale P. Tetrahedron Lett 2005;46:2259.
- 20. This is in contrast to the analogous acyloxy system where attempts at isolation or independent preparation of the allenes failed.
- 21. Triflic acid was tested as a potential catalyst in the base-free reactions depicted in Schemes 1 and 2, but failed to result in product formation.
- 22. There has been a recent discussion on the role of Brønsted acids in homogenous transition metal-catalyzed reactions. For the most relevant references, see:a Hashmi ASK. Catal. Today 2007;122:211. b Li Z, Zhang J, Brouwer C, Yang C-G, Reich NW, He C. Org. Lett 2006;8:4175. [PubMed: 16956180] c Rosenfeld DC, Shekhar S, Takemiya A, Utsunomiya M, Hartwig JF. Org. Lett 2006;8:4179. [PubMed: 16956181] d Rhee JU, Krische MJ. Org. Lett 2005;7:2493. [PubMed: 15932231]
- 23. The authors wish to acknowledge one of the referees, who brought our attention to this possibility.
- For discussions on propargyl/allenyl cations, see:a Olah GA, Krishnamurti R, Prakash GKS. J. Org. Chem 1990;55:6061. b Olah GA, Spear RJ, Westerman PW, Denis J-M. J. Am. Chem. Soc 1974;96:5855.
- 25. For Lewis acid-catalyzed allylation of propargyl acetates with allylsilanes, see: refs <sup>18b,c</sup>.

 $26. \ For mechanistic \ discussions \ on \ regions electivity \ of \ nucleophilic \ attacks \ on \ proparyl \ cations, \ see: \ ref \ 18c$ 

- 27. Due to severe signal overlap, quantification of isotopomers was not possible.
- 28. For Kumada cross-coupling of enol phosphates, see:a Kobayashi Y, Takeuchi A, Wang Y-G. Org. Lett 2006;8:2699. [PubMed: 16774235] b Larsen US, Mariny L, Begtrup M. Tetrahedron Lett 2005;46:4261. c Verboom RC, Persson BA, Baeckvall J-E. J. Org. Chem 2004;69:3102. [PubMed: 15104449] d Hayashi T, Konishi M, Okamoto Y, Kabeta K, Kumada M. J. Org. Chem 1986;51:3772. e Sahlberg C, Quader A, Claesson A. Tetrahedron Lett 1983;24:5137. f Hayashi T, Fujiwa T, Okamoto Y, Katsuro Y, Kumada M. Synthesis 1981:1001.
- 29. For Kumada cross-coupling of aryl phosphates, see:a Huang WG, Jiang YY, Li Q, Li J, Li JY, Lu W, Cai JC. Tetrahedron 2005;61:1863. b Hayashi T, Katsuro Y, Okamoto Y, Kumada M. Tetrahedron Lett 1981;22:4449.
- 30. (R)-(-)-Di-*tert*-butyl-{1-[(*S*)-2-(dicyclohexylphosphanyl)ferrocenyl]ethyl}phosphine [158923–11 –6], Strem catalog number 26–0975. For examples on use, see:a Fernández-Rodríguez MA, Shen Q, Hartwig JF. Chem. Eur. J 2006;30:7782. b Shen Q, Shekhar S, Stambuli JP, Hartwig JF. Angew. Chem., Int. Ed 2005;44:1371.
- 31. Mikami K, Yoshida A. Tetrahedron 2001;57:889.

EtO 
$$P = O$$

O  $R^2$ 

OP(O)(OEt)<sub>2</sub>

OP(O)(OET)

OP(O)(OET)

OP(OET)

O

 $^aRun$  with 5 mol%  $AgBF_4$  in dichloroethane at  $80^{\circ}C$ 

## Scheme 1.

Cascade [3,3]-migration/cycloisomerization of phosphatyloxy alkynyl ketones into furans

EtO 
$$P=0$$
  $CuCl (5 mol\%)$   $DMA, 110-130 °C$   $P=0$   $P=$ 

**Scheme 2.** Cascade [3,3]-migration/cycloisomerization of phosphatyloxy alkynyl pyridines into indolizines

**Scheme 3.** Cyloisomerization of labeled phosphatyloxy alkynyl pyridine **18** 

Scheme 4.

prototopic 
$$1,3-H \sim$$
 $R^1$ 
 $R^2$ 
 $R^3$ 
 $R^4$ 
 $R^2$ 
 $R^3$ 
 $R^4$ 
 $R^4$ 

**Scheme 5.** Proposed mechanisms for cycloisomerization of acyloxy alkynyl ketones into furans via dioxolenylium intermediates

prototopic 
$$1,3-H \sim$$
 $R^1$ 
 $R^2$ 
 $R^3$ 
 $R^3$ 
 $R^4$ 
 $R^2$ 
 $R^3$ 
 $R^3$ 

## Scheme 6.

Alternative mechanisms for cycloisomerization of acyloxy alkynyl ketones into furans via oxirenium intermediates

**Scheme 7.** Cycloisomerization of <sup>17</sup>O-labeled acyloxy alkynyl ketone **33** 

**Scheme 8.** Sequential phosphatyloxy migration/cycloisomerization

**Scheme 9.** Sequential tosyloxy migration/cycloisomerization

Scheme 10.

Proposed mechanisms for sequential migration/cycloisomerization of labeled 47

$$R^{1} \xrightarrow{O} R^{2} \xrightarrow{\text{ionization}} R^{1} \xrightarrow{\text{55}} R^{2} \xrightarrow{R^{3}} R^{1} \xrightarrow{\text{37}} R^{3}$$

Scheme 11.
Proposed mechanisms for formation of allene 37 from 36

Scheme 12. Allylation of acetate 39b

Scheme 13. Sequential migration/cycloisomerization of phosphoryl oxygen-labeled **59** 

EtO OEt EtO OEt 
$$R^1$$
 OEt  $R^2$   $R^3$  OEt  $R^$ 

Scheme 14.

Mechanisms for formation of phosphatyloxy-substituted allene 67

EtO

EtO

$$R^{2}$$
 $R^{1}$ 
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{1}$ 
 $R^{2}$ 
 $R^{2}$ 

Scheme 15.
Proposed mechanisms for conversion of allene 67 to furans 72 and 74

$$[3,3] \sim Y=O$$

$$R^{2}$$

$$R^{1}$$

$$37$$

$$R^{1}$$

$$36$$

$$R^{2}$$

$$R^{3}$$

$$R^{1}$$

$$37$$

$$R^{2}$$

$$R^{3}$$

$$R^{3}$$

$$R^{4}$$

$$R^{2}$$

$$R^{3}$$

$$R^{4}$$

$$R^{2}$$

$$R^{3}$$

$$R^{4}$$

$$R^{5}$$

$$R^{4}$$

$$R^{5}$$

$$R^{5}$$

$$R^{6}$$

$$R^{7}$$

Scheme 16.
Generalized mechanism for migration/cycloisomerization of 36 to 38

Table 1
Cascade 1,2-migration/cycloisomerization of acyloxy alkynyl ketones 22 into furans 23

		yield, % <sup>a</sup>	82	8	69	06	98	08	80	83
	${\nearrow}_{R^3}$	product 23	Phcoo							Phedo O TBSO
CuCl (5 mol%)  R <sup>3</sup> Et <sub>3</sub> N (20 mol%)	DMA, 130 °C R <sup>2</sup>	time, h	c 22	- P	6.	<b>,</b> 2	on 17	<b>h</b> 23	i 32	j 46
R <sup>1</sup> 0	R <sup>2</sup> <b>22</b>	substrate 22	PhCOO Ph	MeCOO Ph	EtCOO Ph O	iPrCOO Me	'Buccoo Ph	Phcoo Ph C <sub>5</sub> H <sub>11</sub>	$^{t}BuCOO$ $C_{5}H_{11}$ $O$	Phcoo Ph OTBS
		entry	-	0	ю	4	N	9	L	∞

Schwier et al.	
	yield, % <sup>a</sup>
	product 23
CuCl (5 mol%)  R <sup>3</sup> Et <sub>3</sub> N (20 mol%)  DMA, 130 °C R <sup>2</sup> DMA, 130 °C R <sup>2</sup> S	time, h
$\begin{array}{c} R^1 \\ \searrow \\ \bigcirc \\ \searrow \\ R^2 \end{array}$	substrate 22
	entry

<sup>a</sup>Run on 1.0 mmol scale

 Table 2

 Optimization of acyloxy migration/cycloisomerization

entry	$\mathbb{R}^1$	[M] (mol%)	solvent	yield 40, % <sup>a</sup>
1	<sup>n</sup> Bu ( <b>a</b> )	Cu(OTf) <sub>2</sub> (5)	PhMe	78
2	<sup>t</sup> Bu ( <b>b</b> )	"	"	66
3	<sup>n</sup> Bu (a)	$PdCl_2(MeCN)_2$ (5)	"	72
4	"	$AgBF_4(5)$	DCM	$50^{b}$
5	<sup>t</sup> Bu ( <b>b</b> )	" <del>"</del>	"	99
6	<sup>n</sup> Bu ( <b>a</b> )	AgOTf (5)	PhMe	60
7	<sup>t</sup> Bu ( <b>b</b> )	u v	"	88
8	<sup>n</sup> Bu (a)	$AgSb_{6}(5)$	"	50
9	"	$PtCl_{2}(5)$	"	43
10	<sup>t</sup> Bu ( <b>b</b> )	AuCl <sub>3</sub> (5)	"	72
11	<sup>n</sup> Bu (a)	AuCl <sub>3</sub> (5)	"	62
12	"	$AuCl(PEt_3)(5) + AgPF_6(5)$	"	complex mixture
13	<sup>t</sup> Bu ( <b>b</b> )	TMSOTf (10)	DCM	63
14	"	$Sn(OTf)_2$ (10)	"	72
15	"	$Sc(OTf)_3(10)$	"	65
16	"	$Zn(OTf)_2$ (10)	"	56
17	"	HOTf (10)	"	80

Remarkably, this cascade method allowed for efficient synthesis of fused furan 40g (entry 7), bicyclic scaffold inaccessible by our previous cycloisomerization techniques. 6n

aNMR yield.

 $<sup>^</sup>b\mathrm{Run}$  at 0°C.

Table 3
Sequential [3,3]/1,2-acyloxy migration/cycloisomerization

			yield, %a	66<	73b	70 <sup>b</sup> 73	84	06	98	<sub>0</sub> 09
		R <sup>2</sup>	product 40	Aco Ph						Aco Ph
oí	(s) R <sup>4</sup> (S)	R1 0 40	time, min	2	180	600	15	15	10	15
	cat. (5 mol%)	CH <sub>2</sub> Cl <sub>2</sub> , π	cat.	$\mathrm{AgBF}_4$	$AgBF_4$	${\rm TMSOTf} \\ {\rm AgBF}_4$	$AgBF_4$	$\mathrm{AgBF}_4$	$\mathrm{AgBF}_4$	Cu(OTf) <sub>2</sub>
PA O		0 0 PR <sup>2</sup>		a a	ပ	יסי	ψ.	<b>-</b>	ත	a
	R.	39	substrate 39	tBu————Ph	$^{OO}_{OPv}$	tBu	tBu	tBu———OAc	tBu	"Bu-"Ph
			entry	-	6	w 4	vo	ø	7	œ

			l	I		
			yield, $%a$	<sub>9</sub> 18	78, <sup>bc</sup>	916
		2)	product 40	Pvo Ph	Aco Ph TMS O Ph	Aco Ph Ph
0	¥ <sup>7</sup>	$R^{1}$ $^{\circ}$ $^{\circ}$ $R^{2}$	time, min	180	1.5	1.5
	cat. (5 mol%)	СН <sub>2</sub> СІ <sub>2</sub> , п	cat.	Cu(OTf) <sub>2</sub>	$AuCl_3$	Cu(OTf) <sub>2</sub>
0 4×4	$\begin{pmatrix} R_3 \\ R_3 \end{pmatrix}$	O R		ਥ		<del></del>
0	R <sup>1</sup>	39	substrate 39	OPv nBu ———Ph Ph	TMS-FPh	Ph———Ph
			entry	6	10	Ξ

<sup>a</sup>Run on 1.0 mmol scale.

bRun on 0.5 mmol scale.

 $^{\mathcal{C}}$ Run in toluene.

Table 4

## Cycloisomerization of carbonyl labeled 52

entry	cat (mol%)	solvent	ratio (53:54)	yield, %
1	CuCl (5) + Et <sub>3</sub> N (20)	DMA	38:62	89, <sup>ab</sup>
2	$AgBF_4(5)$	DCM	0:100	69
3	AuCl <sub>3</sub> (10)	PhMe	0:100	64
4	Cu(OTf) <sub>2</sub> (10)	PhMe	0:100	84
5	$Sn(OTf)_2$ (10)	DCM	38:62	42
6	TMSOTf (10)	"	42:58	51 <sup>a</sup>
7	HOTf (10)	"	26:74	80
8	$AuCl_3(5) + TTBP(10)$	"	0:100	84

 $<sup>{}^{</sup>a}{\rm NMR}~{\rm of~recovered~starting~material~showed~no~formation~of~another~isotopomer.~See~Supporting~Material~for~details.}$ 

 $<sup>^</sup>b$ Run at 130°C.

 Table 5

 Effect of Brønsted base on cycloisomerization with different catalysts

entry	cat (mol%)	% conversion of 39b <sup>a</sup> (no base)	% conversion of $39b^a$ (with TTBP $^b$ )
1	TfOH (10)	100 <sup>c</sup>	<1 <sup>d</sup>
2	$Cu(OTf)_2$ (5)	$100^{e}$	$100^{e}$
3	$PdCl_2(MeCN)_2$ (5)	$100^{e}$	$100^{e}$
4	AgOTf (5)	$100^{e}$	$100^{e}$
5	PtCl <sub>2</sub> (5)	$100^f$	$100^{f}$
6	AuCl <sub>3</sub> (5)	$100^{e}$	$100^{e}$

 $<sup>\</sup>ensuremath{a}\xspace$  based on GC analysis. Yields comparable to that reported in Table 2.

 $<sup>^</sup>b 1.5 \ \mathrm{eq}$  (vs catalyst) 2,4,6-tri-tertbutylpyrimidine (TTBP) added.

c reaction complete after 6 hours.

 $<sup>\</sup>frac{d}{d}$  reaction checked after 24 hours.

 $<sup>^{\</sup>it e}$  reaction complete within 30 minutes.

 $f_{
m reaction}$  complete within 3.5 hours.

yield, %

72

97

99

74

 $94^a$ 

Cross-coupling of phosphatyloxy-substituted heteroaromatics

76a 76b **76c 76d** 77a  $^{77}$ 'n product 76 or 77 76 or 77 CyPF-tBu (5 mol%) Pd<sub>2</sub>dba<sub>3</sub> (5 mol%) toluene, 110 °C R³MgX (2 eq) Me Me Bn Ph Bn ĸ ቯ OP(O)(OEt)<sub>2</sub> 14c 17b 14 or 17 OP(O)(OEt)<sub>2</sub> OP(O)(OEt)<sub>2</sub> substrate 14 or 17 Ч entry 7 9

	3	chwier e	t al.			
				yield, %	78	84
					77c	77d
83	7	X-2	<b>76</b> or <b>77</b>	product 76 or 77	-A	C <sub>5</sub> H <sub>11</sub>
26		CyPF-¹Bu (5 mol%) R³MaX (2 ea)	toluene, 110 °C	$\mathbb{R}^3$	Ph	${\sf Me_3SiCH_2}$
OP(O)(OEt) <sub>2</sub>		R1/X/ 	14 or 17		3	:
		_		substrate 14 or 17	3	3
				entry	7	∞

MMD wield