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## Carbocyclization Cascades of Allyl Ketenimines via *Aza-*Claisen Rearrangements of *N-*Phosphoryl-*N-*allyl-ynamides

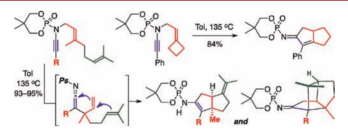
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## **ABSTRACT**



A series of carbocyclization cascades of allyl ketenimines initiated through a thermal aza-Claisen rearrangement of N-phosphoryl-N-allyl ynamides is described. Interceptions of the cationic intermediate via Meerwein—Wagner rearrangements and polyene-type cyclizations en route to fused bi- and tricyclic frameworks are featured.

We recently reported a new class of ynamides<sup>1-3</sup> bearing a phosphoryl group as the required electron-withdrawing component.<sup>4</sup> Most notably, it was demonstrated that N-phosphoryl-N-allyl ynamides 1 [for EWG = PO(OR)<sub>2</sub>] could undergo a thermal 3-aza-Claisen rearrangement<sup>5,6</sup> to generate allyl ketenimine<sup>7</sup> intermediates 2 in situ without suffering a subsequent facile 1,3-shift observed in related N-sulfonyl systems [ $1 \rightarrow 4$  when EWG = ArSO<sub>2</sub>], leading to an effective formation of structurally unique quaternary nitriles  $5^{8,9}$  (Scheme 1). While such a 1,3-sulfonyl shift can be of immense interest, <sup>10</sup> it precluded us from developing a useful carbocyclization of allyl ketenimines  $4^{8,9}$  Consequently, in lieu of a 1,3-phosphoryl shift, we envisioned that carbocyclizations of 2 could take place to afford

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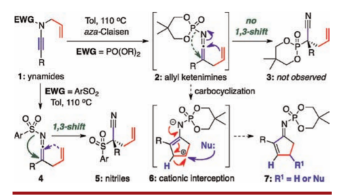
cyclopentenyl zwitter ionic intermediates **6**, which would allow us to construct cyclopentenimine derivatives **7** via a

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Scheme 1. A Carbocyclization of Allyl Ketenimines



1,2-H shift, or more powerfully, nucleophilic trapping<sup>11–13</sup> of **6**. We report herein our success in intercepting these intermediates.

Scheme 2. Feasibility of the Carbocyclization

Our first success with an ynamide-initiated thermal carbocyclization was the rearrangement of ynamide  $\mathbf{8a}$  to  $\alpha,\beta$ -unsaturated cyclopentenimine  $\mathbf{9a}$  in 50% yield (Scheme 2). We were fascinated with this discovery, as it implied that a 1,2-H shift through zwitter ionic intermediate  $\mathbf{6}$  may be in operation. Furthermore, ynamide  $\mathbf{8b}$  also

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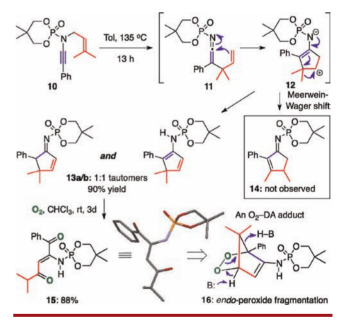
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underwent the tandem *aza*-Claisen rearrangement—carbocyclization to give **9b**, presumably through a tertiary carbocation intermediate.

While it is also possible that deprotonation could lead to 1-amido dienes that then tautomerized to the observed cyclopentenimines, the idea of a 1,2-H shift was enticing. We therefore wondered if other Meerwein—Wagner shifts could occur following the initial *aza*-Claisen rearrangement and carbocyclization (Scheme 3). To explore this possibility, N-prenyl ynamide 10 was heated to 135 °C, with the hopes of demonstrating a 1,2-methyl shift through the formation of cyclopentenimine 14. Unfortunately, 14 was not observed. Instead, a 1:1 tautomeric mixture of 13a and 13b was isolated in 90% yield caused by deprotonation  $\alpha$  to the enamide instead of the desired methyl shift. Remarkably, when air was bubbled through the tautomeric mixture at rt in CHCl<sub>3</sub>, a [4 + 2] cycloaddition of 2-amido diene 13b with O<sub>2</sub> ensued to first give endoperoxide

**Scheme 3.** Attempts at a 1,2-Alkyl Shift: An Unexpected [4 + 2]



16 that subsequently fragmented to the isolated ene-dione 15. While this could be a radical fragmentation, although the reaction conditions involved no base, it is very likely another example of a Kornblum—DeLaMare process. 14,15

The idea of pursuing [4 + 2] cycloadditions with *in situ* generated 2-amido dienes was intriguing; however we were still very interested in intercepting the zwitter ionic intermediates through either Meerwein-Wagner rearrangements or nucleophilic trappings. To explore the former, we prepared ynamides 17 and 21 bearing a tethered

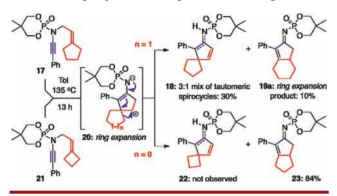
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methylcyclopentylidine and methylcyclobutylidine, respectively, reasoning that the added ring strain should favor ring expansion through zwitter ions 20 (Scheme 4). When 17 was heated to 135 °C, spirocycles 18 resulting from elimination dominated; however bicycle 19a was also isolated in 10% yield representing a successful ring expansion. Moreover, ynamide 21 with increased ring strain yielded ring-expansion product 23 in 84% yield and spirocycle 22 was not observed.

Scheme 4. Ring Expansion through Meerwein-Wagner Shift



We also, rather unexpectedly, discovered a Meerwein—Wagner ring contraction when pursuing the carbocyclization of ynamide **24a** bearing a tethered methylcyclohexene (Scheme 5). In addition to the anticipated 5,6-fused bicycle **19a** resulting from a 1,2-*H* shift, 5,5-spirocycle **26a** was isolated as the major product in 53% yield. It is possible that A<sup>1,2</sup> strain promoted the C–C bond of the fused cyclohexane ring to adopt a pseudoaxial position in

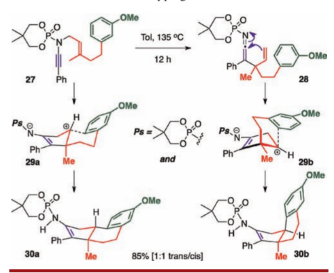
Scheme 5. Ring Contraction through Meerwein-Wagner Shift

**25-CC-ax**, thereby allowing the 1,2-alkyl shift to compete. This notion was furthered by our experimentation with methyl-terminated ynamide **24b**, where the expected 1,2-*H* shift dominated to give **19b** in 75% yield and only 3% of the spirocycle **26b**.

After successfully completing several examples of intercepting the zwitter ionic intermediates via Meerwein—

Wagner rearrangements, we wanted to explore the possibility of using the *aza*-Claisen rearrangement to initiate a carbocyclization cascade with tethered carbon nucleophiles. Gratifyingly, ynamide **27** featuring a *m*-methoxyphenyl moiety tethered to the allyl fragment cleanly underwent the required 3-*aza*-Claisen rearrangement followed by carbocyclization and Friedel—Craft electrophilic aromatic substitution to give **30a** and **30b** as a 1:1 mixture of trans and cis isomers without any competing alkyl shifts (Scheme 6).

Scheme 6. Friedel-Craft Trapping of Cationic Intermediate



The ability to intercept these cationic intermediates with aryl nucleophiles was exciting and propelled us into exploring other nucleophilic trappings. Inspired by the abundance of beautiful work using terpenes in cationic polyene cascades<sup>16,17</sup> we decided to investigate the possibility of an ynamide-initiated carbocyclization cascade with terpene-derived ynamides (Scheme 7).

Starting from commercially available geranylamine 31, a simple two-step protocol involving phosphorylation and Cu-catalyzed amidative cross-coupling gave ynamide 32 in 82% overall yield. When 32 was heated to 135 °C for 12 h, 5,5-cis-fused bicycle 36 bearing an exocyclic olefin was isolated in 55% yield as a single diastereomer. Of note, the cis-fused bicyclic core in 36 is prominent in triquinane 18 natural products. In addition to 36, tricycle 37 featuring four contiguous stereocenters and three all-carbon quaternary centers was isolated in 38% yield as a single diastereomer. The geometry of both products was determined by NOE analysis.

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<sup>(16)</sup> For reviews on polyene cyclizations, see: (a) Johnson, W. S. *Acc. Chem. Res.* **1968**, *1*, 1. (b) Johnson, W. S. *Angew. Chem., Int. Ed.* **1976**, *15*, 9. (c) For a key account, see: Johnson, W. S.; Telfer, S. J.; Cheng, S.; Schubert, U. *J. Am. Chem. Soc.* **1987**, *109*, 2517.

<sup>(17)</sup> Also see: (a) Stork, G.; Burgstahler, A. W. *J. Am. Chem. Soc.* **1955**, 77, 5068. (b) Eschenmoser, A.; Ruzicka, L.; Jeger, O.; Arigoni, D. *Helv. Chim. Acta* **1955**, 38, 1890. (c) Stadler, P. A.; Eschenmoser, A.; Schinz, H.; Stork, G. *Helv. Chim. Acta* **1957**, 40, 2191.

<sup>(18)</sup> For a review on triquinane synthesis: Mehta, G.; Srikrishna, A. Chem. Rev. 1997, 97, 671.

Scheme 7. Ynamide Initiated Cationic Polyene Cascade

The divergence in reaction pathways leading to the biand tricylic products was interesting. As one might imagine, after the initial carbocyclization, the second olefin may add to the carbocation through the facial approach shown in either 34a or 34b, which followed by elimination would give 36. Mechanistically more intriguing is the formal [4+2] cycloaddition to afford 37, which must arise through the olefin approach shown in 34a, followed by enamide addition to the resulting tertiary carbocation.

The described carbocyclization cascade worked equally well with methyl-terminated ynamide **38**. After 12 h, 5, 5-*cis*-fused bicycle **39** was isolated as a 9:1 mixture of endo and exo olefin isomers, as well as tricycle **40** as a single diastereomer.

We have showcased here a tandem aza-Claisen–carbocyclization for the synthesis of  $\alpha,\beta$ -unsaturated cyclopentenimines from N-phosphoryl-N-allyl ynamides. Furthermore, we have demonstrated the ability to intercept the zwitter ionic intermediates through Meerwein–Wagner ring expansions and contractions, as well as with tethered carbon nucleophiles to afford bi- and tricyclic scaffolds. Applications of these carbocyclization cascades in total synthesis are currently underway.

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**Supporting Information Available.** Experimental procedures as well as NMR spectra, and characterizations. This material is available free of charge via the Internet at http://pubs.acs.org.

The authors declare no competing financial interest.

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