## **Reactive Intermediates from DMDO Oxidation of Ynamides. Trapping of a de novo Chiral Push**−**Pull Carbene via Cyclopropanation**

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## **ABSTRACT**



**The reaction profile of DMDO oxidations of ynamides is described. This work illustrates the first examples of highly diastereoselective intramolecular cyclopropanations of a push**−**pull carbene derived from alkyne oxidation. In addition, the ynamide oxidation provides facile access to ketoimides and reveals mechanistic insights into the chemistry of electronically biased oxirenes.**

Oxocarbene intermediates derived from epoxidations of alkynes have been strongly suggested to arise from the rearrangement of oxirenes.<sup>1</sup> Subsequent transformations of the oxocarbenes include Wolff rearrangement and further reaction of the resultant ketene, C-H insertion, hydrogen or alkyl migration, secondary oxidation, and formation of stable metalla-keto carbene complexes.2 Despite unleashing a remarkable array of transformations through a simple  $\pi$ -bond oxidation, little has been reported on engaging this potential toward precise ends.<sup>1</sup> Ynamides,<sup>3</sup> a class of electron-rich and electronically biased alkynes, are uniquely positioned for the tuning of such reactivity.

(2) For an example of metal keto carbenoid formation by DMDO oxidation of alkynes, see: Sun, S.; Edwards, J. O.; Sweigart, D. A.; D'Accolti, L.; Curci, R. *Organometallics* **1995**, *14*, 1545.

The postulated oxirenes **2** derived from an oxidation of ynamides **1**<sup>4</sup> could rearrange to amido-oxocarbene **3** or push-pull carbene **<sup>4</sup>**, the latter experiencing dual stabilization5 (Scheme 1). While amido-oxocarbenes **3** have been represented by related diazo compound decomposition reactions,6 push-pull carbenes **<sup>4</sup>** represent a novel species that is intriguing both in reactivity and in synthetic potential.<sup>7</sup>

<sup>(1)</sup> For reviews on oxirenes, see: (a) Zeller, K. P. *Sci. Synth.* **2002**, *9*, 19. (b) Lewars, E. G. *Chem. Re*V*.* **<sup>1983</sup>**, *<sup>83</sup>*, 519. (c) Torre, M.; Lown, E. M.; Gunning, H. E.; Strauzz, O. P. *Pure Appl. Chem.* **1980**, *52*, 1623.

<sup>(3)</sup> For a review, see: Zificsak, C. A.; Mulder, J. A.; Hsung, R. P.; Rameshkumar, C.; Wei, L.-L. *Tetrahedron* **2001**, *57*, 7575.

<sup>(4)</sup> For oxidations of ynamines employing ozone and oxygen, see: (a) Foote, C. S.; Lin, J. W.-P. *Tetrahedron Lett.* **1968**, *9*, 3267. (b) Schank, K.; Beck, H.; Himbert, G. *Synthesis* **1998**, 1718. (c) During our submission of this work, an account on epoxidation of ynamides appeared. See: Couty, S.; Meyer, C.; Cossy, J. *Synlett* **2007**, *18*, 2819 (received July 28, 2007). (d) For our preliminary disclosure of this work, see: Al-Rashid, Z. F.; Hsung, R. P.; Antoline, J. E.; Ko, C.; Wei, Y.; Yang, J. 40th ACS National Organic Symposium, Durham, NC, June 3, 2007; Abstract No. A-11.

<sup>(5)</sup> For push-pull carbene stability and reactivity, see: (a) Buron, C.; Gornitzka, H.; Romanenko, V.; Bertrand, G. *Science* **2000**, *288*, 834. (b) Moss, R. A.; Zdrojewski, T.; Ho, G.-J. *J. Chem. Soc., Chem. Commun.* 1991, 946. For elegant work on donor-acceptor metal carbenoids, see: (c) Hedley, S. J.; Ventura, D. L.; Dominiak, P. M.; Nygren, C. L.; Davis, H. M. L. *J. Org. Chem.* **2006**, *71*, 5349. (d) Davis, H. M. L.; Hedley, S. J. *Chem. Soc. Re*V*.* **<sup>2007</sup>**, *<sup>36</sup>*, 1109.



Were such a carbene to arise, one may expect promiscuity with respect to the reactive partners it may choose (electronrich versus electron-deficient).5b We report here the first epoxidation of ynamides **1** and trapping of novel chiral push-pull carbenes **<sup>4</sup>** via intramolecular cyclopropanations.8

Our efforts commenced with dimethyldioxirane (DMDO) oxidation of terminally unsubstituted ynamide **7** (Scheme 2).



While initial reaction analysis was complicated by partitioning of the resulting amidoglyoxal **8**<sup>9</sup> with its hydrate (confirmed by X-ray), it became clear that (1) ynamide oxidation could be achieved and is relatively faster than reported alkyne oxidations;<sup>10</sup> (2) oxidation of the proposed oxocarbenes proceeded at a faster rate than cyclopropanation of cyclohexenone, even when DMDO concentrations were limited by syringe pump addition; (3) electron-rich olefin

(8) For leading reviews, see: (a) Lebel, H.; Marcoux, J.-F.; Molinaro, C.; Charette, A. B. *Chem*. *Re*V. **<sup>2003</sup>**, *<sup>103</sup>*, 977. (b) Gnad, F.; Reiser, O. *Chem. Re*V*.* **<sup>2003</sup>**, *<sup>103</sup>*, 1603.

(9) See Supporting Information.

cyclopentene out competed the ynamide for DMDO. While not informative concerning the parent carbenes, amidoglyoxal **<sup>8</sup>** and ketoimides **<sup>9</sup>**-**<sup>12</sup>** could be obtained in good yields and represent potentially useful synthons.<sup>11</sup>

To continue probing for the proposed oxocarbenes, we examined intramolecular systems. As shown in Scheme 3,



oxidation of ynamide **13** provided amido-cyclopropane **14** in a 3:1 isomeric ratio with **14**-*major* being confirmed by X-ray crystal analysis. Ketoimide **15** was also obtained, but its formation appears to decrease at elevated temperatures. The competing intermolecular DMDO oxidation of the proposed oxocarbenes was further suppressed by dilution. Oxidation of 4,5-diphenyl oxazolidinone-substituted ynamide **16** provided amido-cyclopropane **17** as a single diastereomer. The respective ketoimide **18**, free auxiliary **19**, and ketoacid **20** were also identified. The formations of **19** and **20** are likely mutually related.

To account for all the observed products, a comprehensive reaction profile for the oxidation of ynamides is shown in Figure 1. Amido-cyclopropanes **14** and **17** unambiguously confirm the presence of push-pull carbenes **<sup>4</sup>**. The major stereoisomer is likely derived from cyclopropanations of oxocarbenes **4** assuming a conformation that accommodates both the nitrogen electron pair donation (pink) to the carbene empty p-orbital (red) and delocalization of the carbene electron pair into the keto carbonyl group (green), while the oxazolidinone carbonyl is *anti* to the carbene lone pair.

Ketoimide formation can be rationalized through a second DMDO oxidation of oxocarbene intermediates  $4$  (or  $3$ ),<sup>10a</sup> although a pathway involving dioxabicyclobutane **21** cannot be ruled out. The stability of ketoimides **15** in acetone/water suggests ketoacids **20** do not arise simply by hydrolysis.

<sup>(6) (</sup>a) Doyle, M. P.; Dorow, R. L.; Terpstra, J. W.; Rodenhouse, R. A. *J. Org. Chem.* **1985**, *50*, 1663. (b) Doyle, M. P.; Kalinin, A. V. *Synlett* **1995**, 1075. (c) Jiang, N.; Ma, Z.; Qu, Z.; Xing, X.; Xie, L.; Wang, J. *J. Org. Chem.* **2003**, *68*, 893.

<sup>(7)</sup> For some examples of related  $\alpha$ -azacarbenoids, see: (a) Schöllkopf, U.; Hauptreif, M.; Dippel, J.; Nieger, M.; Egert, E. *Angew. Chem., Int. Ed. Engl.* **1986**, *25*, 192. (b) Rigby, J. H.; Cavezza, A.; Heeg, M. J. *Tetrahedron Lett.* **1999**, *40*, 2473. (c) Wurz, R. P.; Charette, A. B. *J. Org. Chem.* **2004**, 69, 1262. (d) Bégis, G.; Sheppard, T. D.; Cladingboel, D. E.; Motherwell, W. B.; Tocher, D. A. *Synthesis* **2005** 3186. For elegant examples of R-azametallo-carbenoids, see: (e) Hegedus, L. S. *Tetrahedron* **<sup>1997</sup>**, *<sup>53</sup>*, 4105. (f) Hegedus, L. S.; Lastra, E.; Narukawa, Y.; Snustad, D. C. *J. Am. Chem. Soc.* **1992**, *114*, 2991. (g) Powers, T. S.; Wulff, W. D.; Quinn, J.; Shi, Y.; Jiang, W. Q.; Hsung, R. P.; Parisi, M.; Rahm, A.; Jiang, X. W.; Yap, G. A.; Rheingold, A. L. *J. Organomet. Chem.* **2001**, *617*, 182.

<sup>(10)</sup> For DMDO oxidation of alkynes, see: (a) Zeller, K.-P.; Kowallik, M.; Haiss, P. *Org. Biomol. Chem.* **2005**, *3*, 2310. (b) Murray, R. W.; Singh, M. *J. Org. Chem.* **1993**, *58*, 5076. (c) Curci, R.; Fiorentino, M.; Fusco, C.; Mello, R.; Ballistreri, F. P.; Failla, S.; Tommaselli, G. A. *Tetrahedron Lett.* **1992**, *33*, 7929.

<sup>(11)</sup> For some examples, see: (a) Kim, S. M.; Byun, I. S.; Kim, Y. H. *Angew. Chem., Int. Ed.* **2000**, *39*, 728. (b) Corey, E. J.; McCaully, R. J.; Sachdev, H. S. *J. Am. Chem. Soc.* **1970**, *92*, 2476.



Figure 1. A mechanistic overview.

Instead, a sequence of Wolff rearrangement of **4** (or **3**) follow by oxidation of the resulting ketene **22**12a has been proposed as the major pathway en route to **20** via  $\alpha$ -lactones **23**.<sup>10a</sup><br>In the context of vnamides **13** and **16** the involvement of

In the context of ynamides **13** and **16**, the involvement of amido-oxocarbenes **3** in cyclopropanation is less likely given the concomitant cyclobutane formation. To probe for **3**, we examined oxidations of homologous ynamides **24a**-**<sup>e</sup>** (Scheme 4) because cyclopropanations of the respective olefin in



**24a**-**<sup>e</sup>** through the amido-oxocarbene intermediate **<sup>3</sup>** would lead to bicyclo[3.1.0]hexanes **25**. To our surprise, the epoxidation of the acrylate containing ynamide **24a** gave neither cyclopropanes **25** nor **26a** but only ketoimide **27a**. This outcome was unaltered by inclusion of a buttressing moiety in the tether (see 24b).<sup>13</sup>

Reasoning that the respective amido-oxocarbene **3** could prefer electron-rich olefins, we prepared ynamide **24c**. Although the olefin of **24c** was not entirely stable to oxidation conditions,12b a single isomer of amido-cyclopropane **26c** was obtained, thereby confirming the promiscuity of push-pull carbenes **<sup>4</sup>**. Attempting to again enhance the cyclopropanation pathway, we examined the oxidation of

(12) (a) Attempts at trapping this ketene afforded ketoaminal **iii** accompanied by free auxiliary. The formation of **iii** suggests the interception of oxocarbene **iv**. (b) Ketoimide **v** was isolated in 8% yield.



ynamide **24d** only to find ketoimide **27d** as the sole product.13 A tether dependence reported in enyne cycloisomerizations<sup>14</sup> prompted our investigation of the oxidation of nitrogentethered ynamide **24e**. In this case, amido-cyclopropane **26e** was isolated in 41% along with ketoimide **27e** with no traces of bicyclo[3.1.0]hexane **25**.

While amido-oxocarbenes **3** cannot be ruled out, these experiments further demonstrated the tendency for ynamide oxidation to yield cyclopropane products exclusively arising from push-pull carbenes **<sup>4</sup>**, thereby suggesting either a strong bias in the rearrangement of oxirenes **2** or a pathway that does not involve an oxirene. Electrophilic substitution of ynamides 28 could occur preferentially at the  $\beta$ -position,<sup>3a</sup> as is conveyed by the electronic distribution in resonance structure **29** (Figure 2). This in turn could lead to the



**Figure 2.** An alternative pathway to push-pull carbenes **<sup>4</sup>**.

oxyketene iminium ion **<sup>30</sup>** giving rise to push-pull carbenes **4**.

We have described here a mechanistic profile of DMDO oxidation of ynamides that led to the first examples of stereoselective intramolecular cyclopropanations of a chiral push-pull carbene derived from alkyne oxidation, a facile access to ketoimides, and insights in the chemistry of electronically biased oxirenes. Further investigations into reactivities and applications are underway.

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**Supporting Information Available:** Experimental procedures as well as <sup>1</sup> H NMR spectral and characterizations are available for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

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<sup>(13)</sup> We attributed this observation as a result of the complexation of the ester carbonyl oxygen with carbene intermediates derived from **24a** and **24b** (or **24d**), leading to carbonyl ylides **vi** and **vii**, respectively, with an enhanced nucleophilicity favoring the second electrophilic addition of DMDO over cyclopropanation. Related discussions were reported (ref 10a) where the solvent acetone (see **viii**) was observed to facilitate the double oxidation.



(14) (a) Fu¨rstner, A.; Szillat, H.; Stelzer, F. *J. Am. Chem. Soc.* **2000**, 122, 6785. (b) Nieto-Oberhuber, C.; López, S.; Jiménez-Núñez, E.; Echavarren, A. M. *Chem.*<sup>-</sup>*Eur. J.* **2006**, *12*, 5916.