

# Asymmetric N-Heterocyclic Carbene Catalyzed Addition of Enals to Nitroalkenes: Controlling Stereochemistry via the Homoenolate Reactivity Pathway To Access $\delta$ -Lactams

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**Supporting Information** 

**ABSTRACT:** An asymmetric intermolecular reaction between enals and nitroalkenes to yield  $\delta$ -nitroesters has been developed, catalyzed by a novel chiral N-heterocyclic carbene. Key to this work was the development of a catalyst that favors the  $\delta$ -nitroester pathway over the established Stetter pathway. The reaction proceeds in high stereoselectivity and affords the previously unreported *syn* diastereomer. We also report an operationally facile twostep, one-pot procedure for the synthesis of  $\delta$ -lactams.

Recently, N-heterocyclic carbenes (NHCs) have been shown to be powerful catalysts for a variety of useful transformations.<sup>1</sup> NHC catalysis traditionally operates via the acyl anion equivalent (Figure 1 A), and this manifold has



received a great deal of attention. When an enal is employed, both the acyl anion and homoenolate pathways (Figure 1 B) become accessible.<sup>2</sup> In 2004, Bode and Glorius independently reported the first NHC generated homoenolate in their annulations between enals and aldehydes to synthesize  $\gamma$ -lactones.<sup>3</sup> Since those initial reports the application of the NHC homoenolate has grown tremendously. A variety of annulations have been developed to synthesize lactones, lactams, and carbocycles.<sup>4</sup> Despite these advances, challenges in stereo-induction and differentiation between the acyl anion and homoenolate pathways remain. Stereoinduction is challenging due to the distal relationship between the NHC and the  $\beta$ -

carbon. Differentiation between the acyl anion and homoenolate is problematic, as the majority of electrophiles used in homoenolate pathways have also been shown to be competent acceptors in the acyl anion pathway.

Nitroalkenes represent attractive electophiles for the homoenolate; the  $\delta$ -nitroesters produced in this reaction are useful synthons for  $\delta$ -lactams and piperidines, common motifs in drug targets and natural products.<sup>5</sup> In 2009, Nair reported the NHC-catalyzed homoenolate reaction between enals and nitroalkenes. With the use of an achiral imidazolium precatalyst, aromatic enals and nitrostyrene derivatives were coupled in good yield and good anti selectivity.<sup>6</sup> Recently, Liu and coworkers rendered the reaction asymmetric by employing an aminoindanol-based triazolium precatalyst. Liu's work showed a variety of enals, both aromatic and aliphatic, to be competent coupling partners for aromatic nitrodienes, nitroenynes, and nitrostryenes in excellent ee and good anti diastereoselectivity (Figure 1).<sup>7</sup> Herein we report our own concurrent studies on this reaction that delivers complementary stereoselectivity and allows coupling with aliphatic nitroalkenes.

During the course of our previous investigations into the asymmetric intermolecular Stetter reaction, we observed homoenolate addition to nitroalkenes as a side product.<sup>8</sup> Intrigued by this reactivity, we began our investigation with the addition of cinnamaldehyde 1a to (E)-1-nitrobut-1-ene 2a. Our exploration of chiral catalysts began with aminoindanol-derived  $5a^9$  (Table 1) and found it provides the desired product with excellent syn diastereoselectivity (17:1), but the yield and ee were unsatisfactory (25% yield, 40% ee). The selective formation of the syn diastereomer was unexpected in light of Nair<sup>6</sup> and Liu's<sup>7</sup> previous reports which delivered the anti diastereomer. The incorporation of an aliphatic nitroalkene in this reaction is also noteworthy, as only activated and aryl nitroalkenes have been previously shown. We continued our investigation by employing fluorinated precatalyst 5b<sup>8a</sup> and were pleased to observe the product in 42% yield, 83% ee, and excellent diastereoselectivity (17:1). Unfortunately this system provides no preference for the homoenolate pathway, and a 1:1 mixture of the Stetter product 4 and desired nitroester 3a is formed. Efforts to improve enantioselectivity with 5b via reaction optimization proved fruitless, and the yield of 3a never surpassed 50%. With these results in hand, we chose to focus

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#### Table 1. Chiral Catalyst Screen<sup>a</sup>



<sup>*a*</sup>Reactions were conducted with 1.5 equiv of **1** and 1.0 equiv of **2**. <sup>*b*</sup>Isolated yield after chromatography. <sup>*c*</sup>Diastereoselectivity determined by <sup>1</sup>H NMR of the unpurified reaction mixture. <sup>*d*</sup>Enantiomeric excess was determined by HPLC analysis on a chiral stationary phase.



on developing conditions to promote the homoenolate pathway over the Stetter pathway.

We hypothesized that bulky substituents in the ortho/ortho' position of the N-aryl ring of the NHC would shift product distribution toward the desired nitroester by partially blocking the acyl anion position. Precatalyst **5c** was synthesized, and a considerable increase in product selectivity (5:1 **3a:4**, 70% yield) was observed, although the enantio- and diastereose-lectivity suffered considerably (2:1 dr, 30% ee). Bistrifluoromethyl triazolium **5d** showed good selectivity for the nitroester (6:1 **3a:4**, 60% yield), and diastereoselectivity was excellent (15:1) but enantioselectivity was low (22% ee).

Given the partial success with larger substituents on the aryl side, we hypothesized that an increase in sterics on the alkyl side of the azolium may deliver similar levels of selectivity favoring the homoenolate product with the potential for improved stereoselectivity. With that in mind, we examined Ye's bis-phenyl O-TMS triazolium precatalyst 5e which provided excellent selectivity for the nitroester (>20:1 3a:4, 17% yield) and excellent stereoselectivity (17:1 dr, 95% ee).<sup>10</sup> The remainder of the mass balance with catalyst 5e was enal, suggesting the catalyst was either prohibitively slow or was undergoing decomposition. We postulated that replacing the bis-phenyl moiety with an aliphatic group may lead to a more efficient catalyst. In pursuit of this belief we synthesized 5f and were pleased to observe a substantial increase in yield while product selectivity (>20:1 3a:4, 49% yield) and stereoselectivity remained excellent (17:1 dr, 93% ee). A brief screen of bases and base equivalents led us to our optimized conditions of 50 mol % sodium acetate (see Supporting Information).

A variety of aliphatic and aryl nitroalkenes are competent in the reaction, Table 2. Aliphatic nitroalkenes typically provide product in excellent stereoselectivity, albeit with moderate yield





<sup>a</sup>See Table 1 footnotes a-d. <sup>b</sup>2.5 equiv of aldehyde 1a used.

(3a-3j). Sterically bulky nitroalkenes such as isopropyl are tolerated, but *tert*-butyl does not participate (not shown). Notably, acetal 3g and terminal olefin 3h are formed with useful selectivities and provide valuable handles for further manipulation. Enantio- and diastereoselectivities are typically lower for aryl nitroalkenes (3k-3o), but the reactions proceed in good yield. Heteroaromatic nitroalkenes participate as well (3k, 3m, 3n).

Electron-withdrawing and -rich aryl enals provide product in modest to good yield, Table 3. In the case of *p*-nitrocinnamaldehyde, the reaction delivers a modest yield of 3q, with the remainder of the aldehyde starting material converted to *p*-nitrodihydroethylcinnamate. In this highly electronwithdrawing system, protonation of the homoenolate outcompetes addition to the nitroalkene.<sup>11</sup> Use of *trans*-2-pentenal in this reaction with catalyst **5f** yields only the Stetter product. This is unusual, as it is the only substrate that gives the Stetter product in greater than trace amounts. However, the use of fluorinated catalyst **5b** at 50 °C delivers the desired product in 25% yield with the remainder of the mass balance as the Stetter product **4**.

Due to the orthogonality of the two ends of products 3, we were motivated to explore manipulation of the nitro-ester product to deliver valuable synthons. We found we could

# Table 3. Enal Scope<sup>a</sup>



 $^a$ See Table 1 footnotes a–d.  $^b$ Catalyst **5b** was used, and the reaction was run at 50  $^\circ\text{C}.$ 

generate the corresponding  $\delta$ -lactams<sup>12</sup> in good yield via a onepot two-step process (Table 4). Addition of zinc dust and acetic

# Table 4. One-Pot Synthesis of $\delta$ -Lactams<sup>a</sup>



acid to the crude reaction mixture after 12 h, followed by heating for an additional 4 h, provides an operationally simple protocol for the one-pot synthesis of  $\delta$ -lactams.<sup>13</sup> The  $\delta$ -lactam product may be further reduced to the piperidine via the action of LiAlH<sub>4</sub> (Scheme 1).<sup>14</sup>

In conclusion we have developed a highly effective catalytic system for the asymmetric and diastereoselective generation of a diverse array of syn  $\delta$ -nitro-esters. Key to our success was the development of catalyst **Sf**, which is highly selective for the homoenolate pathway over the established acyl anion (Stetter) pathway. Our system also allows the previously unreported

#### Scheme 1. Lactam Reduction



coupling with aliphatic nitroalkenes and provides access to the syn diastereomer. We also report the operationally facile onepot two-step reaction sequence to arrive at synthetically useful  $\delta$ -lactams.

# ASSOCIATED CONTENT

#### **S** Supporting Information

Full experimental details, spectroscopic data for all new compounds, and crystallographic data (CIF). 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.

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