# Chemoselective silver-catalyzed nitrene transfer: Syntheses of azepines and cyclic carbamimidates

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#### Abstract

Azepines and their saturated azepane counterparts are important moieties in bioactive molecules but are underrepresented in current drug screening libraries. Herein, we report a mild and efficient azepine formation via silver-catalyzed dearomative nitrene transfer. A 2,2,2-trichloroethoxysulfonyl (Tces)-protected carbamimidate nitrene precursor, coupled with the appropriate ligand for silver, is essential for achieving the unexpected chemoselectivity between arene dearomatization and benzylic  $C(sp^3)$ –H amination. Potential applications in the late-stage diversification of azepines to complex molecular scaffolds and diastereoselective hydrogenations to high Fsp<sup>3</sup> azepanes are also highlighted.

## Introduction

*N*-heterocycles are common motifs in natural products and drugs; approximately 60% of FDA approved small-molecule drugs contain at least one heterocycle.<sup>1</sup> Of these, seven-membered rings containing one nitrogen atom, including azepines and azepanes, are unrepresented in current drug libraries. However, this is not due to a lack of bioactivity, as there are reported examples of azepanes that function as glycosidase inhibitors, anti-diabetics, anticancer agents, and anti-viral compounds (Figure 1A).

Common strategies for the syntheses of substituted azepanes (Figure 1B) include various types of intramolecular cyclizations, [4+3] and related cycloadditions, expansion of strained ring intermediates, and ring-closing metathesis.<sup>2</sup> However, these *de novo* approaches largely require pre-installation of desired functionality and stereochemistry into the precursors and are not particularly modular; thus, there is intense interest in new methods for the synthesis of seven-membered *N*-heterocycles that employ readily available precursors. Based on our group's long-

standing interest in transition metal-catalyzed nitrene transfer (NT), we were intrigued by an underexplored strategy involving nitrene insertion into an aromatic ring to form the sevenmembered heterocycle. While the Büchner reaction to insert a carbene into an arene to form the seven-membered carbocycle is well-known,<sup>3a-h</sup> the analogous 'aza-Büchner' involving insertion of a nitrene into an aromatic precursor is underexplored. Early examples employed sulfonylazide<sup>4a</sup> or carbamate<sup>4b</sup> precursors and high temperatures and/or pressures to generate the key free nitrene intermediate. As a result of these harsh conditions, reactions were generally low-yielding and exhibited poor chemoselectivity between competing reactions of the free nitrene. More recently, the Wei group reported a Rh<sub>2</sub>(esp)<sub>2</sub>-catalyzed insertion of a nitrene formed from a carbamoyl azide precursor to furnish azepine scaffolds.<sup>5</sup> However, inconsistent selectivity between nitrogen insertion and C(sp<sup>2</sup>)–H amination, coupled with the need for high temperatures, Rh loadings, and prolonged reaction times limited the versatility of this chemistry. Luan and coworkers similarly reported the synthesis of azepinones via the aminative dearomatization of naphthols,<sup>6</sup> although



**Scheme 1.** Previous approaches to azepine synthesis and chemoselective silver-catalyzed dearomative ring expansion.

high temperature and arene functionality at the *ortho* position were required. The Leonori group reported an elegant synthesis of azepines via a singlet aryl nitrene generated from photolysis of a nitroarene (Figure 1C), followed by reduction of the azepines to their respective azepanes.<sup>7</sup> A large excess of  $P(O'Pr)_3$  in the presence of blue LEDs was required to generate the nitrene intermediate, with the amine reagent required to stabilize the resultant ketimine en route to the 1H-azepine product. Thus, functionality at the *ortho* position of the nitroarene is not tolerated and high loadings of Pd and/or Pt catalysts were typically needed for the subsequent reduction to the saturated azepane.

Our group has reported several examples of chemo- and site-selective silver-catalyzed nitrene transfer (NT), where the identity of the nitrene precursor and the ligand can be fine-tuned to control the reaction outcome.<sup>8</sup> We envisioned that the combination of a suitable nitrene precursor and an appropriate ligand would enable us to selectively toggle between either benzylic C–H amination or insertion of the nitrene into the aromatic ring in an aza-Büchner-type reaction (Scheme 1D). Herein, we report a chemoselective method for synthesis of either azepines or cyclic carbamimidates using silver-catalyzed nitrene transfer. Highlights of our method include the replacement of expensive Rh complexes with cheaper silver salts, excellent chemoselectivity for either C–H insertion or aziridination, mild reaction conditions, the ability to tolerate substitution at any carbon of the arene and the ability for diverse post-functionalizations of the bicyclic azepine products.

#### **Results and Discussion**

Our group recently reported a highly chemo- and enantioselective intramolecular aziridination catalyzed by silver catalyst supported by bis(oxazoline) (BOX) ligand.<sup>9</sup> Interestingly, we noted differences in the asymmetric bis(oxazoline) (BOX) ligand that provided the highest yields and *ee* for a carbamate versus a carbamididate.<sup>9,10</sup> In the case of carbamates, (*S*,*S*)-'BuBOX (**A**, Scheme 2A) provided the optimal *ee*, with (*S*,*S*)-indanBOX (**B**, Scheme 2B) leading to significantly lower *ee*.<sup>10</sup> The converse proved true for the carbamimidate, highlighting our ability to 'match' the steric bulk of the nitrene precursor with the ligand to influence reaction outcome.<sup>9</sup> Carbamimidates are underexplored as nitrene precursors; indeed, prior to our work, only one example showing aziridination was reported by the Dauban group.<sup>11</sup>

Inspired by these results, we were curious if we might harness the unusual versatility offered by silver to achieve an unprecedented and challenging example of chemoselective NT. Our goal was to identify the best combination of nitrene precursor and ligand to selectively tune for either amination at a benzylic C–H bond or to override this preferred reactivity in lieu of dearomative NT via an aza-Büchner reaction (Scheme 2C). The intermediate aziridine could then undergo an



Scheme 2. Tunable control of silver-catalyzed chemo- and enantioselective NT by matching the nitrene precursor to the ligand.

electrocyclic ring-opening along Path A to furnish the azepine or rearomatize via Path B to deliver a product representing a formal amination of a  $C(sp^2)$ –H bond. Two main elements of the reaction design were used to guide our hypothesis for catalyst-controlled chemoselectivity (Scheme 2D). First, we proposed the sp<sup>2</sup>-hybridized nitrogen of the carbamimidate would behave similarly to the carbamate oxygen in our previous work, binding to the Lewis acidic silver and enforcing a squareplanar geometry in the transition state of the NT.<sup>8i</sup> Second, we reasoned that combining the bulky Tces-protected carbamimidate with a sterically undemanding ligand would favor the typical benzylic C–H amination. In contrast, employing a bulky ligand would lead to clashing between the ligand and the Tces group and might favor competing aziridine, despite the energy penalty associated with disrupting aromaticity.

The impact of diverse bidentate ligands on the chemoselectivity of silver-catalyzed amination of a model Tces-protected carbamimidate **1** was investigated (Scheme 3). A 2:1 silver/ligand ratio was initially employed for two reasons: firstly, it is expected to preferentially form a monomeric silver complex in solution to promote reactivity, and secondly, we have previously noted that excess silver salt aids in breaking down the polymeric PhIO.<sup>8a,12</sup> A series of substituted phenanthroline ligands **L1-L5** favored the C–H insertion product **1b**; more electron-rich ligands **L2** and **L3** gave slightly lower yields compared to electron-neutral **L1** and **L4**. Excitingly, installing an electron-withdrawing chlorine on **L5** gave the first indication of the desired azepine **1a**, albeit in a 3.6:1 ratio of **1b:1a**. This result highlights the ability of electronic effects to exert an impact on chemoselectivity. Ultimately, more sterically demanding ligands, as exemplified by the neocuproine ligands **L6-L8** and the dmBOX ligand **L10**, strongly favored formation of the azepine **1a**, with minimal-to-no formation of **1b**. Previous density functional theory (DFT) calculations on the transition state of silver-catalyzed NT catalyzed by (dmBOX)AgClO<sub>4</sub> showed that a seven-



Scheme 3. Ligand screening for silver-catalyzed chemoselective nitrene transfer.

membered transition state was required to accommodate the near-linear geometry needed for the hydrogen atom transfer step;<sup>8b-c</sup> coordination of carbonyl oxygen of the carbamate to silver was key to favoring the larger transition state. The results in Scheme 3 support our hypothesis that the imidate nitrogen binds the Ag similarly, with the increased steric congestion of the bulky Tces protecting group further destabilizing the smaller transition state that leads to the competing C–H insertion.

With the identification of L10 as the best ligand for formation of the azepine 1a, further optimization was carried out prior to exploring the scope of the chemistry. Decreasing both the catalyst loading and the concentration to minimize product degradation resulted in an improved yield of **1a** to 87% (see the SI for details). The optimal conditions were applied to a range of substituted arenes to ascertain the scope of the azepine formation (Table 1), with excellent chemoselectivity observed across the majority of substrates. A variety of para substituents on the arene were tolerated, including a methyl in 2a, a phenyl in 3a, protected amine and alcohol functional groups in 4a-5a and 8a, halogens in 6a and 7a, an ester in 10a, and a trifluoromethoxy group in 9a. Excellent-to-good yields were observed with electron-neutral and donating substituents, while diminished yields were observed with the electron-withdrawing ester group in 10a. As expected, no formation of the azepine 11a was noted with a highly electron-deficient arene; competing C-H insertion to 11b was observed instead. Modifications to the carbamimidate tether to furnish 12a and 13a were well-tolerated. A carbamimidate derived from ibuprofen delivered 14a in 96% yield, while we were pleased to see that both ortho and para substitution precursors 24 and 15 successfully furnished a 78% yield of trisubstituted azepine 24a and a 90% yield of the tetrasubstituted azepine 15a. Due to the high propensity of the intermediate aziridine to rearomatize, naphthalene-derived carbamimidate 16 did not furnish the azepine, but instead yielded the formal C(sp<sup>2</sup>)-H insertion products **16a'** in a 2.6:1 *rr* through Path B in Scheme 2C; efforts are ongoing to identify conditions tunable for both Paths A and B.

The ring expansion was next investigated with a series of *o*-substituted carbamimidates **17-20**. An electron-donating OMe group in **17** resulted in a single regioisomer **17a**. The *o*-Me substituted carbamimidate **18** favored aziridination at the less substituted alkene of the arene, presumably due to increased steric hindrance. The presence of the electronegative, albeit small F group in carbamimidate **19**, resulted in a 1.6:1 mixture of isomers of **19a**, where aziridination was favored



Table 1. Scope of chemoselective silver-catalyzed ring expansion of carbamimidates to azepines.

<sup>a</sup> Yields determined by <sup>1</sup>H NMR using trimethylphenyl silane as an internal standard. <sup>b</sup> Heated at 50 °C for 4 h.

at the less substituted C=C bond of the arene, likely due to electronic effects. In contrast, the increased steric demand of the bromine in 20 significantly improved the regioselectivity for the

aziridination at the less substituted double bond, resulting in the observation of only one regioisomer **20a**. These results provide insight into the likely regiochemical behavior of arenes containing substituents that vary in their steric and electronic properties.

The electronic effects of the *meta* substituents in **21-23** was diminished compared to their *ortho*substituted counterparts, and essentially a 1:1 mixture of regioisomeric products were observed in **22a** and **23a**. However, a moderate regioselectivity of 3.6:1 in favor of initial aziridination of a less substituted alkene of the precursor was observed in the reaction of the *m*-F carbamimidate **21** to **21a**. This improved regioselectivity suggested that the steric and electronic effects of various substituents can be biased toward different regioisomers. The small, electronegative fluorine promotes formation of the electronically favored azepine isomer, while bulkier electron-neutral to slightly donating substituents show little bias. The potential for the ring expansion strategy to be employed in late-stage functionalizations of complex molecules is highlighted by reaction of the estrone-derived carbamimidate **25** under mild conditions to furnish the azepines **25a** in 78% yield. This result also showcases the ability of the chemistry to accommodate substituents at both the *meta* and *para* positions of the arene.

The scope of the benzylic  $C(sp^3)$ -H bond amination was examined next using either 3,4,7,8tetramethylphenanthroline L2 or bathophenanthroline L4 as the ligand (Table 2). Similar to the azepine formation, precursors 1-3 bearing electron-neutral substituents gave excellent yields of the C-H amination products 1b-3b. Strongly electron-donating groups in the p-position, such as the -OMe group of 4, resulted in amination of the benzylic C-H bond, but the product 4b rapidly decomposed under the reaction conditions. Halogens in 6-7 gave the corresponding benzylic amines in good yields. Interesting, replacing the -OMe group of 4 with a -OCF<sub>3</sub> in 9 furnished 9b in 79% yield with no decomposition. Compared to the azepine formation, the precursors 10-11, bearing electron-withdrawing substituents, furnished better yields and chemoselectivity for the C-H amination, including a 61% yield of 11b Substituents in the carbamimidate tethers of 12 and 13 successfully delivered 12b and 13b, respectively. The stereochemistry in 13b was validated through the collection of an X-ray crystallographic structure (see the Supplementary Information for details). However, precursors bearing electron-donating groups on the arene showed competing azepine formation. For example, 14 derived from ibuprofen gave a 1:1 mixture of 14a:14b; sterics at the benzylic carbon may also play a role in controlling the reaction outcome. In the case of the mesityl carbamimidate 15, no 15b was observed and an 81% yield of the azepine 15a was obtained,





<sup>a</sup> Yields determined by <sup>1</sup>H NMR using trimethylphenylsilane as internal standard. <sup>b</sup> Me<sub>4</sub>phen was used. <sup>c</sup> bathophen was used. <sup>d</sup> Decomposition upon work-up.

even using the typical C(sp<sup>3</sup>)–H amination conditions; a similar result was noted for 17, with decomposition of 17b upon purification. Halogens and alkyl groups at both the *ortho* and *meta* positions of the arene in 18-23 gave the desired products 18b-24b in good yields. Finally, the estrone-derived 25 furnished the C–H amination product 25b in moderate yield, highlighting the

ability of this chemistry to be employed for chemoselective nitrene transfer in more complex molecule settings.

The azepine scaffolds could be post-functionalized in diverse ways to generate high Fsp<sup>3</sup> scaffolds and unusual nitrogen-containing molecular architectures (Scheme 4). Hydrogenation of azepine **3a** gave **26** in good yield and 10:1 diastereoselectivity, highlighting the promising potential of this approach for modular and stereocontrolled syntheses of highly substituted azepanes. The Tces protecting group of **1a** was also easily removed under mild conditions in essentially quantitative yield to give **27**. The bicyclic azepines serve as convenient precursors for the construction of more complex scaffolds. For example, irradiation of **1a** in degassed CHCl<sub>3</sub> using a 370 nm Kessil lamp furnish the [4.5.6] tricyclic product **28** in 90% yield.<sup>13</sup> Furthermore, the unusual, bridged tricycle **29** could be prepared in good yield via a regio- and stereoselective intermolecular hetero-Diels-Alder reaction.<sup>14</sup>



Scheme 4. Selective diversifications of azepine scaffolds.

## Conclusion

In conclusion, we have developed a highly chemoselective NT of aryl-substituted carbamimidates where the nature of the ligand controls the reaction outcome. More sterically hindered bidentate ligands promote dearomative ring expansion to deliver substituted azepines in good yields, while less congested ligands yield the C–H amination products. The interaction of the steric bulk of the Tces protecting group with the ligand controls the favored transition state in the NT to achieve the observed chemoselectivity. Arenes bearing a wide range of functional groups and varying electronics were tolerated in the chemistry. Simple post-functionalizations of the

azepine products produced substituted azepanes with good yield and dr, as well as complex fused and bridged compounds. Ongoing efforts are focused on expanding this chemistry to heteroarenes and achieving intermolecular aza-Büchner reactions selectivity on a broad range of substituted arenes.

# ASSOCIATED CONTENT

Characterization data, optimization tables, additional substrates/catalysts, and details of computational methods are included in the supplementary materials, which are available free of charge via the Internet at http://pubs.acs.org.

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## **Author Contributions**

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. <sup>‡</sup>E.Z.S. and C.L. contributed equally.

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