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Bridged bicyclic building block upgrading: Photochemical synthesis of bicyclo[3.1.1]heptan-1-amines

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ABSTRACT: Compounds containing bridged bicyclic carbon skeletons are desirable building blocks for medicinal chemistry. However, as a result of their inefficient, linear syntheses, commercially available compounds of this sort are plagued by high costs and/or a lack of diversity in substitution patterns. Herein we report the conversion of the readily available bicyclo[1.1.1]pentan-1-amine substructure to a wide range of poly-substituted bicyclo[3.1.1]pentan-1-amines using imine photochemistry. To our knowledge, this is the first reported method to convert the bicyclo[1.1.1]pentane skeleton to the bicyclo[3.1.1]heptane skeleton. Hydrolysis of the imine products gives complex, sp³-rich primary amine building blocks.

MAIN TEXT: The fraction of carbon atoms that are sp³-hybridized in a drug candidate is positively correlated with the molecule's advancement though development and clinical trials.¹ Growing cognizance of this trend among medicinal chemists has increased interest in building blocks containing small, bridged bicyclic carbon skeletons. These substructures, like aromatic rings, can provide a well-defined, rigid framework upon which to append substituents, but retain the pharmacological benefits of being aliphatic.² However, most common building blocks containing such ring systems are mono- or "*para*"-disubstituted as a result of synthetic limitations. Consequently, they are restricted to use as linear spacers or as isosteric replacements for monosubstituted or *para*-disubstituted benzene rings.³ Furthermore, existing building blocks in this class are generally achiral, despite the positive correlation of stereochemical complexity with drug candidate success.⁴ The invention of new synthetic methods providing easy access to poly-substituted bicyclic cores will allow for a more complete biological evaluation of low molecular weight, sp³-rich chemical space.

To this end, our group has developed new methods to synthesize 1-aminonorbornanes, potential aniline bioisosteres with a high degree of substitutional variability. In particular, our group has reported an iridium-catalyzed photoredox strategy and a metal-free, imine-based photochemical strategy toward these structures (Figure 1).^{5,6} In both cases, a species with *N*-centered radical character facilitates the homolytic cleavage of a bond in an adjacent cyclopropane ring, triggering an intramolecular radical ring closure cascade. In the photoredox case, the initial *N*-centered radical species is an amine radical cation, whereas in the imine-based case, it is the imine S₁(n, π^*) excited state diyl.

A. Literature Precedent

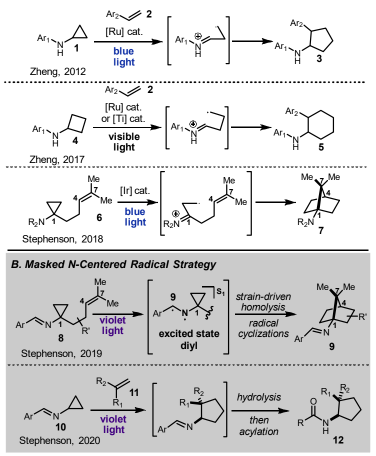


Figure 1. Selected inter- and intramolecular reactions featuring strain-driven homolysis of small-ring amines and subsequent ring formation via addition to an alkene

The photoredox strategy towards strain-driven homolysis and subsequent alkene addition is well-precendented. Zheng and coworkers developed intermolecular variants of the photoredox approach, reacting alkenes with cyclopropyl- or cyclobutylanilines to form 5- and 6-membered rings, respectively (Figure 1).^{7a-e} This annulation chemistry was extended to cyclopropenes by Waser.⁸ However, the imine-based strategy had not been explored in the intermolecular context. Our group has recently disclosed work in this area.⁹ Though cyclopropane ring opening was observed in all cases, we have not observed cyclobutane ring opening under these conditions. This is consistent with the approximately 10³ difference in rate constants determined by Newcomb for the ring opening of cyclopropylaminyl radicals and cyclobutylaminyl radicals.¹⁰ Thus, we propose that the necessary imine excited state lifetime is sufficiently long to achieve cyclopropane ring opening but too short to realize the analogous reactivity with cyclobutanes.

Previous kinetic studies

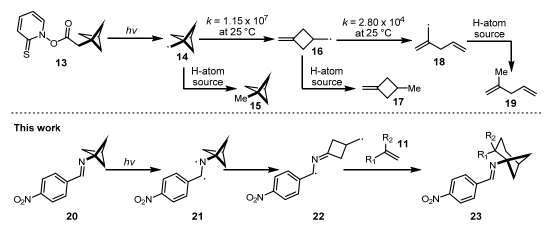
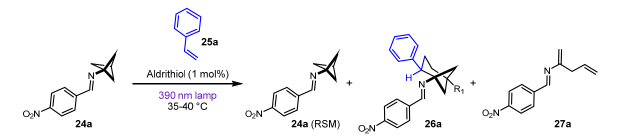


Figure 2. Rate constants from published studies of the bicyclo[1.1.1]pentylcarbinyl radical and our idea of trapping of the intermediate primary radical with an alkene

Intrigued by this difference in selectivity from that observed in the photoredox-based methods, we hypothesized that an imine derived from an abnormally reactive cyclobutylamine, namely a bicyclo[1.1.1]pentan-1-amine, may undergo the desired ring cleavage, generating primary radical intermediate **22**, and then react with an alkene to afford an inert cyclobutylamine, namely a bicyclo[3.1.1]heptan-1-amine like **23**. Though the risk of a second ring opening of the intermediate radical **22** was apparent, previous kinetic studies suggested that radicals of this sort (**16**) are long-lived enough to participate in intermolecular chemistry (Figure 2).^{11,12}



| Trial | 24a (mmol) | Styrene 2a (equiv) | Solvent | 24a (RSM) | 26a | 27a |
|-------|------------|-----------------------|-----------------------------|-----------|-----|-----|
| 1 | 0.2 mmol | 7.5 equiv | 0.65 mL EtOAc | 6% | 15% | 59% |
| 2 | 0.2 mmol | 7.5 equiv | 0.65 mL CH ₃ CN | 5% | 16% | 57% |
| 3 | 0.2 mmol | 7.5 equiv | 0.172 mL EtOAc | 7% | 29% | 40% |
| 4 | 0.2 mmol | 7.5 equiv | 0.172 mL CH ₃ CN | 6% | 26% | 37% |
| 5 | 0.4 mmol | 7.5 equiv | neat | 18% | 38% | 23% |
| 6 | 0.2 mmol | 15 equiv | neat | 6% | 47% | 28% |

Table 1: Optimization of reaction of the parent imine **1a** and styrene **2a**. Yields were determined by ¹H NMR spectroscopy of the crude reaction mixtures using 1,3,5-trimethoxybenzene as internal standard. RSM: recovered starting material.

Indeed, irradiating **24a** and styrene with 390 nm light in ethyl acetate gave the desired product **26a** (Table 1, Trial 1). However, **26a** was the minor product of the reaction, and skipped diene **27a**, which results from the undesired background reaction, was the major product. Changing the solvent from ethyl acetate to acetonitrile had little effect on the reaction profile (Table 1, Trial 2). Increasing the concentration led to a drastic improvement in yield and selectivity, comparable for ethyl acetate and acetonitrile (Table 1, Trials 3 and 4). However, running the reaction in neat styrene gave the best results. Even in 7.5 equivalents of styrene, the imine dissolved, and the reaction proceeded to high conversion within 3 hours. The selectivity was comparable to when using 15 equivalents of styrene (Table 1, Trials 5 and 6). As a precaution, the additive Aldrithiol (2,2'-dipyridyl disulfide) was included in all cases since it was shown to reduce alkene polymerization in our intermolecular [3+2] work.⁹

Despite the inclusion of the additive, prolonged reaction times led to an increasing amount of colored impurities and solid deposits inside the reaction vessel. However, these reaction times also sometimes led to decomposition of **27a** through an unknown mechanism, thereby simplifying purification of the desired products. Reaction times of approximately 12 hours typically led to complete conversion and an acceptable impurity profile.

With suitable conditions in hand, the reaction scope was studied using the paren "*para*"-unsubstituted imine as well as the corresponding imine bearing a "*para*" methyl ester group (Figure 3). Gratifyingly, a wide range of styrenes showed the desired reactivity, albeit with modest yields. Reaction of **24b** with styrene gave product **26b** in 41% yield on a 10 mmol scale (1.56 g product). Alpha-methyl styrene was well-tolerated, giving **26g** and **26h** in 32 and 30% yields, respectively. 1,1-Diarylethylenes were able to undergo the desired ring closure to form sterically hindered bicyclo[3.1.1]heptan-1-amines but also produced a significant amount of a cyclobutene-containing class of byproducts. Cyclobutene **28l** was isolated in 11% yield alongside a 31% yield of compound **26l**. We propose this byproduct forms via an intramolecular hydrogen atom abstraction proceeding through a 6-membered cyclic transition state (Figure 4). Cyclobutenes of this type (**31**) and the ketones resulting from their hydrolysis (**32**) were observed as byproducts for several other alkene substrate classes. A number of examples of such byproducts were isolated and characterized.¹³

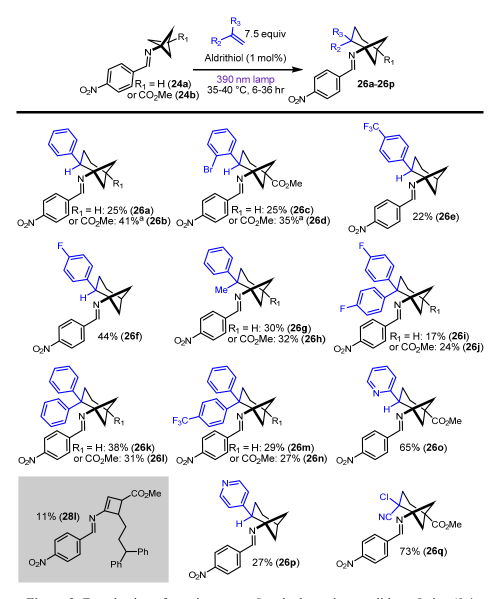


Figure 3. Examination of reaction scope. Standard reaction conditions: Imine (0.4 mmol), alkene (3.0 mmol), 2,2'-dipyridyl disulfide (0.004 mmol) in 1-dram vial with Teflon septum. Degassed through 3 freeze-pump-thaw cycles. Sealed with ParafilmTM and irradiated. Isolated yields reported. Deviations from standard reaction conditions: a) Performed on 10 mmol scale in 6-dram vial. External temperature stabilized at 50 °C as measured by an infrared thermometer. Monitored by ¹H NMR. Starting material consumed after 36 hours. b) Two identical 0.4 mmol runs were combined for purification.

We were also pleased to see alkene classes besides styrenes undergo this chemistry. For example, vinyl pyridines gave modest to moderate yields of the desired products (Figure 3, Compounds **26n** and **26o**). Pyridines commonly act as H-bond acceptors in drugs and their incorporation into building blocks is appealing to for medicinal chemistry applications. Certain unsaturated carboxylic acid derivatives also participate in this reaction. Reaction of **24b** with 2-chloroacrylonitrile gave compound **26q** in 73% yield, the best performance we have observed yet for an alkene undergoing this reaction. Furthermore, the background reaction (Figure 4, "Undesired Reaction Pathway One") was completely suppressed by 2-chloroacrylonitrile, suggesting that certain alkenes may be reactive enough to compete with the background reaction even in the solution phase.

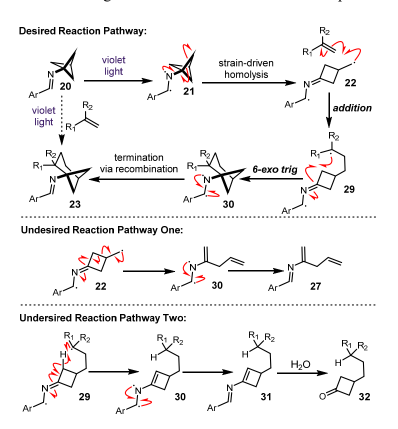


Figure 4. Proposed mechanisms for the formation of the desired product and the observed byproducts.

Imine **26b** was elaborated to desirable protected gamma amino acid building blocks (Figure 5). First, **26b** was hydrolyzed in 83% yield to give **33b**. This material can be protected with di-*tert*-butyl dicarbonate to give carbamate **34b**, which in turn can be saponified to give amino acid building block **35b** containing a protected amine and free carboxylic acid. Amine **33b** can also be protected with benzyl chloroformate and subsequently saponified to give building block **36b**.

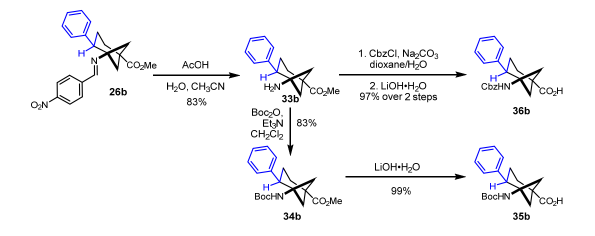


Figure 5. Conversion of 26b to several protected gamma-amino acid building blocks.

In conclusion, we have developed a method of upgrading bicyclo[1.1.1]pentan-1-amines, an increasingly popular class of building blocks, to bicyclo[3.1.1]heptan-1-amines using imine photochemistry. Several classes of alkenes known to react rapidly with nucleophilic radicals are competent in this transformation. The imine products are easily hydrolyzed to give primary amine building blocks. Efforts are underway in our laboratory to further examine the scope of this reaction.

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- 13. See the Supporting Information for further information on byproducts and their characterization.