A General Iridium-Catalyzed Reductive Dienamine Synthesis Allows a Five-Step Synthesis of Catharanthe via the Elusive Dehydrosecodine

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ABSTRACT: A new reductive strategy for the stereo- and regioselective synthesis of functionalized isoquinuclidines has been developed. Pivoting on the chemoselective iridium(1)-catalyzed reductive activation of β,γ-unsaturated δ-lactams, the efficiently produced reactive dienamine intermediates readily undergo [4 + 2] cycloaddition reactions with a wide range of dienophiles, resulting in the formation of bridged bicyclic amine products. This new synthetic approach was extended to aliphatic starting materials, resulting in the efficient formation of cyclohexenamine products, and readily applied as the key step in the shortest (five-step) total synthesis of vinca alkaloid catharanthe to date, proceeding via its elusive biosynthetic precursor, dehydrosecodine.

Saturation and semisaturation nitrogen-containing heterocycles are prevalent structures in bioactive natural products and pharmaceutical compounds, and accordingly, new strategic approaches for their efficient and selective synthesis are important. In parallel, Diels–Alder reactions have been— for nearly a century—one of the most powerful tools for the construction of cyclic and polycyclic products, allowing the disconnection of six-membered rings to a four-electron diene component and a two-electron dienophile. In the normal electron-demand Diels–Alder reaction, electron-rich dienes locked in the reactive s-cis conformation are exceptionally reactive. As such, 1,2-dihydropyridines are a class of compounds particularly poised for cycloaddition reactions, producing the 2-azabicyclo[2.2.2]octane ring system, also called isoquinuclidine (Scheme 1a). This bridged nitrogen-containing bicycle is a familiar structural feature in a range of alkaloid natural products, for instance, catharanthe (3), cononosine (4), and caldaphinidine D (5) (Scheme 1b). Additionally, isoquinuclidines have been used as intermediates toward octahydroisoquinolines in drugs and natural products, such as pseudotabersonine (6) and oseltamivir (7) (Scheme 1c).

To date, because of their inherent instability, the selective and efficient generation of electron-rich 1,2-dihydropyridines has been challenging, and in most cases the presence of a carbamoyl, or similar, electron-withdrawing group on the nitrogen atom is required to make them sufficiently stable for downstream manipulation, albeit at the expense of further deprotection steps or functional group manipulation. Other methods rely on the partial reduction of, or nucleophilic addition to, pyridinium species (Scheme 1d1), but indirect strategies are often required to circumvent the undesired or imperfect regioselectivity in the borohydride-mediated reduction or nucleophilic addition. More recently, highly substituted (and inherently more stable) 1,2-dihydropyridines such as 12 have been generated via Rh-catalyzed C–H activation of α,β-unsaturated imines as well as via multistep cascade reactions involving proline-catalyzed Mannich cyclization followed by oxidation and reduction (Scheme 1d3). Notwithstanding these elegant reports, only specific substitution patterns are currently accessible, and a general strategy for the controlled synthesis of electron-rich 1,2-dihydropyridines currently remains elusive.

Because of the important role of these compounds, and the challenges associated with their generation, we recognized that a mild and general reductive functionalization approach to access 1,2-dihydropyridines using readily available lactam starting materials could be of high synthetic value. Mechanistic studies from our group on the iridium-catalyzed reductive nitro-Mannich reaction revealed that tertiary lactams have a strong propensity to form enamines from the silylated hemiaminal intermediates via their corresponding iminium species. Aware of this, and the tolerance of alkene moieties to the reductive activation conditions, we reasoned that in the presence of suitably placed β,γ-unsaturation in the lactam ring of 15 (Scheme 1e), the 1,2-dihydropyridine species would likely arise from iminium ion via silylated hemiaminal 16. Reactive conjugated dienamine intermediates such as 18 are primed for downstream cycloaddition reactions with various dienophiles, and granting new access to them via a reductive manifold would provide a wealth of opportunities in both library generation, and natural product synthesis alike; herein we wish to report our findings.

We began our studies with a 1H NMR experiment to assess the feasibility of formation of the desired dienamine from...
Scheme 1. (a) Diels–Alder Cycloadditions of 1,2-Dihydropyridines; (b) Isoquinuclidine-Containing Natural Products; (c) Use of Isoquinuclidines in Synthesis; (d) Existing Methods (and Limitations) toward the Synthesis of 1,2-Dihydropyridines and Downstream Isoquinuclidines; and (e) This Work

Figure 1. $^1$H NMR spectra of the reduction of lactam 15a to the dienamine 18a and downstream cycloaddition with N-phenylmaleimide. Reaction performed in $d_8$-toluene, in an NMR tube; 1,3,5-trimethoxybenzene (TMB) was used as internal standard.
lactam precursors (Figure 1). We subjected the model N-benzyl \(\beta,\gamma\)-unsaturated \(\delta\)-lactam substrate \(15a\) to standard reduction conditions in \(d_8\)-toluene (0.1 mol % of Vaska’s complex and 2 equiv of TMDS),\(^{12}\) and very pleasingly, after 20 min, we observed a clean \(^1H\) NMR spectrum fully assignable to dihydropyridine \(18a\).\(^{13}\) Because of the expected instability of this intermediate, we chose to add in one portion the reactive dienophile \(N\)-phenylmaleimide \(21a\) directly to the reaction mixture, and indeed the desired \([4 + 2]\) cycloadduct \(19a\) was formed as the major reaction product (along with TMDS-derived side-products) in 93% NMR yield and as the \(endo\) diastereoisomer.

Encouraged by these preliminary data, we began investigating the scope of this reaction by varying the substituents and substitution patterns on the lactam substrate (Scheme 2). These substrates were accessible via \(\alpha\)-functionalization of the parent lactam (15b, 15d), already known in the literature (15c, 15e),\(^{14}\) or synthesized using a recently developed three-component reaction (15f, 15g).\(^{15}\) We were pleased to find that, when used in conjunction with \(N\)-phenylmaleimide (1.05 equiv) as the dienophile, the corresponding cycloadducts of increasing complexity \(19a\)–\(19g\) could be isolated in good to excellent yields and with essentially complete diastereoselectivity.

Modification of the substitution on the nitrogen atom showed that reactivity was not diminished when using linear (19f) or alicyclic side-chains (19g–19l). Keeping 15g as the parent lactam, we also explored the range of dienophiles that could be successfully deployed in the cycloaddition step. Pleasingly, the use of maleimide \(21h\) as the dienophile resulted in a smooth reaction, providing \(19i\) in excellent 85% yield and \(>95:5\) dr, while oxazolidinone \(21i\) reacted similarly, forming \(19i\) in 90% yield and \(>95:5\) dr. Methyl acrylate (21j), dimethyl fumarate (21k), and acrylonitrile (21l) also led to the formation of the respective cycloadducts 19j, 19k, and 19l, albeit with imperfect diastereoselectivity (85:15, 91:9, and 64:36 dr, respectively).

Having successfully established a scope for the formation of isoquinuclidines from unsaturated \(\delta\)-lactams, we turned our attention to acyclic systems. Simple \(\beta,\gamma\)-unsaturated amides are indeed readily available from secondary amines via coupling with 3-butenolic acid. Our hope was that our newly developed methodology could be extended to the generation of acyclic dienamine species that, in turn, could be valuable intermediates for the formation of tertiary amine-appended cyclohexene architectures, with potential control of up to four newly formed stereocenters.\(^{16}\)

Although the reduction step required longer reaction times than for cyclic systems (3 h, see Scheme 3), we were pleased to find that but-3-enamides \(22a\)–\(c\) did indeed form the desired products.
dienamines 23a–c and the downstream cyclohexene structures 24a–f with complete diastereocorelution upon reaction with N-phenylmaleimide or other dienophiles in good to excellent yields. Moving away from simple but-3-enamides, indole substrate 25a,b, where the βγ-unsaturation is an integral part of the heteroaromatic ring, also produced the desired cycloadducts 26a,b. For ease of isolation, these were further oxidized by addition of DDQ at the end of the reaction and isolated as the aromatized β-carbolines 27a and 27b in 77% and 89% yield, respectively. Finally, both amide functional groups within succinamide 28 could be reduced to their respective enamine intermediates, forming overall a symmetric bisamino-diene species 29 that underwent cycloaddition to furnish symmetric tetrasubstituted 30 as a single isomer. Remarkably, during the course of this reaction, all six carbons contained within the final cyclohexene product saw their hybridization state change from sp3 to sp2 (or vice versa), resulting in a relatively complex architecture arising in a single-pot transformation from a simple building block.

To firmly establish this reductive dienamine generation strategy in complex natural product total synthesis, we set our sights on one of the most important yet elusive intermediates in monoterpene indole alkaloid natural products chemistry, dehydrosecodine (20). Since the pioneering studies of Wenkert in 1962,17 Scott,18a and recently De Luca18b and O’Connor,18c–e this functionally rich molecular entity has been putatively identified as the common precursor to a wide variety of skeletally varied Vinca, Iboga, and Aspidosperma alkaloids.18d Possessing a 1,2-dihydropyridine motif capable of meeting either the electronic demands of a diene (normal electron demand Diels–Alder cycloaddition toward catharanthine 3; see Scheme 4a) or a dienophile (inverse electron demand Diels–Alder cycloaddition toward tabersonine 31),19 dehydrosecodine (20) has remained elusive due to its high reactivity and inherently redox-sensitive functionalities, in particular 1,2-dihydropyridine and indole-2-acrylate.18e–i Not unsurprisingly, nature’s way has inspired the approaches of many synthetic chemists over the years;21 in fact, more than half of the total and formal syntheses of catharanthine published to date have indeed relied on a Diels–Alder approach to the isoquinuclidine core.21a–k Interestingly, however, not one proceeded directly via dehydrosecodine. This is partly due to the difficulty of accessing the 5-ethyl-substituted 1,2-dihydropyridine motif (because of undesired regioselectivity in the reduction of pyridinium ions; see Scheme 1d), particularly in the presence of the sensitive/reactive indole-2-acrylate fragment.20

Recognizing that our reductive strategy offers reliable regiocontrol in 1,2-dihydropyridine synthesis, as well as notable and well-documented chemoselectivity for the reduction of the lactam carbonyl over other functional groups, including alkenes, we set on a journey to access catharanthine (3) via its elusive biosynthetic precursor dehydrosecodine (20).

Our synthesis began with the formation of the α-substituted, βγ-unsaturated δ-lactam 35 in a two-step sequence from commercially available starting materials (Scheme 4b). At high temperatures, tryptamine (33) and dihydropropyne (34) reacted to form the unsaturated lactam as a mixture of constitutional isomers in 51% yield.22 Subsequent double deprotonation of the mixture with 2 equiv of LDA and α-alkylation with ethyl iodide resulted in the formation of desired 35 in 83% yield. After extensive investigations (see the Supporting Information), and taking inspiration from Stephenson’s photoredox-catalyzed C2-functionalization of unprotected indoles,23 we were able to introduce a phosphonooester group at the C2 position of indole 35, resulting in isolation of 37 in 54% yield. The phosphonooester 37 could in turn be used to install the terminal methylene group of 38 via the Rathke modification of the Horner–Wadsworth–Emmons reaction by using paraformaldehyde, in 83% yield.24,25

Having established a four-step route to the precursor of dehydrosecodine 20, the stage was set for the final reductive [4 + 2] cycloaddition sequence. Pleasingly, upon submission of 38 to the newly developed reaction conditions, catharanthine (3) was indeed produced, albeit in trace amounts as determined by 1H NMR analysis of the crude reaction mixture. Extensive optimization of the reductive activation step led to an improved isolated yield (11%) of 3 when TMDS was slowly added to a solution of precursor 38 and Vaska’s complex, thus completing the fully biomimetic total synthesis of the alkaloid and establishing the intermediacy of its evasive and intriguing biosynthetic precursor, dehydrosecodine.

Attempts to isolate byproducts in the final reaction, to understand the low mass return, were unfruitful. Consequently, the reaction was performed in deuterated solvent in an NMR tube, in the hope of observing transient species.26 Upon slow addition of TMDS to a solution of 38 and Vaska’s complex in d8-toluene, catharanthine was immediately produced in 15% NMR yield, alongside reduced species 40 (85% NMR yield, as a mixture of isomers at the dihydropyridine), arising from the apparent hydridic reduction of the indole-2-acrylate in dehydrosecodine (20) (Scheme 5).27 Attempted purification via flash column chromatography on silica gel failed to provide...
dehydrosecodine will likely always induce that any chemical synthesis of dihydropyridine-triggered hydride reduction of the pendant environment.

to give catharanthine and the undesired rearranged product, suggests—hydride scavengers did not change the ratio between any of these experiments.

40,28 while 3 could be isolated in 11% yield. Interestingly, no reaction product arising from the other intramolecular Diels–Alder (IMDA) pathway (see 31, Scheme 4) was observed in any of these experiments.

Further efforts to improve reaction efficiency by introducing hydride scavengers did not change the ratio between catharanthine and the undesired rearranged product, suggesting an intramolecular hydride transfer, followed by protonation and hydridic reduction of the resulting pyridinium species 39 to give 40.29 Although not completely unprecedented,30 this dihydropropyridine-triggered hydride reduction of the pendant indole-2-acrylate suggests that any chemical synthesis of dehydrosecodine will likely always suffer from this undesired internal redox adjustment outside of the exquisitely controlled environment offered by nature’s optimized enzymatic pathways.

In conclusion, an iridium(I)-catalyzed reductive activation of β- unsaturated δ-lactams and amides allows efficient and controlled access to cyclic and acyclic dienamines, delivering—after [4 + 2] cycloaddition—a range of bridged bicyclic and cyclohexene-substituted amine products. This robust approach proceeds with high stereocontrol, low catalyst loading, from readily available starting materials, and has enabled a short and protecting group-free total synthesis of catharanthine via its biosynthetic precursor, dehydrosecodine. Further work to uncover new reactivity of common functional groups through reductive activation approaches is ongoing in our laboratory, and the results will be disclosed in due course.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c04980.

Experimental procedures and characterization data (PDF)

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NOTES

The authors declare no competing financial interest.

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