## Total Syntheses of $(\pm)$ - $\alpha$ -Lycorane and $(\pm)$ -1-Deoxylycorine

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New synthetic routes to  $(\pm)$ - $\alpha$ -lycorane and  $(\pm)$ -1-deoxylycorine were exploited. The *endo*-cycloadduct of 3,5-dibromo-2-pyrone with styrene-type dienophile provided the pivotal intermediate for the syntheses of the titled natural products.

Lycorine (1) is a toxic crystalline alkaloid present in a number of Amaryllidaceae plant species that include *Lycoris, Pancratium, Narcissus, Galanthus, Zephyranthes,* and *Haemanthus.*<sup>1</sup> Bearing a pyrrolo[*de*]phenanthridine common framework (Figure 1),<sup>2</sup> lycorine and its congeneric natural compounds have many important biological activities ranging from the inhibition of ascorbic acid biosynthesis to the prevention of cyanide-insensitive respiration

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to the inhibition of growth and cell division in higher plants.<sup>3</sup> Because of their potentially useful bioactivities, lycorine alkaloids have been the targets of interest, along with other Amaryllidaceae small molecule constituents such as *trans*-dihydronarciclasin and pancratistatin.<sup>4</sup>



Figure 1. Selected pyrrolo[de]phenanthridine natural alkaloid.

Many synthetic studies were then followed,<sup>5</sup> leading to the development of various innovative synthetic strategies

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and structural analogues.<sup>3a</sup> As a part of our ongoing studies exploring the utility of 3,5-dibromo-2-pyrone in target-oriented synthesis,<sup>6</sup> we have envisioned that the pyrrolo[*de*]phenanthridine skeleton of lycorine and its congeners could be rapidly assembled from lactam **8** readily accessible from bicyclolactone **10**, the cycloadduct of 3,5-dibromo-2-pyrone **11**, and dienophile **12** (Scheme 1). Reported herein are our path-finding efforts that led to the total syntheses of  $(\pm)$ - $\alpha$ -lycorane (**3**) and  $(\pm)$ -1-deoxylycorine (**2**). The key features are the formation of the lactam **8** from allylic alcohol **9** and subsequent oxidative functionalization of the cyclohexene subuit (**8**  $\rightarrow$  **7**).

R Pictet-Spenale reduction 1 (R = OH): lycorine 2 (R = H): deoxylycorine Curtius rearrangement Claisen CO-Me rearrangement 8 с 00 B Diels-Alde ъ 10 12 11 Ŕη

Scheme 1. Retrosynthesis of Lycorine and 1-Deoxylycorine

The synthesis began with the Diels-Alder reaction of 3,5-dibromo-2-pyrone (11) with styrene dienophile 12. Heating in benzene afforded endo/exo-cycloadducts as a readily separable mixture (10:1) in 83% total yield (Scheme 2). The isolated endo-adduct 10 was subjected to the Zn-mediated reductive debromination process to give lactone 13 (82% yield). Acid-catalyzed methanolysis of the lactone delivered ester 9 in 83% yield. In order to effectuate the Eschenmoser-Claisen rearrangement reaction,<sup>7</sup> the resulting allylic alcohol 9 was heated with 1,1-dimethoxy-N, N-dimethylethanamine under microwave irradiation (300 W) in xylene. After 8 min, dimethylamide 14 was obtained in 91% yield. Hydrolysis of the methyl ester and subsequent Curtius rearrangement gave isocyanate 16 in 84% total vield over two steps. Successive treatment with LiOH and HCl (aq) gave rise to bicylic lactam 8, which upon a reaction sequence involving a Pictet-Spengler reaction and reductions completed the synthesis of  $(\pm)$ - $\alpha$ -lycorane 3.

After accompishing the synthesis of  $\alpha$ -lycorane **3**, we have mulled over ways to manipulate the cyclohexene double bond toward the synthesis of more decorated 1-deoxylycorine. For this purpose, we examined the oxidative functionalizations of the intermediates **17** and **8** (Scheme 3).





Scheme 3. Oxidative Functionalization of Various Intermediates



Similar to Tossell's intermediate 18, lactam 17 did not give acceptable stereoselectivity or chemical yield when subjected into the typical epoxidation or dihydroxylation reaction. The nearby pyrrolidine ring provided little steric bias for the  $\pi$ -facial discrimination as it occupies a pseudoequatorial position with respect to the C4. Bicyclic lactam 8 gave a higher chemical yield but only marginal selectivity (55% total yield, 3:1 selectivity for epoxidation; 95% total yield, 2:1 selectivity for dihydroxylation). Further study revealed that monocyclic amide 14 is the system of choice as its dihydroxylation reaction produced diol 19 as single isomer in 98% yield. With diol 19 in hand, we explored the synthesis of 1-deoxylycorine (2). At this point, we realized the installation of the double bond between C3 and C4 could be problematic and must be settled before a full-fledged investigation. Using xanthate 23 as a model system, we opted to study the viability of the Chugaev elimination approach. Toward this end, ester 19

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was hydrolyzed into acid **20** after protection of the *cis*-1, 2-diol function (Scheme 4). Subsequent Curtius rearrangement followed by sequential treatments with base and acid effected both lactam formation and acetonide deprotection to afford lactam diol **21** in 51% total yield over three steps.



Scheme 4. Installation of C3–C4 Double Bond: A Model Study

Despite our initial concern, simple treatment with TBSCl allowed selective protection of the lactam **21** C2-OH to afford mono-TBS ether **22** in 82% yield. Sequential treatments with NaH, CS<sub>2</sub>, and MeI gave rise to xanthate **23**. During the reaction, the lactam nitrogen was inevitably methylated. After screening various reaction conditions in the literature, we learned that the elimination reaction is most effective when performed under microwave irradiation.<sup>8</sup> After 5 h at 200 °C, we obtained elimination product **24** in 55% isolation yield. We then decided to construct the B ring lactam before the installation of xanthate group in order to avoid the unnecessary protecting group manipulation (Scheme 5).

Scheme 5. Attempted Installation of C3-C4 Double Bond



Acetylations followed by a Pictet–Spengler cyclization and deprotection furnished diol **26** (78% total yield over three steps from **21**). Despite the possible change in the conformation of the cyclohexane, the silylation reaction occurred predominantly at C2-OH to provide TBS ether **27** in 70% yield. The xanthate group was then introduced for the ensuing Chugaev elimination reaction. Unfortunately, xanthate **28** did not give the anticipated pyrolysis product **29** but a complex mixture of decomposition products. This failure forced us to come back to our original plan and reshuffle the reaction sequence once again, accordingly, i.e., installing the double bond before the assembly of the B ring lactam as illustrated in Scheme 6.

Scheme 6. Attempted Synthesis of  $(\pm)$ -1-Deoxylycorine 2



For easier workup and isolation, the diol function of lactam 21 was masked as an acetonide prior to the LiAlH<sub>4</sub> reduction. The resultant bicyclic pyrrolidine 30 was converted into ethyl carbamate. Treatment with 1 N HCl in THF, during the workup, afforded diol 32. Similar to diol 21, selective protection of the C2-OH required no special reagent or conditions; simple treatment with TBSCl afforded mono-TBS ether 33 in 85% yield. Installation of xanthate group and subsequent pyrolysis under the microwave irradiation gave the elimination product 35 (53%)yield, 40% recovered starting xanthate, 93% yield based on recovered starting material) after 5 h at 200 °C. Removal of TBS group followed by the Mitsunobu reaction with benzoic acid afforded benzoate 36. However, the B-ring lactam formation using the Bischler-Napieralski protocol did not give the expected lactam 38, but diene 37 in 72% yield. Unable to resolve the above elimination problem during the lactam formation, we decided to

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reinvestigate the Chugaev elimination approach at the fully functionalized tetracyclic lactam stage as a last venture. At this point, we came to believe that the location of lactam carbonyl function may affect the elimination process. For this, we prepared tetracyclic lactam **39** with the lactam carbonyl group on the left six-membered ring side (Scheme 7).

Scheme 7. Attempted Synthesis of  $(\pm)$ -1-Deoxylycorine



Thus, carbamate 31 obtained from 30 in 94% yield was treated with Tf<sub>2</sub>O and DMAP to construct the lactam B ring. After concomitant removal of the acetonide group under the conditions, diol 39 was obtained in 81% total vield. Although not unexpected, silvlation of diol 39 was much less selective than that of diol 26, affording both C2-OTBS 40 and C3-OTBS 41 in the ratio of 2:1. Since the undesired 41 can be recycled, the low selectivity of the monosilulation would not be a serious issue at the moment. What mattered was that the xanthate forming process with the isolated 40 gave not only 42 but also the regioisomeric 43 as an inseparable 2:1 mixture. Evidently, the TBS group in 40 was migrated onto the C3-OH during the installation of xanthate group. The use of bulkier silicon-based protecting group may suppress the silvl group migration. Also expected would be a higher selectivity in the C2-OH protection of diol 39. When employed for the selective protection, BPSCl(tert-butyldiphenylsilylchloride) indeed gave much better results, providing the corresponding BPS ethers 46 and 47 in 75% and 20% isolation yield, respectively (Scheme 8). Moreover, the xanthate forming reaction of the BPS protected 46 gave desired xanthate 48 in Scheme 8. End-Game Synthesis of  $(\pm)$ -1-Deoxylycorine



higher selectivity and total yield (48:49 = 3:1, inseparable, 92% total yield). Interestingly, the same reaction with the isolated regioisomeric BPS ether 47 also produced a mixture of 48 and 49 in the same ratio (3:1), indicating the reversible nature of the silyl group migration. Thus, the mixture of BPS ethers 46 and 47 were directly subjected into the xanthate installation process without separation to afford the same product mixture (48 and 49, 3:1) in 92% total yield. The resultant mixture was heated under the microwave irradiation conditions to afford elimination product 50 in 60% overall yield from diol 39 (three steps). After the removal of the BPS group, the configuration of the allylic OH group was inverted under the Mitsunobu protocol. The reduction of amide 51 finally deconvoluted our synthesis of ( $\pm$ )–1-deoxylycorine 2.

In summary, new synthetic routes to  $(\pm)$ - $\alpha$ -lycorane and  $(\pm)$ -1-deoxylycorine were developed. The cycloadduct of 3,5-dibromo-2-pyrone with styrene type dienophile provided the pivotal intermediates for the syntheses of the titled natural alkaloids.

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**Supporting Information Available.** Details of experimental procedures and characterization data. This material is available free of charge via the Internet at http://pubs.acs.org.

The authors declare no competing financial interest.