Catalytic dearomatization approach to quinolizidine alkaloids: Five step total synthesis of (±)-lasubine II

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Supporting Information Placeholder



ABSTRACT: A series of high-yielding silver(I)-catalyzed cyclization reactions of pyridine-, isoquinoline- and pyrazine-ynones are described. The operationally simple and mild reaction conditions are a significant improvement over previously reported thermal cyclizations. The quinolizinone products were also used in a novel dearomatization strategy to prepare 0.53 g of the alkaloid lasubine II in five steps and 36% overall yield.

Pyridine and piperidine are the two most prevalent N-heterocycles used in medicinal chemistry, with a recent study showing their presence in 12% of all US FDA approved drugs.1 New methods for the synthesis of these heterocycles and their derivatives, such as quinolizines/quinolizidines, are therefore of high value. The saturated quinolizidine framework is particularly notable for its presence in a number of bioactive natural products, such as **1**–**3** (Figure 1A),2 which makes them highly attractive synthetic targets.



**Figure 1.** A) Natural products containing the saturated quinolizidine framework; B) Proposed dearomative retrosynthesis.

An unexplored and expedient strategy in which to access these bicyclic frameworks is by the dearomatization of a 2*H*-quinolizin-2-one system **4** (Figure 2B). Dearomatization reactions are important transformations as they enable high value spiro-3 or bridged-compounds4 to be formed from inexpensive and readily available aromatic feedstocks.5 However, current methods for the synthesis of 2*H*-quinolizin-2-ones are limited, relying primarily on harsh thermal conditions,6 with catalytic examples being rare.7 Seeking to address this, an opportunity to build upon our recent work on the catalytic dearomatization/cyclisation of aromatic alkynes was indentified.8 This previous work is based on the activation of alkynes with π-acidic catalysts9 to promote cyclization to generate spirocyclic/annulated products, and based on this, it was considered that pyridine-ynones would serve as useful precursors to 2*H*-quinolizin-2-ones. The viability of a related cyclization protocol had already been briefly demonstrated by both Katritzky and Natarajan (Scheme 1A);6d,e however, in this work the pyridine-ynone species is formed and cyclized in situ via the acylation of 2-picoline under relatively harsh, thermal conditions and the reported modest yields are modest. Herein, we describe a simple and scalable alternative approach, in which pyridine-, isoquinoline- and pyrazine-ynones **5** can all be cyclized into annulated products **6** at room temperature using mild silver(I)-catalyzed conditions (Scheme 1B).

Scheme 1. Pyridine-ynone cyclizations.



Using this new method, a diverse array of quinolizinone products have been prepared in high yield, including reactions performed on gram-scale. The methodology is likely to be of high value in natural product synthesis, and to demonstrate this, an efficient five-step synthesis of 0.53 g of the alkaloid lasubine II has also been developed, including a catalytic dearomatization of a quinolizinone product as a key step.

Our studies began with the preparation of pyridine-ynone **5a** via the deprotonation and acylation 2-picoline with methyl phenylpropiolate (Table 1). Ynone **5a** was then reacted with four π-acidic catalysts in dichloromethane (Table 1, entries 1–4), of which only AgOTf was effective for the formation of quinolizinone **6a**. This prompted further scrutiny of other silver(I) catalysts, and of those tested, AgNO3 provided the best reactivity (entries 7 and 8).

Table 1. Optimization of the pyridine-ynone cyclization.



|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| entry | [catalyst]*a* (equiv) | [solvent] | Time  (h) | conv.  (%)*b* |
| 1 | Cu(MeCN)4PF6 (0.1) | CH2Cl2 | 16 | 0 |
| 2 | Cu(OTf)2 (0.1) | CH2Cl2 | 16 | 0 |
| 3 | Ph3PAuNTf2 (0.1) | CH2Cl2 | 16 | 0 |
| 4 | AgOTf (0.1) | CH2Cl2 | 16 | >95 |
| 5 | AgOTf (0.02) | CH2Cl2 | 2 | 90 |
| 6 | AgSbF6 (0.02) | CH2Cl2 | 2 | 25 |
| 7 | AgNO3 (0.02) | CH2Cl2 | 2 | >95 |
| 8 | AgNO3 (0.01) | CH2Cl2 | 0.5 | 50 |

a Reactions performed with 0.2 mmol of **5a** and catalystin the stated solvent (0.1 M) at RT; b Calculated using the 1H NMR spectrum of the unpurified reaction mixture, rounded to the nearest 5%.

The silver(I) catalyst is proposed to catalyze this transformation as depicted in Scheme 2. The ynone starting material, which exists as an equilibrium of *keto*–*enol* tautomers (with the *enol* tautomer believed to be an unproductive resting state), is presumably activated by the π-acidic silver(I) catalyst (**A**), promoting nucleophilic attack from the pyridine lone pair to form pyridinium species **B** (which is likely a reversible process).10 Deprotonation of the pyridinium species **B** at the acidic α-keto position then forms vinyl silver intermediate **C**; subsequent protodemetalation of this species then affords the final quinolizinone product **6** and regenerates the silver(I) catalyst.

Scheme 2. Proposed mechanism for the silver(I)-catalyzed cyclization.



The scope of this transformation was then examined. Minor modifications made to the conditions shown in Table 1 were to switch to a 1:1 mixture of ethanol:1,2-dichloroethane as the reaction solvent12 and to use 2 mol% of AgNO3 as the catalyst in all of the examples for consistency (Scheme 3).

Scheme 3. Substrate scope for the silver(I)-catalyzed cyclization.



First, the electronics of the aryl ynone subunit were varied to afford quinolizinones **6a**–**c** in quantitative/near-quantitative yield. Pleasingly, these reactions were equally effective on gram-scale; for example, 2.09 g of quinolizinone **6b** was prepared in a single reaction in 99% yield. The aliphatic quinolizinone **6d** was also afforded in near-quantitative yield. Thiophene-substituted and methylated-quinolizinones **6e** and **6f** were also prepared, albeit in lower yield, which is believed to be a direct consequence of the instability of the ynone precursors (see Supporting Information for details). Substituents on the pyridine ring were also well tolerated, with quinolizinones **6g**–**i** bearing cyano, bromo and methyl substituents, all being furnished in excellent to quantitative yield. The structure of the bromide **6h** was also confirmed by X-ray crystallography.11 Finally, this methodology was also demonstrated on other heteroaromatic systems, to afford isoquinoline and pyrazine derived products **6j** and **6k** in excellent yield. The fully unsubstituted quinolizinone **6l** was also synthesized in excellent yield from TMS-ynone **5l** by using 20 mol% AgNO3 and acetone to promote a one-pot desilylation-cyclisation sequence (Scheme 4).13,14 This reaction is particularly pleasing as the TMS ynone **5l** is completely unreactive under the previously reported thermal conditions.6e

Scheme 4. Tandem Ag(I)-catalysed desilylation-cyclisation.



The ease of formation of these quinolizinone products is likely to be of significant value in target synthesis projects, especially given the prevalence of saturated quinolizidine frameworks in Nature.2 This was demonstrated in the five-step dearomative synthesis of (±)-lasubine II (Scheme 5).

Scheme 5. Five step total synthesis of (±)-lasubine II



The synthesis began with the LDA-mediated deprotonation and acylation of 2-picolinewith methyl ester **7**15 to form ynone **5b** in good yield. Next, the silver(I)-catalyzed cyclisation afforded the quinolizine **6b** in near-quantitative yield. Interestingly, the two ring systems of **6b** could then be selectively hydrogenated with either Pd/C or PtO2 to form products **8** and **9** respectively.16 The unpurified quinolizidine **9** was then oxidized under Swern conditions to form ketone **10** in excellent yield over the two-step sequence.17 Finally, the known L-Selectride® reduction of ketone **10** afforded 0.53 g of (±)-lasubine II **3** in 36% overall yield.18,19

In summary, we have developed a mild and operationally simple protocol for the high-yielding catalytic synthesis of quinolizinones. Furthermore, we demonstrated the synthetic utility of the products by preparing 0.53 g of (±)-lasubine II in just five steps and 36% overall yield from 2-picoline. The development of a protocol for the asymmetric hydrogenation of quinolizinones remains the focus of future work,20 which it is hoped will enable the enantioselective synthesis of lasubine II and other related alkaloids.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Experimental procedures and compound characterization data (PDF).

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All authors have given approval to the final version of the manuscript.

Notes  
The authors declare no competing financial interest.

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(11) CCDC 1507022(**6h**) contains the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data\_request/cif.

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