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Prajesh S. Volvoikar, Santosh G. Tilve

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A simple approach for the synthesis of azocine alkaloids: The total synthesis of megallanesine.

Prajesh S. Volvoikar^a and Santosh G. Tilve^{a,b}*

^a Department of Chemistry, Goa University, Taleigao Plateau, Goa, 403206 (India) ^b Organic Chemistry Department, RUDN University, 6 Miklukcho-Maklaya str., Moscow 117198, Russian Federation

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ABSTRACT

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1. Introduction

Nitrogen heterocycles are most abundantly found in nature.¹Azepine and azocine are seven and eight membered ring system containing a nitrogen atom, forms an important class of organic compounds showing interesting conformational structure and wide range of biological activities.² For example isoindolobenzapines alkaloids like chilenine **1**, lennoxamine **2** and deoxychilenine **3** (Fig. 1) are isolated from genus *Berberis* are known for their potent activities against cancerous cells of the lungs, colon, prostrate, etc.³ Magallanesine **4a** a tetracyclic fused ring system is the first isoindolobenzazocine class alkaloid isolated from *Berberis darwinii.*⁴



Fig. 1. Biologically important azepine and azocine ring system.

It is often challenging to synthesize a medium ring size molecule and only a few reports are available for the synthesis of magallanesine or its azocine containing analogues.⁵⁻⁹ Shamma *et al.*⁶ achieved the synthesis of magallanesine prior to its isolation from plant *Berberis darwinii* from a natural product oxyberberine (Scheme 1). Danishefsky *et al.*⁷ achieved first synthesis of **4a** after its isolation using intramolecular aldol condensation in 6 steps with a 56 % overall yield. Kurihara *et al.*^{8a-b} synthesized magallanesine by [1,2]-Meisenheimer rearrangement to construct an azocine ring and modified intramolecular Heck cyclisation as

A new three step synthetic route to construct azocine ring system using anion chemistry and intramolecular Friedel-Craft reaction of an ester is presented. This method allows synthesis of azocine ring analogues in excellent overall yields. The designed strategy was applied for the synthesis of naturally occurring magallanesine.

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the key steps over 11 synthetic steps in 24 % overall yield. The same group ^{8c} later extended the above strategy to synthesize the indole analogue of magallanesine using methyl 2-(9-(phenylsulfonyl)-2,3,4,9-tetrahydro-1*H*-pyrido[3,4-*b*]indol-1-yl)acetate which was synthesized from tryptamine. Recently, Kim *et al.* ⁹ synthesized **4a** using intramolecular Heck and

Kim *et al.* ⁹ synthesized **4a** using intramolecular Heck and Friedel-Crafts cyclisation reaction in seven steps with a 33 % overall yield. Our aim was to synthesize this azocine ring system efficiently in fewer steps.



Scheme 1: Reported route for synthesis of 4a.

In continuation of our interest in synthesis of nitrogen heterocycles¹⁰ we report herein a general route for the construction of the azocine ring system. Our retrosynthetic pathway is depicted is Scheme 2. The dihydro benzoazocine derivative **4a** could be obtained from its tetrahydro derivative **5a**. The construction of eight membered tetrahydro benzoazocine ring B of **5a** could be accomplished from the corresponding ester **6** by intramolecular Friedel-Craft acylation reaction.¹¹ The ester **6**

could be obtained by C-C bond formation via anion chemistry of isoindolinone derivative **7a** which in turn could be prepared by a one pot alkylation-amidation step from intermediates **9** and **10a**.



Scheme 2. Retrosynthesis for construction of azocine ring.

2. Results and Discussion

We started with the synthesis of dimethoxy azocine derivative **5b** (Scheme 3). The required ethyl 6-(chloromethyl)-2,3dimethoxybenzoate **9a** was synthesized according to the procedure described by Rapoport et al. ^{12a}



Scheme 3: Synthesis of azocine derivative 5b.

On heating 9a with 2 equiv of commercially available 2-(3,4dimethoxyphenyl)ethan-1-amine 10b in THF gave the isoindolinone 7b by one-pot alkylation-amidation sequence. ^{12b} Isoindolinone 7b was deprotonated with NaHMDS at -95 °C and then alkylated with ethyl bromoacetate 8a to get the key intermediate ethyl ester 6b.¹³ Only few reports are available in which esters are used as acylating agent.¹¹ The all-important intramolecular Friedel-Craft acylation with PPA (polyphosphoric acid) 110 °C for 16 h failed to deliver the expected eight membered ring compound. Hence, ethyl ester 6b was reacted with 1:10 mixture of P2O5/methanesulphonic acid (Eatons reagent), which is known to be advantageous over using PPA.¹⁴ No change in starting was observed at room temperature even after stirring for 16 h. Hence the reaction mass was heated at 70 °C for 10 h, which gave the desired product tetrahydro benzoazocine 5b in only 25% yield. To improve the yield the corresponding benzyl ester of 6c was prepared which gave 5b in 51% yield at room temperature in 12 h. Further, treating tertbutyl ester 6d with Eaton's reagent gave 99% yield of 5b. To understand the reaction pathway, 6d was treated with Eaton's reagent for 1 h at room temperature. The unreacted 6b and product 5b was seen in small amount along with the corresponding hydrolysed product acid 13a. This suggested that the intramolecular Friedel-Craft reaction of ester 6d is taking place via its acid. The compound 5b was then converted to magallanesine analogue 4b by dehydrogenation with DDO (Scheme 4).⁹

Having successfully synthesized the azocine ring containing 4b, a synthetic derivative of magallanesine, we then completed synthesis of 4c unsubstituted in ring D using starting 10b and

ethyl 2-(bromomethyl)benzoate ¹⁵ **9b** following the same protocol as shown in Scheme 4. Kim *et al.* ⁹ synthesised same compound **5c** using palladium mediated Heck coupling reaction over 6 steps in 57 % overall yield, whereas our method gave the same compound in just 3 steps with an overall yield of 82 % using same starting material **10b**. After this success, we then undertook the synthesis of naturally occurring magallanesine starting from 2-(benzo[*d*][1,3]dioxol-5-yl)ethan-1-amine **10a**¹⁶ which on treatment with **9a** in refluxing THF to furnish the isoindolinone derivative **7a**. The intermediate **7a** was then alkylated with *tert*-butyl bromoacetate **8c** to give ester **6a** in 84 % yield. The ester **6a** was then subjected to the Eaton's reagent in order to get cyclised compound **5a** (Table 1).



Scheme 4: Synthesis of magallenisine and its analogues Table 1: Intramolecualr Friedel-Craft's cyclisation of **6a**

Entry	Reaction condition	Results
1	Eaton reagent, rt, 2 h	Decomposition
2	1:9 (P ₂ O ₅ /methanesulphonic acid),	Decomposition
	DCM, rt, 2 h	
3	1:9 (P2O5/methanesulphonic acid),	Decomposition
	DCM, -20 °C , 6 h	
4	TFA, rt, 2 h, then Eaton reagent, 1 h	Decomposition
5	P ₄ O ₁₀ , DCE, rt, 10 h	5a(26%) + 13b(20%)

However, the only decomposition of the starting was seen at room temperature (Table 1, entry 1). We felt that use of a solvent to dilute the reaction mixture may help. Hence the reaction was attempted in dichloromethane, but similar results were obtained (entry 2). Lowering the reaction temperature also did not help (entry 3). Ester 6a was then hydrolyzed with TFA and treated with Eaton's reagent in a one pot condition which again resulted in decomposition of the starting material. The decomposition of the starting material was attributed to the strong acidic nature of the Eaton's reagent which may be breaking the methenedioxy group. Hence, the reaction was carried out in the presence of excess of P₄O₁₀ in dichloroethane during which two products, the required dihydromagallanesine 5a and its corresponding acid 13b were obtained. Further dehydrogenation of 5a with DDQ provided magallanesine, the structure of which was confirmed by matching with the literature spectral data.^{7, 8b} It is to be noted here that the reported ⁹ synthesis of magallanesine uses BF₃.Et₂O, TFAA for the intramolelcular Friedel-Craft reaction while PPA is used for the other analogues and no reason is attributed to the choice.

Having successfully synthesized magallanesine **4a** and two of its analogues we decided to extend this protocol for indole analogue **4d** (Scheme 4). Thus 2-(1-methyl-1*H*-indol-3-yl)ethan-1-amine **10c**¹⁷ was condensed with **9b** to give 86 % yield. Isoindolinone derivatives **7d** was then successfully alkylated to get **6f** in 96% yield. The ester **6f** when subjected to Eaton's

reagent gave two products 5d and 5e in 30 and 23 % yield respectively. IR spectra of these two compounds were almost similar. But the ¹H NMR spectrum of **5e** showed an extra peak at 1.33 ppm accounting for 9 protons. Also a substitution at 5position of indole ring was observed based on the splitting pattern of the aromatic protons. The compound 5d was identified to be expected tetrahydro azocine based on the spectral data. The mass spectrum of **5e** showed additional 56 a.m.u $([M + Na]^+)$ 409.1892) as compared to 5d, which shows $[M+Na]^+$ peak at m/z. 353.1266. This suggested the presence of an additional C_4H_9 group which may be arising from tert-butanol, which is formed by the hydrolysis of the ester. The tetrahydro azocine derivative 5d was then subjected to the dehydrogenation with DDQ. However, it gave a complex mixture. To account for the for the formation of 5e we conducted a controlled experiment by treating 5d with t-butanol in Eaton's reagent at room temperature. On work-up 62 % of 5e was isolated as expected, thereby confirming the formation of **5e** from **6d** during cyclisation.



Scheme 4. Synthesis of indole fused azocine ring.

3. Conclusion

We have successfully designed a new strategy to construct an azocine ring containing alkaloids using anion chemistry on isoindolinone and intramolecular Friedel-Craft reaction of an ester. The designed strategy was applied for the synthesis of naturally occurring magallanesine. The present method is simple, shorter and has a comparable overall yield in comparison to the reported methods.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://

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- A short and efficient route for synthesis of eight membered azocine ring.
- Synthetic method is extended for synthesis of naturally occuring magallanesine and its derivatives.
- Accepter