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N-Methoxy-N-acylnitrenium lons: Application to the Formal Synthesis of (±)-Desmethylamino FR901483

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ABSTRACT

The formal synthesis of (\pm) -desmethylamino FR901483 (2) is described. Construction of the unique azatricyclic skeleton of 2 was accomplished by a sequence which involved (i) preparation of dienone 7 by an *N*-methoxy-*N*-acylnitrenium ion-induced spirocyclization, (ii) formation of 2-azabicyclo[3.3.1]nonane 5 by the $6-(\pi-exo)-exo-trig$ radical cyclization of 1,7-enyne 6, and (iii) installation of the C-5 *p*-methoxybenzyl side chain by Lewis acid-mediated alkylation of silyl enol ether 18.

The immunosuppressive natural products cyclosporin A and FK506 have played a key role in the advancement of transplant surgery and the treatment of autoimmune diseases. 1,2 However, these drugs have serious side effects and as a result, there is a need to develop less toxic immunosuppressants that selectively inhibit organ rejection but leave the native immune system able to respond to viral, fungal, and tumor antigens. FR901483 (1) (Figure 1), a secondary metabolite of *Cladobotryium* sp. No. 11231 isolated by a group at Fujisawa, is a potent immunosuppressant which significantly increases the survival time of grafts in the rat allograft model. 4,5 This intriguing alkaloid contains 2-aza-

Figure 1. 1-Azaspiro[4.5]decane alkaloids: FR901483 (1), desmethylamino FR901483 (2), and TAN1251A (3).

bicyclo[3.3.1]nonane and pyrrolidine rings spiro-fused to form a azatricyclic skeleton previously unprecedented in Nature. It is not surprising then that synthetic interest in this target has been considerable. To date, syntheses of **1** have been reported by Snider, ^{6a,b} Sorenson, ^{6c} Ciufolini, ^{6d} and, most recently, by Funk. ^{6e} In addition, a number of approaches to the azatricyclic core of **1** have also been published. ⁷ We

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recently reported the synthesis of (+)-TAN1251A (3),⁸ which shares a common 1-azaspiro[4.5]decane skeleton with FR901483, utilizing the Ar₁-5 spirocyclization of *N*-methoxyl-*N*-acylnitrenium ion **8** (Scheme 1).⁹ Having completed the

Scheme 1. Retrosynthetic Analysis for Desmethylamino FR901483 (2)

first leg of our divergent study, we now report the application of this chemistry to the formal synthesis of (\pm) -desmethy-lamino FR901483 (2).

As indicated in Scheme 1, the ultimate goal of the study reported here was tricycle 4, an advanced intermediate in Snider's pioneering synthesis of desmethylamino FR901483 (2).6a We viewed 4 as being accessible from tricycle 5 through a sequence involving alkylation of the corresponding C-6 ketone. In common with other groups,⁶ we deemed formation of the C-6/C-7 bond as the most convenient strategy to access the 2-azabicyclo[3.3.1]nonane ring system.¹⁰ However, while all reported syntheses of **1** have achieved this bond formation through internal aldol reactions (or variants thereof), we were somewhat concerned about both the issue of C-6 stereocontrol and the likelihood of epimerization at the adjoining benzylic stereocenter. To side step these issues, we opted to prepare 5 from enol ether 6 via a 6- $(\pi$ -exo)-exo-trig radical cyclization, ¹¹⁻¹³ mediated by the addition of a stannyl radical, 14 and then capitalize on

the facial dissymmetry of **5** to install the C-5 *p*-methoxybenzyl side chain and C-6 stereocenter at a late stage in the synthesis. The cyclization precursor **6** could be obtained from dienone **7** which, in turn, would be accessible through the spirocyclization of the *N*-acyl-*N*-methoxynitrenium ion **8**, as reported in our synthesis of TAN1251A.

Our route to **4** commenced from commercially available **9** which was coupled with methoxylamine hydrochloride (DCC, Et₃N) to provide **10** in excellent yield (Scheme 2).

^a Reagents and conditions: (a) MeONH₂·HCl, DCC, Et₃N, CH₂Cl₂, 0 °C → rt, 21 h; (b) i. PhI(OCOCF₃)₂, CH₂Cl₂, MeOH, 0 °C, 15 s; ii. NaHCO₃, H₂O, 0 °C, 5 min; (c) H₂ (1 atm), Pd/C, EtOAc, rt, 20 h; (d) (CH₂OH)₂, PPTS, PhH, reflux, 3.5 h; (e) i. Na, NH₃, THF, −78 °C, 30 min; ii. NH₄Cl, 0 °C → rt, 3 h.

Treatment of **10** with bis(trifluoroacetoxy)iodobenzene resulted in rapid spirocylization to furnish **7**. While this material proved prone to decomposition under acidic conditions, by reducing the reaction time to a mere 15 s and immediately quenching the reaction with aqueous NaHCO₃ we were able to prepare multigram quantities of **7** in reasonable yield. Dienone hydrogenation (H₂, Pd/C), protection of the resulting ketone as the 1,3-dioxolane acetal, and reductive cleavage of the *N*-methoxyl amide under Birch conditions¹⁶ now provided pyrrolidone **11**. *N*-Alkylation with propargyl bromide¹⁷ and acetal hydrolysis furnished **12** which

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was converted to the corresponding trimethylsilyl enol ether using trimethylsilyl iodide and (TMS)₂NH (Scheme 3). ¹⁸ This

^a Reagents and conditions: (a) i. NaH, DMF, 0 °C, 3 h; ii. BrCH₂C≡CH, 0 °C, 4 h; (b) 0.5 M HCl, acetone, 50 °C, 18 h; (c) i. (Me₃Si)₂NH, CH₂Cl₂, rt, 30 min; ii. Me₃SiI, -20 °C, 10 min then rt, 3 h; (d) i. Bu₃SnH (1.2 equiv), AIBN (0.1 equiv), PhH (0.08 M), 80 °C, 4.5 h; ii. MeOH, HCl, H₂O, rt, 2 h.

material was found to be of sufficient purity, by ¹H NMR spectroscopy, to allow direct submission to the radical cyclization.

After screening a number of radical initiators and solvents, we found that slow addition of a mixture of *n*-Bu₃SnH and AIBN in benzene to a solution of **6** in the same solvent at reflux over 4.5 h followed by protodestannylation (HCl, MeOH) of the crude reaction mixture gave the desired cyclization product **5** together with tricycle **13**¹⁹ and a small amount of reduction product **14**. Both **5** and **13** where isolated as single diastereomers whose relative stereochemistries were determined through a combination of COSY, NOESY, and HMQC experiments.

Interestingly, the outcome of the cyclization reaction displayed a significant temperature dependency, e.g., use of toluene as the reaction medium favored the translocation—cyclization pathway and led to the formation of 13 and 5 in a 2:1 ratio. As illustrated in Figure 2, we have rationalized these observations in terms of conformers 15 and 16. While 15 can adopt the geometry necessary for 6-exo-trig cycliza-

Figure 2. Competing radical pathways: $6-(\pi-exo)-exo-trig$ cyclization vs 1,5-hydrogen atom transfer/ $5-(\pi-exo-)-exo-trig$ cyclization.

tion to take place,²⁰ **16** is unable to cyclize. However **16** does meet the stereoelectronic requirements for 1,5-transfer of the adjacent allylic hydrogen atom.^{21,22} The allylic radical thus generated then undergoes diastereoselective cyclization with the pendant vinyl stannane to form **13**.²³

Having established a protocol for formation of the 2-azabicyclo[3.3.1]nonane core of the target, we now proceeded to install the C-5 side chain. Protection of the hydroxyl group of **5** as the benzyl ether and oxidative cleavage of the *exo*-olefin gave ketone **17** (Scheme 4). Our

^a Reagents and conditions: (a) BnBr, NaH, Bu₄NBr, DMF, rt, 6 h; (b) i. OsO₄, py, *tert*-BuOH, THF, H₂O, 30 min, rt; ii. NaIO₄, 6 h, rt; (c) i. KHMDS, THF, −50 °C, 15 min; ii. Et₃SiCl, −50 °C, 40 min; (d) **18**, *p*-MeOBnBr, ZnCl₂·Et₂O, Et₂O, −78 °C → −25 °C, 16 h; (e) SmI₂, THF, H₂O, rt, 5 min; (f) LiAlH₄, THF, −78 °C → rt, 22 h; (g) H₂ (1 atm), Pd(OH)₂/C, MeOH, rt, 3 h; (h) see ref 6a.

initial attempts to alkylate various metal enolates of **17** with *p*-methoxybenzyl bromide were unsuccessful, with low yields of **19** and significant amounts of the dialkylated product and starting material being obtained. Encouragingly, we found

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that this obstacle could be overcome by Lewis acid-mediated α -alkylation of enol ether **18**. ²⁴ Thus, sequential treatment of **17** with KHMDS and Et₃SiCl furnished **18** which, after isolation, was treated with *p*-methoxybenzyl bromide and ZnCl₂ to give **19** as a single stereoisomer. Reduction of the C-6 ketone with samarium diiodide now cleanly generated the desired *exo*-alcohol **20** in good yield. ²⁵ The relative stereochemistry of **20** was confirmed by a NOESY experiment which revealed correlations between H-6 and H-7 and the axially positioned proton at C-9.

Reduction of lactam **20** with LiAlH₄ in THF now gave the desired pyrrolidine **21** (28%) and, rather unexpectedly, diol **4** (39%), the product of benzyl ether cleavage.²⁶ Catalytic hydrogenolysis of **21** over Pd(OH)₂/C now proceeded smoothly to give **4** in 99% yield. The combined overall yield for the conversion of **20** to **21** was 66%. As illustrated in Figure 3, the relative stereochemistry of **4** was confirmed by measurement of NOESY correlations. In addition, a comparison of the spectroscopic data collected for **4** with that reported by Snider indicated a close match.^{6a}

Figure 3. Selected NOESY correlations for 2-azabicyclo[3.3.1]-nonane **4**.

In summary, we have developed a synthetic route to **4** which Snider has previously carried to **2** in six steps with an overall yield of 38%. ^{6a} Accordingly, the work reported here represents a formal synthesis of desmethylamino FR901483 (**2**). Efforts to complete the asymmetric synthesis of FR901483 (**1**) are now underway and will be reported in due course.

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Supporting Information Available: Full experimental procedures and spectral data for compounds 4–21. This material is available free of charge via the Internet at http://pubs.acs.org.

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