

Total Synthesis of Rapamycin via a Novel Titanium-Mediated Aldol Macrocyclization Reaction

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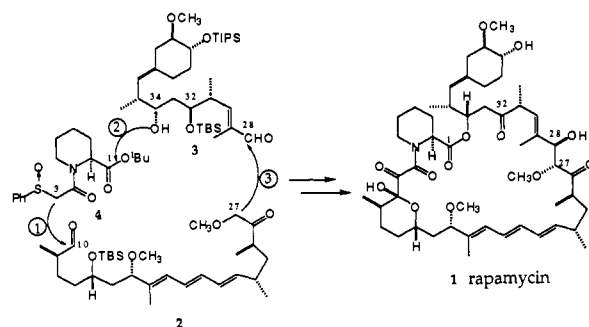
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Received July 14, 1993

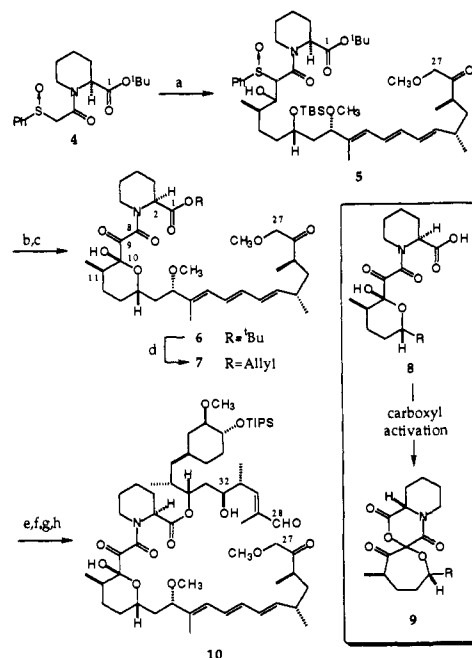
The advent of the two "tricarboxyl" macrolactam-macrolactone immunosuppressive agents rapamycin (**1**)^{1,2} and FK-506³ has provided exciting multidisciplinary scientific challenges.⁴ Indeed, by drawing upon the powerful resources of chemistry (both synthetic and structural)^{5,6} in conjunction with those of molecular and cell biology, Schreiber and associates are in the process of defining the mechanisms of action of these drugs in striking detail.⁷ Total^{8,9} as well as formal¹⁰ syntheses of FK-506 have been achieved. The total synthesis of rapamycin was first reported by Nicolaou *et al.*¹¹ and has also been accomplished by Schreiber and associates.¹²

Our laboratory has been engaged in degradative¹³⁻¹⁵ as well as synthetic^{16,17} studies centered around FK-506 and rapamycin. The long-term aim of both efforts is to pinpoint the substructural features which are critical for binding to natural receptors and for immunosuppressive function.¹⁸ In the case of rapamycin, we recently described investigations which led to the syntheses of compounds **2**¹⁹ and **3**²⁰ (Scheme I). Another building block, **4**, was readily obtained from L-pipecolic acid.²¹ Herein, we disclose the total synthesis of rapamycin featuring as a key step the highly

Scheme I



Scheme II^a



^a (a) LDA, THF, -78 °C; **2**, 57%. (b) Dess-Martin periodinane, CH₂-Cl₂, pyridine, 25 °C. (c) HF·pyridine, THF, 25 °C, 32% from **5**. (d) (i) HCO₂H, CH₂Cl₂, 25 °C; (ii) allyl bromide, K₂CO₃, catalytic *n*-Bu₄N⁺I⁻, DMF, 25 °C, 66%. (e) TMS-imidazole, catalytic DMAP, DMF, 25 °C, 94%. (f) 20 mol % Pd(PPh₃)₄, catalytic PPh₃, CH₂Cl₂, 25 °C, 70%. (g) 3, DCC, catalytic DMAP, CH₂Cl₂, -20 °C, 85%. (h) *n*-Bu₄N⁺F⁻, HOAc, THF, 50 °C, 50%.

novel macroaldolization of *seco* intermediate **11** via its titanium enolate to establish the 31-membered ring.

We combined the synthetic building blocks **2**, **3**, and **4** in the sequence shown in Scheme I (see arrows) and by the methods specified in Scheme II. We defined as an interim goal compound **6**, which contains the sensitive tricarboxyl and triene sectors of **1** as well as the enolate source for a projected macroaldolization (*vide infra*). For this purpose, **4** was converted to its lithio derivative, which condensed with **2**, thereby producing **5** in 57% yield. Upon oxidation with Dess-Martin periodinane²² followed by desilylation, the desired **6** was obtained in 32% overall yield. We emphasize the simplicity of this new method of tricarboxyl synthesis²³ and its compatibility with the rather fragile functionality of this system.²⁴

At this stage, we were in a position to exploit another piece of chemistry developed in our laboratory, i.e., that of generating a competent acylating agent from the pipecolinyl section of a

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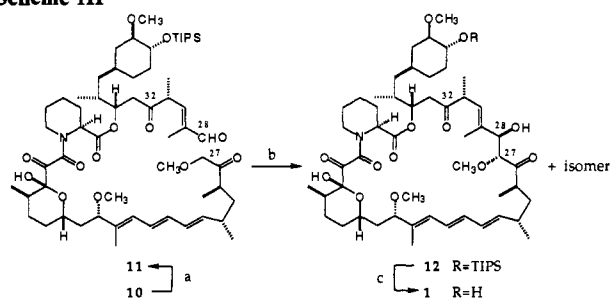
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(21) Reaction of *tert*-butyl *N*-[2-(phenylthio)acetyl]-L-pipecolate (see ref 24) with NaIO₄ in MeOH/H₂O produced **4** in 93% yield.

Scheme III^a

^a (a) Dess–Martin periodinane, CH_2Cl_2 , pyridine, 25 °C, 89%. (b) $\text{TiCl}_3(\text{OiPr})_3$, CH_2Cl_2 , -78 °C; Et_3N , 11%. (c) HF-pyridine, THF, 25 °C, 85%.

secorapamycin which, though truncated, does contain the hemiketal and triene linkages.²⁵ The difficulty that had to be overcome was the tendency of systems bearing the free hemiketal hydroxyl to undergo lactonization (see transformation $8 \rightarrow 9$)²⁶ upon attempted esterification with various external alcohol constructs.

Accordingly, the *tert*-butyl ester function of **6** was cleaved, and the resulting acid was converted to its allyl ester **7**. After protection of the tertiary hemiketal hydroxyl group as its trimethylsilyl ether,²⁵ the allyl ester was cleaved through the agency of $\text{Pd}(\text{PPh}_3)_4$ ²⁷ to afford an acid. This acid was esterified with the C-34 hydroxyl (rapamycin numbering) of compound **3**²⁰ using DCC–DMAP⁸ in 85% yield. Selective twofold desilylation of the C-10 and C-32 silyloxy groups afforded **10**. Finally, oxidation of the C-32 hydroxyl of **10** with Dess–Martin periodinane led to the C-40-silylated secorapamycin **11**, the substrate for the projected intramolecular macroaldolization (Scheme III).²⁸

With compound **11** in hand, an extensive investigation of macroaldol cyclization methods was initiated. The first sets of conditions surveyed were those which were expected to generate metalloenolates of the type well known to participate in aldol condensations. These efforts were unsuccessful. For instance, attempts to generate either lithium²⁹ or cerium³⁰ enolates of **11** by standard protocols did not lead to detectable cyclization products. Similarly, recourse to reagents such as (cyclohexyl)₂ BCl ,^{31a} $\text{Sn}(\text{OTf})_2$,^{31b} Bu_3SnCl ,^{31c} SnCl_4 ,^{31d} ZnCl_2 ,^{31e} and ZrCp_2Cl_2 ^{31f} each in the presence of prescribed amines also led to no detectable cyclization products.

The first success that was realized resulted from recourse to a presumed titanium enolate generated under conditions (**11** +

TiCl_4 , CH_2Cl_2 , -78 °C; Et_3N) described by Evans.³² There were thus isolated a 7% yield of C-40 TIPS rapamycin **12**, a 14% yield of an apparent stereoisomer of **12**,³³ and a 35–45% yield of recovered **11**. The use of isopropoxytitanium trichloride as the promoter³⁴ afforded a 33% yield (42% based on recovered **11**) of a 1:2.3 ratio of **12** to the same stereoisomer that was observed in the reaction mediated by TiCl_4 . The identity of synthetic **12** was established by comparison of its NMR, infrared, and FAB mass spectra as well as its chromatographic mobility to those of **12** prepared from silylation of native rapamycin. The structure of **12** was further proven by desilylation (HF-pyridine) to afford rapamycin itself.³⁵

Many features of the fascinating formation of the C-27–C-28 bond of rapamycin by intramolecular aldolization are being explored (including the use of biomimetic schemes for cyclization). This work will be described in due course. Also awaiting full clarification are issues relating to the thermodynamic stabilities and kinetic connectivities among these and possibly other C-27–C-28 stereoisomers. Finally, we note that the capacity to interpolate compound **3** between the pipercolinic acid (C-1) and carbon-27 in the manner described might well be exploited to create new analogs of rapamycin for biological investigation.

Acknowledgment. This work was supported by NIH Grant AI 16943. NMR spectra were obtained through the auspices of the Northeast Regional NSF/NMR Facility at Yale University, which was supported by NSF Chemistry Division Grant CHE7916210. We also thank Drs. Craig E. Caufield and Guy Schiehsler of Wyeth-Ayerst Research, Inc. for generous samples of rapamycin, for invaluable insights regarding the degradation of rapamycin, and for their continued encouragement of our program.

Supplementary Material Available: Analytical data for compounds **7**, persilylated **10**, **11**, and **12**, and synthetic **1** (6 pages). Ordering information is given on any current masthead page.

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(33) The major product of the macroaldol cyclization was degraded according to the procedure of Goulet (cf. ref 15b: ozonolysis of bisilylated **1**). A product was obtained which is very similar to but not identical with the rapamycin-derived degradation fragment. Given the virtually identical multiplicities with the rapamycin-derived Goulet fragment, it is tempting to assign the major aldol to be in the C-27–C-28 *anti* series. However, this point has not been proven. Also, the possibility of additional differences elsewhere in the molecule arising from epimerization under the reaction conditions is not excluded.

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