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Allylic Alcohol Transposition by Ortho Ester-Initiated Carbonate Extension. Synthesis of the Vasodilator 11(*R*),12(*S*),15(*S*)-Trihydroxyeicosa-5(*Z*),8(*Z*),13(*E*)-trienoic Acid

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ABSTRACT

The title compound 1 was obtained via methyl ester 2, which was synthesized in four steps from an isomeric 11,14,15-triol ester 5. In the key step, Boc orthoformate 9 was treated with TMS triflate to initiate intramolecular nucleophilic substitution with allylic transposition, forming cyclic carbonates 10 and 11.

In the course of research on endogenous mammalian vaso-modulators in the laboratories of Falck and Campbell, a potent vasodilator was discovered in rabbit aortic endothelium and identified as 11(R), 12(S), 15(S)-trihydroxyeicosa-5(Z), 8(Z), 13(E)-trienoic acid (1, Figure 1). Compound 1 is

$$CO_2R'$$
 CO_2Me
 C

Figure 1. Synthetic plan.

a member of a class of triol acids, designated THETAs, that are biosynthesized from arachidonic acid by a sequence of lipoxidation to the hydroperoxide 15-HPETE, isomerization to an epoxy alcohol, and hydrolysis. The 11,12,15-triol pattern of **1** was deduced by mass spectral analysis, but because of the scarcity of the natural product, assignment of stereochemistry awaited comparison with synthetic material. To that end, Falck and co-workers synthesized the four C11–C12 diastereomers of structures **1** and **2** from stereodefined vicinal diol subunits.¹ Herein is reported a concise synthesis of **1** from the known 11,14,15-triol ester **5**,² in which transposition of the C12–C14 allylic alcohol array was accomplished by a new cationic ScN' reaction³ initiated by a cyclic ortho ester⁴ and terminated by a *t*-butyl carbonate.^{5,6}

Triol ester **5** was obtained in two steps from 15(S)-HETE methyl ester (**3**) as described earlier by Falck (Scheme 1).²

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Scheme 1. Preparation of Boc Orthoformate 9

R

$$CO_2Me$$
 CO_2Me
 CO_2Me

Epoxidation of **3** on a 3.5-g scale⁷ afforded the anti (erythro) epoxy alcohol **4** (49%) plus its syn (threo) diastereomer (21%). Hydrolysis of **4** provided triol **5** and its more polar epimer **6** (\sim 1:1). Treatment of the separated triols with excess HC(OMe)₃ and catalytic PPTS in dichloromethane gave orthoformates **7** and **8**, plus bisorthoformates from which the 11-OH group could be released by titration of the reaction mixtures with methanol.⁸ In this way, **7** and **8** were obtained reproducibly in 77–78% yield from **5** and **6**. The C11 configurations were assigned at this stage by *O*-methylmandelic ester analysis.⁹ The 11*R* epimer **7** was converted to *t*-butyl carbonate **9** in 94% yield by treatment with Boc₂O, *i*-Pr₂NEt, and DMAP in 2:1 hexane—diethyl ether.¹⁰

Reaction of **9** (275 mg) with TMSOTf in EtCN at -20 °C gave trans cyclic carbonate **10** and its cis epimer **11** (4.5:1) in 62% yield (Scheme 2). The useful desformyl congeners were also obtained in 6% yield. The Boc derivative of **8** likewise delivered a 5.7:1 trans/cis product mixture in

Scheme 2. Ionization of 9 Leading to Cyclic Carbonates and Methanolysis Affording Transposed Triols

comparable yield. Propionitrile (mp -93 °C) was chosen instead of acetonitrile to enable experimentation at lower temperature without resorting to the use of mixed solvents.^{5b} In trials at -78 °C, little formation of **10/11** was observed. The ring stereochemistry was evident from the chemical shifts of H11 and H12 (**10**: δ 4.32, 4.70; **11**: 4.71, 5.13).¹¹ The H13 and H14 signals at δ 5.7–5.9 were well isolated at 600 MHz and showed J = 15.6 Hz, characteristic of the E configuration.

Exposure of the separated carbonates **10** and **11** to methanolic NaOMe afforded 12-*epi*-**2** and **2** (86–90%), whose 1 H and 13 C NMR spectra agreed with those of the 11*R* product pair synthesized by Falck. The H11 resonances occurred at δ 3.53 and 3.71, respectively, consonant with the assigned syn and anti C11–C12 relationships. Saponification of **2** yielded acid **1**, which by negative-ion ESI/MS/MS showed a diagnostic m/z 197 ion arising from C11/C12 cleavage; in the spectrum of the acid corresponding to **5**, there appeared instead an m/z 223 ion attributable to C13/C14 cleavage.

The genesis of 1,3-dioxolan-2-yl cations by ionization of ortho esters, as depicted in Scheme 2 for the reaction $\mathbf{9} \rightarrow \mathbf{A}$, is well documented.⁴ Subsequent nucleophilic capture with ring opening typically leads to vicinally functionalized products such as bromohydrin esters. An orthoformate linked to a phenolic group gave the bromo formate when treated with AcBr but cyclized to the chromane under catalysis by PPTS, demonstrating the influence of the counterion on intervs intramolecular reactivity.^{4c} Here, the latter mode prevails by isomerization of \mathbf{A} to \mathbf{B} , which is formulated as an

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2442 Org. Lett., Vol. 8, No. 11, 2006

equilibrium driven forward by charge dispersal. Related iodocyclizations of homoallylic *t*-butyl carbonates have shown evidence of stereochemical equilibration prior to departure of the *t*-butyl cation, as the less sterically hindered diastereomer predominated in all cases.⁵ The same products also appear to be kinetically favored on the basis of results of low-temperature experiments.⁵ The preponderant formation of trans carbonates in the present work accords with these precedents.

In Bartlett's studies on carbonate extension, acetonitrile was found to play a critical role by trapping the t-butyl cation as the Ritter intermediate; in turn, this species proved able to transfer t-Bu $^+$ to dimethylformamide. Flausibly, similar transfers could impact the reaction of $9 \rightarrow 10/11$, e.g., by participation of the solvent as a cation shuttle; adjuvant trapping agents such as anisole might therefore be enlisted to assess the nature and extent of such effects.

To augment the supply of the anticonfigured triol **2**, a method was devised to utilize the major carbonate **10** by inversion at C12 (Scheme 3). Syn diol **12** was obtained from **10** by a sequence of deformylation, silylation, and carbonate cleavage. Exposure of **12** to the oxidant formed from 4-acetamido-TEMPO and p-TsOH· H_2O^{14} then delivered enone **13** in 75% yield. The compatibility of this oxidizing system with acid-sensitive functionality can be attributed to the low solubility of p-TsOH in CH_2Cl_2 . 14b Luche reduction of **13** at -50 °C was followed by cleavage of the t-butyl-dimethylsilyl ether to provide triols **2** and 12-epi-**2** in a 3.5:1 ratio. 15 No attempt was made to identify optimal conditions for this reduction.

In summary, the key features of this synthesis include utilization of the differential solvolytic lability of acyclic and cyclic orthoformates⁸ to access intermediate **9**, deployment

Scheme 3. Inversion at C12 1. i-Pr2NEt 0.01M in MeOH, 0 °C осно ΗÕ ĎН OTBS 2. TBSCI, Im, DN 3. NaOMe 0.2M TBSCI, Im, DMF 10 12 78%, 3 steps CH₂Cl₂, 0 °C 1. NaBH₄ CeCl₃ PPTS 1M OTBS 'nΗ НÕ ÓН 2. in MeOH

of a new 1,3-dioxolan-2-yl cation-mediated ScN' reaction to effect allylic transposition, and processing of the diaster-eomeric carbonates 10 and 11 to converge on the penultimate target 2.

87%, 2 steps

13

2

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Supporting Information Available: Experimental procedures and NMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

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Org. Lett., Vol. 8, No. 11, 2006

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