First Total Synthesis of Paracaseolide A

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ABSTRACT

The first total synthesis of the paracaseolide A, a very unusual tetraquinane oxa-cage bislactone recently isolated from the mangrove Sonneratia paracaseolaris, has been achieved. The final step and culmination of the eight step synthetic sequence is a [4+2] dimerization of a 4-hydroxybutenolide, generated by singlet oxygen-mediated oxidation of a furan precursor.

An investigation into the stem bark of Chinese mangrove plants, belonging to the genus Sonneratia, resulted recently in the isolation by Guo and his coworkers1 of a very unusual α-alkylbutenolide dimer which was named paracaseolide A (1, Scheme 1). Structural elucidation techniques revealed the presence of a rare tetraquinane bislactone skeleton bearing two adjacent linear alkyl chains on the convex face of its oxa-cage-like structure. Natural product 1 showed significant inhibitory activity against dual specificity phosphatase CDC25B, with an IC50 value of 6.44 μM.1 CDC25B is a key enzyme required for cell cycle progression and has been observed in a variety of cancers and is associated with tumor aggressiveness and poor prognosis.2 Herein, we report the first total synthesis of the highly complex molecular architecture that is paracaseolide A.

The primary goal of our research is to develop methods that provide more efficient and effective responses to synthetic challenges, particularly in the context of traditional intransigent polyoxygenated target structures.3 Thus, in any given piece of work, we are aiming to achieve as many of the recently delineated4 and exacting criteria for “ideal synthesis”5 as possible. Our synthesis of paracaseolide A was developed with these ideas in mind and with the goal of rapidly increasing molecular complexity with minimal use of concessionary steps.6 Even the use of the TBS-group is non-traditional as it serves several constructive purposes alongside its more conventional protecting role (vide infra).

The isolation team had proposed a very reasonable biosynthetic path for the assembly of paracaseolide A with a [4+2] dimerization of the α-alkenyl-γ-hydroxybutenolide 4 at its heart (Scheme 1). We would like to add to the proposal that a concerted Diels–Alder6 reaction is a "ideal synthesis" concept, see: Hendrickson, J. B. J. Am. Chem. Soc. 1975, 97, 5784.

5 For the first introduction to the "ideal synthesis" concept, see: Hendrickson, J. B. J. Am. Chem. Soc. 1975, 97, 5784.
starting \( E,E \)-diene 4 would be expected to afford the C7 diastereoisomer of the natural product, compound 2 (Scheme 1). Although a stepwise Diels–Alder reaction cannot be excluded, we believe that a concerted Diels–Alder reaction might be occurring followed by a fast epimerization of the readily epimerizable C7-centre of the initially formed diastereoisomer 2, leading to the formation of the natural compound 1 (Scheme 1). This type of epimerization should be thermodynamically favored since it repositions the long linear alkyl chain (appended at C7), from the concave, to the convex face of the cage-like structure. Following these considerations, our synthetic design (Scheme 1) hinged upon the use of this bioinspired \([4+2]\)-Diels–Alder dimerization of \( \alpha \)-alkenyl-\( \gamma \)-hydroxybutenolide 4, and, as such, it followed in the footsteps of other successful and elegant syntheses that have utilized Diels–Alder-dimerizations to access complicated architectures.\(^7\) It was anticipated that the \( \alpha \)-
alkenyl-γ-hydroxybutenolide 4 could be accessed through the singlet oxygen photooxygenation of an appropriately substituted furan precursor.

With this in our minds our investigation began with synthesis of the requisite furan photooxygenation precursor 11 which was accomplished as shown in Scheme 2. Lithium-halogen exchange of 1-iodotridecane (6) followed by addition to 3-furaldehyde resulted in the formation of furanol 7. The resultant hydroxyl functionality then served another useful purpose when it was utilized to effect a regioselective silylation of the furan (at the 2-position as opposed to the less hindered 5-position, 7 → 9) Specifically, (3-α-trialkylsililoxy)alkylfuran 8 underwent a [1,4] O→C silyl migration⁹ to furnish 2-trialkylsilyl-3-(α-hydroxy)alkyl-furan 9 (Scheme 2). After acidic dehydration using p-TsOH, the remaining unsubstituted ortho-position of the furan was methylated using standard conditions to afford the photooxygenation precursor furan 11. The presence of the TBS group at the 2-position of the furan not only guaranteed the regioselective methylation of position-5, it also directed the regioselectivity of the singlet oxygen (²¹Ο₂) oxidation⁹ such that it yielded exclusively the requisite α-substituted γ-hydroxybutenolide 4. In contrast to this clear and desired regioselective outcome, it is well known that γ-hydroxybutenolides prepared by photooxygenation of 3-alkyl furans are mixtures of α- and β-alkyl γ-hydroxybutenolides,¹⁰αβ or solely the β-regiosomers when Hunig’s base is included.¹⁰c Thus, through a short and high yielding synthetic sequence, α-alkenyl-γ-hydroxybutenolide 4 was synthesized.

With the monomer 4 in-hand, the stage was now set to explore the ambitious bioinspired [4+2] dimerization/ketalization/epimerization sequence to reach our target, paracaseolide A. Unfortunately, heating of a toluene solution of 4 at 110 °C in a sealed tube for 12 h, resulted in formation of the α,γ-disubstituted butenolide 13 in 42% isolated yield (Scheme 3). A mechanistic explanation for the formation of compound 13 invokes an intramolecular oxa-Michael type addition of the open keto-acid form 12 is shown in Scheme 3. In order to avoid this unwanted reaction (4 → 13) and favor the desired bimolecular [4+2] dimerization, we opted to investigate increasing the concentration. To our satisfaction when the reaction was performed neat in a sealed tube at 110 °C for 12 h, formation of 3:1 mixture of the natural product paracaseolide A (1) vs the α,γ-disubstituted butenolide 13 was observed by ¹H NMR analysis of the crude reaction mixture. Chromatographic separation of the two products afforded the clean natural product 1 in 59% isolated yield.

In summary, an extremely short (8 steps in total, starting from commercially available compounds), fast (executed easily in 4-5 days) and efficient protocol (15% overall yield) has been developed for the first total synthesis of paracaseolide A (1). The synthetic sequence involves transformation of a simple and readily accessible furan substrate into the desired 4-hydroxybutenolide 4 using a singlet oxygen-mediated oxidation.¹¹ This 4-hydroxybutenolide is the precursor for the final bioinspired [4+2]-dimerization/ketalization/epimerization that gives rise to the natural product paracaseolide A. This atom¹² and step-economic¹³ protocol, which utilizes

\[ \text{Scheme 3. Bioinspired [4+2] Dimerization of 4-Hydroxybutenolide 4 to Paracaseolide A} \]

\[ \text{10a} \]

\[ \text{10c} \]

\[ \text{11} \]

\[ \text{12} \]

\[ \text{13} \]
environmentally-benign oxygen from the air as the oxidant, and, which uses a bioinspired [4+2]-cycloaddition to accomplish rapid increase of molecular complexity in one synthetic operation, represents an achievement that takes us a step, albeit a small one, towards achieving the prized goal of an ideal synthesis.\textsuperscript{4}

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**Supporting Information Available:** Experimental procedures, full spectroscopic data and copies of $^1$H and $^{13}$C-NMR spectra for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.