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Toward the Synthesis of Reidispongiolide A: Stereocontrolled Synthesis of the $C_{17}-C_{22}$ and $C_{23}-C_{35}$ Degradation Fragments

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ABSTRACT

Reidispongiolide A

By relying on the asymmetric aldol reactions of chiral ketones, a highly stereocontrolled synthesis of each of the C_{17} – C_{22} and C_{23} – C_{35} degradation fragments of reidispongiolide A has been achieved. This permits a configurational assignment of the complete C_{17} – C_{36} region of this antimitotic macrolide, along with providing advanced intermediates for a projected total synthesis.

The discovery of new cytotoxins that retain activity toward multidrug resistant (MDR) cancer cell lines continues to fuel the burgeoning field of cancer chemotherapy. Recently, a number of actin-binding macrolides of marine origin have attracted attention as novel antimitotic agents that cause rapid loss of microfilaments in cells, without affecting microtubule organization. In particular, the scytophycins, aplyronines, sphinxolides, and reidispongiolides demonstrate pronounced antimicrofilament activity in vitro and potently inhibit the

growth of MDR cancer cells. While these preliminary findings highlight their potential, both as candidates for new anticancer drugs and versatile molecular probes of the organization and function of the actin cytoskeleton, the scarcity from the natural sources has generally hampered the biological evaluation and preclinical development of these stereochemically complex macrolides.

The sphinxolide/reidispongolide family of 26-membered macrolactones are prominent members of this emerging class of actin-binding cytotoxic macrolides (Scheme 1). Originally isolated from an unidentified Pacific nudibranch,^{5a} these polyketide metabolites have more recently been obtained from the marine sponges *Neosiphonia superstes* and *Reidispongia coerulea*,^{5b-d} collected off the coast of New Caledonia. Extensive analysis of sphinxolide A by NMR methods, based largely on the interpretation of proton—carbon ^{2,3}*J* couplings, has recently enabled an assignment of the *relative* configurations in the five isolated stereoclusters, as indicated in the boxed regions of structure 1.⁶ Additionally, controlled ozonolysis of the closely related macrolide reidispongiolide A (having the proposed stereostructure in 2), followed by

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reductive workup with NaBH₄, provided three distinct polyol fragments, assigned as structures 3 (mixture of C_5 epimers) and 4 and 5 (mixture of C_{31} epimers), whose spectroscopic data appeared consistent with that of the corresponding segments in $1.^7$ While the full configuration of the sphinx-olide/reidispongiolide macrolides could be determined by completing a total synthesis, which remains our ultimate objective, the preparation of these three degradation fragments (or stereoisomers thereof) should simplify the stereochemical quandary and establish the interconnections between the isolated stereoclusters.

6: $R^1 = Me$, $R^2 = R^3 = H$, $R^4 = OMe$

We now report an expedient stereocontrolled synthesis of the reidispongiolide fragments **4** and **5**, as well as the diastereomer **6** having the opposite configuration at C₃₂ and C₃₃, following the retrosynthetic analysis outlined in Scheme 1. On the basis of our reservations regarding the relationship between the isolated stereoclusters, we designed a flexible and modular synthetic approach using appropriate aldol reactions of the chiral ketones **7**, **8**, **9**, and *ent-***9**. The present work leads to a configurational assignment for 10 out of the 15 stereocenters in reidispongiolide A, and also sets a solid foundation for ongoing total synthesis efforts.

First, an asymmetric synthesis of the C₁₇–C₂₂ fragment **4** of reidispongiolide A using a convenient one-pot aldol/reduction sequence⁸ with methyl ketone **7**⁹ and crotonaldehyde was developed (Scheme 2). In previous work with such ketones,^{8–11} we had shown that high levels of diastereoselectivity can be obtained for 1,4-*syn* adducts by appropriate choice of Ipc ligand chirality in boron aldol reactions. By using our standard conditions with (—)-Ipc₂BCl/Et₃N, the resulting boron aldolate was reduced^{8b} in situ with LiBH₄ to provide the 1,3-*syn* diol **10** with high diastereoselectivity (91%, 91:9 dr). Conversion into the bis-methyl ether (NaH, MeI) was then followed by ozonolysis and in situ reduction (NaBH₄) of the ozonide to give the corresponding alcohol. Finally, TBS ether removal (TBAF) afforded the

ORTEP drawing derived from the X-ray analysis of 1

^a Conditions: (a) (−)-Ipc₂BCl, Et₃N, Et₂O, −78 °C; crotonal-dehyde; LiBH₄; (b) NaH, MeI, THF; (c) O₃, CH₂Cl₂, −78 °C; MeOH, NaBH₄, −78 °C; (d) TBAF, THF; (e) *p*-nitrobenzoyl chloride, Et₃N, DMAP, CH₂Cl₂.

diol **4**, obtained in four synthetic transformations and 27% yield from **7**.

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The ¹H and ¹³C NMR data exhibited by this diol **4** were consistent with that reported by D'Auria and co-workers for both the C₁₇-C₂₂ fragment obtained from the chemical degradation of natural reidispongiolide A and that of a synthetic sample of *ent*-**4**.⁷ The absolute configuration of the degradation fragment was further confirmed as (18*S*,19*S*,-21*R*) through comparison of the specific rotation values.¹² Additionally, treatment of the diol **4** with *p*-nitrobenzoyl chloride (Et₃N, DMAP) gave the bis-ester **11**, where single-crystal X-ray analysis confirmed the *all-syn* stereochemistry.

The convergent assembly of the C_{23} – C_{35} fragments commenced with a substrate-controlled, boron-mediated aldol reaction (c-Hex₂BCl, Et₃N)^{8a,10} between the ethyl ketone $\mathbf{8}^{13}$ and aldehyde $\mathbf{12}$ (Scheme 3). Upon standard oxidative

^a Conditions: (a) c-Hex₂BCl,Et₃N,Et₂O, −78 °C; (b) Me₄NBH(OAc)₃, MeCN/AcOH; (c) DDQ, 4 Å MS, CH₂Cl₂; (d) NaH, MeI, THF; (e) TBAF, THF.

workup, the expected anti-anti adduct 13, resulting from the high level of π -face discrimination exercised by the intermediate (E)-enolate,8 was obtained in 95% yield (95:5 dr). This adduct provided a suitable substrate for hydroxyldirected reduction to set up the C23-C29 stereopentad. By employing Me₄NBH(OAc)₃,¹⁴ the desired 1,3-anti diol 14 was obtained cleanly (90%, >99:1 dr). With the five contiguous stereocenters now secured in a concise manner, the differentiation of the two hydroxyl groups was required. Treatment of the diol 14 under DDQ-mediated oxidative cyclization conditions¹⁵ resulted in the exclusive formation of the corresponding six-membered PMP acetal, as a single diastereomer. Finally, methylation of the free hydroxyl (NaH, MeI) and removal of the TIPS ether (TBAF) afforded the crystalline alcohol 15 in 80% overall yield from 14. At this point, the relative stereochemistry of this C₂₃-C₂₉ subunit was confirmed by X-ray crystallographic analysis of 15.

With the required stereopentad **15** in hand, we turned to preparing the methyl ketones **9** and *ent-***9** to access both possible *anti*-relationships at C_{32} and C_{33} in the extended C_{23} – C_{35} fragment (Scheme 4). By using Brown's methodology, ¹⁶ the aldehyde **16** was treated with the (*E*)-crotylborane reagent derived from *trans*-butene and (–)-Ipc₂BOMe to

afford the *anti*-adduct **17** in 81% yield (95:5 dr, 95% ee).¹⁷ Following methyl ether formation (NaH, MeI), the terminal alkene was oxidized to give **9** (71%)¹⁸ under modified¹⁹ Wacker conditions. By using (+)-Ipc₂BOMe, the enantiomeric methyl ketone *ent-***9** was prepared from **16** in an analogous manner.

We next examined the coupling of the ketone **9** and the aldehyde **18**, obtained by Dess—Martin oxidation of **15** (99%). This pivotal aldol coupling step was best achieved by generation of the lithium enolate of **9** (LDA, THF, -78 °C) and addition of a solution of **18** (0.5 equiv, 30 min) to give the β -hydroxy ketone **19** in 85% yield, as an inconsequential 4:1 mixture of diastereomers. Subsequent elimination, through formation of the corresponding mesylate (MsCl, Et₃N) and in situ treatment with DBU, provided (*E*)-enone **20** (65%). Subjection of **20** to standard hydrogenation conditions (H₂, Pd/C) then effected both clean hydrogenolysis of the PMP acetal and reduction of the alkene. Finally, ketone reduction (NaBH₄) and TBS ether removal (TBAF) provided the alcohols **5**, obtained as a 4:1 mixture of epimers at C₃₁, which were separated by flash chromatography.

Notably, the ¹H NMR data (500 MHz, CD₃OD) of the alcohols 5 were in close agreement²⁰ to that reported by D'Auria and co-workers for the C₂₃-C₃₅ fragment obtained in their degradation work,7 thus supporting this relationship between the two remote stereoclusters at C_{24} – C_{28} and $C_{31}-C_{33}$ in reidispongiolide A. At the time, inconsistencies between our ¹³C NMR data and that reported by the Naples group led us to prepare 6 having the other anti stereorelationship at C_{31} – C_{32} . This involved the analogous aldol coupling between 18 and ent-9 to give 21 followed by elaboration into 6, obtained as a 4:1 mixture of epimers at C₃₁. Spectroscopic analysis²⁰ of the separated alcohols **6** revealed that these were clearly diastereomers of the C₂₃-C₃₅ fragment obtained by ozonolysis of reidispongiolide A. Subsequently, comparison of the ¹³C NMR data for synthetic fragments 5 and 6 with the revised data provided²¹ for material obtained by chemical degradation allowed the confident assignment of the *relative* stereochemistry for the C₂₃-C₃₅ sequence of reidispongiolide A.

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⁽¹²⁾ The low value of $[\alpha]^{20}_D$ recorded for synthetic 4 (+4.1, MeOH) was in accord with that reported (ref 7) for the corresponding degradation fragment. The bis-(S)-MTPA ester was also prepared for spectroscopic comparison.

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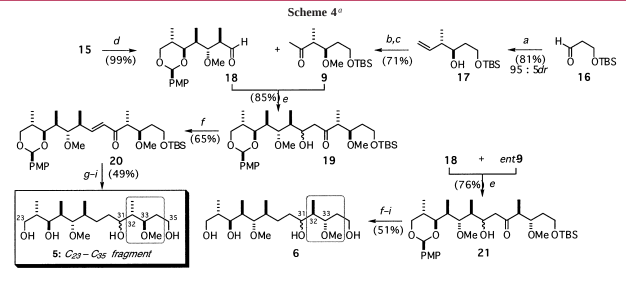
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⁽²¹⁾ Professor D'Auria has since provided us with revised 13 C NMR data for their C_{23} – C_{35} degradation fragment which are in complete agreement with the data we have acquired for the alcohols 5.



^a Conditions: (a) *trans*-butene, *t*-BuOK, THF, *n*-BuLi; (−)-Ipc₂BOMe; BF₃•OEt₂; **16**, −78 °C; (b) NaH, MeI, THF; (c) O₂, Cu(OAc)₂ (20 mol %), PdCl₂ (10 mol %), AcNMe₂/H₂O (4:1); (d) Dess−Martin periodinane, NaHCO₃, CH₂Cl₂; (e) LDA, THF, −78 °C; (f) MsCl, NEt₃, CH₂Cl₂; DBU; (g) H₂, cat. Pd/C, EtOH; (h) NaBH₄, MeOH; (i) TBAF, THF.

In conclusion, on the basis of the established absolute configuration in the C_{17} – C_{22} sequence and the relative configuration at C_{23} – C_{35} , and relying on a common biogenesis for the sphinxolide/reidispongioloide family and the structurally related scytophycin and aplyronine macrolides, ^{1a} a full assignment of the complete C_{17} – C_{36} region of reidispongiolide A, as indicated in **22**, is strongly suggested. Confirmation of this proposal will rely on completing the total synthesis of reidispongiolide A, work that is ongoing in our laboratory.

22: C₁₇-C₃₆ sequence of reidispongiolides

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Supporting Information Available: Physical and spectroscopic data for new compounds and CIF files for **11** and **15**. This material is available free of charge via the Internet at http://pubs.acs.org.

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