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Progressive-Convergent Elucidation of Stereochemistry in Complex Polyols. The Absolute Configuration of (–)-Sagittamide A

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Abstract

The absolute stereostructure of sagittamide A (1), a O-hexacetyl long-chain hexahydroxy- α , ω -dicarboxylic acid, was assigned using a progressive-convergent approach that integrates three powerful regimens for stereochemical analysis of acyclic natural products: J-based analysis, 13 C NMR universal database comparisons and exciton coupling circular dichroism.

The structure of (–)-sagittamide A ($\mathbf{1}$)¹–an unprecedented polyacetoxy, long-chain α , ω -dicarboxylic acid isolated from a tropical didemnid tunicate—was solved by application of conventional 2D NMR spectroscopic methods, however, only partial stereochemistry could assigned. Although configurations of the terminal amino acids (L-ornithine and L-valine) were determined readily by conventional methods, the contiguous 5,6,7,8,9,10-hexol hexaacetate in $\mathbf{1}$ represented a significantly more complex NMR problem, in part, because of isolated stereohexad C5–C10 flanked by CH₂ groups² and equivocal interpretations of J coupling.

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We now report the complete stereostructure of **1** using a *progressive-convergent* approach that integrates three powerful regimens for stereochemical analysis of natural products: use of Murata's *J*-based analysis (${}^{3}J_{HH}$ and ${}^{2,3}J_{HC}$), application of Kishi's Universal Database⁴ (pairwise-comparison of ${}^{13}C$ NMR chemical shifts with stereo-defined models) and highly sensitive exciton coupling circular dichroism (ECCD). The integrated approach rapidly converges upon a unique stereochemical assignment for **1** with internal validation.

A basis set of ${}^3J_{\rm HH}$ and ${}^{2,3}J_{\rm CH}$ values were obtained by 2D heteronuclear 2D NMR experiments of **1** (COSY and HSQMBC, respectively, see Supporting Information) and used to predict an all *anti*-relative configuration for C6–C9 for **1**. Consequently, the number of remaining possible diastereomers of **1** was reduced from N=32 to 4. A synthetic route to six model compounds, representing permutations of the six stereocenters C5–C10 congruent with those proposed for **1**, was conceived and executed starting with D-xylose (see Supporting Information). In order to address an equivocal C8 ${}^3J_{\rm CH}$ value in **1**, a parallel set of models **2–9** was also prepared from D-ribose as described below (Scheme 1).

Indium-promoted Barbier reaction of D-ribose with allyl bromide gave a 2:1 mixture of epimeric homoallylic alcohols⁷ **10** and **11** after protection. Each acetonide was deprotected and hydrogenated (Pd/C, CF₃CH₂OH, 1 atm H₂)⁸ followed by Swern oxidation to the corresponding C9 aldehydes and homologation using two stereocomplementary methods (*Z*-selective Wittig olefination using phosphonium salt **14** and *E*-selective Julia-Kocienski olefination with tetrazole **15**⁹) to give **12** and **13**.

Stereoselective OsO₄ dihydroxylation¹⁰ of **12** gave diols **16** and **17**. In this manner, *E*- and *Z*-olefins were converted to diol diastereomers and purified by HPLC, prior to deprotection to the hexaols. Peracetylation of each hexaol furnished the eight C7–C9 *ribo*-model compounds **2–9** and six *xylo*-models (Supporting information). The correct relative configuration of **1** emerged from ¹³C NMR comparisons with the model compounds (Figure 1).⁴

The 1 H and 13 C NMR spectra of each peracetate model were carefully assigned from COSY and HMBC spectra. Pairwise comparisons of the differences of the 13 C chemical shifts ($\Delta\delta$) for C4–C11 in model compounds and **1** clearly showed an excellent match for the C8 epimer **6** obtained from D-ribose, but an mismatch for the corresponding *xylo*-C8 epimer (e.g. C8: $\Delta\delta = +0.05$ and -3.93 ppm, respectively, see Supporting Information). A valuable object lesson is revealed here that promotes a progressive-convergent approach to stereochemistry. Although anomalous $^{3}J_{\text{CH}}$'s in **1** predicted an erroneous *xylo*-configuration during *J*-based analysis, 11 this was readily rectified in the progressive 13 C $\Delta\delta$ analysis allowing reassignment of C8 configuration to that of **6**.

The absolute stereochemistry of **1** was secured by transformation of the natural product, and hexaol diastereomers corresponding to **6** and **7**, to the per-benzoate ester derivatives, **18**, **19** and **20**, respectively, ¹² and comparison of their corresponding CD spectra (Figure 2). Since the fingerprint Cotton effects observed in the CD spectra of **18** and **19** were equal in magnitude but opposite in sign, the absolute configuration of **1** corresponds to *ent-***19** and is related to L-ribose. ¹² Thus, the complete configuration of sagittamide A (**1**) is depicted as (5*S*,6*S*,7*S*,8*R*,9*R*,10*S*).

In summary, we have deployed an integrated approach to solve the configuration of sagittamide A (1). The power of this triple-combination of methodologies lies in judicious interpretation of homonuclear 3J and heteronuclear ${}^{2,3}J$ to provide *partial* stereochemical information which is then used to inform correct choices for synthesis of model compounds to be used in the next stage: ${}^{13}C$ NMR comparative analysis.

A significant advantage is gained by a requirement for only a limited sub-set of stereo-model compounds without the necessity for synthesis of all 64 possible permutations. The progressive-convergent approach succeeds where other singular methods based on NMR may become irreducibly complex ¹³ or rendered equivocal by second-order effects that militate against reliable stereochemical assignments.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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- 6. The carbons numbered C7, C8 and C9 in 1 map to C4, C3 and C2 of ribose or xylose, respectively. Thus, the stereochemical descriptors 'xylo-' and 'ribo-' in the context of this work refer to C7–C9 of 1.
- 7. The configuration of the major isomer was assigned by analogy with the well-known 1,2-syn-stereopreference for In°-promoted allylation of aldohexoses [Kim E, Gordon DM, Schmid W, Whitesides GM. J Chem Org. 1993; 58:5500–5507.Kobayashi S, Nagayama S. J Org Chem. 1996; 61:2256–2257.] and subsequent conversion to the acetonides 10 and 11.
- 8. Deprotection of **10** and **11** to the corresponding primary alcohols was rapidly effected when CF₃CH₂OH was used as solvent for hydrogenolysis. No reaction was observed in ethanol, even after several days at 3 atm H₂.
- 9. Blakemore PR, Cole WJ, Kocienski PJ, Morley A. Synlett. 1998; (1):26–28.Both **14** and **15** were prepared from δ-valerolactone in three and four steps, respectively (see Supporting Information).
- 10. Diastereomeric assignments of 5,6-diols were based on the expectation of anti-selectivity of OsO₄ addition to allylic alcohols and confirmed by the outcomes from double-diastereoselection using the Sharpless asymmetric dihydroxylation (Kolb HC, VanNieuwenhze MS, Sharpless KB. Chem Rev. 1994; 94:2483–547.) and observed pseudo-C₂ symmetry in the ¹H and ¹³C NMR spectra of 2 and 8. See Supporting Information.
- 11. This observation suggests caution in using J-based methodology and over-reliance on the underlying assumption of all-staggered conformations and the accuracy of J's measured in strongly coupled contiguous polyols that may not be amenable to first-order spin analysis.
- 12. The lactam-mono methyl ester that formed spontaneously upon treatment of 1 (CH₂N₂, MeOH-ether, ref. 1) and the hexaols corresponding to 6 and 7 were each converted (excess BzCl, pyridine, 40 °C) to hexabenzoates 17, 18, and 19, respectively, after HPLC purification. Benzoylation at higher temperatures (60–90 °C) lead to significant formation of tetrabenzoyloxy-tetrahydrofuran.
- 13. The similarity of CD spectra of diastereomeric **19** and **20** reflect the dominance of the C7–C10 configuration on the Cotton effects.

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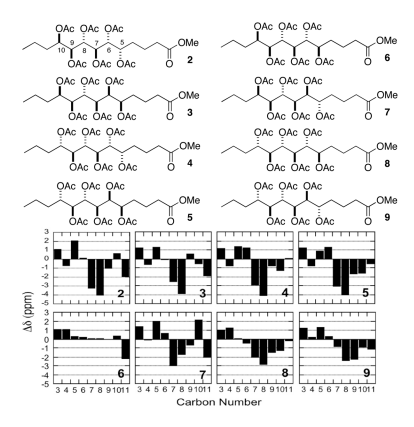


Figure 1. ^{13}C NMR (125MHz, d₆-DMSO, T=298 K) $\Delta\delta$ values (δ_C 1– δ_C model) of ribo-model compounds 2–9

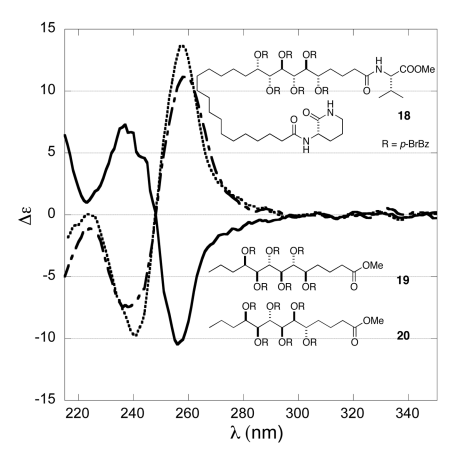


Figure 2. CD spectra of sagittamide A derivative 18 (—), together with models 19 (…) and 20 (— -), (CH₃CN, $c=10~\mu$ M).

Scheme 1.

a) In°, allyl bromide, H₂O; b) TrCl, pyridine, reflux 53% (2 steps); c) CSA, acetone, CH₃C(OCH₃)₂CH₃ 58%, **10:11** dr 2:1); d) SiO₂-HPLC 1:19:EtOAc hexanes; e) H₂, 1 atm, Pd/C, CF₃CH₂OH, 35–69%; f) i. (COCl)₂, DMSO, CH₂Cl₂, -78 °C ii. Et₃N; g) i. **15**, DME, NaHMDS, -78 °C, ii. aldehyde, 25%, dr 3:1 (2 steps); h) i. **14**, THF, NaHMDS, -78 °C, ii. aldehyde, dr>19:1,16% 2 steps; i) OsO₄, NMO, acetone, H₂O; dr 1.7:1, 93%; j) K₃Fe(CN)₆, K₂OsO₄, K₂CO₃ (DHQ)₂PHAL, t-BuOH, H₂O, CH₃SO₂NH₂, dr 3.8:1, 86%; k) 2% TMS-Cl, MeOH, ii. CH₂N₂, ether/MeOH, iii. Ac₂O, pyridine 6h: 22% **3** (3 steps), 48%; **2** (3 steps), 44%, **6** (4 steps), 26%, **7** (4 steps).