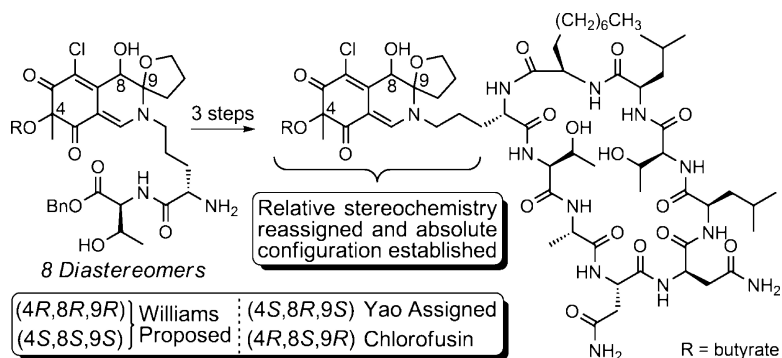


Total Synthesis of Chlorofusin, Its Seven Chromophore Diastereomers, and Key Partial Structures

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J. Am. Chem. Soc., **2008**, 130 (37), 12355-12369 • DOI: 10.1021/ja8012819 • Publication Date (Web): 20 August 2008

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Total Synthesis of Chlorofusin, Its Seven Chromophore Diastereomers, and Key Partial Structures

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Abstract: Chlorofusin is a recently isolated, naturally occurring inhibitor of p53–MDM2 complex formation whose structure is composed of a densely functionalized azaphilone-derived chromophore linked through the terminal amine of ornithine to a nine residue cyclic peptide. Herein we report the full details of the total synthesis of chlorofusin, resulting in the assignment of the absolute stereochemistry and reassignment of the relative stereochemistry of the complex chromophore. Condensation of each enantiomer of an azaphilone chromophore precursor with the N^3 -amine of a protected ornithine–threonine dipeptide, followed by a one-step oxidation/spirocyclization of the most reactive olefin provided all eight diastereomers of the fully elaborated chromophore–dipeptide conjugate. Comparison of the spectroscopic properties for these eight compounds and those of simpler models with that reported for the natural product allowed the full assignment of the (4*R*,8*S*,9*R*)-stereochemistry of the chlorofusin chromophore. The natural, but stereochemically reassigned, diastereomer of the dipeptide conjugate was incorporated in a convergent total synthesis of chlorofusin confirming the stereochemical reassignment and establishing its absolute stereochemistry. Similarly and enlisting the late stage convergent point in the total synthesis, the remaining seven diastereomers of the chromophore–dipeptide conjugates were individually incorporated into the nine-residue cyclic peptide of chlorofusin (4 steps each) providing all seven remaining possible chromophore diastereomers of the natural product.

Introduction

The tumor suppressor p53 is an important part of an innate cancer defense mechanism and acts as a transcription factor that initiates cell cycle arrest and apoptosis in response to stress such as DNA damage.^{1–4} The activity of p53 is modulated by MDM2 (HDM2), which tightly binds p53 preventing it from acting as a regulator of cell division^{5–7} and targeting it for nuclear export and degradation.^{8,9} Overexpression of MDM2 has been implicated in many cancers,^{10–16} defining the p53–MDM2 interaction

as an attractive target for therapeutic intervention.^{4,17} An X-ray crystal structure of the N-terminal domain of MDM2 bound to a 15-residue transactivation domain of p53 revealed the structural details of their complex that is mediated by the interaction of three hydrophobic residues of a p53 α -helix with a hydrophobic cleft of MDM2.¹⁸ Molecules that bind the hydrophobic cleft of MDM2 disrupt this protein–protein interaction with p53, restoring its regulatory function and inhibiting tumor growth.^{4,19–22}

Chlorofusin (**1**, Figure 1) was isolated from the fungal strain *Microdochium caespitosum*²³ and found to disrupt the p53–MDM2 interaction by directly binding to the N-terminal domain of MDM2 ($IC_{50} = 4.6 \mu\text{M}$, $K_D = 4.7 \mu\text{M}$).^{23,24} Thus, chlorofusin represents an exciting lead for antineoplastic intervention that

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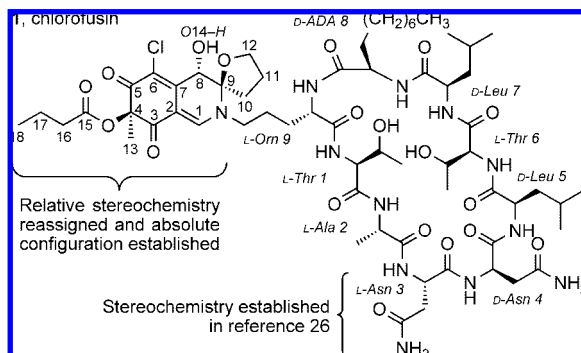


Figure 1. Chlorofusin.

acts by a rare disruption of a protein–protein interaction although the structural details of this inhibitory interaction derived from MDM2 binding have yet to be established.^{20,21,25} On the basis of extensive spectroscopic and degradation studies, the chlorofusin structure was proposed to consist of a densely functionalized, azaphilone-derived chromophore linked through the terminal amine of ornithine to a 27-membered cyclic peptide composed of nine amino acid residues.²³ Two of the cyclic peptide amino acids possess a nonstandard or modified side chain, and four possess the D-configuration. Although the spectroscopic and degradation studies of chlorofusin permitted the identification of the cyclic peptide structure and connectivity, the two asparagine residues Asn3 and Asn4 were only established to have opposite stereochemistries (L and D) and their respective assignments were not possible. In earlier studies, we reported the synthesis of the two cyclic peptide diastereomers bearing either the L-Asn3/D-Asn4 or D-Asn3/L-Asn4 stereochemistry and correlation of the former with the spectroscopic properties (¹H and ¹³C NMR) of the natural product.²⁶ In that work, four key subunits were assembled, sequentially coupled, and cyclized to provide the nine residue cyclic peptide. The coupling and macrocyclization sites were chosen to minimize the use of protecting groups and maximize the convergency of the synthesis, and a deliberate late-stage incorporation of the subunit bearing the two asparagine residues allowed convenient access to both diastereomers required to assign the absolute stereochemistry. Concurrent with this disclosure, Searcey reported the synthesis of the L-Asn3/D-Asn4 diastereomers incorporating either a D-ADA8 or L-ADA8 residue,²⁷ and recently Nakata²⁸ has also reported a synthesis of this cyclic peptide.

Similarly, the chlorofusin spectroscopic studies conducted by Williams provided the structure and an assigned relative stereochemistry for the unusual azaphilone-derived chromophore, but did not permit an assignment of its absolute stereochemistry (Figure 2). The relative stereochemistry was assigned using gradient 1D NOE studies in which irradiation of the proton attached to C8 with a mixing time of 50 ms resulted in NOEs to the protons attached to C10 as well as C12, whereas a shorter mixing time of 25 ms resulted in NOEs to

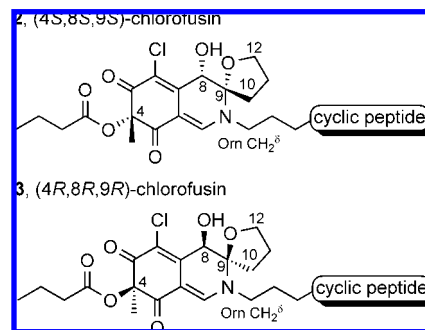


Figure 2. Original chromophore relative stereochemical assignment.²³ No assignment of absolute stereochemistry was made.

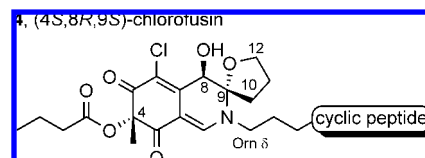


Figure 3. Stereochemical assignment made by Yao and co-workers.³⁰

only the protons attached to C10. In addition, irradiation of the C4–Me protons provided a long-range NOE to the proton attached to C8, but only if conducted with very extended mixing times (500 ms). These results suggested that the protons attached to C8, C10 and C4–Me all lie on the same face of the chromophore and led to the relative stereochemical assignment depicted in Figure 2.

Herein, we report the full details of studies leading to a reassignment of this relative stereochemistry for the chlorofusin chromophore that is still, but less obviously, consistent with these experimentally observed NOEs, as well as the assignment of the chromophore absolute configuration (4*R*,8*S*,9*R*). A total synthesis of this revised chlorofusin structure provided material displaying spectroscopic properties indistinguishable from that reported for the natural product confirming the new chromophore structural assignment and establishing the accuracy of our earlier L-Asn3/D-Asn4 cyclic peptide stereochemical assignment. A preliminary account of the first stage of these studies was disclosed²⁹ immediately following the report of Yao³⁰ which claimed to have completed a total synthesis of the natural product. Although this group similarly reassigned the chromophore relative stereochemistry, they misassigned the absolute stereochemistry overlooking key diagnostic spectroscopic differences that distinguish their unnatural (4*S*,8*R*,9*S*)-diastereomer **4** (Figure 3) from the natural (4*R*,8*S*,9*R*)-diastereomer. In extending our previously disclosed studies which provided all four 4*R*-diastereomers of chlorofusin, we have now additionally incorporated the four 4*S*-diastereomers of the chlorofusin chromophore (enantiomeric series) into fully elaborated chlorofusin diastereomers. Thus, in addition to natural chlorofusin, the remaining seven chlorofusin chromophore diastereomers [(4*R*,8*R*,9*R*), (4*R*,8*S*,9*S*), (4*R*,8*R*,9*S*), (4*S*,8*S*,9*S*), (4*S*,8*R*,9*R*), (4*S*,8*S*,9*R*), and (4*S*,8*R*,9*S*)] including the (4*R*,8*R*,9*R*)/(4*S*,8*S*,9*S*)-diastereomers proposed by Williams as well as the

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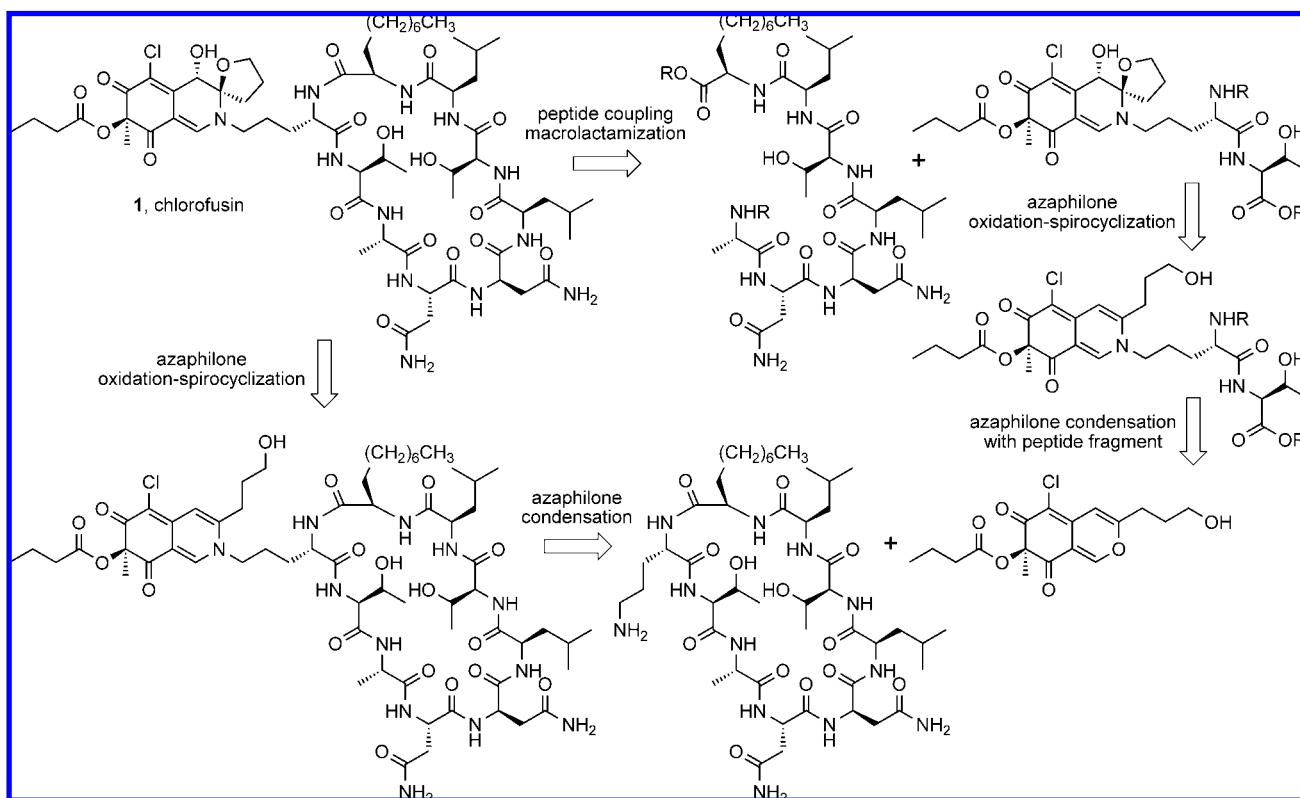
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Scheme 1



Yao misassigned (4*S*,8*R*,9*S*)-diastereomer were synthesized as key analogues of the natural product and the comparison of their spectral properties with that reported for chlorofusin provide unambiguous support for the structural reassignment.

The chlorofusin chromophore is derived from a class of compounds referred to as azaphilones, so named for their ability to readily condense with ammonia or primary amines.³¹ In light of this, at least two potential routes to chlorofusin could be envisioned: (1) condensation of an appropriate azaphilone with the full cyclic peptide followed by chromophore oxidation and spirocycle formation, or (2) construction of the fully functionalized chromophore appended to a smaller peptide fragment followed by peptide coupling and macrocyclization to afford chlorofusin (Scheme 1). The former requires the conduct of an azaphilone chromophore oxidative spirocyclization on a complex, late stage synthetic intermediate with control of the resulting relative and absolute stereochemistry or effective separation and characterization of the resulting diastereomers, whereas the latter requires that the chromophore *N,O*-spiroketal in the smaller peptide fragment not unravel during the late stages of the ensuing cyclic peptide synthesis. Both routes require the same precursor azaphilone allowing us to potentially explore either route as the work progressed.

Because the absolute configuration of the chromophore was unknown and the relative stereochemical assignment came under question as the work progressed, the routes pursued in our first generation synthesis permitted access to both enantiomeric series and to all possible diastereomers, albeit optimized to access the original diastereomer assigned by Williams. As discussed herein, confidence in a requisite reassignment of the relative stereochemistry emerged with the observation of diagnostic spectro-

scopic distinctions (¹H and ¹³C NMR) made possible by access to all diastereomers. Due to concerns of an unambiguous stereochemical assignment as the studies progressed, the strategy of early stage versus late stage chromophore oxidative spirocyclization with clear stereochemical assignments for all diastereomers became the focus of our efforts and was effectively conducted on the azaphilone adduct with an ornithine–threonine dipeptide. Beautifully, even an assignment of the chromophore absolute configuration emerged from diagnostic distinctions observed in the spectroscopic properties of the series of all eight diastereomers derived from the chromophore adduct (four diastereomers of the two enantiomeric series) that was confirmed by the total synthesis of the natural product. In retrospect, a late stage chromophore elaboration, like that of Yao,³⁰ may have failed to produce sufficient quantities of the natural product for detection and may not have allowed an unambiguous stereochemical assignment given that the natural product was found to possess a contrathermodynamic *N,O*-spiroketal stereochemistry.

Results and Discussion

Chromophore Azaphilone Precursor: Route 1. The initial approach to the key intermediates **17** and **18** used to explore the C8–C9 oxidative spirocyclization followed established routes to access azaphilones bearing a range of C9 substituents,^{32–34} and began with the one carbon homologation of commercially available 3,5-dimethoxy-4-methylbenzoic acid (**5**) (Scheme 2). The crude acid chloride derived from **5** was alkylated with TMSCH₂Bt to afford the ketone adduct (SOCl₂, reflux;

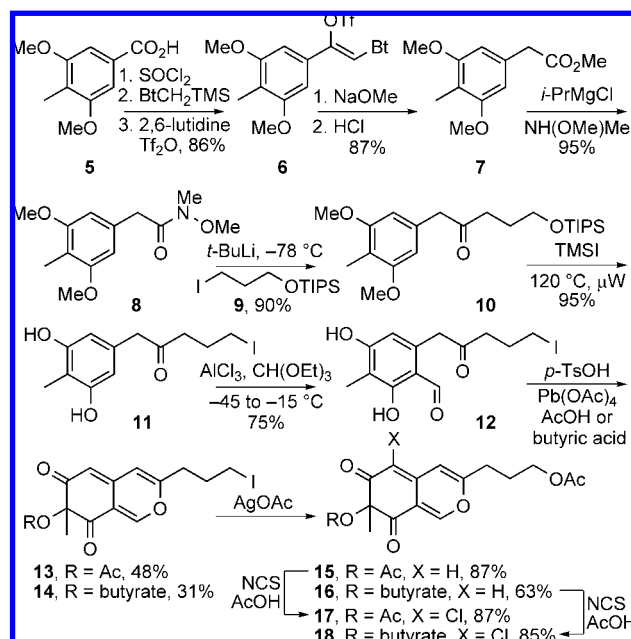
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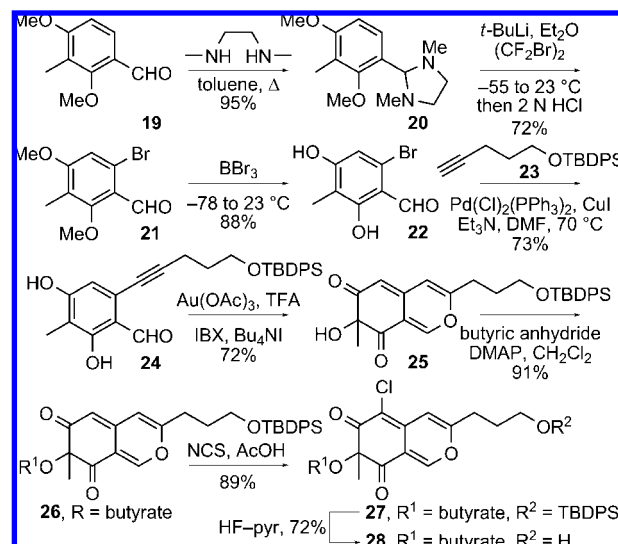
Scheme 2



TMSCH₂Bt, THF, reflux, 6 h)³⁵ and subsequent treatment with 2,6-lutidine and Tf₂O provided **6** (2,6-lutidine, Tf₂O, CH₂Cl₂, 0 to 23 °C, 2 h, 86%, three steps). The enol triflate **6** was converted to the methyl ester **7** by treatment with NaOMe followed by acidic hydrolysis of the resulting ynamine (NaOMe, MeCN, reflux, 12 h; 12 N HCl, MeOH, reflux, 24 h, 87%, two steps). After conversion to Weinreb amide **8** (NHMe(OMe), *i*-PrMgCl, THF, -20 °C, 30 min, 95%),³⁶ the lithium reagent derived from **9** was added to **8** to provide the ketone **10** (**9**, *t*-BuLi, -78 °C, 2 h, Et₂O-THF, 90%). Selective deprotection of the two methyl ethers of **10** in the presence of the silyl ether using BBr₃, BF₃·Et₂O, AlCl₃, LiI, TMSI, or PhSH was not successful. In all cases, the silyl ether was found to preferentially or nonproductively undergo cleavage. However, the two methyl ethers were cleaved and the TIPS-protected primary hydroxyl group was displaced cleanly with iodide to provide **11** in remarkably high yields when **10** was treated with TMSI under microwave irradiation (TMSI, MeCN, 120 °C, 50 min, normal absorption, 95%). Formylation of **11** provided **12** (AlCl₃, CH(OEt)₃, toluene, -45 to -15 °C, 1 h, 75%)³⁴ and one-pot oxidative cyclization of **12** in acetic acid gave the azaphilone **13** while conducting the same reaction in butyric acid provided **14** (*p*-TsOH, AcOH or butyric acid, 95 °C, 2 h; Pb(OAc)₄, 15 °C, 40 min, 48% for **13**, 31% for **14**).³⁴ Displacement of the primary iodide of **13** and **14** with acetate (AgOAc, AcOH, 55 °C, 2 h, 87% for **15**, 63% for **16**)³⁷ gave **15** and **16** whose regioselective C6-chlorination (NCS, AcOH, 23 °C, 24 h, 87% for **17**, 85% for **18**)^{38,39} afforded the chloroazaphilones **17** and **18**.

Chromophore Azaphilone Precursor: Route 2. The analogous intermediate **28** was also prepared by a more concise route based

Scheme 3



on the work of Porco.³⁸ Although a one-step conversion^{40,41} of commercially available 2,4-dimethoxy-3-methylbenzaldehyde to 6-bromo-2,4-dimethoxy-3-methylbenzaldehyde suffered from poor conversions, a more controlled two-step procedure^{42,43} was adopted beginning with the condensation of 2,4-dimethoxy-3-methylbenzaldehyde (**19**) with *N,N'*-dimethylethylenediamine (toluene, 130 °C, 3 h, 95%) (Scheme 3). The resulting dimethylimidazolidine **20** was regioselectively lithiated (*t*-BuLi, Et₂O, -55 to 0 °C, 1 h) and subjected to electrophilic bromination followed by amination hydrolysis (CF₂Br₂, -78 to 23 °C, 18 h; 2 N HCl (aq), 30 min, 72%) to provide 6-bromo-2,4-dimethoxy-3-methylbenzaldehyde (**21**) in good yield. This two-step procedure offers more expeditious access to **21** than a previously reported four-step synthesis.³⁸ Removal of the aryl methyl ethers proceeded in high yield to give **22** (BBr₃, CH₂Cl₂, -78 to 23 °C, 88%). Sonogashira coupling of aryl bromide **22** and alkyne **23** provided the azaphilone precursor **24** (Pd(Cl)₂(PPh₃)₂, CuI, Et₃N, DMF, 70 °C, 16 h, 73%). Oxidative cyclization of **24** using Porco's methodology³⁸ generated the azaphilone **25** in good yield (Au(OAc)₃, TFA, dichloroethane, 4 min; IBX, Bu₄NI, 4 h, 72%). Acylation of the tertiary alcohol of **25** to give **26** (butyric anhydride, DMAP, CH₂Cl₂, 24 h, 91%), regioselective C6-chlorination yielding **27** (NCS, AcOH, 36 h, 89%) and removal of the TBDPS protecting group (HF-pyridine, THF, 2 h, 72%) proceeded smoothly to provide azaphilone **28**. The racemic azaphilones **25**, **26**, **27** and **28** could be chromatographically separated into their constituent enantiomers, and the most effective chiral phase resolution was achieved with **27** ($\alpha = 1.33$). The absolute stereochemistry of **25**–**28** was assigned based on the sign of their optical rotation (α_D) and the sign of the longest wavelength Cotton effect (350–370 nm) in their CD spectra (Figure 4).^{44–46} This assignment was further confirmed by application of Porco's asymmetric oxidative

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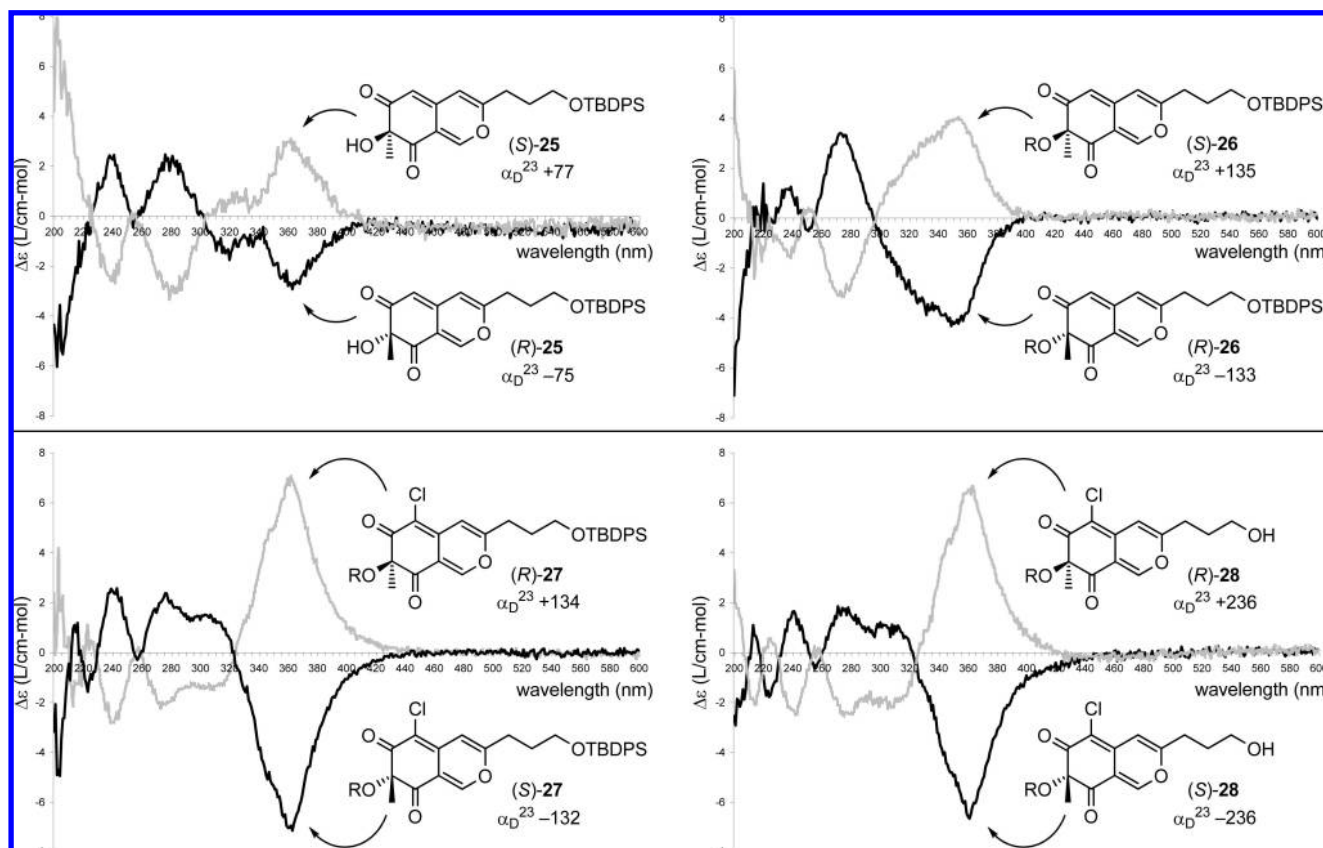
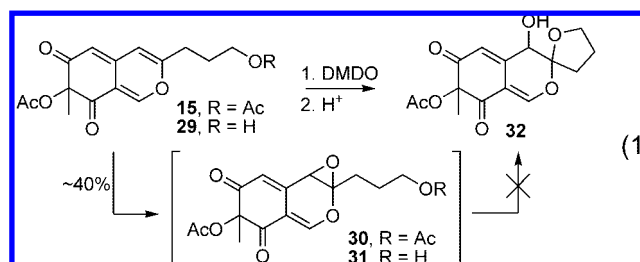


Figure 4. CD spectra (0.2 mM in MeOH) and α_D^{23} for azaphilones **25–28** (R = butyrate).

cyclization⁴⁷ of the OTIPS variant of **24** conducted using the catalyst generated from Cu(I) and (–)-sparteine to provide the corresponding azaphilone (94% ee), matching the assignment made based on CD. As fate would dictate and since (+)-sparteine is not yet readily accessible, this asymmetric route currently only provides access to the chromophore unnatural enantiomer series.

Oxidative Spiroketalization. With azaphilones **15** and **17** in hand, we briefly examined their direct oxidative spiroketalization. Thus, selective epoxidation or dihydroxylation of the C8–C9 double bond followed by oxonium ion formation upon epoxide opening or water elimination was anticipated to install the C8 alcohol and lead to subsequent C9-spiroketalization with the pendant side chain alcohol. Confident that the Williams proposed C8–C9 stereochemistry with the two oxygen substituents syn would be the thermodynamically most stable, conducting the reaction under conditions of reversible ketal formation would favor generation of the assigned C8–C9 syn versus anti diastereomers. Thus, a variety of oxidizing agents were explored to effect this transformation. Whereas subjecting azaphilone **15** to the Sharpless asymmetric dihydroxylation protocol⁴⁸ or that of oxidative spiroketalization with Re_2O_7 ⁴⁹ did not lead to identifiable products, treatment of **15** with DMDO⁵⁰ led to epoxidation of the desired C8–C9 double bond in ca. 40% yield (eq 1). However, the labile epoxide product⁵¹

quickly decomposed and proved challenging to work with or even purify. In hopes of trapping the labile epoxide, the analogous azaphilone **29** possessing the unprotected primary alcohol was treated with DMDO alone or followed with addition of a catalytic amount of acid (HCl, HOAc, PPTS, CSA).⁵² Although the formation of the intermediate epoxide was confirmed by ¹H NMR,⁵¹ the subsequent spiroketal formation was not observed and the intermediate epoxide was nonproductively consumed. Consequently, we focused our efforts on conducting the oxidative spirocyclization following amine introduction with substrates such as **34** avoiding the issue of C9-isomerization during amine conjugation.



Chromophore Model Studies. We explored a variety of conditions for condensation of the azaphilones **15** and **17** with

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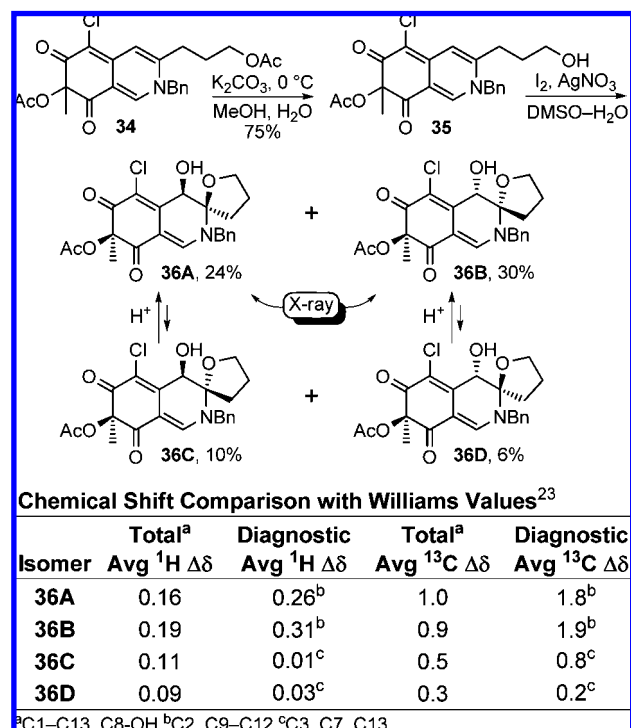
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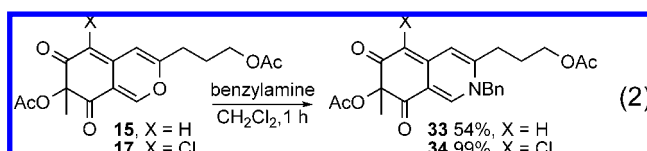
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(51) For **30**: ¹H NMR (500 MHz, CDCl_3) δ 7.64 (s, 1H), 6.31 (s, 1H), 5.72 (s, 1H), 4.14 (t, $J = 6.3$ Hz, 2H), 2.46 (t, $J = 7.5$ Hz, 2H), 2.16 (s, 3H), 2.07 (s, 3H), 1.96 (m, 2H), 1.54 (s, 3H); HR ESI-TOF m/z 351.1072 ($M + H^+$, $\text{C}_{17}\text{H}_{19}\text{O}_8$ requires 351.1074) (Representative Diastereomer). Treatment of **29** with DMDO followed by concentration provided a crude ¹H NMR indicating the consumption of starting material and appearance of signals corresponding to epoxide **31** by analogy to **30**.

Scheme 4

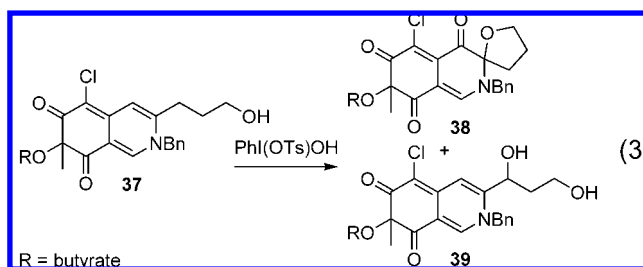


benzylamine to probe the compatibility of the condensation with solvents that may be required for effective dissolution of the cyclic peptide or a smaller peptide fragment. Although condensation of azaphilone **15** with benzylamine did provide **33** in modest yield (CH₂Cl₂, 1–16 h, 54%), condensation of chloroazaphilone **17** provided **34** in excellent yield (CH₂Cl₂, 1 h, 99%) establishing the productive role of the electron-withdrawing C6-chlorine substituent (eq 2). Selective hydrolysis of the primary acetate (K₂CO₃, H₂O, MeOH, 0 °C, 30 min, 75%) followed by oxidative spirocyclization of **35** with I₂ assisted by AgNO₃ in the presence of H₂O (I₂, AgNO₃, DMSO–H₂O, 48 h) allowed access to all four diastereomers (**36A–36D**) of the simplified chlorofusin chromophore (Scheme 4).



The development of these conditions followed unsuccessful efforts to effect a direct oxidative spirocyclization enlisting DMDO, *m*-CPBA, peroxyimide, peroxydicarbonate, peroxyurea, singlet oxygen, TMSOOTMS, as well as Shi, Sharpless, and Davis asymmetric epoxidation methods or osmium (OsO₄), rhenium (Re₂O₇), and ruthenium (RuO₄) based dihydroxylation reactions. Hypervalent iodine oxidizing reagents including Koser's reagent provided mixtures of **38**,⁵³ the oxidized form of the desired product, and **39**,⁵³ representing the allylic oxidation product (eq 3) although the yields of **38** were consistently low (10–15%) and the use of alternative reagents has not yet significantly improved on this. Nonetheless, the modest success with such reagents initiated studies with related

reagents that first activate the C8–C9 double bond by reversible bromonium or iodonium ion formation.

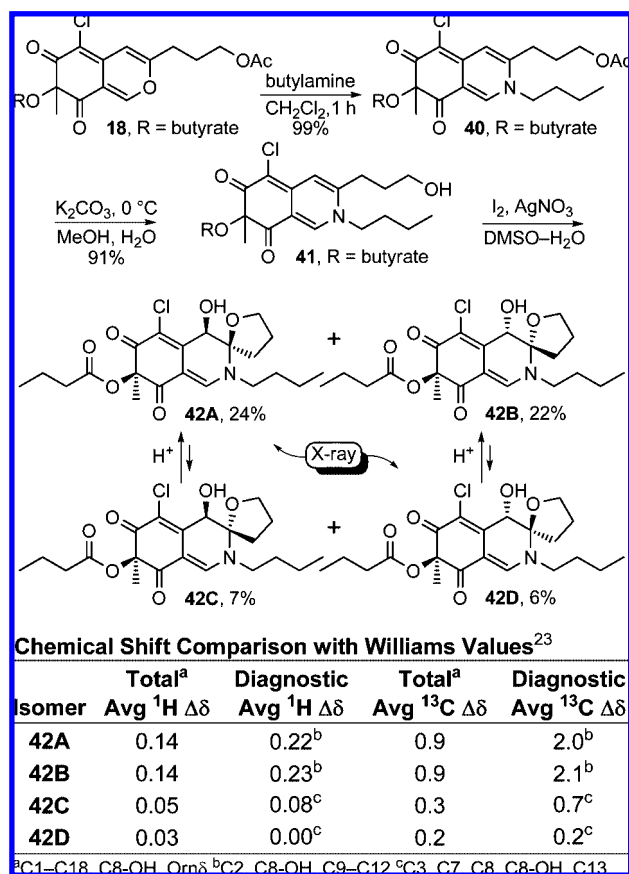


Thus, early efforts exploring a bromonium ion initiated spirocyclization, induced upon treatment of **35** with NBS under anhydrous conditions, provided the α-bromo *N,O*-spiroketal as a 1:1 mixture of C4-diastereomers resulting from clean trans addition across the C8–C9 double bond. Treatment of **35** with NIS under such conditions failed to provide analogous products upon workup despite the slow disappearance of starting material under the reaction conditions. However, inclusion of water in the reaction mixture led directly to the desired syn α-hydroxy *N,O*-spiroketal presumably resulting from water displacement of the iodide while premature workup of the reaction allowed isolation of two isomers of the unstable intermediate α-iodo *N,O*-spiroketal. Switching from NIS to I₂ served to increase the yield of the syn products and significantly, also led to the production of trans α-hydroxy *N,O*-spiroketal products. Although a range of solvents support the reaction (e.g., CH₃CN–H₂O, DMF–H₂O), DMSO–H₂O provided the syn cycloadducts in the best yield along with fewer undesired side products. Finally, the use of Ag(I) salts (AgOTf, AgNO₃, AgClO₄, AgOTFA) in the mixture accelerated the reaction, improved the conversions and minimized a competitive oxidation reaction. Thus, the reaction appears to be initiated by reversible iodonium ion formation and subsequent iodoetherification with *N,O*-spiroketal formation followed by slow Ag(I)-assisted iodide displacement by H₂O providing **36A–36D** directly. Notably, the lone pair of the endocyclic amine is tied up in the π-system of the chromophore as a vinylogous amide, and accounts for the relative stability of the iodonium intermediate, the predominance of C8/C9 syn products arising from S_N2-like trans intramolecular opening of the iodonium ion, and the high degree of planarity at the nitrogen center observed in the X-ray crystal structures (Figure 5). The major products (54%), in which the C8 and C9 oxygen substituents are syn possessing the C8/C9 stereochemistry found in the Williams assignment,

(53) For **38**: ¹H NMR (600 MHz, CD₃OD) δ 7.94 (s, 1H), 7.38 (m, 5H), 4.83 (d, *J* = 15.1 Hz, 1H), 4.75 (d, *J* = 15.3 Hz, 1H), 4.21 (dd, *J* = 14.7, 6.9 Hz, 1H), 4.07 (dd, *J* = 14.9, 7.0 Hz, 1H), 2.53 (td, *J* = 14.1, 7.1 Hz, 1H), 2.38 (t, *J* = 7.2 Hz, 2H), 2.21 (m, 1H), 2.01 (m, 2H), 1.63 (m, 2H), 1.53 (s, 3H), 0.97 (t, *J* = 7.4 Hz, 3H); ¹³C NMR (CD₃OD, 150 MHz) δ 190.6, 190.0, 189.4, 172.9, 148.6, 134.9, 134.3, 129.5 (2C), 129.0, 128.2 (2C), 120.4, 102.1, 97.2, 84.5, 71.2, 54.4, 35.2, 33.0, 25.1, 23.1, 18.4, 13.7; IR (film) ν_{max} 2984, 1737, 1374, 1242, 1046, 917, 734 cm⁻¹; HR ESI-TOF *m/z* 458.1383 (M + H⁺, C₂₄H₂₄ClNO₆ requires 458.1365) (Representative Diastereomer). For **39**: ¹H NMR (500 MHz, CDCl₃) δ 7.80 (s, 1H), 7.37–7.45 (br m, 3H), 7.19 (d, *J* = 7.0 Hz, 2H), 7.10 (s, 1H), 5.26 (d, *J* = 16.3 Hz, 1H), 5.10 (d, *J* = 16.4 Hz, 1H), 5.00 (dd, *J* = 9.1, 4.7 Hz, 1H), 3.82 (m, 1H), 3.75 (m, 1H), 2.44 (dt, *J* = 7.3, 3.3 Hz, 2H), 2.28 (m, 1H), 2.20 (m, 1H), 1.67 (m, 2H), 1.56 (s, 3H), 0.97 (t, *J* = 7.4 Hz, 3H), the two OH signals were not observed; ¹³C NMR (CDCl₃, 125 MHz) δ 193.5, 185.6, 173.3, 148.8, 143.4, 142.4, 134.1, 129.9 (2C), 129.5, 126.6 (2C), 115.1, 114.1, 105.0, 85.0, 77.2, 58.4, 57.1, 39.2, 35.4, 23.1, 18.4, 13.8; MALDI-TOF *m/z* 460.15 (M + H⁺, C₂₄H₂₆ClNO₆ requires 460.15) (Representative Diastereomer).

(52) Corey, E. J.; Kang, M.; Desai, M. C.; Ghosh, A. K.; Houpis, J. N. *J. Am. Chem. Soc.* **1988**, *110*, 649.

Scheme 5



represent those that formally arise from a trans iodoetherification reaction followed by water S_N2 displacement of the iodide. The structures of **36A** and **36B**, the major products of the oxidative spirocyclization reaction, were established by X-ray crystallography⁵⁴ and the C8 and C9 oxygen substituents of both compounds were found to be oriented syn with respect to one another with the C8–OH occupying an axial orientation. *N,O*-Ketal equilibration studies unambiguously established the structure of the remaining two minor anti diastereomers. The ¹H NMR signals of the tetrahydrofuran ring were found to be diagnostic of the relative orientation of the C8 and C9 substituents. Accordingly, chemical shifts of the C10–H signals for the two anti isomers **36C** (m, 2.25 ppm) and **36D** (m, 2.26 ppm) are similar to each other and to the C10–H signal reported for chlorofusin (br m, 2.38 ppm), but are distinct from the C10–H signals for **36A** (m, 1.89) and **36B** (m, 1.81). Similarly the C11–H signals for **36C** (m, 2.01 ppm) and **36D** (m, 2.00 ppm) possess chemical shifts that are closer to that of chlorofusin (m, 2.0–2.2 ppm) than the two syn isomers **36A** (m, 1.79 ppm; m, 1.96 ppm) and **36B** (m, 1.76 ppm; m, 1.82 ppm). The C12–H signals for **36C** (m, 3.80 ppm; m, 3.87 ppm) and **36D** (m, 3.77 ppm; m, 3.83 ppm) were also similar to one another and chlorofusin (q, 3.78 ppm; m, 4.02 ppm), and different from those of the syn isomers **36A** (m, 3.97 ppm; m, 4.18 ppm) and **36B** (m, 3.97 ppm; m, 4.18 ppm), but the correlation with the anti isomers was more tentative. The ¹³C NMR data exhibited similar trends, the most striking difference appearing with C10 signals for **36C** (30.6 ppm) and **36D** (30.5 ppm) being very similar to each other and to chlorofusin (30.3 ppm), but 4 ppm farther upfield than the C10 signals for the two syn isomers **36A** (34.5 ppm) and **36B** (35.0 ppm). A complete tabular comparison of the spectroscopic properties of **36A–36D** (Tables

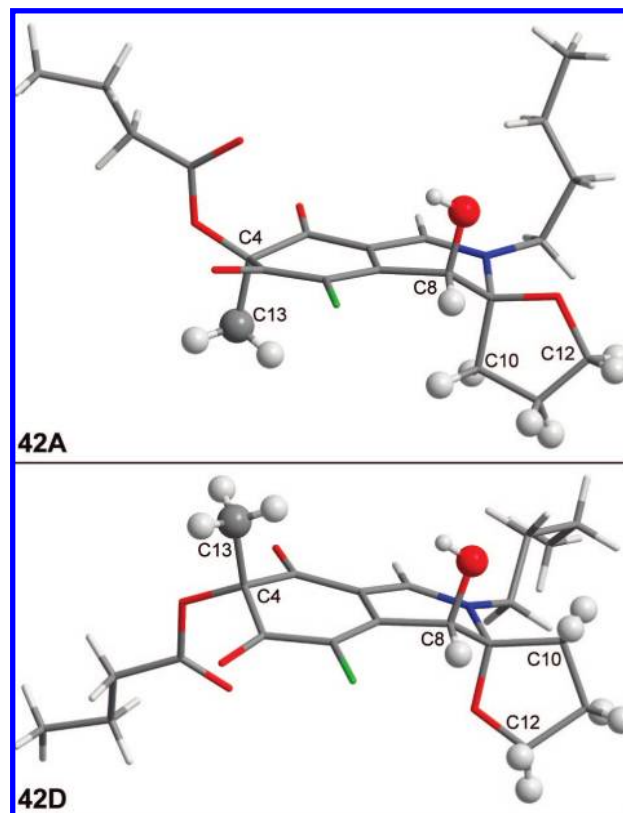
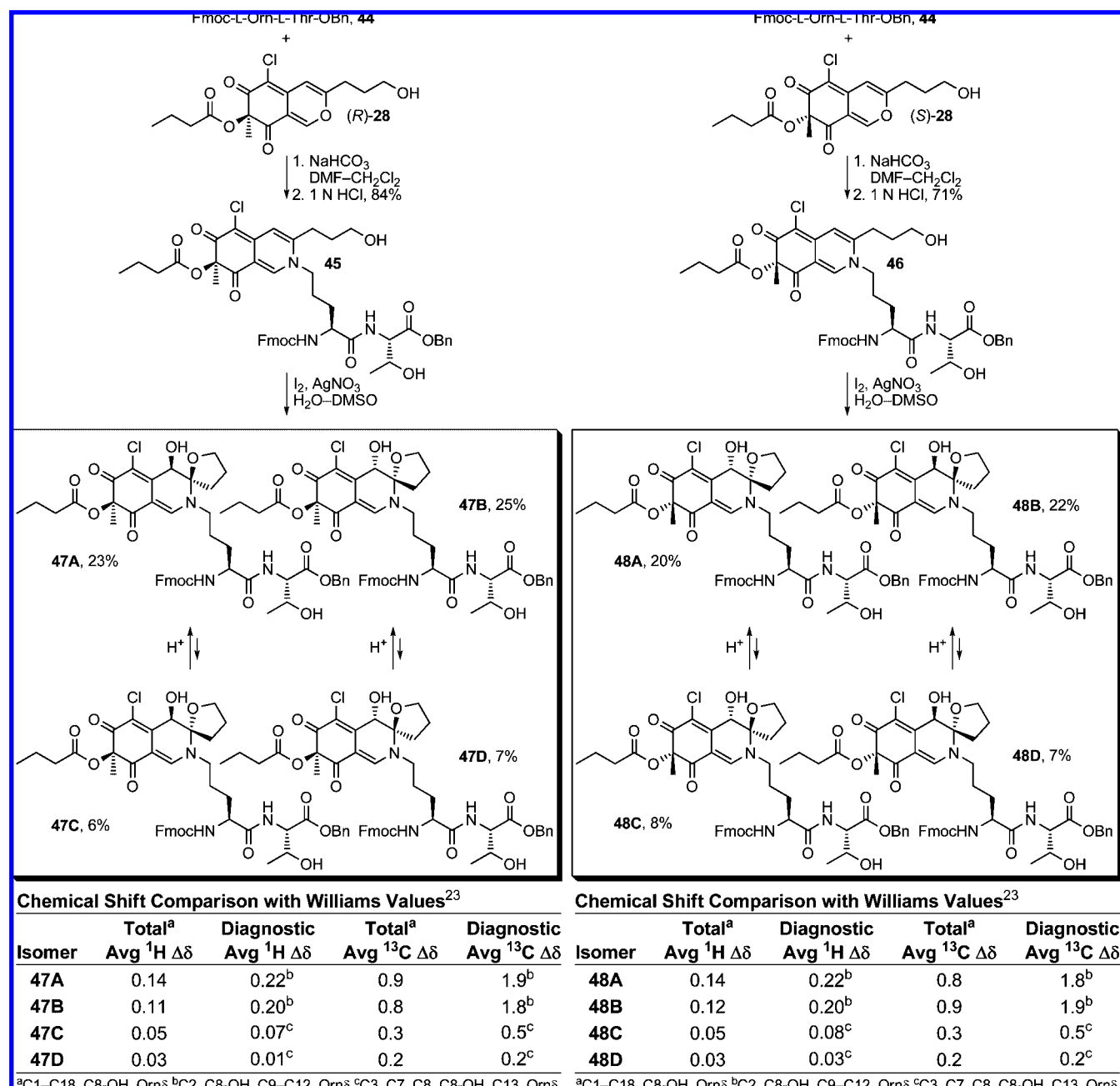


Figure 5. X-Ray structures of **42A** and **42D**.

S1 and S2, Supporting Information) and bar graph representations of the chemical shift differences ($\Delta\delta$, Figures S4 and S5, Supporting Information) relative to chlorofusin are provided in the Supporting Information and they, including the diagnostic distinctions highlighted above, were the first indication that the C8 and C9 oxygen substituents of chlorofusin may be oriented anti with respect to one another, and not syn as originally reported. These may also be summarized more simply as observed chemical shift differences averaged over the entire chromophore or restricted to the more diagnostic comparisons as presented in Scheme 4, both of which indicate that the anti diastereomer **36D** provides the best match with the chlorofusin chromophore. However, we were concerned about potential perturbation of the chromophore NMR signals by the appended benzyl group and elected to examine the additional, more comparable system **42** incorporating a *N*-butyl substituent as well as a C4 butyrate versus acetate. Significantly, the observations with **36A–36D** initiated a switch in our strategy from targeting the total synthesis of chlorofusin to one that could also unambiguously establish the chromophore stereochemistry. As such, the efforts were refocused on preparing and spectroscopically characterizing all possible chromophore diastereomers rather than exclusively targeting the Williams syn isomer.

Thus, butylamine was condensed with chloroazaphilone **18** to give **40** in excellent yield (CH₂Cl₂, 1 h, 99%). Hydrolysis of the primary acetate (K₂CO₃, H₂O, MeOH, 0 °C, 30 min, 91%) followed by the single step oxidative spirocyclization of **41** (I₂, AgNO₃, DMSO–H₂O, 48 h) provided **42A–42D** possessing the *n*-butyl substitution (Scheme 5). As with **36A–36D**, the NMR data collected from the anti isomers **42C** and **42D** provided a better match with chlorofusin than the syn isomers **42A** or **42B** (Supporting Information Tables S3 and S4 and Figures S8 and S9). An X-ray

Scheme 6



crystal structure of **42A**, one of the major products of the spirocyclization reaction representing the Williams assigned diastereomer and the closest match by NMR to **36A**, confirmed that the C4, C8 and C9 oxygen substituents were all syn while an X-ray of **42D**, the closest match with chlorofusin by NMR, confirmed that the relative orientation of the C8 and C9 oxygen substituents is anti and that the C4 methyl group is cis to the C8–OH and trans to the C9 oxygen of the spirotetrahydrofuran (Figure 5).⁵⁵ *N,O*-Ketal equilibration studies established the related syn/anti pairs and completed the unambiguous stereochemical assignments for

the entire series. As the similarity of the model to the chlorofusin chromophore increased, the trends in the NMR data distinguishing the C8/C9 syn diastereomers from the anti diastereomers became more predominant. For C10–H, **42A** (m, 1.85 ppm; m, 1.94 ppm) and **42B** (m, 1.82 ppm; m, 2.00 ppm) each display two signals, neither of which is within 0.3 ppm of chlorofusin (br m, 2.38 ppm), whereas the signals for the two anti isomers **42C** (m, 2.37 ppm) and **42D** (m, 2.39 ppm) differ from chlorofusin by 0.01 ppm. Similarly, the C12–H signals for the syn isomers **42A** (dd, 4.03 ppm; m, 4.22 ppm) and **42B** (dd, 4.04 ppm; dd, 4.22 ppm) are much farther downfield than the analogous signals for **42C** (dd, 3.82 ppm; m, 3.95 ppm), **42D** (dd, 3.80 ppm; dd, 3.95 ppm) and chlorofusin (q, 3.78 ppm; m, 4.02 ppm). An additional and important trend emerged linking **42B** with **42D**, which share the same relative stereochemistry at C8.

(54) Atomic coordinates for the two C8/C9 syn spirocycles **36A** (CCDC646443) and **36B** (CCDC646442) have been deposited with the Cambridge Crystallographic Data Center.

(55) Atomic coordinates for the C8/C9 syn spirocycle **42A** (CCDC678265) and the C8/C9 anti spirocycle **42D** (CCDC646441) have been deposited with the Cambridge Crystallographic Data Center.

The signals for C8–H observed in the spectra of **42A** (d, 4.48 ppm) and **42C** (d, 4.49 ppm) are similar to one another, whereas the signals for **42B** (d, 4.54 ppm) and **42D** (d, 4.53 ppm) are not only similar to one another, but also to that of chlorofusin (d, 4.53 ppm). This, along with other more subtle differences, distinguishes the two anti diastereomers with **42D**, not **42C**, being representative of the stereochemistry found in chlorofusin. Finally, the C8–OH signal in this series appears to distinguish **42A** (d, 6.18 ppm), **42B** (d, 6.11 ppm), **42C** (d, 6.41 ppm), and **42D** (d, 6.25 ppm) as well as link the relative stereochemistry of **42D** with that of chlorofusin (d, 6.26 ppm).

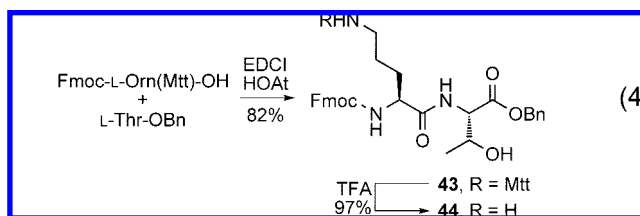
The ^{13}C NMR show similar trends to the observations made with **36A–36D**. However, the differences between the syn and anti compounds become even greater in this series for C10, C11, C12 and C9 in the absence of a proximal phenyl group. With X-ray structures of representative C8/C9 syn and anti diastereomers in hand, the diagnostic differences seen in their C10 ^{13}C NMR chemical shifts can be attributed to its axial (syn) or equatorial (anti) orientation. For C10, the chemical shifts of **42C** (30.2 ppm) and **42D** (30.2 ppm) became even closer to that of chlorofusin (30.3 ppm), but remained over 4 ppm farther upfield than the C10 signals for the syn isomers **42A** (34.6 ppm) and **42B** (35.0 ppm). The butylamine replacement for the Orn side chain also allowed a comparison of the chemical shift of the methylene adjacent to the chromophore ring system, with the anti isomers **42C** (50.5 ppm) and **42D** (50.6 ppm) matching that of chlorofusin (50.5 ppm) as opposed to **42A** (49.7 ppm) and **42B** (49.8 ppm).

Although intuitively surprising but which now may be expected from the X-ray crystal structure of **42D** (Figure 5) that shows an unobstructed path between C8–H and C4–Me (4.761 Å, C8–H to C13), the ROESY NMR for **42D** exhibited a weak cross-peak between C4–Me and C8–H as observed by Williams as a long-range NOE in chlorofusin. Also observed in its X-ray structure is the diaxial orientation of the C8 and C9 oxygen substituents. In this conformation, the equatorial C8–H can exhibit NOEs to both C10–H and C12–H as observed by Williams for chlorofusin. Although these NOEs were not quantitated, it is notable that C8–H is closer to C10–H (2.484 Å) than C12–H (2.592 Å) in this X-ray consistent with such implications in the Williams work. In fact, the C8–OH is axial and the C8–H is equatorial in the X-ray structures for **36A**, **36B**, **42A**, and **42D** indicating that the C8–H NOEs observed by Williams would be possible with all four diastereomers. This dominance of the C8–OH axial orientation arises presumably from the A(1,3)-strain an equatorial C8-alcohol would experience with the C6-chloride which is further destabilized by the electronegative nature of the two interacting substituents (Cl/OH). With this understanding of the conformational properties of the chromophore diastereomers and with the characterization data from both the **36** and **42** model systems in hand, the stereochemistry of the chlorofusin chromophore was confidently reassigned as either the (4*S*,8*R*,9*S*) or (4*R*,8*S*,9*R*) anti diastereomer (**42D** enantiomers).

L-Orn-L-Thr Azaphilone Conjugation and Elaboration to All 8 Chromophore Diastereomers. At this stage, we were convinced that a final stage chromophore coupling to the intact cyclic peptide followed by oxidative spirocyclization would be difficult to implement. Not only would this require the separation, characterization, and unambiguous stereochemical assignments of the four diastereomers generated in each enantiomeric series, but the reassigned anti isomer represented a minor,

contrathermodynamic product of the cyclization reaction in our model systems. As such, we elected to take the less ambitious, more manageable step of linking and elaborating the chromophore on a smaller dipeptide constituting L-Orn9–L-Thr1 of chlorofusin examining the two enantiomeric series and their four possible diastereomers. Not only would this provide intermediates with more manageable physical and spectroscopic properties and a third model on which to carefully reassess the chlorofusin stereochemical assignments, but it provided an opportunity to probe the chromophore absolute stereochemical assignment.

To this end, dipeptide **44** was prepared from the commercially available amino acid derivatives Fmoc-L-Orn(Mtt)-OH and L-Thr-OBn using a standard coupling protocol (EDCI, HOAt, DMF, 24 h, 82%) followed by deprotection of the *N*^δ-amine of ornithine (TFA, CH₂Cl₂, 1 h, 97%). Condensation of **44** with either (*R*)-**28** or (*S*)-**28** (NaHCO₃, DMF, CH₂Cl₂, 24 h; 1 N HCl, 4 h, 84% for **45**, 71% for **46**) provided **45** and **46**, respectively (Scheme 6). While incorporation of simple amines into the azaphilone core proceeded in high yield with simply CH₂Cl₂ as the reaction solvent, condensation with **44** required more specialized conditions. Solubility considerations led to the use of DMF as a cosolvent and additive studies revealed that the early stages of the incorporation were accelerated by addition of NaHCO₃ and subsequent treatment with 1 N HCl (aq) drove the reaction to completion. Oxidative spirocyclization of **45** (I₂, AgNO₃, DMSO–H₂O, 72 h) provided **47A–47D** and similar treatment of **46** provided **48A–48D**. The ^1H and ^{13}C NMR spectra of all eight diastereomers were fully assigned (Tables S5–S8, Figures S11–S14, Supporting Information) using COSY, HMQC, HMBC, and ROESY NMR data and *N,O*-ketal equilibration studies defined the related syn/anti pairs in each enantiomeric series. Notably, the equilibration studies requiring 5% TFA–HOAc (25 °C) revealed that the interconversions, like those in the **36** and **42** series, are slow even under these strongly acidic conditions and that the *N,O*-spiroketals are remarkably robust. Additionally, resubjecting **47B** or **47C** to the oxidative spirocyclization reaction conditions did not provide isomerized products suggesting that the syn products are both thermodynamically and kinetically favored and that the anti products generated are stable to the reaction conditions.



Stereochemical Assignment of Chlorofusin. By using the diagnostic spectroscopic distinctions observed in the **47** and **48** series, having assigned the C4 absolute stereochemistry by CD (see Figure 4),^{44–46} and enlisting *N,O*-ketal equilibrations to define the syn/anti pairs in each enantiomeric series, the complete relative and absolute stereochemical assignments for all eight diastereomers were unambiguously established. Moreover, only diastereomer **47D** (4*R*,8*S*,9*R*) matched all the spectroscopic properties reported for the chlorofusin chromophore. In short, the two syn and two anti diastereomers in each enantiomeric series are readily distinguishable by the diagnostic ^1H NMR (C10–H, C11–H, C12–H, and C8–OH) and ^{13}C NMR (C2 and C9–C12) chemical shifts and supported by kinetic and thermodynamic predominance of the syn isomers.

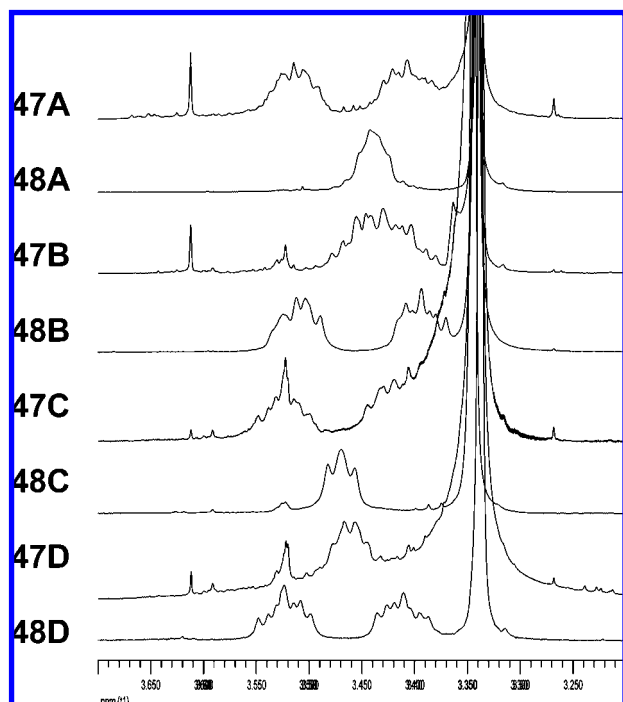


Figure 6. Ornithine CH₂^δ ¹H NMR signals for 47A–47D and 48A–48D.

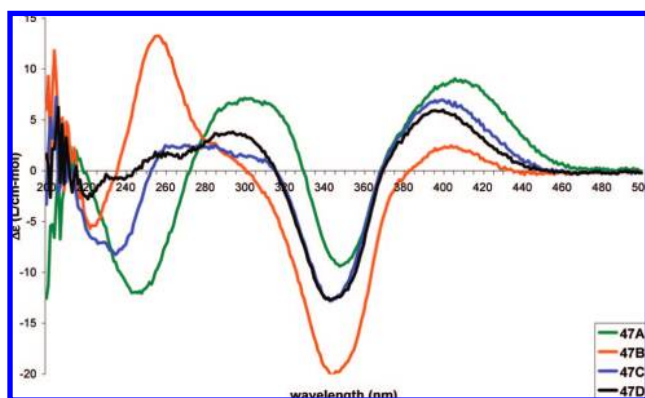


Figure 7. CD spectra (0.2 mM in MeOH) of 47A–47D.

The two anti diastereomers in each enantiomeric series are most readily distinguished by the diagnostic ¹H NMR (C8–H, C13–H, and C8–OH) and ¹³C NMR (C3, C7, C8, and C13) chemical shifts although there are more subtle spectroscopic distinctions as well. These are discussed in detail in the Supporting Information where Figures S11–S14 graphically quantitate these distinctions. In both series, it was the anti diastereomers **47D** and **48D** (chromophore enantiomers) that most closely matched chlorofusin. Additionally, the (4*R*,8*S*,9*R*)-diastereomer **47D** provided a near perfect match with the spectroscopic properties reported for the chlorofusin chromophore, whereas the (4*S*,8*R*,9*S*)-diastereomer **48D** (chromophore enantiomer) proved readily distinguishable by both the ¹H NMR chemical shift and multiplicity of the ornithine CH₂^δ adjacent to the chromophore (δ 3.45, m, 2H for **47D** vs δ 3.41 and 3.52, two m, 1H each for **48D**; chlorofusin = δ 3.42, t, 2H). This final multiplicity distinction between **47D** and **48D** allowed the absolute stereochemical assignment for the chlorofusin chromophore (Figure 6). This assignment of the absolute configuration for chlorofusin is especially significant

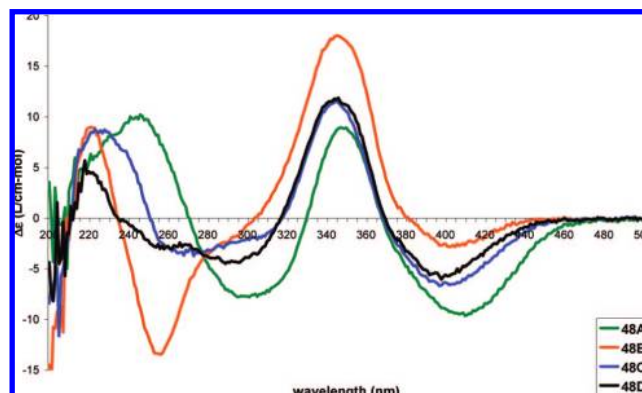
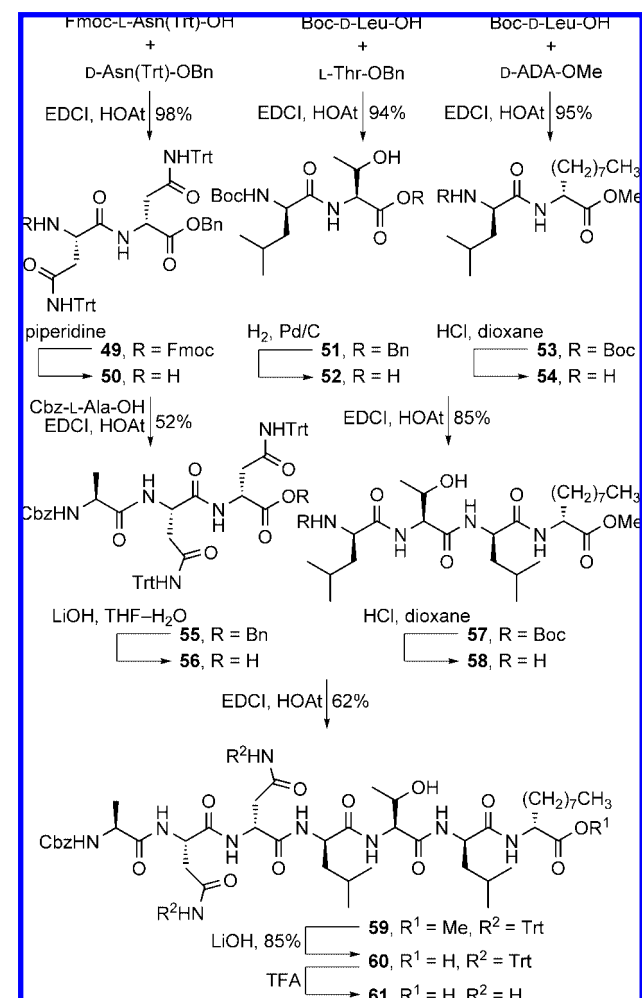


Figure 8. CD spectra (0.2 mM in MeOH) of 48A–48D.

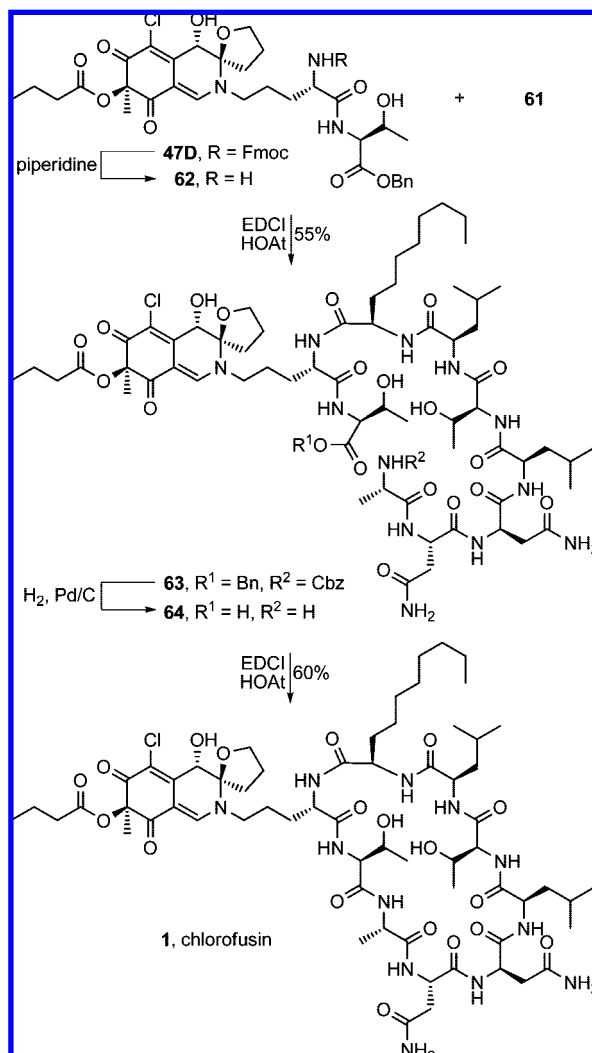
Scheme 7



given that our examination of the extensive series of known azaphilones revealed that both C4 configurations are found naturally with roughly equivalent representations.

As previously discussed, the anti diastereomers adopt a conformation with both the C8 and C9 oxygen substituents axial (see Figure 5) and their equatorial C8–H can exhibit NOEs to both C10–H and C12–H consistent with data for chlorofusin. The syn diastereomers adopt a conformation in which the C8–OH is axial, the C9 oxygen substituent is equatorial, and the C8–H is also equatorial. These isomers similarly exhibit C8–H to C10–H and C12–H NOEs. However, an important

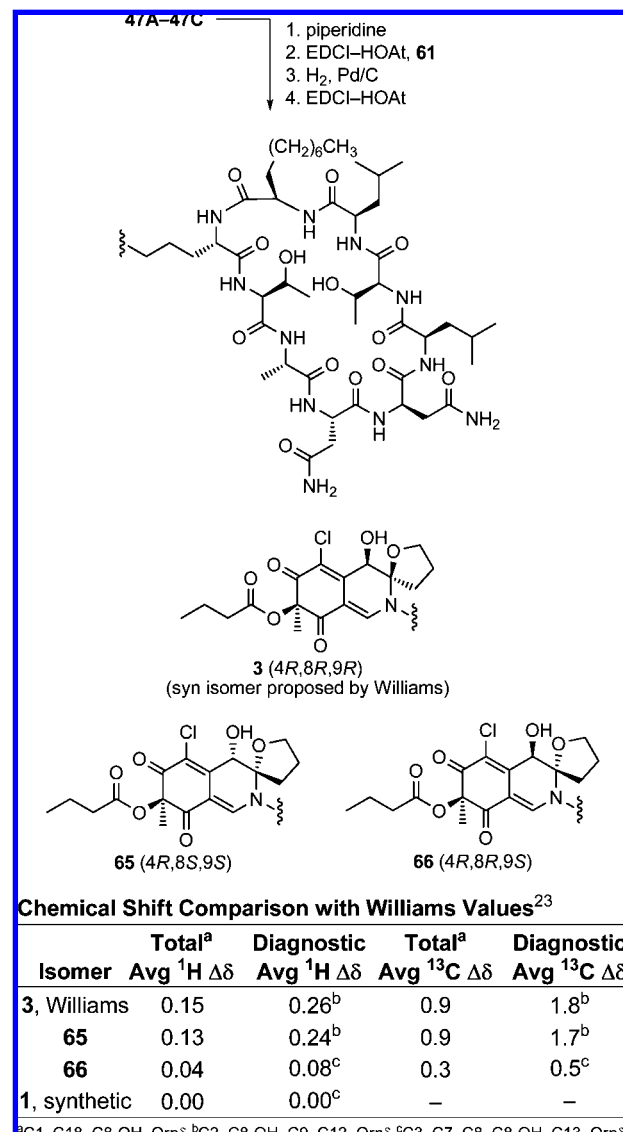
Scheme 8



ramification of the conformational features of the syn and anti diastereomers where each possess an equatorial C8–H is the counterintuitive observation that all four diastereomers possess nearly equivalent C4–Me to C8–H distances (5.42–5.55 Å) and that it is actually closest (5.42 Å) for the anti isomer we assigned to chlorofusin. The ROESY NMR data (300 ms mixing time) for these eight diastereomers reveals an NOE only between C8–H and C4–Me (and not C4–Me/C8–OH) when those groups are cis with respect to one another (47A, 47C, 48A, and 48C) and a correlation between C4–Me and both C8–H and C8–OH when C4–Me is trans with respect to C8–H (47B, 47D, 48B, and 48D). Thus, regardless of the relative stereochemistry of the chromophore, a NOE is seen between C4–Me and C8–H making the reassignment of the chromophore relative stereochemistry consistent with Williams' spectroscopic characterization of the natural product. Importantly, the NOE data collected and reported by Williams is impeccable including that of the long-range C4–Me/C8–H NOE, but it did not prove diagnostic of a C4–Me/C8–H cis stereochemistry.

Comparison of the CD spectra of all eight diastereomers shows that the region between 250–500 nm is dependent solely on the stereochemistry of the chromophore with nearly equal and opposite spectra observed for 47A and 48A, for 47B and 48B, for 47C and 48C, and for 47D and 48D (Figures 7 and 8). Of particular note is the sign of the longest wavelength Cotton effect (395–410 nm).⁵⁶ As with azaphilones 25–28, a

Scheme 9



positive longest wavelength Cotton effect is diagnostic of the C4–R stereochemistry and a negative Cotton effect is diagnostic of the C4–S stereochemistry.^{44–46}

Total Synthesis of Chlorofusin and its Three 4R-Diastereomers. With the chromophore stereochemistry confidently assigned, the (4R,8S,9R)-diastereomer 47D was incorporated into

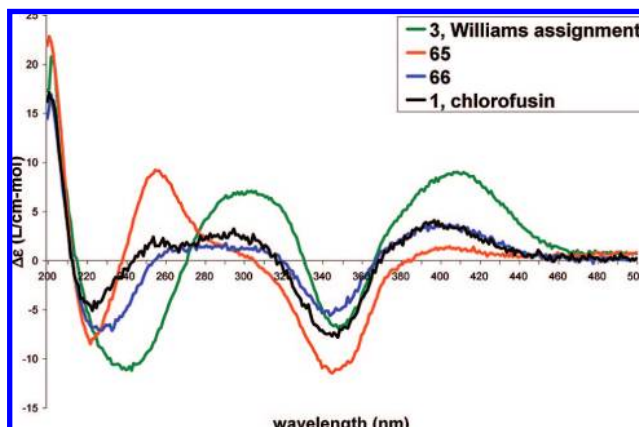


Figure 9. CD spectra (0.2 mM in MeOH) of 3, 65, 66, and 1.

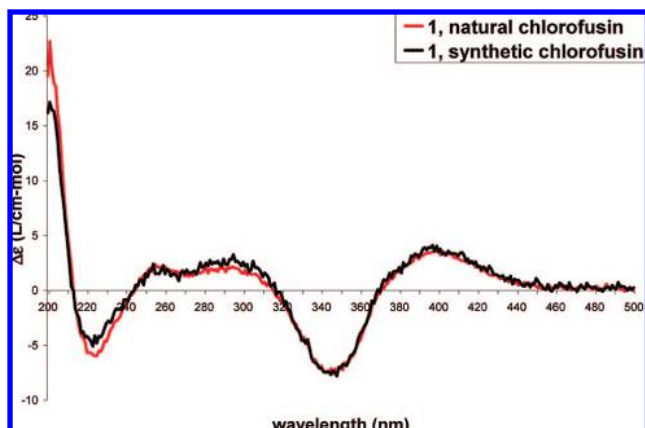


Figure 10. CD spectra of synthetic versus natural chlorofusin.

a total synthesis of chlorofusin. As a prelude to our initial assemblage of chlorofusin, one of the two anti diastereomers (**47C**) in the natural *4R*-enantiomeric series was exposed to many of the reaction conditions and recovered unchanged. Its treatment with EDCI–HOAt and NaHCO₃ in DMF for 16 h followed by exposure to workup conditions (dilute with EtOAc, wash with 1 N HCl (2×), aq. NaHCO₃ (2×), saturated aq. NaCl, and drying over Na₂SO₄) provided a quantitative recovery of **47C** with no detectable isomerization to its more stable syn isomer (**47A**). This would be representative of the peptide

coupling and macrocyclization reaction conditions and their workup. Extended treatment (16 h) with H₂/Pd–C does slowly hydrogenate the material, but it does not isomerize it. We minimized this potentially competitive reaction during the benzyl ester and Cbz deprotections using short reaction times (4 h). The Fmoc deprotection with piperidine (40 min) proceeded uneventfully providing the free amines of **47A**–**47D**, each of which was isolated by chromatography, without detection of isomerization or other conceivable problems. As such, the stability of the *N,O*-ketal is greater than one might intuitively suspect and this may be attributed to the fact that the nitrogen is tied up as a vinylogous amide. Two empirical observations additionally highlight this. First, the isolation of chlorofusin was conducted with TFA (0.01%) in the reverse phase HPLC solvent system.²³ Second, the conditions required for *N,O*-ketal isomerization are strongly acidic (e.g., 5% TFA–HOAc or 1:1 TFA–CH₂Cl₂, 5 h) and they do not isomerize under the conditions of a mild aqueous acid workup. Thus, a projected concern over the chromophore stability did not materialize, simplifying the path forward.

Peptide **61**, synthesized in a convergent sequence (Scheme 7), was protected in a manner that avoids treatment of late stage chromophore-containing intermediates with strong acid that might promote C9 *N,O*-spiroketal isomerization. Fmoc deprotection of **47D** (piperidine, DMF–CH₂Cl₂, 40 min) and coupling of the free amine **62** with heptapeptide **61** (EDCI, HOAt,

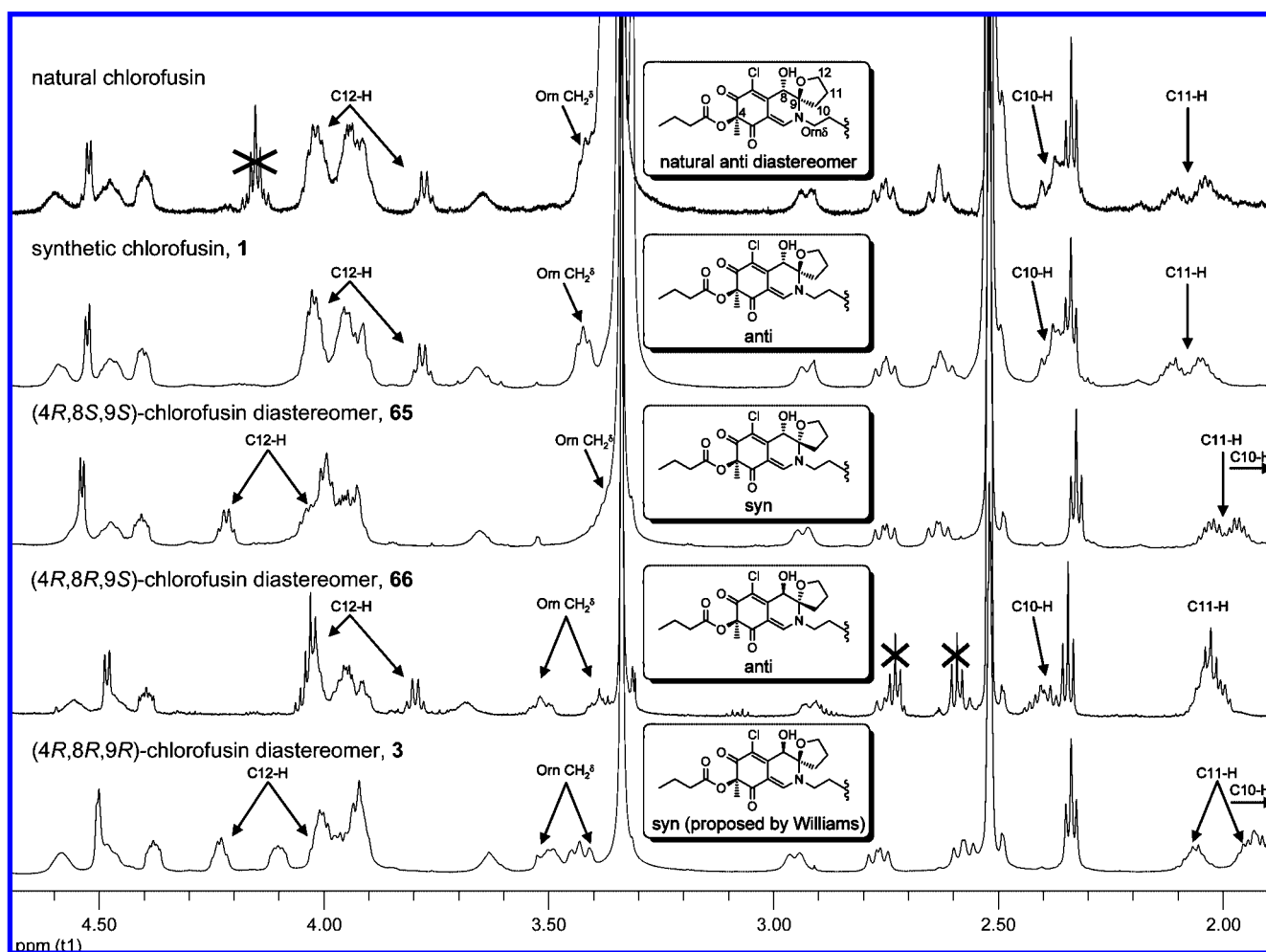
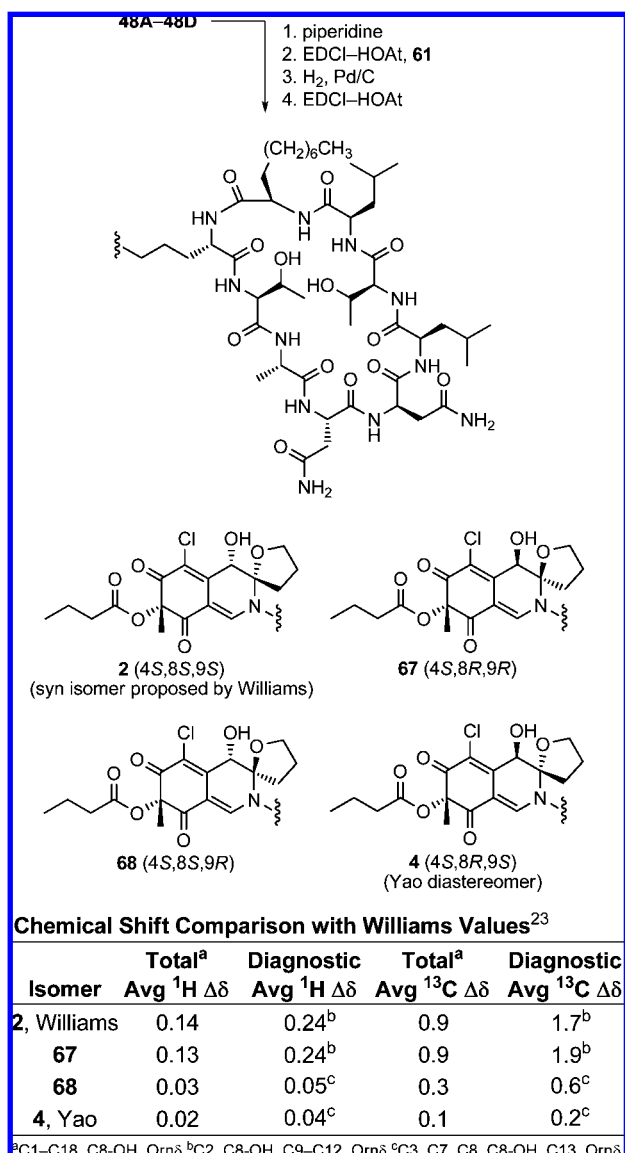


Figure 11. ¹H NMR overlay of natural chlorofusin with all four C4-*R* chlorofusin chromophore diastereomers.

Scheme 10



NaHCO₃, DMF, 0 to 23 °C, 24 h, 55%) provided **63** (Scheme 8). Concurrent benzyl ester deprotection and Cbz removal (H₂, Pd-C, THF-DMF, 4 h) followed by macrocyclization of **64** (EDCI, HOAt, NaHCO₃, DMF, 0 to 23 °C, 40 h, 60%) afforded synthetic chlorofusin (**1**), which displayed spectroscopic properties indistinguishable from those reported for natural chlorofusin. Among the most notable of these spectroscopic correlations was the single signal for the ornithine CH₂^δ (δ 3.42, t, 2H for natural and synthetic chlorofusin) that readily distinguishes the natural diastereomer from the (4*S*,8*R*,9*S*)-diastereomer incorporating the chromophore enantiomer (δ 3.42 and 3.50, two m, 1H each) that was prepared and miscorrelated by Yao.³⁰

In addition to the natural (4*R*,8*S*,9*R*)-diastereomer constituting chlorofusin, the (4*R*,8*R*,9*R*)-diastereomer **3** proposed by Williams as well as the (4*R*,8*S*,9*S*)- and (4*R*,8*R*,9*S*)-diastereomers of chlorofusin (**65** and **66**) were also prepared by this route from **47A–47C**. Notably, this entailed simply the four steps of Fmoc

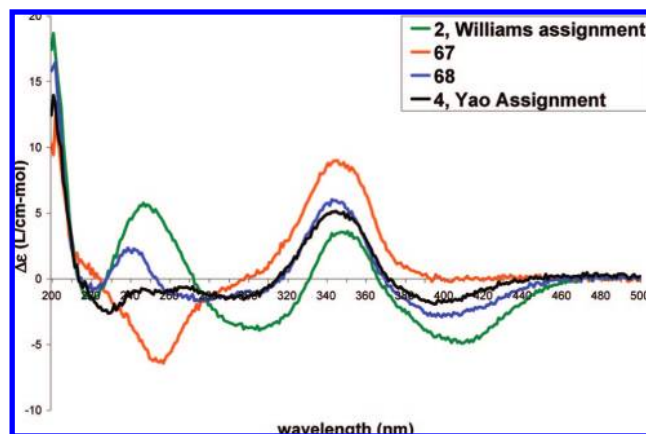


Figure 12. CD spectra (0.2 mM in MeOH) of **2**, **67**, **68**, and **4**.

deprotection of the chromophore dipeptide precursor and EDCI-promoted coupling with the same heptapeptide **61**, concurrent Cbz and benzyl ester deprotection, and a final macrolactamization to provide diastereomers **3**, **65**, and **66** with purification of a single intermediate in addition to the final products (Scheme 9). This late stage divergence in a convergent total synthesis facilitated the assemblage of the collective set of diastereomers and the anticipated noncorrelation of their spectroscopic properties with that reported for the natural product provided further firm support for the new structural assignment. The spectroscopic distinctions in these isomers mirrored those observed with the dipeptide conjugates **47A–47D** with the two syn and two anti diastereomers in each enantiomeric series readily distinguishable by the diagnostic ¹H NMR (C10-H, C11-H, C12-H, and C8-OH) and ¹³C NMR (C2 and C9–C12) chemical shifts. The two anti diastereomers in each enantiomeric series are most readily distinguished by the diagnostic ¹H NMR (C8-H, C13-H, and C8-OH) and ¹³C NMR (C3, C7, C8, and C13) chemical shifts along with more subtle spectroscopic distinctions.

Following the completion of this work and our chlorofusin structural reassignment and upon the Yao report³⁰ that claimed to have prepared chlorofusin (4*S*,8*R*,9*S*-diastereomer), we re-examined a sample of authentic chlorofusin provided by Dr. Stephen Wrigley in 2003 (aged and of unknown quality) that failed to provide a discernible ¹H NMR spectrum in 2003. Now, with an intimate knowledge of the chromatographic and physical properties of such compounds in hand, the processing of the remaining material (<1 mg) provided a sample that exhibited a CD spectrum indistinguishable (sign and magnitude) from synthetic **1** confirming the absolute configuration assignment (Figure 10). Moreover, its CD spectrum was readily distinguishable from the two possible syn diastereomers (4*R*,8*R*,9*R* and 4*R*,8*S*,9*S*) and more subtly distinguishable from the alternative anti (4*R*,8*R*,9*S*)-diastereomer **66** (Figure 9). Like the simple azaphilone chromophore precursors **25–28**, the dipeptide azaphilone conjugates **45** and **46**, as well as the fully elaborated chromophore dipeptides **47** and **48**, the longest wavelength (395–415 nm) CD Cotton effect is positive for all four 4*R* chlorofusin diastereomers as well as the natural product firmly establishing the absolute configuration assignment.^{44–46}

Just as significantly, the sample provided an ¹H NMR spectrum of a sufficient quality to confirm that it represented the authentic natural product (Supporting Information). Illustrated in Figure 11 is the spectral overlay of a diagnostic region of the ¹H NMR spectra of this sample of natural

(56) There is an interesting shift in the longest wavelength CD Cotton effect from 360 nm for the azaphilones **25–28** (Figure 4), to 375–380 nm for the (dipeptide) amine conjugates **45** and **46**, to 395–410 nm for **47A–47D** and **48A–48D**.

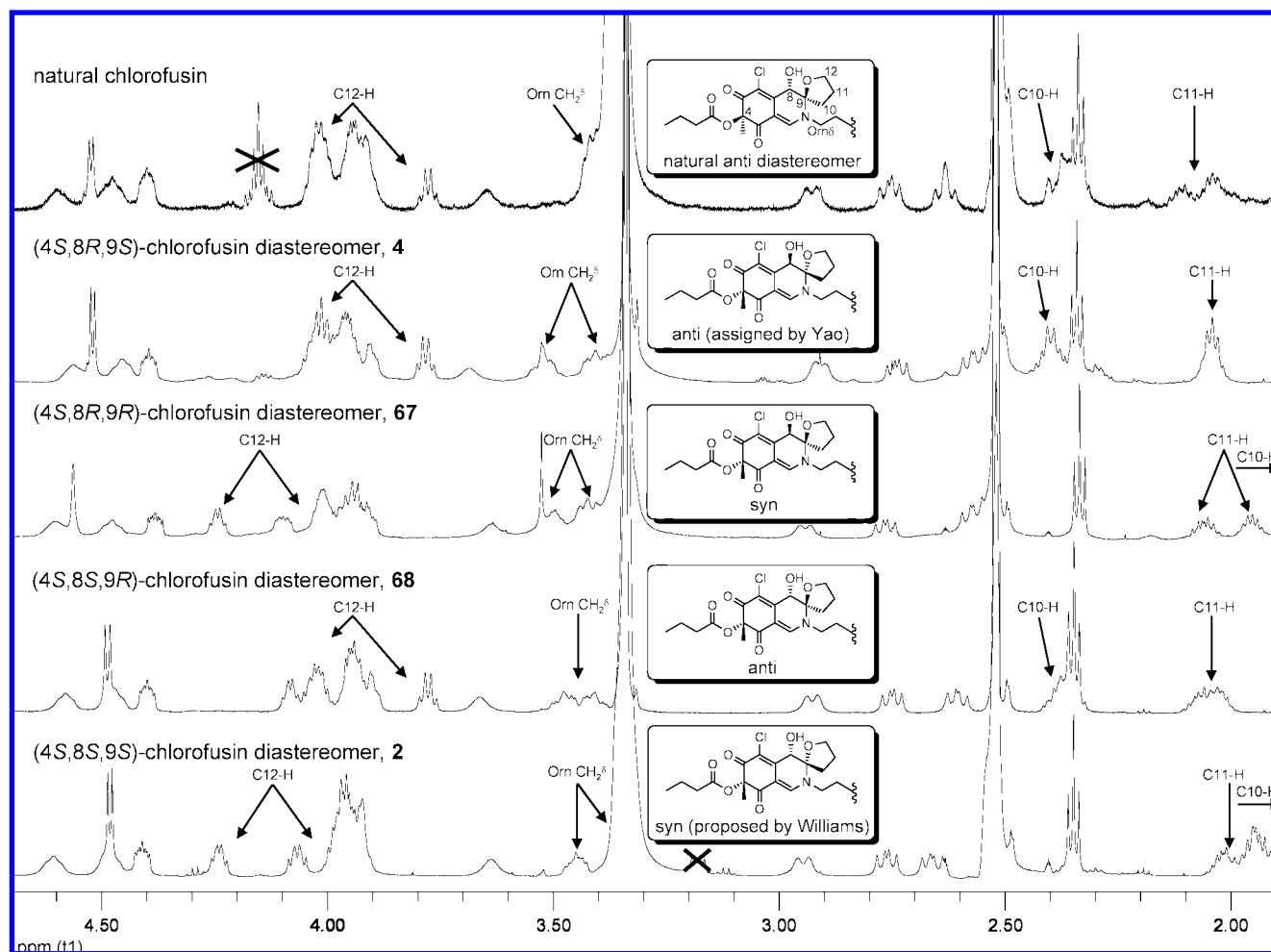


Figure 13. ^1H NMR overlay of natural chlorofusin with all four C4-*S* chlorofusin chromophore diastereomers.

chlorofusin with the four *4R* diastereomers illustrating the clear correlation with only the synthetic (*4R,8S,9R*)-diastereomer **1** and the clear distinctions with any of the alternative *4R* diastereomers in this enantiomeric series. Significantly, the syn diastereomer **3** proposed by Williams is readily discernible from natural chlorofusin not only by CD (Figure 9), but also by ^1H NMR (Figure 11). Additionally, the comparisons in this region of the ^1H NMR spectra clearly confirm the stereochemical purity and integrity of each isomer indicating that each, including the less stable anti diastereomers, is unaffected by the synthetic sequence required for their incorporation into the final products.

Synthesis of All Four *4S*-Diastereomers of Chlorofusin. With the Yao disclosure of the synthesis and misrelation of **4** with chlorofusin,³⁰ we elected to extend our studies to include the preparation and characterization of all four *4S*-chromophore diastereomers of chlorofusin including **4**. The preparation of this enantiomeric series simply required the late-stage individual incorporation of **48A–48D** into the full chlorofusin structure and again benefited from the convergent nature of the total synthesis providing diastereomers **2**, **67**, **68**, and **4** in four steps of which **4** corresponds to the Yao structure (Scheme 10). Though operationally short, the final materials are technically challenging to handle (SiO_2 and C18-reverse phase chromatography) making their synthesis more challenging than our expression of access in four steps might inadvertently convey.

With **2**, **67**, **68**, and **4** in hand, their spectroscopic properties proved readily distinguishable from the natural product. Each

displayed an anticipated longest wavelength negative Cotton effect opposite that observed with the natural product (Figure 12). Moreover, each *4S*-diastereomer provided an essentially identical, but of opposite sign, CD spectrum (250–500 nm) to the corresponding *4R*-diastereomer indicating that it is the chromophore, and not the cyclic peptide, that establishes the sign and magnitude of the CD spectrum in this region. Notably, that of **4** which bears the chlorofusin chromophore enantiomer is nearly identical to that of chlorofusin, but is clearly of the opposite sign.

Similarly, the ^1H NMR and ^{13}C NMR spectroscopic properties of each diastereomer **2**, **67**, **68**, and **4** were distinguishable from those of chlorofusin (see Supporting Information Table S11 and S12). This is most apparent in the region represented with the spectral overlay with the natural product in Figure 13. The Yao diastereomer **4**, bearing the enantiomer of the chromophore found in natural chlorofusin, is most readily distinguishable by its ornithine CH_2^δ signal which appears as two multiplets of 1H each (δ 3.41 and 3.52), but also exhibits more subtle differences including C8–OH (δ 6.22 vs 6.26), Orn9 α -CH (δ 4.56 vs 4.59), Leu5 α -CH (δ 4.45 vs 4.48), Thr1 α -CH (δ 3.68 vs 3.66), Asn3 β - CH^2 (δ 2.57, dd vs 2.62, app t), C10–H (δ 2.40, m vs 2.38, br m), and C11–H (δ 2.04, m vs 2.0–2.2, br m). Similarly, the alternative syn diastereomer **2** proposed by Williams and in this enantiomeric series also exhibited spectroscopic properties readily distinguishable from the natural product. Interestingly, while the ornithine CH_2^δ

signals for **1**, **65**, **66**, and **3** follow the trend observed with **47A–47D** appearing as a single multiplet of 2H in the ^1H NMR when the stereochemistry at C8 is *S* and as two multiplets of 1H each when the stereochemistry at C8 is *R*, it was observed that the ornithine CH_2^δ signal for **2**, **67**, **68**, and **4** appeared as distinguishable multiplets of 1H each independent of the C8 stereochemistry.

Conclusion

Full details of the first total synthesis of chlorofusin are reported resulting in the reassignment of its chromophore relative stereochemistry and the establishment of its chromophore absolute stereochemistry. Careful spectroscopic characterization of each of the four chlorofusin model chromophore diastereomers in two distinct series suggested the initial Williams assignment might not prove accurate and revealed that the key spectroscopic feature leading to the original assignment does not distinguish the four possible diastereomers. Subsequent preparation and characterization of all four diastereomers of a dipeptide conjugate with both enantiomers of the chlorofusin chromophore confirmed the required relative stereochemistry reassignment and permitted an assignment of its absolute configuration. This was unambiguously established with the total synthesis of not only chlorofusin, which displayed spectroscopic and physical properties indistinguishable from the natural product, but also all seven alternative chlorofusin chromophore diastereomers, each of which displayed spectroscopic properties

readily distinguishable from the natural product. This included the diastereomer advanced by Yao³⁰ as being identical with spectroscopic properties reported for natural chlorofusin as well as both enantiomeric chromophores of the original Williams assignment.²³ This study with the comprehensive characterization of all possible isomers was accomplished enlisting a convergent synthetic strategy with a key late-stage divergent introduction of the eight chromophore–dipeptide conjugates requiring only four steps for completion of the synthesis and facilitating the total synthesis of the full series of eight chromophore diastereomers. Currently, efforts are underway to synthesize the chromophore stereoselectively as well as to probe the mechanism by which chlorofusin disrupts the interaction of p53 and MDM2.

Acknowledgment. Dedicated to Professor E. J. Corey on the occasion of his 80th birthday. We gratefully acknowledge the financial support of the National Institutes of Health (CA41101) and the Skaggs Institute for Chemical Biology. We are especially grateful to Dr. S. S. Pfeiffer for initial studies on the chromophore synthesis (Scheme 2) and both Dr. S. S. Pfeiffer and Dr. P. Desai for initial studies on the chlorofusin cyclic peptide.²⁶

Supporting Information Available: Full experimental details are provided. This material is available free of charge via the Internet at <http://pubs.acs.org>.

JA8012819