

Accepted Article

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This manuscript has been accepted after peer review and appears as an Accepted Article online prior to editing, proofing, and formal publication of the final Version of Record (VoR). This work is currently citable by using the Digital Object Identifier (DOI) given below. The VoR will be published online in Early View as soon as possible and may be different to this Accepted Article as a result of editing. Readers should obtain the VoR from the journal website shown below when it is published to ensure accuracy of information. The authors are responsible for the content of this Accepted Article.

To be cited as: Angew. Chem. Int. Ed. 10.1002/anie.201712369 Angew. Chem. 10.1002/ange.201712369

Link to VoR: http://dx.doi.org/10.1002/anie.201712369 http://dx.doi.org/10.1002/ange.201712369

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Total Synthesis of (±)-Phomoidride D

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Dedicated to Professor Amos B. Smith, III in appreciation of his mentorship.

Abstract: Described herein is a synthetic strategy for the total synthesis of phomoidride D. This highly efficient and stereoselective approach provides rapid assembly of the carbocyclic core by way of a tandem phenolic oxidation/intramolecular Diels-Alder cycloaddition. A subsequent Sml₂ mediated cyclization cascade delivers an isotwistane intermediate poised for a Wharton fragmentation that unveils the requisite bicyclo[4.3.1]decene skeleton and sets the stage for synthesis completion.

Since the isolation and structural elucidation of the two fungal secondary metabolites, phomoidride A (1, CP-225,917) and phomoidride B (2, CP-263, 114) (Figure 1), by researchers at Pfizer, ¹ numerous groups have devoted efforts toward developing new synthetic strategies to construct these natural products. ² While the cholesterol-lowering and anticancer properties displayed by 1 and 2 certainly motivated synthetic efforts, there is little doubt that the unique and complex

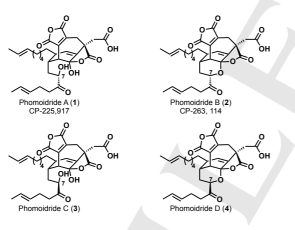


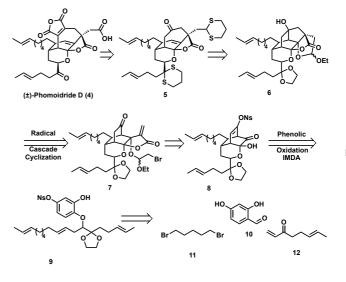
Figure 1. The Phomoidride Family.

 * Dr. J. C. Leung, Dr. A. B. Bedermann, Prof. J. T. Njardarson, Prof. D. A. Spiegel, Prof. G. K. Murphy, Dr. N. Hama, Dr. B. M. Twenter, Dr. P. Dong, Prof. T. Shirahata, Dr. I. M. McDonald, Dr. M. Inoue, Dr. N. Taniguchi, Dr. T. C. McMahon, Dr. C. M. Schneider, Dr. N. Tao, Prof. B. M. Stoltz, Prof. J. L. Wood Department of Chemistry and Biochemistry Baylor University One Bear Place 97348, Waco, Texas 76798, United States E-mail: john _L wood@baylor.edu

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architecture inherent to the phomoidrides has served as the primary inspiration to what has proven to be a variety of markedly creative approaches,^{2, 3} and four completed total syntheses.⁴ Studies toward these targets also led to discovery of two other congeners, phomoidrides C and D (**3** and **4**, respectively, Figure 1),^{4f, 5} which differ in the relative stereochemistry at C7. While previous synthetic efforts have been primarily directed toward the densely functionalized carbocyclic phomoidride core, biosynthetic work has focused on biogenesis and congener interconversion.⁶ Herein we describe a novel synthetic strategy that employs two cascade sequences en route to a successful synthesis of (±)-phomoidride D (**4**).⁷

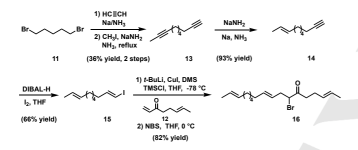
As illustrated retrosynthetically in Scheme 1, our plan for accessing **4** called for late stage introduction of the maleic anhydride moiety, an endgame akin to those reported by Fukuyama⁸ and Shair^{3a}. In contrast to the latter efforts, our strategy employs a regioisomeric β -ketoester that derives from ketone **5**, which was seen as arising from Wharton fragmentation of isotwistane **6**. Although increasing structural complexity in a retrosynthesis appears counterintuitive from a strategic planning perspective, we envisioned accessing **6** via a ketyl-initiated cascade cylclization wherein an exomethylene lactone serves as a lynchpin and bromide as the nucleofuge. The cyclization cascade precursor (**7**) would arise from [2.2.2]



Scheme 1. Retrosynthetic Analysis of Phomoidride D.

bicycle **8**, the product of a tandem phenolic oxidation/inverse electron-demand intramolecular Diels-Alder (IMDA) cycloaddition wherein phenol **9** serves as substrate. This highly efficient combination of two cascade reactions allows for global control of relative stereochemistry and introduces all but five carbons present in phomoidride D. Phenol **9** would arise from the readily available precursors **10**, **11** and **12**.^{4c,7d,8,9}

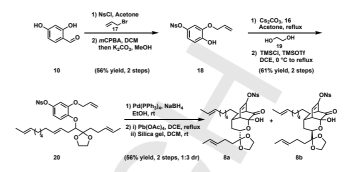
In accord with our synthetic plan, 1,5-dibromopentane (11) was homologated to 13 by exposure to sodium acetylide followed by monomethylation of the intermediate diyne (Scheme 2). Deprotonation of 13 at the terminal alkyne allowed for selective reduction to enyne 14. Subsequent hydroalumination/iodination of 14 delivered vinyl iodide 15,¹⁰ which following conversion to the corresponding cuprate was advanced via conjugate addition to known α , β -unsaturated ketone 12.⁸⁻⁹ Under the illustrated TMSCI-accelerated conditions,¹¹ this latter reaction furnishes an intermediate silyl enol ether which, upon in situ exposure to *N*-bromosuccinimide (NBS), delivers 16.



Scheme 2. Synthesis of α -bromoketone 16.

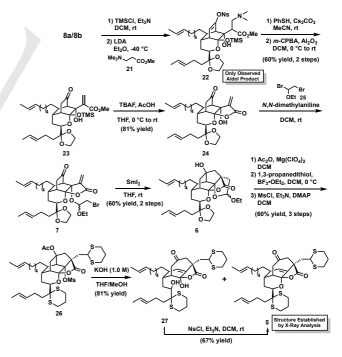
Having developed an efficient five-step sequence to abromoketone 16, we next focused on the aromatic coupling partner. It is worth noting that a considerable number of experiments over many years indicated that the planned IMDA would likely be successful if the diene component were sufficiently electron poor. In efforts to satisfy this electronic requirement the o-nitrobenzenesulfonyl (nosyl) moiety was discovered to be sufficiently electron withdrawing and stable to subsequent synthetic steps. Thus, as illustrated in Scheme 3, commercially available 1,2-dihydroxybenzaldehyde (10) was sequentially nosylated, allylated, and exposed to Dakin oxidation conditions to furnish 18. Alkylation of 18 with 16 furnished an intermediate ketone that was protected under modified Noyori conditions.¹² Exposure of the derived acetal (20) to palladiummediated allyl deprotection followed by a Pb(OAc)₄ induced tandem aryl oxidation cycloaddition sequence, provided the ahydroxy ketones 8a and 8b as an inseparable 1:3 mixture of diastereomers, respectively.13

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Scheme 3. Tandem Phenolic Oxidation/Diels-Alder Cascade.

Interestingly, treating the derived mixture with TMS-CI followed by the lithium enolate of methyl 3-(dimethylamino)propionate results in conversion of only the major diastereomer (**8b**) to the corresponding aldol product **22** (Scheme 4).¹⁴ Subsequent nosyl deprotection, Cope elimination, and desilylation provides an intermediate (**24**) containing an *exo*-methylene lactone poised to serve as a lynchpin in the second cascade reaction. Effecting this latter event begins by condensing tertiary alcohol **24** with dibromide **25** to afford Stork/Ueno bromoacetal **7**.¹⁵ Exposure of **7** to freshly prepared Sml₂ promotes a smooth 5-*exo*-trig/5-*exo*tet cyclization cascade¹⁶ that delivers the key isotwistane **6** in excellent yield. Importantly, this sequential C-C bond forming event sets stereochemistry at the imbedded quaternary center and positions the core structure for fragmentation to the bicyclo[4.3.1]decene. To this end, we first converted **6** to the

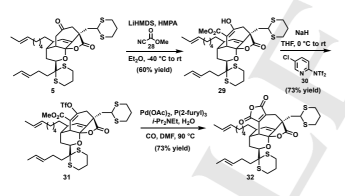


Scheme 4. Synthesis of the [4.3.1]-Bicyclic Core via Radical Cyclization Cascade and a Wharton Fragmentation.

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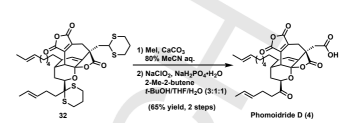
corresponding acetate and then unveiled the latent tertiary alcohol via transacetalization with 1,3-propanedithiol. Introduction of the requisite nucleofuge was accomplished via mesylation of the derived alcohol to provide fragmentation substrate 26. To our delight, exposure of 26 to KOH in a THF/MeOH solvent mixture, resulted in Wharton fragmentation and installation of the bridgehead olefin to furnish 5. Unsurprisingly, the conditions required for fragmentation also resulted in varying amounts of a diol (27) derived from ring opening of the spiroacetal, an unwanted side-reaction that could be reversed by treating the reaction mixture with nosyl chloride and triethylamine. The illustrated stereochemical outcome of the two cascade sequences was firmly established via single crystal X-ray analysis.

With ready access to the bicyclic carbon framework, we turned attention to constructing the maleic anhydride moiety and began exploring conditions for the regio- and chemoselective acylation of **5**. After some experimentation, we were gratified to discover that Mander's reagent (**28**),¹⁷ employed with Et₂O as solvent under thermodynamic deprotonation conditions, not only minimized *O*-acylation but led to predominately the desired regioisomer **29** as a mixture of keto-enol tautomers, favoring the latter.¹⁸ Subsequent transformation of **29** to the corresponding enol triflate **31** using Comins' reagent (**30**)¹⁹ enabled maleic anhydride installation via a palladium mediated carbonylation (Scheme 5). At this point, all that was required to complete phomoidride D was deprotection of **32** followed by oxidation of the derived aldehyde.



Scheme 5. Preparation of the Maleic Anhydride via Palladium-Mediated Carbonylation.

In the latter events, numerous methods to remove the dithiane moieties, including both alkylation and oxidation processes, were attempted but often led to the formation of complex mixtures instead of the desired keto-aldehyde. Eventually, we found that exposing **32** to excess iodomethane (80 equiv) in the presence of calcium carbonate (CaCO₃) resulted in clean conversion to an intermediate keto-aldehyde which, upon Pinnick oxidation using sodium chlorite (NaClO₂), sodium dihydrogenphosphate monohydrate (NaH₂PO₄•H₂O) as buffer, and 2-methyl-2-butene as hypochlorous acid scavenger, furnished phomoidride D (**4**) in excellent yield (Scheme 6).



Scheme 6. Completion of the Phomoidride D Total Synthesis.

In conclusion, a total synthesis of phomoidride D has been achieved by employing a novel strategy that requires 26-steps in its longest linear sequence. From the outset, the primary motivation for pursuing a synthesis of this intriguing molecule was the challenge of developing a non-obvious yet efficient approach, an endeavor that invariably advances the forefront of strategies and tactics in the science of synthesis.

Acknowledgements

The authors gratefully acknowledge support of this work from Bristol-Myers Squibb, Eli Lilly, Glaxo-Wellcome, Yamanouchi (now Astellas), AstraZeneca, and Amgen through their faculty award programs and the Camille and Henry Dreyfus Foundation for a Teacher Scholar Award to J.W. J.T.N. thanks the Helgu and Jónsdóttur and Sigurliða Kristjánsson Memorial Foundation, G.K.M thanks the NSERC of Canada, T.S. thanks Prof. Satoshi Ōmura and the Kitasato Institute, and N.T. thanks Yamanouchi (now Astellas). The authors thank Prof. Kevin Klausmeyer and Ms. Sam Yruegas for obtaining and analyzing X-ray crystallographic data. We also gratefully acknowledge financial support from Baylor University, the Welch Foundation (Chair, AA-006), and the Cancer Prevention and Research Institute of Texas (CPRIT, R1309).

Keywords: Phomoidride • Synthesis • Cascade • Diels-Alder • Aryl-Oxidation

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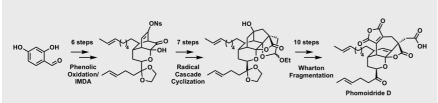
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