

Total Synthesis of Selaginpulvilin C and D Relying on in Situ Formation of Arynes and Their Hydrogenation

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Supporting Information

ABSTRACT: The total syntheses of selaginpulvilins C and D is described. The key strategy for the construction of the core fluorene moiety involves in situ formation of an aryne intermediate followed by its formal hydrogenation. The precursor tetraynes that undergo aromatization via hexadehydro Diels—Alder reaction were prepared from readily available building blocks through typical alkyne-coupling reactions.

S elaginpulvilins A–D that contain a novel 9,9-diphenyl-1-(phenylethynyl)-9H-fluorene framework were reported by Yin and co-workers in 2014. The structural assignment of these tetraphenolic compounds was secured by spectroscopic, chemical, and single-crystal X-ray diffraction analyses. These unprecedented natural products were isolated from *Selaginella pulvinata* (Hook. et Grev.) Maxim. (Selaginellaceae), which has been widely used in traditional Chinese medicine. The activity-guided bioassays of ethanolic extracts of *S. pulvinata* containing selaginpulvilins A–D and other constituents showed significant inhibitory activities (IC $_{50}$ values of 0.11–5.13 μ M) against phosphodiesterase-4 (PDE4).

The important biological profiles of selaginpulvilins in combination with their novel structural features make them highly attractive targets for total synthesis, which will also provide an opportunity to develop a range of related structures for further biological activity-relationship studies. In terms of designing an effective synthetic approach, we envision that 9,9diphenyl substituents on the central fluorene substructure³ of selaginpulvilins can be installed via double Friedel-Crafts arylations with precursor 1 (Scheme 1). In turn, the phenylethynyl-9H-fluorenone structure can be constructed from tetrayne 2 via a hexadehydro Diels-Alder reaction⁴⁻⁶ (HDDAR) to form an aryne⁷ intermediate followed by its formal hydrogenation via either intramolecular hydride transfer from a trialkylsilyl group followed by protonation^{5c} or hydrogen transfer from cyclooctane.^{4h} Finally, tetrayne **2** would be accessed from readily available arene and alkyne building blocks.

The preparation of tetrayne **2** commenced with a Sonogashira coupling of commercially available 2-bromo-5-methoxybenzaldehyde with (trimethylsilyl)acetylene to generate **3** (Scheme 2). ⁸ 4-(Methoxyphenyl)-1,3-diyne **4** was also prepared through a merger of 4-iodoanisole and (trimethylsil-

Scheme 1. Selaginpulvilin Retrosynthesis

yl)acetylene, desilylation, and a Cadiot—Chodkiewicz coupling with 4-bromo-2-methylbut-3-yn-2-ol followed by removal of the acetone moiety to liberate a terminal alkyne under basic conditions. Generation of an acetylide from 4 and its addition to aldehyde 3 afforded alcohol 5.

Removal of the trimethylsilyl group from 5 followed by a copper-catalyzed coupling with bromoalkynes delivered tetraynes 7a-e.

With these substrates for HDDAR in hand, we explored their conversion to the projected fluorenones. On the basis of the established protocol^{5c} for intramolecular hydride transfer to an

Received: October 28, 2016
Published: November 18, 2016

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Scheme 2. Synthesis of Tetraynes

aryne species catalyzed by a silver catalyst, tetrayne 7a was treated with MnO2 at 25 °C in CH2Cl2 in the presence of AgSbF₆ (10 mol %) (Scheme 3). Unfortunately, substrate 7a decomposed upon addition of AgSbF₆. Next, 7b was oxidized with MnO₂ in cyclooctane as a source of hydrogen. 41 Gratifyingly, desired compound 9b was generated in 34% overall yield as a mixture of two isomers in a 6:1 ratio, the minor isomer of which was formed via the alternative mode of HDDAR. On the other hand, the corresponding trimethylsilylcontaining 7c provided 9c in 54% yield along with another isomer in a 5:1 ratio. Addition of anisole-based Grignard reagent 10 to ketone 9b provided an adduct tert-alcohol 11b, which was directly subjected to the known Friedel-Crafts arylation conditions to afford 12b in 51% overall yield from 9b. 9,10 Subsequent removal of the triethylsilyl group from 12b was found to be recalcitrant. Treating 12b with an excess amount of Bu₄NF at 90 °C afforded the corresponding desilylated product 13b, O-trimethylselaginpulvilin D, in only 34% yield along with 22% recovered starting material 12b. In stark contrast, the Friedel-Crafts arylation of trimethylsilylcontaining compound 11c directly generated protodesilylated compound 13b in 38% yield devoid of 12c under otherwise identical conditions. The final cleavage of the methyl ethers with Me₃SiI¹¹ or BBr₃¹² afforded a mixture of several products of incomplete demethylation After much experimentation, it was found that treating 13b with MeMgI in neat at 160 °C under reduced pressure delivered selaginpulvilin D.¹³

With the successful route in hand, synthesis of another congener of the same family, selaginpulvilin C, was attempted. The (trimethylsilyl)methyl-substituted tetrayne 7d was proposed to be a good choice for this purpose. Treating 7d with MnO₂ in cyclooctane at room temperature afforded fluorenone 9d, which was reacted with Grignard reagent 10 to generate 11d (Scheme 4). Subsequently, with the established Friedel—Crafts arylation protocol, 11d was converted to 12d in 86% yield. Desilylation of 12d with TBAF quantitatively afforded 13d, a trimethyl-protected form of selaginpulvilin C. Finally,

Scheme 3. Total Synthesis of Selaginpulvilin D

Scheme 4. Total Synthesis of Selaginpulvilin C

global cleavage of the methyl ethers with methylmagnesium iodide resulted in the formation of selaginpulvilin C in 69% yield, 13 which constitutes a short total synthesis of the natural product.

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In summary, we have accomplished total syntheses of selaginpulvilins C and D. The construction of the core fluorene moiety of these natural products relies on a strategy that involves in situ formation of an aryne intermediate via the hexadehydro Diels—Alder reaction (HDDAR) followed by formal hydrogenation of the aryne moiety. The required tetrayne substrates for HDDAR were prepared efficiently via repetitive use of alkyne-coupling reactions starting from readily available building blocks. We are pursuing the synthesis of selaginpulvilins A and B and other structurally diverse analogues for structure—activity relationship studies on the basis of HDDAR-based approaches, which will be reported in due course.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.orglett.6b03241.

Detailed experimental procedures, characterization data, ¹H and ¹³C NMR spectra (PDF)

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Notes

The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

We thank the NSF (CHE-1361620) and TACOMA Technology for financial support of this work and the high school intern Ms. Muskaan Gupta from Adlai E. Stevenson High School for her contribution in preparing compounds 7b, 11b, and 12b. The mass spectrometry facility at UIUC is greatly acknowledged.

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