
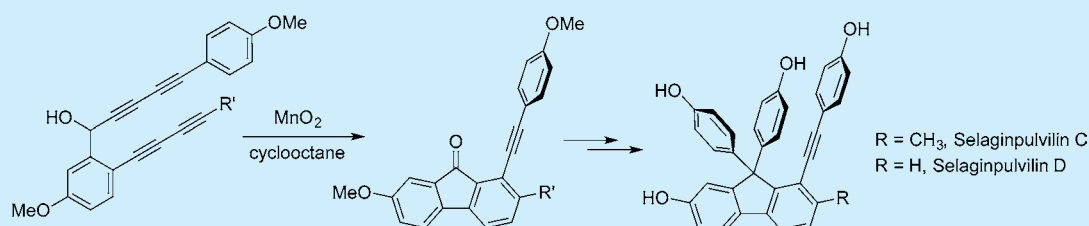


Total Synthesis of Selaginpulvilin C and D Relying on *in Situ* Formation of Arynes and Their HydrogenationRajdip Karmakar and Daesung Lee^{*†}

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 Supporting Information

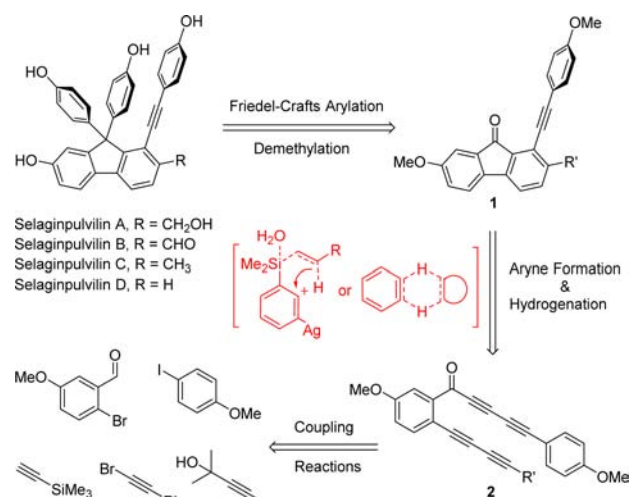
ABSTRACT: The total syntheses of selaginpulvilins C and D is described. The key strategy for the construction of the core fluorene moiety involves *in situ* formation of an aryne intermediate followed by its formal hydrogenation. The precursor tetraynes that undergo aromatization via hexadehydro Diels–Alder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

Selaginpulvilins A–D that contain a novel 9,9-diphenyl-1-(phenylethynyl)-9*H*-fluorene framework were reported by Yin and co-workers in 2014.¹ 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.² The activity-guided bioassays of ethanolic extracts of *S. pulvinata* containing selaginpulvilins A–D and other constituents showed significant inhibitory activities (IC₅₀ values of 0.11–5.13 μM) 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 9,9-diphenyl 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-9*H*-fluorenone structure can be constructed from tetrayne **2** via a hexadehydro Diels–Alder reaction^{4–6} (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**.

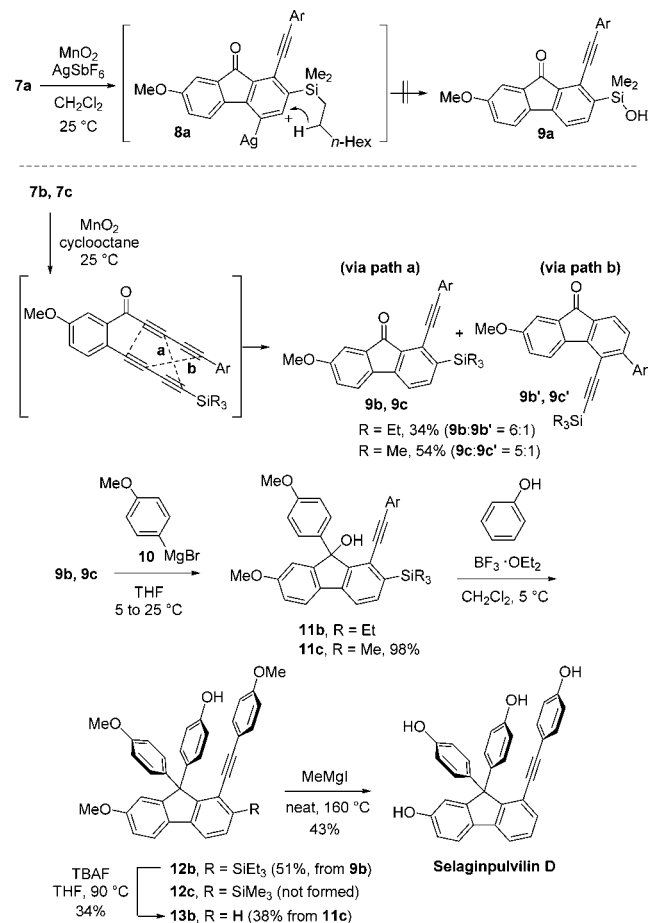
Removal of the trimethylsilyl group from **5** followed by a copper-catalyzed coupling with bromoalkynes delivered tetraynes **7a–e**.

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

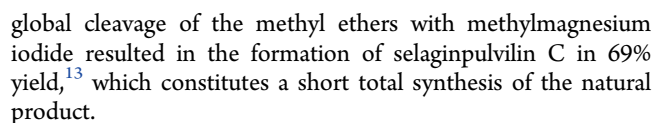
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Scheme 3. Total Synthesis of Selaginpulvinin D



Scheme 4. Total Synthesis of Selaginpulvinin C



In summary, we have accomplished total syntheses of selaginulvilins 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 selaginulvilins 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 ^{13}C NMR spectra (PDF)

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Notes

The authors declare no competing financial interest.

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