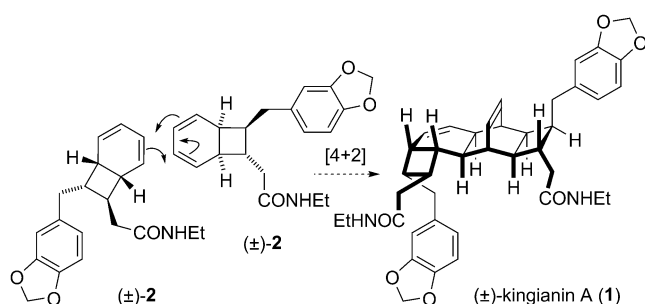


Total Synthesis of Kingianins A, D, and F**

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In Memory of Rodney W. Rickards

The kingianin natural products are a unique group of complex racemic bicyclo[4.2.0]octadiene dimers, isolated from the bark of *Endiandra kingiana* (Lauraceae) by Litaudon and co-workers.^[1] The first reported kingianin, (±)-kingianin A (**1**),^[1a] formulates as a dimer of bicyclo[4.2.0]octadiene **2**, and the Litaudon group proposed a biosynthesis involving

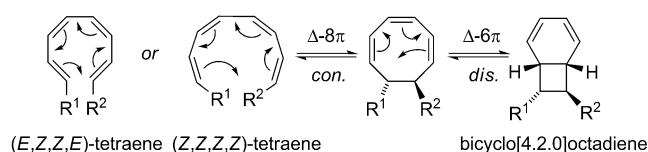


Scheme 1. Diels–Alder biosynthetic pathway to (±)-kingianin A (**1**), as proposed by Litaudon et al.^[1]

spontaneous (non-enzyme-mediated) Diels–Alder dimerization (Scheme 1).^[1] Several reports, however, describe the need for temperatures in excess of 150 °C for Diels–Alder dimerization of 1,3-cyclohexadiene.^[2] The notion that a structural feature within compound **2** may lower the barrier to thermal Diels–Alder dimerization was investigated by Moses and co-workers in 2011.^[3] An elegant synthesis of monomer **2** was achieved by the Moses group, but all attempts to induce thermal dimerization failed.^[3] Inspired by the pioneering work of Bauld and co-workers,^[4] we hypothesized that

a radical cation Diels–Alder dimerization could explain the formation of the kingianins in nature.

The bicyclo[4.2.0]octadiene framework present within Litaudon's proposed biosynthetic monomer **2** is a skeletal feature found in several natural products.^[5–9] The endiandric acids, which were isolated in racemic form in the early 1980s by Black and colleagues, were the first reported examples.^[5]



Scheme 2. The 8π–6π biosynthesis of bicyclo[4.2.0]octadiene structures, as proposed by Black et al.^[5b–f]

Black proposed that the bicyclo[4.2.0]octadiene structure was formed through a spontaneous 8π–6π domino electrocycloaddition of either an (*E,Z,Z,E*)-tetraene or a (*Z,Z,Z,Z*)-tetraene (Scheme 2).^[5b–f] Beautiful biomimetic syntheses of various bicyclo[4.2.0]octadiene natural products by Nicolaou,^[10] Trauner,^[11] Baldwin,^[12] Parker,^[13] and Moses^[14] have successfully utilized the proposed (*E,Z,Z,E*)-tetraene precursors. Evidently, the difficulty associated with preparing conjugated all-(*Z*)-polyenes has precluded their use in synthesis. In fact, (2*Z*,4*Z*,6*Z*,8*Z*)-decatetraene is both the highest all-(*Z*)-conjugated polyene and the only (*Z,Z,Z,Z*)-tetraene synthesized thus far.^[15]

Given the unprecedented structure and puzzling biosynthetic origin of the kingianin natural products,^[1] we decided to embark upon efforts towards their synthesis. The wealth of synthetic work in the literature utilizing (*E,Z,Z,E*)-tetraene precursors to access bicyclo[4.2.0]octadiene structures^[3,10–14] convinced us that we should take this opportunity to investigate the alternative biosynthetic precursor, namely the (*Z,Z,Z,Z*)-tetraene (Scheme 2).^[5b–f] Although initially drawn to the sp²–sp² cross-coupling strategy utilized by Negishi for the synthesis of (*Z,Z,Z*)-trienes,^[16] we elected instead to investigate the feasibility of a four-fold stereoselective partial reduction of a conjugated tetrayne. We anticipated that if this unprecedented^[17] and highly challenging^[18] synthetic transformation were realized then a remarkably short synthesis of the kingianins could be achieved.

The application of previously reported methods^[19] for the synthesis of unsymmetrical tetraynes was met with great difficulties. The instability of the requisite intermediates and problems associated with scaling up these approaches led us to develop a new scalable synthesis of unsymmetrical tet-

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failed to dimerize under these reaction conditions (Scheme 4).

The synthesis of ($\pm$)-kingianins A (**1**) and D (**14**) was eventually optimized to a sequence involving oxidation of alcohol **11** using the tetrapropylammonium perruthenate/*N*-methylmorpholine-*N*-oxide (TPAP/NMO) conditions of Stark et al.,^[34] with the product directly subjected to radical cation Diels–Alder dimerization using the Ledwith–Weitz salt (**13**; 5 mol %).^[4,33] The resultant mixture of diastereomeric diacids was directly converted into the corresponding diamides. Column chromatography afforded a mixture of three dimeric diamides in 17% yield over the three steps from alcohol **11**. Reverse-phase preparative HPLC allowed the isolation of analytically pure samples of ($\pm$)-kingianin A (**1**), ($\pm$)-kingianin D (**14**) and a third, as yet undetermined, structure.^[1] This radical cation Diels–Alder dimerization is a remarkably selective reaction, with only three of the potential thirty-two isomeric products isolated. Both ($\pm$)-kingianin A (**1**), a homochiral dimer, and ($\pm$)-kingianin D (**14**), a heterochiral dimer, are the result of *endo*-selective formal Diels–Alder reactions occurring at the convex faces of both diene and dienophile. Previous studies have shown that the radical cation Diels–Alder dimerization of 1,3-cyclohexadiene is *endo* selective,^[4] however, a full explanation of the site and orientational regioselectivity observed in the present study will require further investigation. The natural product ($\pm$)-kingianin F (**15**) was similarly obtained by dimerization of the other bicyclo[4.2.0]octadiene diastereomer **12**, followed by double oxidation and diamide formation.^[35]

In summary, our highly divergent biomimetic strategy has resulted in the total synthesis of ($\pm$)-kingianins A (**1**), D (**14**) and F (**15**), in a longest linear sequence of ten steps. The noteworthy synthetic aspects of our successful approach include the gram-scale preparation of an unsymmetrical tetrayne, the unprecedented reduction of a conjugated tetrayne to a (*Z,Z,Z,Z*)-tetraene, and radical cation Diels–Alder dimerization of functionalized bicyclo[4.2.0]octadienes. From these studies, we conclude that the kingianins are not formed through spontaneous Diels–Alder dimerization. Instead, we propose that nature uses a SET-mediated cycloaddition analogous to the approach described herein.^[36,37] Our results, in conjunction with previous biomimetic syntheses,^[3,10–14] demonstrate that (*E,Z,Z,E*)-tetraenes, and not their all- (*Z*) congeners,^[32] are the likely biosynthetic precursors to bicyclo[4.2.0]octadiene natural products.^[38]

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