Synthesis of (+)-Chatancin

**Significance:** Platelet-activating factor (PAF) antagonist (+)-chatancin was isolated from the soft coral *Sarcophyton* species. Its unique carbon skeleton features a fused ring system bridged by a hemiketal, and six contiguous stereocenters. Bio-synthetically, the natural product is hypothesized to originate from the macrocyclic diterpene cembrene A by a sequence of oxidations and a transannular Diels–Alder cycloaddition. Avoiding the lengthy synthesis of such a macrocycle, Zhao and Maimone employed a related Diels–Alder reaction prior to cyclizing the final ring. As a result, their route is remarkable for its conciseness, giving the natural product in only eight linear steps and 13% overall yield.

**Comment:** Starting from farnesal, a short sequence was devised to give key intermediate C. Quick access to pyrone C enabled the probing of the transannular Diels–Alder reaction as a means to construct the first decalin motif. Thus, palladium-catalyzed carbonylative esterification and subsequent heating in toluene gave diastereomers E and F exclusively, creating four new stereocenters in the process. Allylic chlorination yielded G, which was directly treated with zinc dust to give the full carbon skeleton as a single diastereomer in a Barbier-type annulation. After palladium-catalyzed regioselective hydrogenation of the exo olefin in H, (+)-chatancin was obtained. The structure was also confirmed by X-ray analysis.