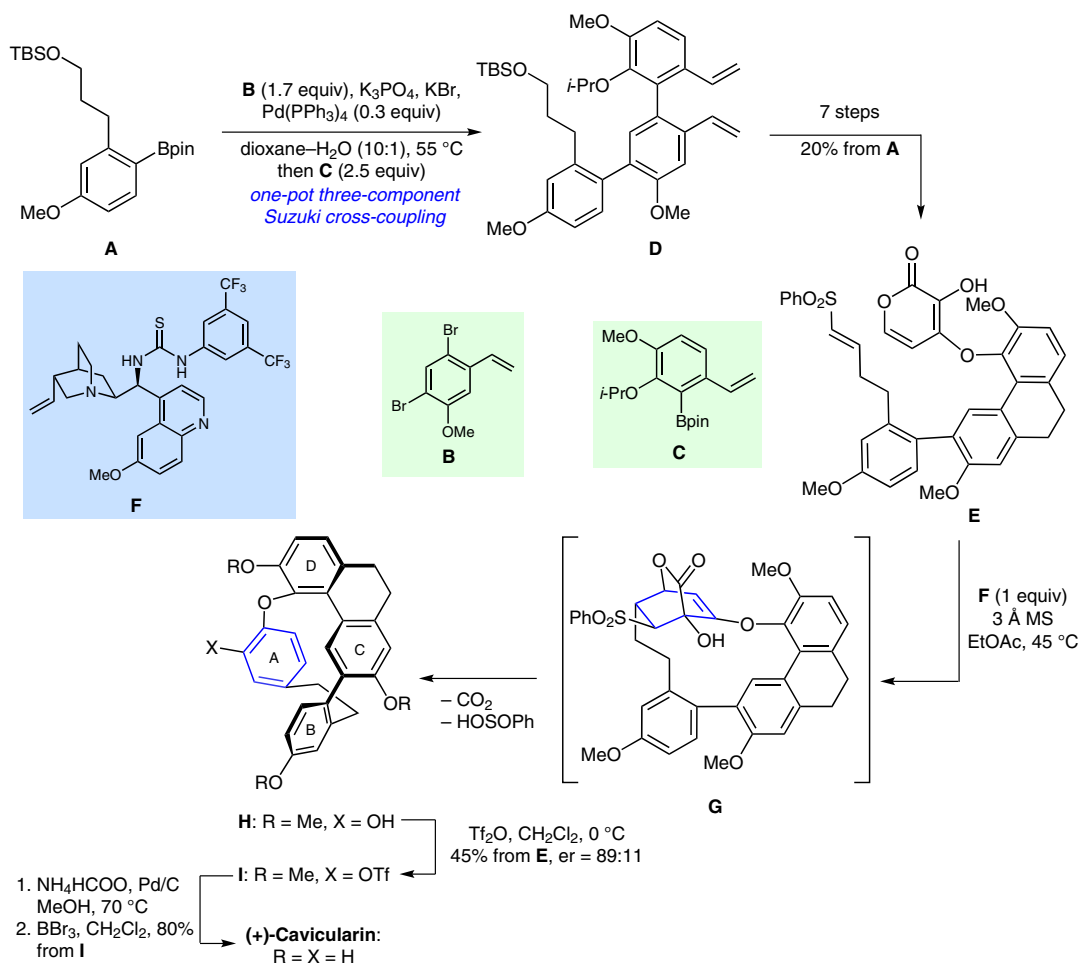


P. ZHAO, C. M. BEAUDRY* (OREGON STATE UNIVERSITY, CORVALLIS, USA)
 Enantioselective and Regioselective Pyrone Diels–Alder Reactions of Vinyl Sulfones: Total Synthesis of
 (+)-Cavicularin
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Total Synthesis of (+)-Cavicularin



Significance: (+)-Cavicularin, isolated from the liverwort *Cavicularia densa*, is a chiral cyclophane natural product. Because of its unusual molecular structure, several total syntheses have been reported to date. Zhao and Beaudry report a conceptionally different approach, which relies on an intramolecular enantioselective pyrone Diels–Alder reaction with subsequent CO_2 extrusion to generate the aromatic A ring of the natural product.

Comment: The synthesis commences with a remarkably selective one-pot three-component Suzuki cross-coupling between dibromide **B** and boronic esters **A** and **C**. Coupling product **D** was further advanced to α -hydroxy pyrone **E**. In the presence of cinchona alkaloid **F**, this material underwent the desired Diels–Alder reaction to yield intermediate **G**, which immediately eliminated CO_2 and phenylsulfonic acid to generate **H** as a single regioisomer. Finally, reduction and protecting group removal yielded (+)-cavicularin in 7.3% overall yield.

SYNFACTS Contributors: Erick M. Carreira, Christian Ebner
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