## Category

Synthesis of Materials and Unnatural Products

## Key words

reductive double cyclization

methylene-bridged stilbenes

dibenzo[a,e]pentalenes



H. ZHANG, T. KARASAWA, H. YAMADA, A. WAKAMIYA, S. YAMAGUCHI\* (NAGOYA UNIVERSITY, JAPAN)

Intramolecular Reductive Double Cyclization of o,o'-Bis(arylcarbonyl)diphenylacetylenes: Synthesis of Ladder  $\pi$ -Conjugated Skeletons

Org. Lett. 2009, 11, 3076-3079.

## Synthesis of Conjugated Ladder Systems by a Reductive Double Cyclization

**Significance:** The intramolecular cyclization of *ortho*-substituted benzene rings is a popular synthetic technique for the formation of benzene-fused five-membered rings. The authors have extended this technique to the synthesis of ladder-type polycyclic compounds by employing diaryl acetylene precursors. In previous reports, reduction of the acetylene moiety led to a double cyclization with heteroatoms. In the current paper, carbonyl substituents on the benzene ring act as the reducible moiety and are used to generate methylene-bridged stilbenes (2) and dibenzo[a,e]pentalenes (3).

**Comment:** The authors used computational studies to determine that the carbonyl groups in compounds of type **1** are more readily reduced than the acetylene moiety. Cyclic voltammetry demonstrated close proximity of the two irreversible one-electron reduction potentials; therefore, the authors propose a mechanism involving synchronous double cyclization through biradical **4**. Compound **3** is formed by the overreduction of compound **2** by lithium naphthalenide. The  $\pi$ -conjugation of these polycyclic hydrocarbons can be extended further via cross-coupling chemistry, presenting interesting candidates for organic electronics.

**SYNFACTS Contributors:** Timothy M. Swager, Rebecca R. Parkhurst Synfacts 2009, 10, 1096-1096 Published online: 22.09.2009 **DOI:** 10.1055/s-0029-1217930; **Reg-No.:** S09909SF