BORON TRIFLUORIDE PROMOTED REACTION OF DITHIO-SUBSTITUTED ALLYLIC ANIONS AND CYCLIC Ethers

Jim-Min Fang* and Ming-Yi Chen
Department of Chemistry, National Taiwan University
Taipei, 10764, Taiwan, Republic of China

Abstract— Allylic anions generated from 2-propenyl-1,3-dithiane and 2-styryl-1,3-dithiane react exclusively at α-carbons with three- to six-membered cyclic ethers in the presence of BF₃·Et₂O.

It has been shown that combination of organolithium and BF₃·Et₂O is superb for the ring-opening reaction of oxiranes and oxetanes. With 2-substituted oxirane, the ring-opening reaction occurs selectively at the less hindered site. The active species in the reaction of alkyllithium is once presumed to be RĊ=C-BF₃, although other evidence indicates the reaction may proceed via a coordinate intermediate of Lewis acid BF₃ and cyclic ether, which is then attacked by alkyl- or vinyl- or phenyllithium. More recently, ring-opening reactions of tetrahydrofurans and tetrahydropyrans, most of them having phenyl group(s) at the C-2 positions, are also realized by treatment with the mixed reagent of allyl- (or alkynyl-)silane and TiCl₄. Since the reaction occurs selectively at the more substituted C-2 site, it may proceed via a cationic intermediate stabilized by phenyl group(s). This method is synthetically useful in extension of four and five carbon units, but it is not applicable to reactions with parent tetrahydrofuran, or tetrahydropyrans, or their 2-alkyl derivatives. We here demonstrate effective ring-opening reactions of various cyclic ethers by allylic anions generated from 2-propenyl-1,3-dithiane and 2-styryl-1,3-dithiane (Scheme 1).

The typical experimental procedure is described as follows. To a solution of 2-propenyl-1,3-dithiane (1 mmol) in ether (10 mL) was added dropwise n-BuLi (1.2 mmol, 1.6 M in hexane) at -40 °C. After stirring for 20 min, the resulting allyllithium solution was cooled to -78 °C, and 1 mmol of BF₃·Et₂O (diluted with 1 mL of ether) was added dropwise, followed by addition of 1.2 mmol of appropriate cyclic ether. The ring-opening reaction completed in 20 to 60 min as revealed by the TLC analysis.

The reaction is highly regioselective. As the hardness of cyclic ether greatly increases when it coordinates with BF₃, it is expected to react at the α-site of dithio-substituted allylic anion. In agreement with precedent, present ring-opening reactions took place exclusively at the less hindered α'-carbons of 2-methyloxirane and 2-methylfuran, while reactions with 2 phenyloxirane showed preference for attack at the benzyl position. This method provides a wide scope of carbon-carbon bond formation with both parent and substituted three- to six-membered cyclic ethers in giving high yields of alchol products (3-18). As the dithiane moiety is readily hydrolyzed with HgCl₂, this method also culminates in conversion of α,β-unsaturated aldehydes to enones and related compounds by the polarity reversal process. Thus, treatment of styryldithiane 6 with HgCl₂ (aqueous MeOH, 80 °C, 0.5 h) afforded 5-hydroxy-1-phenyl-1-hexen-3-one (19) as the sole product. Hydrolysis of 11 (HgCl₂, aqueous MeOH) followed by demercuration (NaOH, NaBH₄) resulted in a single product of ketone 20.
Scheme 1.

\[ \text{BuLi} \quad \text{R S} \quad \xrightarrow{1)} \quad \text{BF}_3\text{Et}_2\text{O} \quad \text{R S} \quad \xrightarrow{2)} \quad \text{cyclic ether} \quad \text{Y= R = Me} \quad \text{R = Ph} \]

1 \( R = \text{Me} \)
2 \( R = \text{Ph} \)

(a) The number in the parenthesis represents isolated yield (%) of products.
(b) 9 is the \( \gamma \)-substitution product consisted of two diastereomers (2:1).

Similar reaction of propenyldithiane 12 in aqueous CH₃CN gave dihydroxy ketone 21 in a high yield. When compound 3 was treated subsequently with HgCl₂, NaOH and excess of NaBH₄, a product of cis 2-methyl-4-hydroxytetrahydropyran (22) was obtained exclusively.

References and Notes:
9. All products had satisfactory elemental analyses and compatible spectra (IR, MS and NMR).

(Received in Japan 25 August 1988)