# Reaction of Benzyne with Butadiene 

By L. F. Hatch* and D. Peter<br>(Department of Chemistry, The University of Texas at Austin, Austin, Texas 78712)

Benzyne has been shown to act as a dienophile with cyclic dienes, but apparently not with acyclic dienes. The reason has been given that benzyne does not survive long enough to permit the open-chain diene to assume the requisite cislike arrangement that the cyclic dienes possess. ${ }^{1}$ We have now been able to effect this type of reaction.
Benzyne and buta-1,3-diene, generated by the concurrent thermal decomposition of benzene-diazonium-2-carboxylate ${ }^{2}$ and 3 -sulpholene ( 2,5 dihydrothiophen 1,1-dioxide) respectively, react in pentan-2-one at $100^{\circ}$ to give a $9 \%$ yield of


1,4-dihydronaphthalene. A temperature of $100^{\circ}$ is necessary to provide a reasonable, continuous concentration of butadiene from the 3 -sulpholene; the method of Friedman and Logullo ${ }^{3}$ was therefore useful for the preparation and decomposition in situ of benzenediazonium-2-carboxylate.

It is suggested that the nascent butadiene from the decomposition of 3 -sulpholene would be in the preferred cis-conformation, thus enabling reaction to take place.

That 3 -sulpholene acts simply as a source of butadiene, and that the $\mathrm{SO}_{2}$ entity is not involved in a transition state is shown by kinetic data ${ }^{4}$ obtained from the thermal decomposition of 3 -sulpholene in the presence and absence of maleic anhydride as the dienophile. The reactions are first order with $k=2.4 \times 10^{-3}\left(\min .^{-1}\right)$. At temperatures below the decomposition temperature of 3 -sulpholene, no addition product is formed with benzyne.

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