

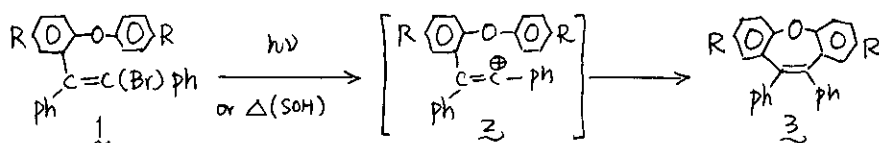
FORMATION OF HETEROCYCLES BY INTRAMOLECULAR CYCLIZATION
OF UNSATURATED CARBENIUM IONS

Tsugio Kitamura, Hironobu Kawasato, Chihiro Hatano, Shinjiro Kobayashi,
and Hiroshi Taniguchi

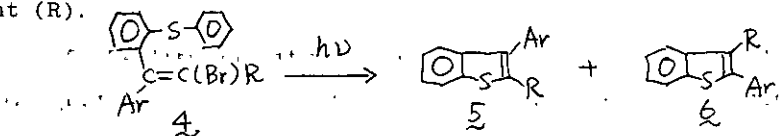
Department of Applied Chemistry, Faculty of Engineering,
Kyushu University 36, Hakozaki, Fukuoka 812, Japan

Intramolecular cyclization of vinyl bromides and allenyl chlorides bearing arylhetero groups was studied.

The reaction of β -(*o*-aryloxyphenyl)vinyl bromides 1 proceeded under both solvolytic and photolytic conditions. The products, dibenzoxepin derivatives 3, are derived from the resulting vinyl cations, followed by electrophilic attack on the aryloxy group.



On the other hand, β -(*o*-arylthiophenyl)vinyl bromides 4 afforded benzothio-
phene derivatives 5 and 6 under photolytic conditions. Vinyl radicals and vinyl
cations are responsible for the formation of 5 and 6, depending on the α
substituent (R).



Intramolecular reaction of γ -(*o*-arylheterophenyl)allenyl chlorides 7 with
 ZnCl_2 -or SnCl_4 gave six-membered heterocycles 8 as the sole products. Cyclization
at the propargyl position on the allenyl cations is explained by the large positive
charge density and the strain of the products derived from cyclization at the
allenyl position.

