

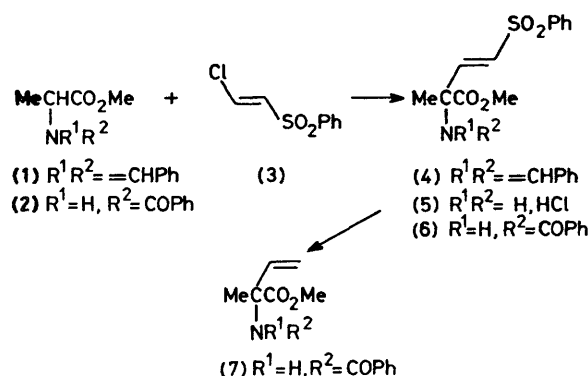
Phenyl *trans*-2-Chlorovinyl Sulphone, a Vinyl Cation Equivalent

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Summary Phenyl *trans*-2-chlorovinyl sulphone is a vinyl cation equivalent useful for the conversion of α -amino acids into α -vinyl α -amino acids.

It is known that phenyl *trans*-2-chlorovinyl sulphone (3) readily undergoes substitution with a variety of heteroatomic nucleophiles,¹ and that vinyl sulphones can be reduced to the corresponding olefins.² A sequential combination of these reactions, using a carbanionic nucleophile would appear, therefore, to offer a means for the electrophilic introduction of a vinyl unit at carbon. Our interest in α -vinyl α -amino acids as potential irreversible enzyme inhibitors³ has led us to investigate, in a model sequence, the transformation of alanine to *N*-benzoyl- α -vinyl-alanine



methyl ester, using phenyl *trans*-2-chlorovinyl sulphone (3) as a vinyl cation synthon.⁴

Thus the ester enolate derived from the ester (1)^{5,6} adds, with concomitant elimination of chloride, to (3) at -78°C to afford the phenyl *trans*-vinyl sulphone (4). Without purification compound (4) was treated with dilute hydrochloric acid, the resulting amine hydrochloride (5)[†] (m.p. 120°C) being isolated in 83% overall yield. Compound (5) was converted into the benzamide (6)[†] (m.p. 143°C) by routine methods, and (6) was readily desulphurized using aluminium amalgam,² to afford *N*-benzoyl- α -vinyl-alanine methyl ester (7)[†] (m.p. 109°C) in 80% yield. Alternatively *N*-benzoylalanine methyl ester (2) can be directly converted,

in 57% yield, into (6), *via* the reaction of its derived dianion⁷ with (3).

Another α -vinyl α -amino acid in which the α carbon is also fully substituted, α -vinyl-3,4-dihydroxyphenylalanine, has been made previously *via* reduction of acetylenic intermediates,^{3,8} while Baldwin⁹ has reported the synthesis of some $\beta\gamma$ -unsaturated glycine analogues *via* reduction of the corresponding nitro derivatives. The use of a vinyl cation synthon offers a complementary strategy, as the α -vinyl α -amino acids may be prepared from the corresponding α -amino acids.

(Received, 3rd July 1978; Com. 698.)

[†] N.m.r. and i.r. spectra and elemental analyses are consistent with the proposed structure.

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