of pentalenolactone as its benzylamine salt (see ref 19) were provided by the Pfizer Laboratories through the courtesy of Drs. Arthur Nagel and Robert Volkmann and by Professor David Cane of Brown University. The authors also acknowledge the help and advice of Dr. Milan Uskokovic of the Hoffmann-La Roche Laboratories in dealing with the Brederick reagent.

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Thione S-Methylides as Quasi-Wittig Reagents

Sir:

Betaines (1) derived from reaction of carbonyl functions with alkylidenesulfuranes (2) undergo an intramolecular displacement reaction leading to oxiranes,1 while alkylidene-





Figure 1. Frontier molecular orbitals involved in betaine decompositions.



phosphoranes (Wittig reagents) select the alternate closure to a hypervalent phosphorus intermediate which is penultimate to the alkene product.² The partition of **1** between two potential surfaces whose maxima are described by transition states 3 or 4 electronically³ depends upon the magnitude of the first-order frontier interaction⁴ (E) between the overlapping oxygen p orbital (p_o) and the terminus of the high lying σ^* orbital involved (S in 4, C in 3; Figure 1). Sulfonium ylide derived betaines might be encouraged to undergo intramolecular closure to yield a "Wittig" reaction if a low-lying π^* orbital was available at sulfur (Figure 1). This requirement would be met by betaines (6) generated from thione methylides⁵ and appropriate carbonyl substrates. We now report that thione



methylides can undergo Wittig-type reactions and stereochemically complement the Wadsworth-Emmons phosphonate modification of the Wittig reaction in substituted acrylic ester syntheses.

The quasi-Wittig reagent, N, N'-dimethylimidazole-2-thione S-carbomethoxymethylide (7), is easily generated in situ from the corresponding salt 8. Specifically, to a solution of 1 equiv



of 8 in dry acetonitrile at 0 °C is added 1 equiv of base and after 5 min the carbonyl reagent is introduced. The resulting mixture is stirred for 10 min at 0 °C to complete the reaction. Table I summarizes the results including overall isolated yield of pure product and the isomer distribution for some representative

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Table I. Products from Quasi-Wittig Reaction

9, R	Z isomer	E isomer	% yield
Ph	75	25	60
CH ₃ CH ₂ CH ₂	71	29	31 <i>ª</i>
$(CH_3)_2C = CHCH_2CH_2CH_2$	75	25	42 <i>ª</i>
CH ₃ CH=CHCH=CH	35	65	12

^a Molar ratio of aldehyde to 8 was 2:1.

aldehydes (9).⁶ Improvement of the overall yield of acrylic ester from enolizable substrates was effected by increasing the ratio of **8** to substrate present in the reaction mixture.

The stabilized thione S-methylide differs from its Wittig counterparts in that the former produces primarily the Z isomer (Table I). With this difference in mind, two pathways via 11 or 12 may be involved in the formation of the observed products. Considering the most favorable conformation, the



erythro-betaine, 11, arises from addition of the ylide to the carbonyl function with the changed centers diametrically opposed while the opposite is true for the development of the *threo*-betaine, 12. If steps $11 + 12 \rightarrow 13 \rightarrow 10$ are considered irreversible or faster than betaine dissociation the isomer distribution of 10 is determined by the ratio of 11 to 12 which, in turn, is a result of this conformational preference, i.e.,



This must be the case as the predominant Z isomer arises from the sterically most encumbered hypervalent sulfurane 13b.⁷ Addition of a metal cation to the reaction should lower the transition state energy leading to 11 relative to 12 by ion pairing at the alkoxide site and increase the proportion of betaine 11 available for Z-alkene production. In fact, addition of LiI (3 equiv) to the ylide reaction with benzaldehyde raised the Z to E isomer distribution of the methyl cinnamate product to 92:8.⁸ Thione S-methylides with sterically less-demanding groups at the carbonionic center yield a more nearly equal distribution of product stereoisomers. For example, 14 provides (50% isolated yield) a ratio of (E)- to (Z)-cinnamonitriles of



59:41. It should be noted also that the electron donating ability of the thione substituents is critical in determining both the partition of the reaction paths and the distribution of geometrical isomers.⁹ The ylide **16** derived from tetramethylthi-



ourea gave a 90:10 ratio of E- to Z-17 along with 18 in an overall ratio of 1:1.

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- (9) The energetic placement of the acceptor orbital of the equatorial substituent governs the bond energy of the hypervalent system (see ref 5).

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Isolation and Characterization of Two C₁₁H₁₁ Cationic Rearrangements

Sir:

Unsaturated substrates rearrange by some mechanisms that are cationic and by others that are not. The distinction is easier to recognize in principle than to resolve in practice. How can one isolate the rearrangement of an ionic intermediate from the competitive rearrangement of its covalent precursor or product?

We encountered this problem, as one might have expected, in trying to generate the bicyclo[4.3.2]undecatetraenyl cation¹ (1, Figure 1) from its covalent precursor under classic S_N1 conditions (2,6-lutidine-buffered 70% aqueous acetone hydrolysis of the 9-syn-p-nitrobenzoate (2)² at 80 °C). The problem was previously recognized in studies of apparently cationic C₉H₉ rearrangements.³ Within the C₁₁H₁₁ series, some ten different rearrangements have already been reported.^{4,5} Most of them appear to be cationic, but some clearly are not.^{5a,b,d,e}

The S_N1 conditions which we selected generated only two products of kinetic control,⁶ both rearranged: the anti-tetracyclic alcohol (4)^{7a,b} and anti-pentacyclic alcohol (5).^{7a,c} We

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