nomenclature system would logically involve changing the ligand name as a function of complex oxidation state. Consequently, we favor using the trivial name diaminomaleonitrile (damn) for the ligand to be consistent with previous literature (ref 11)

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Synthesis of New β -Lactam Antibiotics¹

Sir:

In a previous paper² we reported the preparations of azetidinone disulfides, 1, and 2β -halomethylpenicillins, 4 (Y = halogen), which are important precursors for the syntheses of 2β -substituted methylpenicillins 4, 3β -substituted cephams 5,3 and desacetoxycephalosporin 11.2 The present communication deals with the isolation of sulfenylanilide 2b $(Y = NHC_6H_5)$, an intermediate involved in the conversion of 1 into 4 and 5, and the synthesis of a new tricyclic β -lactam antibiotic 6a via intramolecular cyclization of 4a (Y =

We have recently established a method⁴ which allows the stereospecific conversion of 1 into 5 as well as 4 (Y = halogen)² by treatment with various nucleophiles under the presence of Ag+. The formation of 4 and 5 presumably proceeds through the sulfenyl derivative 2 which is then transformed into episulfonium ion 3; we have now secured corroborative evidence for this mechanism by isolation of the sulfenylanilide 2b (Y = NHC₆H₅). Thus treatment of 1b with aniline at room temperature in CH2Cl2 under the presence of AgBF₄ yielded 3β -anilinocepham $5b^4$ (Y = NHC₆H₅), mp 129-130°, $[\alpha]D$ +58.5° (EtOH), 45% yield. On the other hand, treatment of 1b with aniline in ethyl acetate under the presence of AgOAc affords 2b (Y = NHC_6H_5): colorless oil; 90% yield; m/e 455; ir 1770 cm⁻¹ (β -lactam); NMR (CDCl₃), 1.97 (s, 4'-H), 4.78 (s, 2'-H), 5.05 and 5.22 (two br s, 5'-H), 5.08 (d, J = 5 Hz, 4-H), 5.53 (dd, J = 5and 8 Hz, 3-H). Reaction of 2b (Y = NHC₆H₅) with BF_3OEt_2 gave 5b (Y = NHC₆H₅) in 40% yield, thus supporting the intermediacy of 2b (Y = NHC₆H₅) (Scheme I).

Furthermore, sulfenyl derivative 2b (Y = NHC₆H₅) may act as an intermediate for introducing other nucleophilic groups. Thus treatment of 2b $(Y = NHC_6H_5)$ with BF₃OEt₂ in MeOH gave in 80% yield a 1:3 mixture of 4b $(Y = OCH_3)$ and 5b (OCH_3) .⁴ Similarly reaction of 2b $(Y = OCH_3)$

= NHC₆H₅) with HCl yielded 4b $(Y = Cl)^2$ in quantitative yield.

We have been able to achieve the conversion of 4a (Y = Br) into the intramolecularly cyclized product 6a. Thus base treatment of 4a (Y = Br) with bases led to the ringexpanded cephem 11a2 in high yield, but formation of a minute amount of a by-product was also observed; silica gel chromatography of this gave colorless crystals, mp 140-143°, $[\alpha]D + 203.2°$ (CHCl₃). The structure **6a** was assigned on the basis of spectral data: ir, 1795 cm⁻¹ (β -lactam); NMR (CDCl₃), 1.70 (s, 2-CH₃), 2.06 and 2.27 (ABq, J = 7 Hz, 3-H), 5.47 (dd, J = 4 and 9 Hz, 7-H),6.19 (d, J = 4 Hz, 6-H). It is reasonable to postulate that in the present case product 6a is formed directly by an intramolecular nucleophilic displacement in 4a (Y = Br) and not through an episulfonium ion 3a, although formation of the latter has been demonstrated in both ring expansion^{2,4,5} and nucleophilic substitution⁴ reactions of 4.

Formation of cephem 11a was avoided by starting from the corresponding sulfoxide 8a (Y = Br) which would not yield the episulfonium ion 3. Thus when 8a (Y = Br) obtained by oxidation of 4a (Y = Br) with m-chloroperbenzoic acid. was treated with Et₃N in acetone at room temperature for 72 hr, a 1:1 mixture of 7a and 9a was obtained in ca. 52% yield. The compounds were separated by silica gel chromatography. The structure of 7a, mp $148-148.5^{\circ}$, $[\alpha]D + 22.8^{\circ}$ (CHCl₃), was established by comparison with a sample synthesized by oxidation of 6a with m-chloroperbenzoic acid. The second product, mp $147-148^{\circ}$, was assigned structure 9a from spectral data; this was substantiated by an independent synthesis from $10a^{7}$ by m-chloroperbenzoic acid oxidation. It is obvious that compound 7a is formed by a nucleophilic attack of the C-3 carbanion on the 2β -bromomethyl group, while formation of compound 9a can be accounted for by the arrows shown in structure 8, i.e., a rearrangement initiated by formation of the C-6 carbanion.

Cyclization leading to 7a was best achieved under the following optimum conditions. Namely, treatment of 8a (Y = Br) with 1,5-diazabicyclo[5.4.0]undec-5-ene (DBU) in DMF at -30° for 1 hr gave 7a in 80% yield. Similarly, sulfoxide 8c (Y = Br) derived from 8a (Y = Br) by a deacylation method⁸ gave 7c, tosylate mp 176-179° dec, in 81% yield. These results indicate that the intramolecular cyclization proceeds faster than the intramolecular rearrangement and also suggest that cyclization of 4a (Y = Br) would take place preferentially without production of the episulfonium ion 3 if a strong base was employed.

This was corroborated as follows. Treatment of 4a with DBU under the conditions as described above yielded 6a in ca. 80% yield; however, this reaction was always accompanied by production of the undesired cephem 11a, 8%. Tricyclic sulfide 6a was also derived from 7a by reduction with PCl₃ in DMF in 76% yield. The synthesis of 6a from 4a represents the first intramolecular nucleophilic displacement in the penam system.^{9,10}

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- (10) Conversion of compound 6 to 2-methylcephem will be reported shortly.

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Evidence for $S(^1D)$ Atom Reactions Involving $^{34}S(n,\gamma)^{35}S$ Nuclear Recoil Generated Sulfur

Sir:

Reactions of recoil sulfur atoms produced by both the $^{35}\text{Cl}(n,p)^{35}\text{S}$ and $^{34}\text{S}(n,\gamma)^{35}\text{S}$ nuclear processes in the gas phase are known to be quite complex due primarily to the

Table I. Data for ³⁵S + CS₂ Exchange Reaction as a Function of Gaseous Additives

Additives		Sample composition (Torr)			Yield CS35S nor-
M	N	P_{CS_2}	$\dot{P_{ m M}}$	P_{N}	malized activitya
		200			490 ± 10
Η,		200	100		350 ± 10
H,		200	200		256 ± 10
H,		200	400		153 ± 5
H_2 H_2 H_2 H_2	Ar	200	100	1077	235 ± 10
C_2H_6		200	400		49 ± 3
C_2H_4		200	400		41 ± 2

a The yield of CS³⁵S is the normalized activity per Torr of S available. Results reported are the average of two or more determinations and the errors are based upon the statistical errors in the aliquots counted.

polyvalent nature of the atom. In addition the propensity for sulfur species to undergo oxidation reduction processes further complicates the identification and characterization of primary reaction channels. Earlier accounts of nuclear recoil sulfur reactions have, in general, ignored the role of electronic excitation, or, as reported in one case, the contribution of $S(^1D)$ has been discounted as being unimportant.²

The differences in reactivity for the low lying electronic states of sulfur atoms have been extensively characterized in systems where the atoms are photochemically generated.³ In particular, changes in both reaction mechanisms and reaction rates are known to be associated with $S(^1D)$ and $S(^3P)$ reactions. Direct use of the techniques employed in photochemical systems, however, has not led to an unambiguous interpretation of nuclear recoil systems.² We have obtained evidence for the gas phase reactions of singlet sulfur atoms, $S(^1D)$, generated by the $^{34}S(\mathbf{n},\gamma)^{35}S$ nuclear recoil reaction in the presence of CS_2 .

The recoil sulfur species were generated by thermal neutron irradiation for 10 min to 1 hr at a flux of 10^{12} (n/cm²)/sec on a rotating multiple sample holder at the Washington State Nuclear Radiation Center. All samples were prepared by standard high vacuum techniques and flame sealed in 15-cm³ cylindrical quartz irradiation vessels. Carbon disulfide (Matheson Coleman and Bell) was thoroughly degassed and vacuum distilled before use. No impurities were detected by gas chromatography. Research grade C_2H_4 and C_2H_6 (Phillips) and H_2 and Ar (Matheson) additives were used directly from gas cylinders. Product analysis was carried out by radio gas chromatography incorporating an internal flow proportional detector.

The reactions between both triplet and singlet sulfur atoms and CS₂ are known to occur in the gas phase and have been discussed previously.⁴ In this study the effect of various gas additives on the total production of CS³⁵S from nonlabeled CS₂ has been determined in order to investigate the role of various forms of recoil atom excitation in driving the exchange reaction. In Table I, the CS³⁵S produced as a function of H₂ dilution and at fixed dilution with C₂H₄, C₂H₆, and Ar is tabulated. The activity of the labeled CS₂ is reported as specific activity per Torr of CS₂ originally present in the sample mixture.⁵

The results indicate that the efficiency of the sulfur atom exchange reaction with CS₂ rises linearly within experimental error as the mole fraction of CS₂ increases. This behavior suggests that the hydrogen additive is competing effectively for the reactive sulfur atoms or sulfur containing intermediates.⁶ Such competition might arise from a moderating effect on a hot reaction, an exchange reaction between an excited intermediate and CS₂, or a quenching effect which eliminates electronically excited states of atomic sulfur. The reaction of ground electronic state sulfur atoms with H₂, if it occurs at all, cannot compete with the reaction