AZAFULVENES 4'. CYCLOADDITION REACTION OF 6-AMINO-1-AZAFULVENE TO ISOCYANATE, KETENE AND SULFENE

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T<u>akashi</u> K<u>obayashi, Shoji</u> K<u>ajigaeshi,</u> and S<u>huji Kanemasa</u>*
<u>Department of Industrial Chemistry, Faculty of Engineering</u> Department of Industrial Chemistry, Faculty of Engineering, Yamaguchi University, Tokiwadai, Ube 755

6-Amino-1-azafulvene and its benzo analog, generated from the iminium perchlorate by deprotonation, reacted with isocyanate, ketene and sulfene affording the corresponding [6+2] cycloadducts.

It was previously reported that 6-amino-1-azafulvene 1 could be regenerated by the pyrolysis of **5,10-dihydro-5,1O-diaminodipyrrolo[1,2-a:1,2-d]pyrazine,** the dimer of **1,** and was captured by isocyanate giving the corresponding [6+2] cycloadduct². In addition, the treatment of the iminium perchlorate 2 , synthesized from pyrrole- and indolecarbaldehyde with perchlorate of secondary amine, with some bases gave the dimer of 1³. This result indicates that deprotonation of 2 would lead to the generation of l , followed by the dimerization.

In order to obtain a corroboratory evidence for the generation of 6-amino-lazafulvene 1 by deprotonation from 2 and also to investigate the reactivity of 1 in its cycloaddition reaction, the reactions of 2 with base in the presence of isocyanate, ketene and sulfene were attempted in this communication.

A mixture of equivalent amount of **N-(2-pyrrolylmethy1ene)morpholinium** per-

chlorate 2a and phenyl isocyanate in benzene was treated with triethylamine at room temperature to give the 1:1 adduct 3a in 72% yield which showed the identical ir and nmr spectrum with **2,3-dihydro-1-morpholino-2-phenyl-1H-pyrrolo-**[$1,2$ -c]imidazol-3-one already prepared by us². After adding triethylamine to the suspension of the iminium salt 2a in methylene chloride at -50°C, the subsequent addition of phenyl isocyanate to this reaction mixture formed the same product 3a in 88% yield. This indicates that the iminium salt 2a liberated 6morpholino-1-azafulvene la on the treatment with base, which is considerably stable at low temperature enough to permit the cycloaddition with phenyl isocyanate into the corresponding [6+2] cycloadduct.

The similar reaction of N-(3-methyl-2-indoly1methylene)morpholinium perchlorate 2b with triethylamine in the presence of phenyl isocyanate gave $2,3$ **dihydro-9-methyl-l-morpholino-2-phenyl-lH-imidazo[l.5-alindol-3-one 3** in 76% yield. The structure of 3b was deduced on the basis of the spectral data shown in Table 1.

As well known, ketene and sulfene can be 1 iberated from acyl ha1 ide and sulfonyl halide by the elimination of hydrogen halide, respectively. The reactions of 1 with ketene and sulfene were therefore carried out by use of the method in which the generation of 1 from 2 was made concurrently with that of ketene from acyl halide and that of sulfene from sulfonyl chloride by the action of base.

Thus the mixture of iminium salt $2a$ and α , α -diphenylacetyl chloride in benzene was treated with an excess of triethylamine and the resultant reaction mixture was chromatographed on alumina to afford the 1:l adduct of la with diphenylketene in 67% yield. This product 5a was identified to be 2,3-dihydro-1-morpholino-**2,2-diphenyl-lH-pyrrolizin-3-one** by its ir and nmr spectrum in which the absorption band at 1740 cm⁻¹ was assigned to the five-membered carbonyl group and three the protons on pyrrole ring appeared as each multiplet. Similarly la reacted with dimethylketene and lb did with dimethyl- and diphenylketene to give the respective [6+2] cycloadducts, 4a, 4b and 5b. The reactions of la and 1b with phenylketene, however, resulted in no isolation of the [6+2] cycloadducts,
<u>6a</u>' and <u>6b</u>', but formation of the deaminated products as red crystals. The ease of deamination would due to the trans conformation of the initially formed cycloadducts **6'.** The comparable deamination reactions have been allowed in the case of the trans $[6+2]$ cycloadducts of $\underline{1a}$ with acrylates and methyl vinyl ketone⁴.

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The reactivity of 6-amino-1-azafulvene 1 in its cycloaddition to ketene is quite distinct from that of an ordinary fulvene which underwent the [2+2] cycloaddition to mono- and dichloroketene at the endocyclic double dond^{5,6}. Only an example of the [6+2] cycloaddition of 1,4-diazafulvene to ketene has been reported by Rohr and his co-workers'.

The first cycloaddition of fulvene to sulfene will be presented as follows: The treatment of the iminium salt 2a and methane sulfonyl chloride with triethylamine gave the $[6+2]$ cycloadduct $7a$ whose structure was confirmed to be $2,3-di$ **hydro-3-morpholinodipyrrolo[l,2-b]isothiazole-l,-dioxide** on the basis of the spectral data shown in Table 1. The yield of $7a$ was depended upon the reaction temperature. Similarly 1b reacted with sulfene to give the corresponding $[6+2]$ cycloadduct 7b. ro-3-morpholinodipyrrolo[1,2-b]isothiazole-1,1-dioxide on the base

ctral data shown in Table 1. The yield of $\frac{7a}{2a}$ was depended upon th

perature. Similarly 1b reacted with sulfene to give the correspon

loadduct <u></u>

With asymmetrically substituted sulfene, phenylsulfene, la afforded the stereospecific [6+2] cycloadduct 8a. This compound 8a should be either cis or trans isomer, while the coupling constant of 7.0 Hz between two the methine protons at 1- and 2-position failed to determine the geometrical structure.

Although the unisolated $[6+2]$ cycloadducts $6'$ from the reactions with phenylketene readily eliminated morpholine to yield the deaminated products **6** as described above, the deamination reaction from the [6+2] cycloadducts with sulfenes *(L* and **8)** could not occur even on heating or treatment with base. An inspection using Dreiding model showed that the eliminating groups in **6'** are

located in nearly eclipsed position which favors to promote the cis elimination, whereas showed the difficulty of the deamination from $\frac{7}{5}$ and $\frac{8}{5}$ with the eliminating groups in staggered positions.

The low yield of the [6+2] cycloadducts from benzoazafulvene lb would owe to the tendency to dimerization under the reaction conditions. In fact, a considerable amount of the dimer of lb was isolated in almost all cases.

Table 1. The Reactions of $\underline{1}$ with Isocyanate, Ketenes and Sulfenes.

Table 1. Continued

	6b 113-115 16 -50 °C			1700 $vC=0$	2.08(3H,s,CH ₃), 7.00(1H,s,-CH=), 6.80-7.80 (9H, m, indolyl and phenyl protons)	259
With Sulfenes						
7a	$133 - 135$ 44 R.T.		$83 - 70^{\circ}$ C	vS0 ₂	1330 2.51, 3.65(each 4H,m,CH ₂), 3.85(2H,m,SO ₂ CH ₂), 242 4.64(1H,dd,J=6.0, 7.0Hz,CH), 6.04, 6.32, 6.85 (each lH,m,pyrrolyl protons)	
7b	156–158 decomp.	36 20	80° C R.T.	1330	2.32(3H,s,CH ₃), 2.59, 3.79(each 4H,m,CH ₂), 950 2 3.89, 3.96(each 1H,m,SO ₂ CH ₂), 4.99(1H,dd, J=5.0, 8.0Hz, CH), 7.26-7.86(4H, m, indolyl protons)	302
8a	163-164	- 14 88	R.T. -40° C	1340 v50 ₂	2.57, 3.63(each 4H,m,CH ₂), 4.94(1H,d,J=7.0Hz, 318 CH-N), 5.15(1H, d, J=7.0Hz, CHPh), 6.15, 6.48, 7.03(each lH,m,pyrrolyl protons), 7.44(5H,m, phenyl protons)	

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