## PL 1

## A NEW SYNTHESIS OF 5-NITROFURYLETHYLENE COMPOUNDS

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The methods of synthesis known up to this date of 5-nitrofurylethylene derivatives give possibilities of preparing only some types of these compounds. The new found ways in this area enabled to synthetise new type of 5-nitrofurylethylene compounds.

The first way is based on the use of the acidic hydrogens of some nitrofurfuryl derivatives in the condensation reactions with carbonyl compounds.

This method is suitable for proparing three and foursubstituted 5-nitrofurylethylene derivatives. The modified Horner-Witting reaction on suitable ketono was used in the case, when X was an electronodonating graup.

The second way is more common and give possibilities to prepare unknown compound of 5-nitrofurylethylene series. This method is based on the use of 5-nitrofurylvinylbromide (1) in the nucleophilic vinylic substitution reaction, where the reagents are compounds containing a free electron pair

nitrogen heterocycles and so on

The elimination of HBr goes by the action of strong bases on I and the 5-nitrofurylacethylene arises,

The use of 5-nitrofurylvinylamonium salts is more advantageous while the reaction proceeds in polar solvent and this compound reacts with more reagents than compound I.

The enamines of 5-nitrofurylethylene type are the potential source for preparation of derivatives of unknown 5-nitrofuryl-acetaldehyde

Pl 2

RINGSCHLUSSREAKTIONEN AN SELENOCARBONAMID--DERIVATIVEN ZU 1,3-SELENAZOLEN, 1,3.4-SELENADIAZINEN UND 1,3.4-SELENADIAZOLEN

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Selenazoles are the selenium analogues of thiazoles, and although the first derivatives of 1,3-selenazole were prepared by Hofmann as early os 1889, only a few publications on this subject have appeared in the literature until now. Hafmann was a student of Hantsch's and transferred his thiazole synthesis to the selenazoles. This most useful and versatile of all the thiazole syntheses is the reaction of an 2-halo ketone or aldehyde with a thioamide. For his purpose Hofmann used selenaurca instead of thioamide. Even nowadays this method is still the most usual one. It permits a wide variety by choosing suitable reaction partners as to be seen in formula 1.

The selenoureas, needed for condensation, can usually be prepared by addition of hydrogen selenide to a solution of the corresponding eyonomides or carbodinimides. They can be less dangerously and quite smoothly synthesized according to Douglass via the isoselenocyanotes by the alkaline hydrolysis of the corresponding ocyl-substituted selenoureas (formula 2).

2-Hydrazinoselenazoles have been prepared by the use of sclenosemicarbazide as selenacarbanamide. Starting from ace-tone selenacemicarbazide an prepare the 2-isopropylidenehydrazinoselenazoles in a smooth reaction by condensation with a-halacarbanyl compounds. Their careful addic hydrolysis leads to the free 2-hydrazino-selenazoles (formula 3).

$$\begin{array}{c} \begin{array}{c} \text{Hol} \\ \\ \text{C}_0 \end{array} + \begin{array}{c} \text{Se} \\ \text{NN-N} = \\ \text{Me} \end{array} \begin{array}{c} \text{NH-N} = \\ \text{Me} \end{array} \begin{array}{c} \text{NH-N} = \\ \text{NH-N} = \\ \text{NH-NH-2} \end{array} \end{array}$$

On the other hand, the  $\alpha$ -holocarbonyl component affers the possibility of variation too. For instance,  $\alpha$ -holocarboxylic acids, or rather their esters, form derivatives of 4-oxoselenazoline (formula 4).

First investigations on the reactivity of the 1,3-selenazoles were undertaken by Hoginiwa. He came to the conclusion that the 5-position of selenazoles is slightly reactive towards electrophilic substitution. Thi3 reactivity is still further increased by substituents in the 2-position, which can exert the + M-effect. We found during our investigations multiple intration depending on the conditions of the reaction and the substrate, whereas Haginiwa only described the formation of the 5-intro derivatives. Thus, the 2-benzamino-4-aryl(alkyl)selenazoles form the corresponding 5-nitro derivatives under mild conditions using the nitrate/sulfuric acid method. The use of cold mixed nitric and sulfuric acids also effects the phenyl groups which may be present, leading to dinitro or trinitro compounds (formula 5).

The bromination is not as complicated as the nitration. Bromination was carried out in several solvents and with various amounts of bromine. In spite of the great variation in conditions, monobrome derivatives containing the bromine atom in the 5-position are always formed (formula 6).

The high reactivity of the 5-position in 1,3-selenazoles towards electrophilic substitution was also to be observed on any coupling. By reacting molar quantities of an equeous solution of a diazonium solt with an ethanolic solution of a 2-arylamino-selenazole, for instance, the corresponding 2-arylamino-5-azoselenazols are formed in a smooth reaction (formula 7).

If the free selenosemicarbazide is used instead of selenosemicarbazones for the condensation with 2-halocarbanyl compounds, as mentioned above, the reaction will be quite different. The sole final product that can be isolated in neutral as well as acidic solution is the 2-p-chloroanilino-5-phenyl-1,3,4-thiadiazine, as illustrated by the 4-p-chlorophenylselenosemicarbazide. In the mother liquor there are, however, traces of the 2hydrozonoselenozoline. The third possible isomer, the 3-amino--2-iminoselenozoline derivative, could not be detected in any case (formula 8).

Starting from selenosemicarbazides one can also arrive at the third substance type, the 1,3.4-selenadiazoles. The first compounds of this type were obtained by Stollé and Gutmann in 1904 on reaction of N.N-diacytyletazines with phosphorus pentoselenide. Lalezari and Shafice synthesized 2-amino-1,3.4-selenadiazoles from selenosemicarbazide and carboxytic acids in the presence of phosphorus oxychloride. They obtained the free 2-amino compounds by hydrolysing the initially formed 2-ocylamino derivatives (formula 9).

$$2R \cdot C_{n}^{OH} + \bigvee_{H_{2}H}^{Se_{n}/H_{2}} \underbrace{(POCl_{3})}_{H-N} \underbrace{R \setminus A_{n-N}^{Se_{n}/H_{2}}}_{N+N}$$

$$(9)$$

We found in connection with the synthesis of substituted selenosemicarbazides than the cyclisation of the acyl derivatives under suitable conditions is a generally applicable method for obtaining the 2-amino-1,34-selenadiozoles. The 2-arylamio-1,34-selenadiazoles could be obtained in the same way by cyclisation of the 1-acyl-4-arylselenasemicarbazides (formula 10).