Transguanylation and Formation of 2-Arninothiazoline from *S-* **(2-Aminoethy1)isothiourea**

By **MICHAEL** D. **HALLAS, PATSY BOSTICK** REED, and R. **BRUCE MARTIN*** *(Chemistry Department, University of Virginia, Charlottesville, Virginia 22901)*

Summary The tripartite pH-rate profile obtained for disappearance of *S*-(2-aminoethyl)isothiourea indicates that a tetrahedral carbon addition intermediate occurs in the transformations of this system.

s- **(2-AMINOETHYL) ISOTHIOUREA (I)** is believed to provide radiation resistance in organisms by means of an intramolecular transguanylation reaction to yield Z-mercaptoethylguanidine **(11),** which has **a** free SH group. In addition

to (11), which is produced in neutral aqueous solutions, dilute acid solutions of **(I)** yield 2-aminothiazoline **(111)** and ammonia as the main products.1 From the results of

quantitative ion-exchange and SH content studies at $0.09M (I)^1$ it can be calculated that (II) and (III) are produced in equal amounts at *ca.* pH 4.2. Due to the appearance of these two products from one starting material, it has been suggested that a cyclic tetrahedral carbon intermediate is formed.¹ Since the possible intermediate has an SH group and three amino-nitrogen atoms joined to the tetrahedral carbon, there is opportunity for a mechanistic study of a tetrahedral carbon intermediate with groupings different from, though related to, those studied previously. Related reactions, the mechanisms of which have been studied, include transacetylation from sulphur to nitrogen in S-acetylmercaptoethylamine,² hydrolysis of thioimidate esters to give thiolesters,³ and the methoxy-aminolysis of methyl thiolformate.* In these three cases a diffusioncontrolled proton transfer reaction is rate determining under some conditions.

The occurrence of a tetrahedral addition intermediate in the transformations of (I) is indicated by the tripartite pH-rate profile shown in the Figure. The profile consists of two straight-line portions of unit slope joined by a bridging region. Rates of disappearance of *S*-(2-aminoethyl)isothiourea, 2HBr (Calbiochem) were followed at 235 nm. Replacement of Br⁻ by the less absorbing BF_4^- did not affect the results. Significant differences in molar extinction coefficients exist at 235 nm where ϵ is 1700 for (I) 500 for **(11),** and 2900 for **(111).** When **(11)** and **(111)** are produced in equal amounts from **(I),** no net change in absorption appears and accurate rate constants are difficult to determine in this region near pH **4-2. A** solution originally containing (I) displays an increase in absorption below pH **4-2** and a decrease above, demonstrating the transformation from **(111)** to **(11)** as the product as the pH increases through the bridging region shown in the Figure. Production of the two products in equimolar amounts occurs near pH **4.2,** consistent with the estimate made from the earlier study, performed at more than 200 times greater concentration. Reverse reactions starting with **(11)** or **(111)** and even an excess of NH, were not detected under

the conditions of this study. Hydrolysis to 2-mercaptoethylamine and dicyandiamide does not occur under the conditions used.1 As indicated by a weak absorption at **265** nm, a minor amount of disulphide formed in some **of** the final, more basic solutions.

FIGURE. *Logarithm of observed first order rate constants extrapolated to zero buffer concentration in min-1* versus **pH** *for dis* a *ppearance of* (I) a *at* 25.0° *and* 0.2M *ionic strength controlled with* **KCl**.

Analysis of the dependence of the reaction rate on buffer concentration reveals that the disappearance of (I) is general base catalysed with respect to free amine(AH+). For 2-(morpholino) ethanesulphonic acid (IV) buffer (p $K_{\mathbf{a}}$ **6.16)** from **0-01-0-20 M** concentration, studies at five **pH**

values where the fraction of buffer in the basic form varied from $0.2-0.8$ give k_{obs} $(H^+)/(B) = (3.6 \pm 0.3) \times 10^{-6}$ min-1. Similar values within a factor **of** *2.5* are obtained for acetic, formic, and chloroacetic acid buffers indicating a low Brønsted exponent in this formulation. This result suggests that hydroxide ion catalysis would not become important until pH > **10.** At a given pH, the higher buffer concentrations yield lower final absorbances, indicating that they promote formation of (11) over (111). At a concentration of 0.01 *M* phosphate buffer is about fifty times more effective as a catalyst than buffer (IV) at pH **6.1-6.7.**

The mechanism shown is the simplest that yields a rate expression which accommodates the tripartite pH-rate profile of the Figure and general base catalysis with respect to free amine. By assuming that the tetrahedral carbon addition intermediate, $(CH⁺)$, exists in a low, steady-state concentration, we obtain equation **(1)** for the observed

$$
k_{\rm obs} = \frac{K_{\rm a}k_1}{\rm (H^+)} \times \frac{k_3\rm(H^+) + \Sigma k_4\rm(B)}{k_2 + k_3\rm(H^+) + \Sigma k_4\rm(B)}\tag{1}
$$

first order rate constant for disappearance of **(I).** Due to the rapidity of the reaction at high pH, the acidity constant K_a cannot be measured, but is less than 10^{-8} . The points of the Figure are fitted with the curve of equation **(1)** drawn with $K_a k_1 = 1.6 \times 10^{-6} \text{ min}^{-1}$, $k_3/k_2 = 360 \text{ m}^{-1}$, and $k_4(H_2O)/k_2 = 0.020$, where water is the only base of consequence. In the case of buffer (IV) $k_4/k_2 = 2.3 \text{ M}^{-1}$. An alternative formulation with general acid catalysis of a neutral tetrahedral carbon addition intermediate $(HB + C)$ is possible. In either case, the amine expulsion reactions k_2 and k_3 appear to be extremely rapid bond-breaking steps.

(Received, Aupst **9th, 1971;** *Corn.* **1387.)**

- ¹ J. X. Khym, R. Shapira, and D. G. Doherty, *J. Amer. Chem. Soc.*, 1957, **79**, 5663.
² R. E. Barnett and W. P. Jencks, *J. Amer. Chem. Soc.*, 1969, **91**, 2358; 1968, **90**, 4199.
-
- K. K. Chaturvedi and G. L. Schmir, *J. Amer. Chem. Soc.,* **1969, 91, 737.**
- G. **31.** Blackburn, *Chem. Comm.,* **1970, 249.**