

RECOIL-ACTIVATED AND THERMAL EXCHANGE REACTIONS BETWEEN SULFUR-35 AND CARBON DISULFIDE

Sir:

Two interesting exchange reactions have been observed in research on the preparation of S^{35} -tagged CS_2 (CSS*). Each reaction presents an interesting chemical phenomenon, namely, the exchange of a free sulfur atom or ion with one bound in the CS_2 molecule, and each is applicable to the preparation of CSS*. Such *atomic* exchange reactions, involving energetic covalent bonds, have been found in the past to be much slower.

The activation energy in the first of these reactions is supplied by the specific nuclear process which results in formation of S^{35} . Two experiments have been performed to date, each utilizing the *n,p* reaction on Cl^{35} to make the 87-day S^{35} . In the first experiment, a solution of CS_2 in CCl_4 (10 volume per cent. CS_2) was placed in the stray neutron field near the Massachusetts Institute of Technology cyclotron for one month. The total S^{35} activity and that present as CS_2 were assayed by Curium analysis. For the CS_2 analysis, exhaustive extraction with Na_2CO_3 solution was followed by distillation of the mixture to remove the other S^{35} -containing compounds ($CSCl_2$ etc. No effort was made to separate CCl_4 and CS_2). About 50% of the S^{35} formed in compounds not volatile below room temperature was present as CSS*. In the second experiment, a solution of one gram of C_2Cl_6 in 1 ml. of CS_2 was sealed in a quartz vial for a thirty-day bombardment in the Oak Ridge pile, and assayed as described above, except that upon receipt, the sample was kept frozen until aliquoted for total S^{35} analysis to avoid loss of the more volatile compounds (of $BaCS_3$). In this case, 12% was recovered as CSS*. The lower value in the second experiment may be attributed (a) to the precaution taken to recover volatile compounds, and (b) to the greater variety and number of radiation-induced side reactions possible in the higher neutron flux of the pile. The specific activity of S^{35} as CSS* in the Oak Ridge sample attained a value of greater than one millicurie per gram.

The second reaction, now being studied, is the exchange of sulfide ion in aqueous solution with CS_2 as a separate phase. The reaction proceeds through sulfide exchange with thiocarbonate ion (CS_3^-), and like the electron transfer reactions of thallium¹ and iron² recently reported appears to be catalyzed by precipitation (of $BaCS_3$). On the other hand, when the CS_3^- is decomposed with acid and the resulting CS_2 extracted with CCl_4 and analyzed, this exchange shows a half-time of about forty minutes (sulfide concentration about 0.5 M, thiocarbonate about 0.15 M, pH 9.5, 30°). Investigation of the kinetics of this reaction continues.

(1) R. J. Prestwood and A. C. Wahl, *THIS JOURNAL*, **70**, 880 (1948).

(2) L. Van Alten and C. N. Rice, *ibid.*, **70**, 883 (1948).

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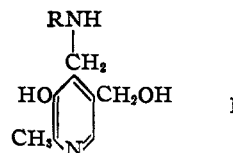
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PYRIDOXYLAMINES

Sir:

Pyridoxal has been reductively coupled with certain amines, including several pressor amines, to give compounds of structure I. For example,



β -phenylethylamine, tyramine, tryptamine, isobutylamine, histamine (amines derived from naturally occurring amino acids), as well as benzylamine reacted with pyridoxal to give yellow Schiff bases, which were hydrogenated over a platinum catalyst to give pyridoxyl- β -phenylethylamine dihydrochloride, II (m. p. 227-228°, dec.), pyridoxyltyramine dihydrochloride, III (m. p. 238-239°, dec.), pyridoxyltryptamine hydrochloride, IV (m. p. 222-223°, dec.), pyridoxylisobutylamine hydrochloride, V (m. p. 204-205°, dec.), pyridoxylhistamine dihydrochloride, VI (m. p. 236-237°, dec.), and pyridoxylbenzylamine dihydrochloride, VII (m. p. 220-221°, dec.). These new compounds as well as the intermediary Schiff bases were also analytically characterized.

These pyridoxylamines, which are derivatives of both pyridoxine and the pressor amines, are being studied for vitamin B₆ activity and for pressor activity.

The tests of these compounds for vitamin B₆ activity in deficient rats were made by Dr. Gladys Emerson and Miss Elizabeth Wurtz of the Merck Institute for Therapeutic Research, who have found that compounds II, III, IV and VII show activities which range between 50 and 100% of the activity of a molar equivalent of pyridoxine. Such high biological activity for these new compounds is in contrast to the low activity which has been found for previous structural modifications of the vitamin B₆ group.¹

(1) Unna, *Proc. Soc. Exptl. Biol. Med.*, **43**, 122 (1940); Harris and Wilson, *THIS JOURNAL*, **63**, 2526 (1941); Harris, *ibid.*, **63**, 3363 (1941).

Extensions of these chemical and biological studies will be detailed later.

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RECEIVED MARCH 20, 1948

EVIDENCE FOR THE INVOLVEMENT OF GLUTATHIONE IN THE MECHANISM OF PENICILLIN ACTION

Sir:

Several authors have suggested the involvement of —SH groups in the antibacterial action of penicillin (see review¹). The similarity in the molecular structure of glutathione and of penicillin^{2,3} suggests the possible involvement of glutathione in the antibiotic action of penicillin. The following experiments (supported partly by the Cutter Laboratories, Berkeley, California) bear on this question.

When standard penicillin assay plates are flooded with a 1% solution of 2,6-dichlorophenol-indophenol in a saturated aqueous solution of sodium bicarbonate the inhibition zones promptly stain intensely blue, and are sharply delineated from the faintly bluish uninhibited background by a narrow colorless rim that locates the ring of enhanced growth that circumscribes each zone. Similar patterns obtain on plates pretreated for five minutes with acetone, which blocks —SH groups from cysteine but not those from glutathione.⁴ However, if —SH groups of glutathione are blocked by flooding the plates for ten minutes with a 10% solution of formaldehyde in saturated sodium bicarbonate the 2,6-dichlorophenol-indophenol is no longer reduced to the colorless form in the ring of enhanced growth, which now stains deep blue.

The reducing activity in the regions of enhanced growth may be strikingly revealed also by flooding plates with a 0.5% aqueous solution of 2,3,5-triphenyltetrazolium chloride, whereupon these regions become intensely red, while the zones of inhibition remain uncolored. Pretreatment of the plates with 10% formaldehyde blocks this reaction. When such plates are subsequently flooded with the tetrazolium reagent, the red color fails to develop, except at the extreme outer margin of the ring of enhanced growth where it is very faint.

Such simple experiments do not themselves afford unequivocal proof of the participation of glutathione in the mechanism of penicillin action. However, it is generally assumed that —SH groups are involved. Our results indicate that some of these —SH groups are less reactive than those of cysteine, and in view of the work on the role of

glutamine revealed by Gale and Taylor^{5,6,7} it seems reasonable to deduce the involvement of glutathione.

(5) E. F. Gale and E. S. Taylor, *Nature*, **158**, 676 (1946).

(6) E. F. Gale and E. S. Taylor, *J. Gen. Microbiol.*, **1**, 314 (1947).

(7) E. F. Gale, *Nature*, **160**, 407 (1947).

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IMPROVED ION EXCHANGE METHOD FOR SEPARATING RARE EARTHS IN MACRO QUANTITIES¹

Sir:

Previous communications from this laboratory² described ion exchange methods by which rare earths were separated from one another in kilogram quantities. The process consisted essentially of absorbing the mixed rare earths on the top of long columns of commercial IR-100 Amberlite resin, in the acid cycle, and then eluting by means of citric acid solutions whose pH had been adjusted to the required value by the addition of ammonium hydroxide. While these processes represented an enormous saving, in man-hours required per gram of pure rare earth produced, over the old processes of fractional crystallization, etc., they were not ideal in the sense that when a mixture of rare earths was present, shapes of the elutions bands were such that there was a slight trailing of the preceding rare earth across the main band of the following one. This cut down the amount of pure rare earth obtained from any one pass of the column and frequently resulted in the necessity of recycling considerable quantities of the material.

Considerable work has been done in this Laboratory concerning the nature of the separation process. Good spectroscopic evidence has been obtained that at least four complexes of the rare earths with citrate solution exist and that each of these in turn becomes important as the pH range and citric acid concentrations are changed. Recently, it has been found that separation of the rare earths in large amounts can be markedly increased by eluting with a 0.1% citric acid solution in the pH range between 5.0 and 5.5. Under these conditions both the front and rear edges of the elution band (amount of rare earth eluted per liter plotted against liters of the eluant passed through the column) are steep and the tops of the eluting bands are flat. Furthermore, the bands separate from each other until the front edge of the one rare earth band is riding on the rear edge of the preceding band. Increasing the length of the column beyond the limit necessary to do this does not separate the bands any further, so there is good evidence that the one rare earth is replacing the

(1) R. Pratt and J. Dufrenoy, *Bact. Rev.*, **12**, 79 (1948).

(2) E. Fischer, *Science*, **106**, 146 (1947).

(3) R. Pratt and J. Dufrenoy, *J. Bact.*, in press (1948).

(4) L. Genevois and P. Cayrol, *Enzymol.*, **6**, 352 (1939).

(1) This document is based in part on work performed under Contract No. W-7405 eng-82 for the Atomic Energy Project.

(2) Spedding, *et al.*, *THIS JOURNAL*, **69**, 2777, 2786, 2812 (1947).