TABLE I

Eu(II)-E	u(III) E	XCHANGE	RATE	S AT	39.4°,	$\mu = 2.0$
Total Eu(II) + Eu(III) concn., f	Eu(II) conen.,	Eu(III) conen., f	H+ concn., f	Cl - concn.,	Half- time, min- utes	k moles ~2-12- min1
0.0653	0.0244	0.0409	1.00	1.86	53	0.108
.0894	.0258	. 0636	1.00	1.82	40	. 107
.1055	. 0683	.0372	1.00	1.84	33	.108
.0677			1.00	.716	132	.108
. 0630			0.30	1.87	58	. 102

The europium used was of 99.9% purity and was loaned to us by Mrs. Ethel Terry McCoy to whom we express our sincere gratitude. We thank Professor Don M. Yost and Dr. David L. Douglas of the California Institute of Technology for the Eu152 activity.

DEPARTMENT OF CHEMISTRY University of California Los Angeles, California

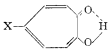
DALE J. MEIER CLIFFORD S. GARNER

RECEIVED MARCH 5, 1951

ON p-AMINOTROPOLONE

Sir:

Very recently, Dewar¹ not only reported a molecular orbital calculation² of tropolone (I, X = H)



but also predicted that paminotropolone (II, X NH₂), which should be obtainable by the reduction of its

azo-compound, may show interesting pharmacological properties as a precursor in vivo of p-aminobenzoic acid.

We have already reported on the syntheses of I³ (independent of other three laboratories⁴), α -aminohinokitiol, 5 i.e., p-amino-m-isopropyltropolone (III), o-bromo-p-aminotropolone⁶ (IV) and other various derivatives^{3,6} of I. We have also recently synthesized II, a brief account of which is given here.

Catalytic reduction of phenylazotropolone³ (m.p. 161–161.5°; anal. Calcd. for $C_{13}H_{10}N_2O_2$: N, 12.39. Found: N, 12.58) or p-tolylazotropolone³ (m.p. 202.5–203°; anal. Calcd. for $C_{14}H_{12}N_2O_2$: N, 11.66. Found: N, 11.42) with Adams catalyst, or their reduction with sodium hydrosulfite, yields yellow scaly crystals (II), m.p. 177-177.5°. *Anal.* Calcd. for C₇H₇O₂N: C, 61.32; H, 5.14; N, 10.21. Found: C, 61.09; H, 5.32; N, 10.02; yield, 30-40%. II is also obtained by the similar methods from p-nitrosotropolone⁷ (X = NO), charring at $180^{\circ} \cdot (Anal. \text{ Calcd. for } C_7H_5O_3N: N, 9.27.$ Found: N, 9.10) in better yield (70-80%). II is

- (1) M. J. S. Dewar, Nature, 166, 790 (1950).
- (2) Similar colculation and measurement of dipole moments of I and its related compounds have already been reported; Y. Kurita, T. Nozoe and M. Kubo, J. Chem. Soc. Japan, 71, 543 (1950); Bull. Chem. Soc. Japan, in press.
- (3) T. Nozoe, S. Seto, Y. Kitahara, M. Kunori and Y. Nakayama, Proc. Japan Acad., 26, (7) 38 (1950); presented at the Annual Meeting of the Chemical Society of Japan in Kyoto, April 2, 1950.
- (4) W. von E. Doering and L. H. Knox, This Journal, 72, 2305 (1950); J. W. Cook and A. R. Gibb, Chemistry & Industry, 427 (1950); R. D. Haworth and J. D. Hobson, ibid., 441 (1950).
- (5) T. Nozoe and E. Sebe, Proc. Japan Acad., 26, (9) 45 (1950); T. Nozoe, S. Ebine, S. Itô and A. Konishi, ibid., 27, 10 (1951).
- (6) T. Nozoe, Y. Kitahara, K. Yamane and A. Yoshikoshi, ibid., 27, 18 (1951); T. Nozoe, S. Seto, T. Ikemi and T. Arai, ibid., 27, 24 (1951). (7) T. Nozoe and S. Seto, to be published soon.

amphoteric and its chemical behaviors are closely analogous to III and IV. Copper complex salt: greenish yellow microcrystals. Picrate: yellow scaly crystals, m.p. $225-226^{\circ}$ (dec.); anal. Calcd. for $C_{13}H_{10}O_{9}N_{4}$: N, 15.30. Found: N, 15.42. Diacetate: colorless scaly crystals, m.p. 180.5-181°; anal. Calcd. for $C_{11}H_{11}O_4N$: N, 6.33. Found: N, 6.02.

Application of the Sandmeyer reaction to II yields the following halogen compounds. p-Bromotropolone (V, X = Br); m.p. 189–190°, alone or in admixture with β -bromotropolone, β obtained as a by-product during the synthesis of I and the position of its bromine atom was later established to be at para,6 so that the amino group in II is also clearly in the para position. p-Chlorotropolone: orange needles, m.p. 147-149°. p-Iodotropolone: orange needles, m.p. 169-170°.

Details of the results of our studies will be reported shortly. The effects of I, II, and some of their allied compounds on Yoshida sarcoma have already been published.8

We are deeply indebted to Dr. R. Majima (Emeritus Professor of this University) for his unfailing encouragement and to the Ministry of Education of Japan for the financial support.

(8) S. Katsura, K. Satô, K. Akaishi, T. Nozoe, et al., Proc. Japan Acad., 27, 31, 36 (1951).

CHEMICAL INSTITUTE FACULTY OF SCIENCE Tôhoku University Sendai, Japan

TETSUO NOZOE Sutichi Seto SEIJI EBINE Shō Itō

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ON A NEW TYPE OF AROMATIZATION BY THE DIAZOTIZATION OF O-AMINOTROPOLONE RIVATIVES

Sir:

Tropolone and its allied compounds, the tropoloids, possess a fair degree of aromatic properties in spite of the unsaturated, seven-membered ring structure. On the other hand, it has also been established that these compounds, when heated with highly concentrated alkalies, undergo benzylic rearrangement to carboxylic acids of benzenoid series. According to Raistrick¹ and Dewar,² tropolones can be taken as precursors, in vivo, of natural benzenoid carboxylic acids, and recently Robinson³ has also discussed on the assumption of a biogenetic relation between the tropolones and various alkaloids or anthocyanines. These certainly seem attractive suggestions but they must be confirmed by future experimental evidence. In this connection, studies on the aromatization of tropoloid series become of great significance.

Previously, we had encountered a notable fact that when o',p-dinitro-m-isopropyltropolone is heated with 50% aqueous ethanol for ten minutes at 100°, or with absolute methanol, ethanol or isopropyl alcohol at 50–60° for a few minutes, it undergoes rearrangement to form o', p-dinitro-m-cumic acid or its respective esters quantitatively.4

- H. Raistrick, Proc. Roy. Soc. (London), A199, 141 (1949).
 M. J. S. Dewar, Nature, 166, 790 (1950).
 R. Robinson, ibid., 166, 930 (1950).

- (4) T. Nozoe, Science of Drugs, 3, 171 (1949) [English translation, Sci. Rep. Tohoku Univ., I, 34, in press (1951)].

In a recent experiment, o'-amino- (I, m.p. 99°) and o-aminohinokitiol (II, m.p. 121°) were submitted to the Sandmeyer reaction, in order to obtain various structurally identified o'- and o-halogenohinokitiols. Unexpectedly, however, p- and o-isopropylsalicylic acid derivatives were obtained in a good yield besides the objective compounds, under certain conditions.

In addition to the o'- and o-chloro and bromohinokitiols, I also yielded colorless scales (III), m.p. 95–96°, and II yielded colorless needles (IV), m.p. 122–123°, both in 25–30% yield. In either case, no iodo derivatives were obtained, differing from p-aminohinokitiol (m.p. 131°), and instead III and IV are generally obtained in a better yield. It was also found that the same compounds were obtained in a good (50–60%) yield by heating the solution of diazonium salt of I and II with diluted sulfuric acid.

III showed no depression of the melting point when fused with a pure specimen of p-isopropylsalicylic acid. IV gives the same reddish violet coloration as III with ferric chloride in methanol solution. Anal. Calcd. for C₁₀H₁₂O₃: C, 66.65; H, 6.71. Found: C, 66.17; H, 6.74. The phenolic substance obtained by its decarboxylation gives phenoxyacetic acid derivative (V), m.p. 64°. Anal. Calcd. for C₁₁H₁₄O₃: C, 68.04; H, 7.21. Found: C, 67.76; H, 7.70. V showed no depression of the melting point when fused with the 3-isopropylphenoxyacetic acid, m.p. 64°, derived from the decarboxylation product of III.

It has been assumed, from the experimental facts, that the mechanism of their rearrangement might be as shown in the scheme. It is naturally possible to assume formation of a carbonium ion as an intermediate during the decomposition of respective diazonium cations (VI and VII). It is interesting to note, that o-, m- and p-cumic acid derivatives are easily formed from the same m-isopropyltropolone.

CHEMICAL INSTITUTE FACULTY OF SCIENCE TÔHOKU UNIVERSITY SENDAI, JAPAN

Tetsuo Nozoe Yoshio Kitahara Kôzô Doi

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THE DEGRADATION OF COLCHICINE TO OCTA-HYDRODEMETHOXYDESOXYDESACETAMIDO-COLCHICINE

Sir:

The synthesis of *dl*-colchinol methyl ether¹ has left ring C as the only part of the colchicine molecule for which absolute structural evidence is lacking. The tropolone formulation proposed by Dewar² has received support in recent publications³; however, direct degradative and synthetic evidence as to the seven-membered nature of ring C and the positions of the oxygen functions in this ring would be desirable. We wish to report the degradation of colchicine (I) to octahydrodemethoxydesoxydesacetamidocolchicine (II). Since this latter compound contains the carbon skeleton of colchicine intact, its synthesis, which appears feasible, would provide definitive proof of the size of ring C.

$$\begin{array}{c} CH_{3}O \\ CH_{3}O \\ CH_{3}O \\ CH_{3}O \end{array} \longrightarrow \begin{array}{c} CH_{3}O \\ CH_{3}O \\ CH_{3}O \\ \end{array}$$

Colchicine (m.p. 154-155°), on heating with methanolic dimethylamine gave N,N-dimethylami-

nocolchicide (replacement of methoxyl by dimethamino) [m.p. $174-176^{\circ 4}$; $[\alpha]^{25}$ D $+69.4^{\circ}$ (c, 1.03, ethanol); Anal. Calcd. for $C_{23}H_{28}N_2O_5$: C, 67.0; H, 6.8; N, 6.8; OCH₃, 22.6. Found: C, 66.9; H, 6.9; N, 7.0; OCH₃, 22.2] which formed a picrate [m.p. $186-188^{\circ}$; $[\alpha]^{25}$ D $+171^{\circ}$ (c, 1.08, chloroform); Anal. Calcd. for $C_{29}H_{31}N_5O_{12}$: C, 54.3; H, 4.9; OCH₃, 14.5. Found: C, 54.3; H, 5.0; OCH₃, 14.3] and on catalytic hydrogenation in glacial acetic acid was converted to the ketone, tetrahydrodemethoxycolchicine [m.p. $143-144^{\circ}$; $[\alpha]^{25}$ D -174° (c, 1.11, ethanol); Anal. Calcd. for $C_{21}H_{27}NO_5$: C, 67.5; H, 7.3; N, 3.8; OCH₃, 24.9. Found: C, 67.5; H, 7.3; N, 3.8; OCH₃, 24.9. The latter formed a soluble bisul-

fite addition compound and on further hydrogenation absorbed one mole of hydrogen to yield the carbinol, hexahydrodemethoxycolchicine [m.p. $168-170^{\circ}$; $[\alpha]^{25}$ D -166° (c, 1.01, ethanol); reported m.p. $171^{\circ 5}$ and $173^{\circ 6}$; Anal. Calcd. for

⁽⁵⁾ T. Nozoe, Y. Kitahara and K. Doi, Proc. Japan Acad., 27, in press (1951).

⁽⁶⁾ T. Nozoe and E. Sebe, ibid., 26, [9] 45 (1950).

⁽⁷⁾ O. Jacobseu, Ber., 11, 1061 (1878).

⁽¹⁾ H. Rapoport, A. R. Williams and M. E. Cisney, This Journal. 72, 3324 (1950).

⁽²⁾ M. J. S. Dewar, Nature, 155, 141, 479 (1945).

^{(3) (}a) H. R. V. Arnstein, D. S. Tarbell, G. P. Scott, and H. T. Huang, This Journal, **71**, 2448 (1949); (b) G. P. Scott and D. S. Tarbell, *ibid.*, **72**, 240 (1950).

⁽⁴⁾ This compound has been previously reported as melting at 204-206° [A. J. Ewins, J. N. Ashley and J. O. Harris, British Patent 577, 606 (1945)]; no other physical properties or analytical data were given. Sublimation, crystallization from various solvents, and chromatographic adsorption on alumina all indicated the homogeneity of our product and failed to alter its melting point.

⁽⁵⁾ K. Bursian, Ber., 71, 245 (1938).

⁽⁶⁾ A. D. Kemp and D. S. Tarbell, This Journal, 72, 243 (1950).