[CONTRIBUTION FROM THE KEDZIE CHEMICAL LABORATORY, MICHIGAN STATE UNIVERSITY, EAST LANSING, MICH.]

Two Unusual Reactions Encountered during the Devious Hydrolysis of 3,3-Dimethyl-1,1,2,2-tetracyanocyclopropane to the Corresponding Tetracarboxylic Acid¹

By HAROLD HART AND FILLMORE FREEMAN

RECEIVED OCTOBER 31, 1962

The structure of the first isolable alkaline hydrolysis product of 3,3-dimethyl-1,1,2,2-tetracyanocyclopropane (I) is confirmed as the acid-amide-imide II. Thermal decarboxylation of II proceeds with cleavage of the cyclopropane ring, and probably involves participation by the neighboring *cis*-carboxamido group; mechanisms which would allow decarboxylation without ring opening would require highly strained transition states. Two additional examples of decarboxylation of a cyclopropanecarboxylic acid with ring cleavage are presented. Compound II was converted in four steps and high yield by alternate methylation with diazomethane and hydrolysis with base to the corresponding cyclopropanetarcarboxylic acid (XIII). One of the hydrolytic steps (X \rightarrow XI) involves a remarkable transformation in which a five-membered imide ring between *cis* substituents is hydrolyzed on one side of the cyclopropane ring plane and formed on the opposite side.

Introduction

The discovery² that a crystalline product deposits rapidly when aqueous potassium iodide is added at room temperature to a solution of bromomalononitrile in acetone led to a rather general synthesis of tetracyanocyclopropanes³ which we propose be called the Wideqvist reaction.⁴ By using tetrahydrofuran as solvent, the reaction has been used to prepare tetracyanocyclo-



propane itself from formaldehyde⁵; and in another modification, alkylidene bismalononitriles were converted with bromine to monoalkyltetracyanocyclopropanes.^{6,7}

The assignment of the correct structure to I depended upon the isolation of certain of its hydrolysis products and their characterization by independent synthesis.⁸ These reactions are summarized in Scheme 1, using structures assigned by Ramberg and Wideqvist. Alkaline hydrolysis of I reportedly liberated one-half of the theoretical nitrogen as ammonia very rapidly; acidification gave a crystalline solid, m.p.



- Paper XII in a series on Cyclopropane Chemistry. For the previous paper, see H. Hart and R. A. Cipriani, J. Am. Chem. Soc., 84, 3697 (1962).
 L. Ramberg and S. Wideqvist, Arkiv. Kemi, Mineral. Geol., 12Å, No. 22, 12 (1937).
- (3) S. Wideqvist, *ibid.*, **20B**, No. 4, 8 (1945).
- (4) The equation is written with iodine as a product, but the initial inorganic product may well be iodine bromide.
- (5) R. M. Scribner, G. N. Sausen and W. W. Prichard, J. Org. Chem., 25, 1440 (1960).
- (6) R. P. Mariella and A. J. Roth, III, ibid., 22, 1130 (1957).
- (7) Our own investigations into the scope of the Wideqvist reaction will be described in a separate paper.
- (8) L. Ramberg and S. Wideqvist, Arkiv. Kemi, Mineral. Geol., 14B, No. 37, 13 (1941).

196–197° with CO₂ evolution, to which the acid-amideimide structure II⁹ was assigned. The same product (m.p. and comparison of the crystals in a polarizing microscope) was obtained from the previously known^{10,11} III. The alternate structure IIa was discarded primarily on the basis of titration data (>1 but <2



equivalents of base required for neutralization to phenolphthalein) and synthesis from III. Earlier, Birch, Gough and Kon¹¹ had isolated a presumably identical compound from III but assigned it structure IIb; this was also discarded⁸ on reasonable grounds (among others, its failure to lactonize). Hydrolysis of I or III with fuming hydrochloric acid gave a single product, assigned structure IV. Finally, not shown in Scheme 1, Ramberg and Wideqvist hydrolyzed I with concentrated ammonia which contained potassium hydroxide, and obtained on acidification (CO₂ evolved) a dibasic acid, m.p. 165°, originally thought⁸ to be impure *cis*-caronic acid, but in fact probably γ, γ -dimethylitaconic acid.¹²

In view of the simplicity and generality of the Wideqvist reaction as a synthesis of cyclopropanes, and considering also the current interest in polycyano compounds,¹³ it seemed worthwhile to re-examine the alkaline hydrolysis¹⁴ of I, not only to establish with modern methods the structure of the first product II, but also to complete the hydrolytic degradation of I. This paper describes a sequence by which I may be converted to the tetraacid in five steps, and discloses two novel reactions discovered along the way.

Results and Discussion

The Structure of II.—The yield of II from I was improved to over 80% by adding methanol to the aqueous potassium hydroxide. Potentiometric titration established that II was dibasic in accord with the assigned structure ($pK_{A^{I}}$ 2.45 and $pK_{A^{II}}$ 8.35). Absence of bands in the 4.4 μ region mitigated against structures

(9) 2,4 - Dioxo-5-carboxamido-6,6-dimethyl-3-azabicyclo [3.1.0]hexane-1carboxylic acid; because the systematic names of most compounds described in this paper are of this length, compounds will be referred to only by number in the text. Systematic names are given in the Experimental part.

- (10) I. Guareschi and E. Grande, Atti R. Accad. Torino, 34, 18/6 (June); Chem. Zentr., 70, II, 439 (1899).
- (11) S. F. Birch, W. H. Gough and G. A. R. Kon, J. Chem. Soc., 119, 1315 (1921).

(12) See ref. 6, footnote 9.

- (13) See particularly the series of ten papers from the du Pont group, J. Am. Chem. Soc., 80, 2775-2844 (1958), and subsequent papers.
 - (14) The acidic hy_rolysis to IV has not yet been reinvestigated.

Vol. 85

containing the cyano group.¹⁵ The n.m.r. spectrum in D_2O containing NaOD showed two singlets at 8.74 and 8.76 τ with equal areas (C-CH₃). In dimethyl sulfoxide, the n.m.r. showed singlets at -1.14 and 2.39 τ , relative areas 1:2, ascribed to the imide and amide hydrogens, respectively (succinimide and phthalimide, in the same solvent, show imide singlets at -1.00 and -1.13τ , respectively). These data are consistent only with the formulation of this product as II, the mechanistically unlikely alternative IIc being eliminated by the position of the methyl resonances in the n.m.r.

The amide group of II resisted reaction with nitrous acid or nitrosyl chloride. But prolonged alkaline hydrolysis led to the diacid V in over 80% yield; V was also obtained directly from I (in nearly quantitative



yield, if one allows for the small amount of II also isolated). Diazomethane converted V to the trimethyl derivative VI.¹⁶

When V was heated above its melting point, carbon dioxide was evolved.¹⁷ Analysis of the crystalline product (obtained in 42% yield) showed that two moles of carbon dioxide had been lost, but its melting point (166-167°) was different from that (120°) reported¹⁸ for caronimide. Furthermore, the infrared showed a carbon-carbon double bond (6.0μ) which was confirmed by a positive Baeyer test. The product is therefore undoubtedly isopropylidenesuccinimide (VII)19; indeed, mechanistic considerations suggest that this is the only plausible decarboxylation product, cleavage of the three-membered ring occurring with loss of the first mole of carbon dioxide. The imide carbonyl is prevented from participating in the decarboxylation because the resulting enolic intermediate would contain a [3.1.0] system with a double bond exocyclic to the threemembered ring. The normal reaction of a cis-1,2-cyclopropanedicarboxylic acid, *i.e.*, anhydride formation, is also mitigated against because of strain in the product,¹⁷ but the carbonyl of the *cis*-carboxyl is in a favorable position to assist the decarboxylation. In

(15) L. J. Bellamy, "Infrared Spectra of Complex Molecules," J. Wiley and Sons, Inc., New York, N. Y., 1958, p. 264.

(16) Imides have previously been converted to N-methyl derivatives with diazomethane; see G. Labruto, Gazz. chim. ital., 63, 2669 (1933).

(17) Formation of the anhydride apparently would result in excessive strain



Fused five-membered rings of this type are known, if the remaining bridging ring is larger, as in the formation of the diimide



See W. J. Middleton, R. E. Heckert, E. L. Little and C. G. Krespan, J. Am. Chem. Soc., **80**, 2783 (1958).

(18) K. N. Menon and J. L. Simonsen, J. Chem. Soc., 302 (1929).

(19) Curiously, this simple compound appears not to have been previously reported.



short, forces conspire against all normal paths and favor the one observed.

Two further examples (one presented now and one reserved for a later section of this paper) show that this novel decarboxylation is general and that ring opening occurs with the loss of the first mole of carbon dioxide. When II was heated for a short time just above its melting point, one mole of carbon dioxide was lost and a crystalline product, m.p. $240-241^{\circ}$, was isolated in 87% yield. Its n.m.r. spectrum in 10% sodium hydroxide showed the two allylic methyl groups (singlets at 7.72 and 8.14τ with equal areas) expected for VIII but at lower field than is permissible for VIIIa (cf. II, vide supra). The n.m.r. of VIII in dimethyl sulfoxide had



three equal singlets at -1.03 (imide), 2.27 and 2.80 τ ; the latter two are amide hydrogens, one of which is apparently hydrogen-bound to the imide carbonyl. The single C-H on the ring carbon holding the carbox-amido group is not visible in either solvent. The carbon-carbon double bond in VIII is also evident from its infrared spectrum (6.05 μ) and positive Baeyer test. Treatment of VIII with diazomethane gave IX which, on hydrolysis, gave two moles of nitrogen bases and iso-propylidenesuccinic acid.

Conversion of II to Methyl3,3-Dimethyl-1,1,2,2-tetracarboxylate (XIV).—Compound I was successfully converted, via II, to the tetracarboxylic acid XIII by the unusual route shown in Scheme 2. Alkaline hydrolysis of II or V proceeded only very slowly, presumably be-



cause the di- or tri-anions are only sluggishly attacked by base.

Compound II was therefore methylated with two moles of diazomethane to give a nearly quantitative yield of X which, on hydrolysis, gave in 90% yield a product corresponding to loss of one of the two methyl groups. The two obvious structures XIa and XIb are immediately eliminated because the compound was dibasic; the values of the pK_A 's (I, 2.65; II, 8.38) suggested a carboxyl and an imide (cf. II, vide supra), leaving XI as the only reasonable structure. The n.m.r. spectrum in D₂O containing NaOD showed two peaks, at 7.36 and 8.84 τ , with relative areas 1:2, assigned to the N-CH₃ and C(CH₃)₂, respectively.



When one considers the multiplicity of hydrolytic reactions which might occur when X is treated with strong base, the formation of XI in excellent yield as the only isolable product is truly remarkable. It clearly arises from opening of the imide ring to produce a carboxyl group and an N-methylamide, followed by reaction of the amide and ester functions intramolecularly on the opposite side of the three-membered ring. Although the mechanistic details are not known, it is possible to rationalize the sole production of XI as



Hydrogen bonding between the amide nitrogen and the imide carbonyl, as shown in Xa, would make that carbonyl the most likely site of attack by base. That such hydrogen bonding can be important has already been shown from the n.m.r. of VIII. Hydrogen bonding between the *cis*-amide and ester functions is less likely than that shown in Xa because it requires a sevenrather than six-membered ring. Similarly, the amide and ester are not free to react intramolecularly for reasons cited above.¹⁷ Because of solubility difficulties, it was not possible to obtain a satisfactory n.m.r. spectrum of X, but in a later paper we will present n.m.r. evidence for hydrogen bonding of the type postulated, in an analog of X. Following the opening of the imide ring (Xb) and equilibration to the carboxylate ion Xc, the amide function is freed from both constraints and reacts intramolecularly via an intermediate of the type Xd to furnish the dianion of XI which, because of the accumulated negative charge, is not susceptible to ready further hydrolysis.

As noted in the Experimental part, XI may also be prepared by the alkaline hydrolysis of N-methyl III; this implies that in the conversion of III to II shown in Scheme 1, the imide function of II need not necessarily arise from the imide function of III.

Decarboxylation of XI again led to opening of the three-membered ring, it being converted just above its melting point smoothly to XV.



Reaction of XV with diazomethane followed by hydrolysis allowed the isolation of methylamine (as its hydrochloride) in over 95% yield, thus adding confirmatory evidence to the structure assigned to XI. Isopropylidenesuccinic acid was also isolated from the hydrolysis.

Attempts to hydrolyze XI with alkali failed. With diazomethane, it was converted to XII, which was rapidly hydrolyzed to the tetracarboxylic acid XIII. Because of difficulties with purification, XIII was converted to its tetramethyl ester XIV for analysis. The n.m.r. spectrum of XIV is decisive regarding its structure, consisting of two sharp singlets at 6.42 (O-CH₃) and 8.56 τ (C-CH₃), relative areas 2:1.

The sequence of reactions just described allows the conversion of I to XIV in over-all 43% yield, despite the number of steps. The scheme is general for tetracyanocyclopropanes derived from ketones, as will be described in a separate paper.²⁰

Experimental²¹

2,4-Dioxo-5-carboxamido-6,6-dimethyl-3-azabicyclo[3.1.0]hexane-1-carboxylic Acid (II).--A solution of 2.8 g. (0.016 mole)of 3,3-dimethyl-1,1,2,2-tetracyanocyclopropane (I),² m.p. 208-209°, in 30 ml. of 25% aqueous potassium hydroxide and 40 ml. of methanol was refluxed for 3 hours, the alcohol removed (Rinco rotary evaporator) and the remaining liquid extracted continuously with ether for 4 hours to remove non-acidic materials. The aqueous solution was then acidified (congo red) with dilute hydrochloric acid and extracted continuously with ether for 24 hours. Removal of the solid from the extract by filtration, and recrystallization from water gave 2.81 g. (80.1%) of white crystals of II, m.p. (dec.) 196-197° (lit. value⁸ 196-197°). In order to be certain of its identity with the product described by Ramberg and Wideqvist,⁸ a sample was analyzed.

Anal. Calcd. for $C_{9}H_{10}N_{2}O_{6}$: C, 47.79; H, 4.46; N, 12.39. Found: C, 47.68; H, 4.49; N, 12.33.

Aqueous potentiometric titration gave a neutralization equivalent of 112.3 (theory 113.1) with pK_A^{I} 2.45 and pK_A^{II} 8.35. The infrared spectrum in Nujol had characteristic bands at 2.85, 2.95 (N—H), 5.50, 5.70, 5.85 μ (C=O) and no absorption from 4.40–4.45 (C=N); the n.m.r. in D₂O (containing NaOD) had singlets at 8.74 and 8.76 τ with equal areas and in dimethyl sulfoxide there were singlets at -1.14 and 2.39 τ , relative areas 1:2. Compound II was recovered in >90% yield from reactions with sodium nitrite and hydrochloric acid (0°, 1 hour) and with isoamyl nitrite, ethanol and sulfuric acid (0°, 12 hours).

suitoxide there were singlets at -1.14 and 2.397, relative areas
1:2. Compound II was recovered in >90% yield from reactions with sodium nitrite and hydrochloric acid (0°, 1 hour) and with isoamyl nitrite, ethanol and sulfuric acid (0°, 12 hours).
2.4-Dioxo-3.3-dimethyl-3-azabicyclo[3.1.0]hexane-1,5-dicarboxylic Acid (V). From II.—A solution of 1.13 g. (5 mmoles) of II in 30 ml. of 10% sodium hydroxide was refluxed for 3.25 hours. During reflux, the evolved ammonia was swept by a stream of nitrogen into standard hydrochloric acid, which, at the end of this period, indicated that nearly one mole of ammonia had been produced. The alkaline solution was acidified with

(20) H. Hart and F. Freeman, unpublished results.

(21) Melting points are uncorrected. Analyses are by Spang Microanalytical Laboratories, P. O. Box 1111, Ann Arbor, Mich. Nuclear magnetic resonance spectra were obtained on a Varian model A-60 or HR-60 instrument at 60 Mc. Band positions are in r-units (G. V. D. Tiers, J. Phys. Chem., 52, 1151 (1958)), and relative areas were obtained by electronic interration.

dilute hydrochloric acid and extracted continuously with ether for 36 hours. The ether extract was dried (sodium sulfate), evaporated, ethyl acetate added and, after treatment with Norit, the product was isolated and, arter treatment with acetate-pentane, yielding 0.92 g. (80.7%) of V, m.p. 168– 169° dec. Potentiometric titration gave pK_A^{II} 2.4, pK_A^{III} 5.15, pK_A^{III} 9.75. The n.m.r. spectrum in D₂O containing Na-OD bod circlet with acutal concert \$ 62 cond 2 containing Na-OD had singlets with equal areas at 8.62 and 8.68 τ .

Anal. Calcd. for C₃H₃NO₆: C, 47.58; H, 3.99; N, 6.16; neut. equiv., 75.7. Found: C, 47.36; H, 4.22; N, 6.05; neut. equiv., 75.1.

From I.—A solution of 1.70 g. (0.01 mole) of I in 40 ml. of 2 N potassium hydroxide was refluxed for 9 hours, acidified (congo red) with hydrochloric acid and extracted continuously with ether for 36 hours. Filtration of the ether layer gave 0.39 g. (17.2%) of II, m.p. 196–197° dec. after recrystallization from The ether solution was then dried over anhydrous sowater. dium sulfate, treated with Norit, and the solvent evaporated (Rinco). Recrystallization of the crude residue from ethyl ace-tate-pentane gave 1.88 g. (82.8%) of white crystals of V, m.p. and m.m.p. 167-168° dec. Methyl 2,4-Dioxo-3,6,6-trimethyl-3-azabicyclo[3.1.0]hexane-1,5-dicarboxylate (VI).—To a solution of 2.27 g. (0.01 mole) of V in 50 ml. of absolute methanol cooled in an ice-bath was added

an ethereal solution of diazomethane in small portions until gas evolution ceased and the solution acquired a pale yellow color. Solvent and excess reagent were evaporated, and the residue recrystallized from methanol, giving 2.49 g. (91.1%) of VI, m.p. 106-107°. Its infrared spectrum (CCl₄) showed carbonyl bands at 5.60, 5.75 and 5.85 μ and no N-H stretching band near 3μ.

Anal. Caled. for $C_{12}H_{16}NO_6\colon$ C, 53.52; H, 5.62; N, 5.20. Found: C, 53.47; H, 5.70; N, 5.16.

Decarboxylation of V.—A flask containing 2.27 g. (0.01 mole) of V was swept with a dry nitrogen stream, and immersed in an oil-bath at 168°, whereupon V melted and evolved carbon dioxide. After 1 minute the flask was removed from the oilbath and cooled to room temperature. The yellow residue was dissolved in methanol, treated with Norit, the solvent evaporated and the residue recrystallized from water, giving 0.59 g. (42.2%) of isopropylidenesuccinimide (VII), m.p. 167–168°. The compound gave a positive Baeyer test for unsaturation and showed bands at 2.95 (N—H), 5.65 and 5.85 (C=O) and 6.0 μ (C=C) in chloroform solution.

Anal. Calcd. for $C_7H_9NO_2$: C, 60.42; H, 6.52; N, 10.07. Found: C, 60.40; H, 6.60; N, 10.00. Decarboxylation of II.—The procedure was the same as that

becarboxylation of 11.—1 the procedure was the same as that just described for decarboxylation of V, except that the oil-bath temperature was 197°. From 2.26 g. (0.01 mole) of II there was obtained 1.58 g. (87.1%) of 3-isopropylidene-4-carboxamidosuccinimide (VIII), white needles from water, m.p. 240–241° dec. Its infrared spectrum had bands at 2.95 and 3.15μ (N—H), 5.70, 5.85, 5.96 μ (C=O) and 6.05 μ (C=C). The n.m.r. spectrum in dimethyl sulfoxide had singlets with equal areas at -1.03, 2.27 and 2.80 τ ; in 10% sodium hydroxide, the n.m.r. spectrum showed singlets with equal areas at 7.72 and 8.14 7.

Anal. Calcd. for $C_8H_{10}N_2O_5$: C, 52.74; H, 5.53; N, 15.37. Found: C, 53.06; H, 5.56; N, 15.27.

Preparation and Hydrolysis of 4-Carboxamido-3-isopro-pylidene-1-methylsuccinimide (IX).—Methylation of VIII with diazomethane (as in the preparation of VI) gave IX in 86.4%yield, m.p. 229-230° from methanol.

Anal. Calcd. for $C_9H_{12}N_2O_8;\ C,\,55.09;\ H,\,6.17;\ N,\,14.28$ Found: C, 55.29; H, 6.26; N, 14.19.

A mixture of 0.70 g. (3.5 mmoles) of IX and 30 ml. of 10%sodium hydroxide was refluxed for 5 hours during which time volatile products were swept by nitrogen into standard hydro-chloric acid. Titration after this time showed nearly two moles of volatile base had evolved. The alkaline solution was cooled, acidified with dilute hydrochloric acid and cooled over-

cooled, acidified with dilute hydrochloric acid and cooled over-night in the refrigerator. The solid was filtered and recrystal-lized from water, giving 0.21 g. (38.2%) of isopropylidenesuc-cinic acid, m.p. 164-165° (lit. value²² 161.5-162°), neut. equiv. 79.15 (theory 79.07). Methyl 2,4-Dioxo-5-carboxamido-3,6,6-trimethyl-3-azabicyclo-[3.1.0]hexane-1-carboxylate (X).—The procedure was identical with that described above for the preparation of VI. From 2.26 g. (0.01 mole) of II there was obtained 2.4 g. (98%) of X, re-crystallized from methanol, m.p. 195-195.5°. It had character-istic infrared bands in chloroform at 2.87 and 2.95 μ (N—H) and 5.63, 5.70, 5.85 (C=O). Anal. Calcd. for CuHuN*Os: C. 51.96: H. 5.55°. N 11.02

Anal. Caled. for $C_{11}H_{14}N_2O_5$: C, 51.96; H, 5.55; N, 11.02. Found: C, 51.93; H, 5.58; N,11.16.

(22) C. G. Overberger and C. W. Roberts, J. Am. Chem. Soc., 71, 3618 (1949).

2,4-Dioxo-5-(N-methyl)-carboxamido-6,6-dimethyl-3-azabicyclo[3.1.0]hexane-1-carboxylic Acid (XI). From X.—A solu-tion containing 2.54 g. (0.01 mole) of X, 35 ml. of methanol and 25 ml. of 10% sodium hydroxide was refluxed for 30 minutes then cooled, the methanol removed (Rinco evaporator), the resithen cooled, the methanol removed (knew evaporator), the resi-due made strongly acidic with dilute hydrochloric acid and placed in the refrigerator overnight. Filtration of the white crystals gave 2.15 g. (90%) of XI, m.p. 204-205° dec. Com-pound XI had, by potentiometric titration in water, pK_A^{II} 2.65 and pK_A^{II} 8.38. Its n.m.r. spectrum in D₂O containing NaOD had singlets at 7.36 and 8.84 τ , relative areas 1:2.

Anal. Calcd. for C₁₀H₁₂N₂O₆: C, 49.99; H, 5.04; N, 11.66; neut. equiv., 120.11. Found: C, 49.98; H, 5.04; N, 11.66; neut. equiv., 118.03.

From III.—3,3-Dimethyl-1,2-dicyanocyclopropane-1,2-carboximide (III)¹¹ (m.p. 242° dec., equal singlets at 8.50 and 8.57 τ in acetonitrile) was methylated with diazomethane by the usual procedure. From 1.9 g. (0.01 mole) of III there was obtained 2.0 g. (98.8%) of 3,3-dimethyl-1,2-dicyanocyclopropane-1,2-N-methylcarboximide, white needles from methanol, m.p. 235-236° dec. Its n.m.r. had three equal singlets at 7.16 (N- CH_3), 8.43 and 8.70 τ (C-methyl's).

Anal. Calcd. for $C_{10}H_9N_9O_2$: C, 59.10; H, 4.46; N, 20.68. Found: C, 59.23; H, 4.54; N, 20.61.

A solution of 2.03 g. (0.01 mole) of this N-methylimide in 15 ml. of 10% sodium hydroxide, after 30 minutes reflux followed by acidification with hydrochloric acid and cooling in the re-frigerator overnight, gave 1.99 g. (83.1%) of XI, m.p. 204-205° dec., identical with material obtained from X. In an attempt to hydrolyze XI further, 2.0 g. (8.3 mmoles) was refluxed for 3 hours with 30 ml. of 10% sodium hydroxide, cooled, acidified to congo red with hydrochloric acid and extracted continu-ously with ether for 24 hours. The extracted mixture was fil-tered to give 1.53 g. (76.5%) of tan crystals which, on recrystal-lization from methanol, gave recovered XI, m.p. and m.m.p. 205-206° dec. No crystalline product could be isolated from the ether extract.

Decarboxylation of XI .- The procedure was analogous to that described above for V. 4-(N-Methyl)-carboxamido-3-isopropylidenesuccinimide (XV), m.p. 131-132° from water, was obtained from XI in 89% yield.

Anal. Calcd. for $C_3H_{12}N_2O_3$: C, 55.09; H, 6.17; N, 14.28. Found: C, 55.18; H, 6.25; N, 14.31.

Compound XV gave a positive Baeyer test and, when treated as above with diazomethane, gave 4-(N-methyl)-carboxamido-3-isopropylidene-1-methylsuccinimide, m.p. 69-70° from methanol, in 85.7% yield.

Anal. Calcd. for $C_{10}H_{14}N_2O_3$: C, 57.12; H, 6.71; N, 13.33. Found: C, 57.19; H, 6.87; N, 13.09.

Quantitative hydrolysis of the latter demonstrated that both nitrogens had methyl groups attached to them. A sample (0.26 g., 1.25 mmoles) in 15 ml. of 10% sodium hydroxide was refluxed for 6 hours, evolved gases being swept by a nitrogen stream into hydrochloric acid. The latter was then evaporated to dryness and the white residue dissolved in 10 ml. of refluxing absolute ethanol. On cooling, there was obtained 0.162 g. (95.4%) of methylamine hydrochloride, m.p. and m.m.p. 226–227°. Con-tinuous extraction of the acidified reaction mixture with ether for 24 hours allowed recovery of 0.066 g. (33.6%) of isopro-pylidenesuccinic acid, m.p. 164–165° from water (lit. value¹⁷ 161.5-162°)

Methyl 2,4-dioxo-5-(N-methyl)-carboxamido-3,6,6-trimethyl-3-azabicyclo[3.1.0] hexane-1-carboxylate (XII) was prepared from XI and diazomethane in the usual way. From 1.20 g. from XI and diazomethane in the usual way. From 1.20 g. (5 mmoles) of XI there was obtained 1.25 g. (93.3%) of XII, m.p. 154-155°, from methanol. Compound XII had an N-H band at 2.95 μ and carbonyl bands at 5.70, 5.88 and 5.95 μ . Its n.m.r. spectrum in chloroform showed singlets at 6.22, 7.05, 8.42 and 8.75 τ and a doublet at 7.13 τ , all with equal areas.

Anal. Calcd. for $C_{12}H_{16}N_2O_6$: C, 53.72; H, 6.01; N, 10.44. Found: C, 53.82; H, 6.15; N, 10.31.

3,3-Dimethyl-1,1,2,2-tetracarboxylic Acid (XIII).-A solution of 1.34 g. (5 mmoles) of XII in 13 ml. of 10% sodium hydroxide and 20 ml. of methanol was refluxed for 30 minutes, then cooled and the alcohol removed by evaporation (Rinco). The remaining solution was extracted continuously for 6 hours with ether. This extract was discarded, the aqueous solution acidified with dilute hydrochloric acid and again extracted continuously with ether (36 hours). The ether extract was dried over anhydrous sodium sulfate and the solvent partially evaporated. To the remaining solution there was added 10 ml. of ethyl acetate; the whole was treated with Norit and the product isolated (0.87 g., 70.7%) by recrystallization from ethyl acetate-pentane; m.p. 166-169°. Further attempts at purification led to de-composition, so the acid was converted to its tetramethyl ester for analysis for analysis.

Methyl 3,3-Dimethylcyclopropane-1,1,2,2-tetracarboxylate (XIV).-Compound XIII (0.98 g., 4 mmoles) was methylated with diazomethane in the usual fashion, yielding 0.96 g. (80%) of XIV, m.p. 154.5-155°. Its infrared spectrum in carbon tetrachloride showed a sharp intense carbonyl band at 5.75 μ and its n.m.r. spectrum in chloroform showed two sharp singlets at 6.42 and 8.56 τ , relative areas 2:1.

Anal. Calcd. for $C_{13}H_{18}O_8$: C, 51.65; H, 6.00. Found: C, 51.81; H, 6.12.

Acknowledgment.—We are indebted to the Office of Ordnance Research (Contract No. DA-20-018-ORD- 16492) and to the Petroleum Research Fund of the American Chemical Society (Grant 488-C) for financial support of this research. H. H. also thanks his hosts at Cambridge University for their hospitality and discussions during preparation of this manuscript, and the National Science Foundation for a Senior Postdoctoral Fellowship. Finally, we thank Mr. John S. Fleming for certain of the n.m.r. spectra.

[CONTRIBUTION FROM THE METCALF CHEMICAL LABORATORIES OF BROWN UNIVERSITY, PROVIDENCE 12, R. I.]

Mechanism of the Oxidation of Nitrosobenzenes by Peroxoacetic Acid

By Khairat M. Ibne-Rasa, Carl G. Lauro and John O. Edwards

Received November 9, 1962

The oxidation of nitrosobenzenes by peroxoacetic acid in aqueous ethanolic media has been found to obey second-order kinetics with first-order dependence on each reactant. The oxidation is not acid catalyzed. The rate is enhanced by a factor of about 22 when the solvent is changed from ethanol to water. Electron-releasing *p*-substituents on nitrosobenzene accelerate the rate of oxidation, while the electron-withdrawing groups have the opposite effect. Peroxochloroacetic acid and Caro acid oxidize nitrosobenzene about 30 and 7 times, respectively, as fast as does peroxoacetic acid in 47% ethanol at 30°. The energy of activation $(\Delta S^{\pm}) - 22$ cal. mole⁻¹ deg.⁻¹. The rate of oxidation is not retarded by nitrobenzene, which is a radical trap. Added perchlorate ions do not have any significant influence on the rate of oxidation. A mechanism consistent with these data is suggested.

Introduction

A kinetic study of the peroxoacetic acid oxidation of some anilines to phenylhydroxylamines and then to nitrosobenzenes was reported in a previous paper.¹ The results of a similar study of the next step, the oxidation of nitrosobenzenes to nitrobenzenes, are now presented. These investigations were undertaken partly because of our general interest in reactions involving nucleophilic displacement on oxygen^{1,2} and partly because very few reactions of nitroso compounds have been the subjects of detailed kinetic studies in the past.³

It has been suggested that the oxidation of nitroso to nitro compounds by peroxy acids involves a nucleophilic attack by the peroxoacid on the nitrogen of the nitroso group.⁴ There is, in principle, at least one more pathway available for the oxidation; the nitroso nitrogen might perform a nucleophilic attack on the outer oxygen of the peroxy acid, the reactants thus acting in a manner reverse to that suggested by Bunton. One of the aims of this work was to discriminate, if possible, between the above two mechanisms.

The present studies indicate that in acidic aqueous ethanolic media the oxidation follows a mechanistic path analogous to that suggested for aromatic amines¹; *i.e.*, the peroxy acid behaves as an electrophile rather than as a nucleophile as suggested by Bunton.

Results

Kinetic Technique and Product Analysis.—The kinetic technique involved measuring the rate of disappearance of the nitrosobenzenes spectrophotometrically. Aqueous ethanolic solutions containing 0.01 to 0.02 M of a nitrosobenzene and 0.130 to 0.260 M peroxoacetic acid were placed in thermostated spectrophotometric cells and the disappearance of the nitroso compound followed as a function of time. The absorbancy at time "t" (A_t) was taken as a measure of the unreacted amount of nitrosobenzene at that time. In most (1) K. M. Ibne-Rasa and J. O. Edwards, J. Am. Chem. Soc., 84, 763 (1962).

(2) J. O. Edwards in "Peroxide Reaction Mechanisms," edited by J. O. Edwards, Interscience Publishers, Inc., New York, N. Y., 1962, pp. 67-106.
(3) B. G. Gowenlock and W. Luttke, *Quart. Rev.*, (London) **12**, 321 (1958).

(4) C. A. Bunton in "Peroxide Reaction Mechanisms," edited by J. O. Edwards, Interscience Publishers, Inc., New York, N. Y., 1962, p. 21.

cases the peroxy acid was present in large excess relative to the nitroso compound to provide pseudo-firstorder conditions. Plots of log A_t vs. time were linear to over three half-lives. Pseudo-first-order rate constants (k_1) were calculated from the slopes of these plots. The k_1 values were then divided by the initial concentration of peroxoacetic acid to get the secondorder constants (k_2) listed in Table I.

Table I

EFFECT OF MEDIUM, TEMPERATURE AND SUBSTITUENTS ON SECOND-ORDER RATE CONSTANTS

Initial [peroxoacetic acid] = 0.260 M, initial [Ar·NO] = 0.01 M

Compound	Solvent ^a	Temp. ±0.1°C.	$k_1 \times 10^4$, 1. mole ⁻¹ sec. ⁻¹
Nitrosobenzene	96% EtOH	50.0	5.15
	56% EtOH	50.0	18.4
	47% EtOH	50.0	27.2 ± 0.4
	36% EtOH	50.0	40.2
	16% EtOH	50.0	78.8
	47% EtOH	40.0	12.1 ± 0.1
	47% EtOH	30.0	5.15
p-Chloronitrosobenzene	47% EtOH	50.0	15.8
p-Bromonitrosobenzene	47% EtOH	50.0	14.7
<i>p</i> -Nitrosotoluene	47% EtOH	50.0	63.4
p-Nitrosoanisole	47% EtOH	50.0	200
4 Pr """ F+OH" in	moont (full	maluman	of athenal con

^a By "x% EtOH" is meant "x" volumes of ethanol contained in 100 volumes of reaction solution.

In a few experiments, which had been started with 0.01 M of a nitrosobenzene, 2 ml. of the reaction solution at infinite time was diluted with water to 250 ml. The ultraviolet spectra of these diluted solutions were studied and compared with those of $8 \times 10^{-5} M$ aqueous solutions of authentic samples of the corresponding nitrobenzene. The yield of nitrobenzene from the oxidation of nitrosobenzene was thus found to be quantitative and the conversion of *p*-chloronitrosobenzene to *p*-chloronitrobenzene was better than 95%.

Kinetic Law.—The linearity of the first-order plots indicates that the reaction is first order with respect to the nitroso compound. The dependence of the pseudofirst-order rate constants on the initial concentrations of the reactants was also studied for the oxidation of nitrosobenzene. The pseudo first-order rate constant