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RUBBER, POLYISOPRENES, AND ALLIED COMPOUNDS

PART VII.
ACTION OF NITRIC OXIDE THEREON

BY

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44. Rubber, Polyisoprenes, and Allied Compounds. Part VII. Action of Nitric Oxide thereon.

By George F. Bloomfield and (in part) G. A. Jeffrey.

The reaction of nitric oxide with the olefins cyclohexene, 1-methylcyclohexene, dihydromyrcene, and rubber presents characteristics of a free-radical, chain reaction. In the products the molecular ratio of combined oxygen to nitrogen is considerably in excess of 1:1, the nitrogen atom being generally directly linked to carbon. In many instances definite nitro-compounds have been isolated, and a considerable part of the attack appears to

proceed at the ethylenic carbon atoms, either by substitution or addition of NO_2 and N_2O_3 groups. The precise reaction mechanism is obscure.

Although the reactions of nitrogen dioxide, dinitrogen trioxide, and nitrosyl chloride with olefins have been studied extensively, the action of nitric oxide has received but little attention and it has in fact been stated (Sidgwick, "The Organic Chemistry of Nitrogen," 1937, p. 213) that it does not combine with a normal ethylenic linkage. Nitric oxide is, however, of well-established utility in the study of free-radical reactions, which it can promote, retard, or inhibit in virtue of its property of starting reaction chains, or stopping them by combination with other free radicals.

General Characteristics of the Reaction of Nitric Oxide with Olefins.—Characteristics of a free-radical mechanism were exhibited. For instance, when a solution of rubber, dihydromyrcene, 1-methylcyclohexene, or cyclohexene was shaken in contact with nitric oxide there was generally an induction period of 15—30 minutes, depending on the intensity of the prevailing light, before any sign of reaction became evident. Then the solution acquired a pronounced green colour, absorption of nitric oxide commenced, and some heat was evolved. The induction period was not confined to the initial starting of the reaction, for if the reaction was stopped by removing the nitric oxide from the system and then, after a short interval, nitric oxide was readmitted, an induction period was again observed before reaction recommenced. Under the conditions employed, the reaction ceased owing to admixture by nitrogen produced in the reaction. When provision was made for removing the nitrogen, no difficulty was experienced in carrying the reaction as far as was desired, and it became obvious that absorption of nitric oxide could proceed far beyond one molecular proportion per double bond, and that the volume of nitrogen formed was generally between a third and a quarter of the volume of nitric oxide which had been absorbed at any given stage of the reaction. The formation of nitrogen in reactions of nitric oxide with free radicals (but not with olefins) has been reported by Nauta and Mulder (Rec. Trav. chim., 1939, 58, 1070) and by Sonneborn and Wiselogle (J. Amer. Chem. Soc., 1942, 64, 860), who attributed its formation to decomposition of an intermediate hyponitrite.

Obviously the reaction of nitric oxide with olefins does not proceed by any simple attachment of nitric oxide itself to the unsaturated molecule, and it is not surprising that a variety of products was isolated contain-

ing nitrogen in various states of combination with oxygen.

Reaction with Rubber.—Gorgas's observation (Ber., 1930, 63, 2700) that an insoluble nitrogenous product was precipitated by the action of nitric oxide on a rubber solution has been confirmed. This occurred at a stage represented by the absorption of approximately 1 mol. of nitric oxide per C_5H_8 unit, and the composition of the product corresponded with the introduction into half of the available C_5H_8 units of groups containing nitrogen and oxygen in the ratio 1:2 to 1:2.7, varying from one preparation to another. At a somewhat earlier stage the product could be separated into two components of approximate compositions $C_{10}H_{15}O_2N$ and $C_{10}H_{15}O_3N$. In the early stages of reaction there was considerable reduction of molecular size, by an obscure mechanism which required the absorption of a proportion of nitric oxide appreciably greater than the proportion of oxygen required to bring about a comparable reduction in molecular size (Farmer and Sundralingam, L. 1943, 125).

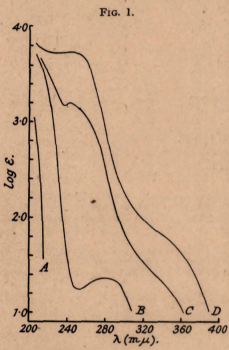
Reaction with Simpler Olefins allied to Rubber.—Dihydromyrcene and 1-methylcyclohexene yielded products which appeared to contain substituent nitro-groups, together with substances of rather lower oxygen: nitrogen ratio. In the formation of the former products the unsaturation was considerably affected, being much lower

than that calculated for simple nitro-substituted olefins.

cycloHexene yielded the following products: (1) the already known crystalline cyclohexene pseudo-nitrosite * (Baeyer, Annalen, 1894, 278, 110; Wieland, ibid., 1921, 424, 71), (ii) a mixture of isomeric nitrocyclohexenes containing a considerable proportion of 1-nitrocyclohexene, and (iii) an unstable, non-distillable, viscous oil constituting the major product. Partial separation of the isomeric nitrocyclohexenes yielded a fairly pure sample of the 1-nitro-isomer but the other isomer, in all probability 3-nitrocyclohexene, could not be isolated in a pure state since it underwent considerable decomposition in the presence of alkali (cf. the instability of α-methylene-substituted halogen; preceding paper). The constitution of the 1-nitrocyclohexene was established by oxidation to adipic acid and by reduction to cyclohexanoneoxime, the latter reaction probably proceeding through the intermediate cyclohexenylhydroxylamine (cf. the reduction of nitrocyclohexane to cyclohexylhydroxylamine), or possibly by direct 1: 4-addition of hydrogen to the C:C·N:O system. As was expected from the presence on the ethylenic carbon atom of a strongly polar group, capable of forming a conjugated system with the ethylenic linkage, addition of iodine chloride to the >C = C< bond was almost completely inhibited, but the additive capacity for hydrogen established the presence of the ethylenic linkage. Spectrographic examination by Dr. H. P. Koch, for which the authors' thanks are due, has confirmed the presence of considerable conjugation in the 1-nitrocyclohexene component of the mixture (Fig. 1). The other nitrocyclohexene which was present exhibited a much more satisfactory additive capacity for iodine chloride. The constitution of the major non-volatile reaction product has not been established; it was a unimolecular substance exhibiting no additive capacity for iodine chloride, with groups containing oxygen and nitrogen in a ratio slightly in excess of 2:1 added or substituted at the original ethylenic linkage of the cyclohexene; treatment with alkali gave a crystalline product of higher nitrogen content than the original oil, suggesting the presence of stable carbon-nitrogen linkages.

^{*} The bimolecular structure proposed by Wieland has been confirmed by X-ray examination (p. 123).

The principal characteristics of the reaction under consideration are therefore: (1) Various products are formed; all contain nitrogen directly linked to carbon, and the nitrogen is combined with oxygen in the pro-



A.—cycloHexene.
B.—Nitrocyclohexane.

-Mixture of isomeric nitrocyclohexenes.

D.-1-Nitrocyclohexene (mainly).

portion 1:2, 2:3, or 1:3; several of the products are identical with those known to be formed by the action of higher oxides of nitrogen on the corresponding olefin. (2) At least 1 mol. of nitrogen is formed for every 4 mols. of nitric oxide reacting. (3) When substitutive reaction is taken into account, the H: C ratio is substantially preserved. (4) The ethylenic carbon atoms are extensively involved in substitutive and additive reactions. (5) The reaction is inhibited by alcohol or acetic acid.

Since most of the reaction products are precisely those obtained by the action of higher oxides of nitrogen or of nitrous acid, it is likely that reaction proceeds by conversion of the nitric oxide into a higher state of oxidation, probably by a free-radical chain mechanism involving the hydrocarbon. The attack at the ethylenic carbon atoms may be closely related to the preferential attack of free halogen radicals at these carbon atoms (preceding paper). Any such reaction must occur in the liquid phase, since brown fumes in the gas phase were not observed; the presence of nitrous acid or, more probably, dinitrogen trioxide in the liquid phase was, however, detected by Sonneborn and Wiselogle's procedure (loc. cit.) during the course of the reaction. If nitrous acid itself is formed, it must become involved in a course of reaction other than the usual formation of \(\psi\$-nitroles with those secondary nitro-groups already present. The formation of some of the reaction products is satisfied by overall reactions of the type represented by the equations

(a) RH + 4NO = RNO₂ + N₂ + HNO₂ or

$$2RH + 8NO = 2RNO_2 + 2N_2 + N_2O_3 + H_2O_3$$

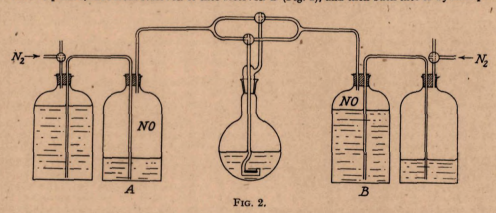
(b) $>C:C < + N_2O_3 = >C(NO) \cdot \dot{C}(NO_2)$

(c)
$$2RH + 10NO = 2RNO_3 + 3N_2 + 2HNO_2$$
 or $2RH + 10NO = 2RNO_3 + 3N_2 + N_2O_3 + H_2O$.

EXPERIMENTAL.

(Microanalyses were carried out by Dr. W. T. Chambers and Miss H. Rhodes.)

Reactions were conducted either in a glass globe or in a small flask fitted with a sintered-glass distributor through which gas could be passed, first from reservoir A into reservoir B (Fig. 2), and then back into A by manipulation of the



stopcocks and application of nitrogen pressure to the reservoirs. The apparatus was first swept out with purified nitrogen, then evacuated until the solvent had boiled vigorously for a few minutes under the reduced pressure; refilled with nitrogen, re-evacuated, and finally filled with nitric oxide obtained from sodium nitrite, potassium ferrocyanide, and acetic acid. The gas remaining at the conclusion of an experiment was displaced into a measuring system and, after removal of any nitric oxide with aqueous potassium permanganate, the volume of residual nitrogen was measured.

Experiments with Rubber.—The hydrocarbon (sol rubber, 5 g.) was dissolved in carbon tetrachloride (300 c.c.); reaction required 30—120 mins. The results are in the table.

Experiments with Dihydromyrcene and 1-Methylcyclohexene.—Dihydromyrcene (15·6 g.) in carbon tetrachloride (150 c.c.) was shaken with nitric oxide (1888 c.c., N.T.P.) in the glass globe during 8 hours in diffused sunlight. The residual gas (588 c.c.) contained a little nitric oxide (38 c.c.), whence the nitric oxide consumed and the nitrogen formed were equivalent to 0·73 mol. and 0·22 mol. respectively per mol. of dihydromyrcene. A little yellow resin was precipitated. The yellow supernatant solution yielded on distillation unreacted dihydromyrcene (10·8 g.), a liquid fraction, b. p. 66—70°/0·001 mm. (0·6 g.), and considerable residue (4·1 g. Found: C, 55·55; H, 8·5; N, 11·6%; I.V.,

Reaction of Nitric Oxide with Rubber.

| NO consumed, | N ₂ formed, | Analysis of product, %. | | | | | |
|---|------------------------|-------------------------|-------------|------|--|---------|---|
| mols. per C ₅ H ₈ unit. | mols. per mol. NO. | c. | H. | N. | 0. | I.V. | Solubility of product. |
| 0.13 | _ | 84.2 | 11.3 | _ | | | Soluble |
| 0.15 | • 0.28 | 83.9 | 11.3 | 1.65 | 3.2 | - | * |
| 0.19 | 0.23 | 83.15 | 11.1 | - | _ | 30000 | |
| 0.48 | 0.25 | 76.0 | 10.15 | 4.2 | 9.65 | 262 | ,, † |
| 0.94 | 0·24 t | _ | 1000 | _ | A STATE OF THE STA | _ | |
| 0.95 | 0.29 | 68.7 | 9.05 | 6.0 | 16.25 | 220 | Partly insol |
| ca. 1·0 | - | 63.25 | 845 | 7.85 | 20.3 | - | Insol. |
| ca. 0.9 | - | {68·0 60·7 | 8·9 7·85 | 6.95 | 16·15 § 24·6 | = | Insol. part, 74% of product Soluble part, 26% of product |
| 1.35 | 0·29 t | OF THE PARTY OF | 372 - 3 S | _ | - | - Total | Insol. |

* Intrinsic viscosity 0.74 in benzene + 15% methanol; mol. wt. 74,000 (original mol. wt. 238,000).
† Intrinsic viscosity 0.57 in benzene; mol. wt. 33,000 (original mol. wt. 220,000).
‡ In these experiments the absence of carbon dioxide in the residual gas was demonstrated by passage of the latter through a weighed CO₂-absorption tube.
§ Empirical formula C₁₀H_{15.4}NO₂.

Empirical formula C₁₀H_{15.4}NO₃.

79, corresponding to $C_{10}H_{18}O_{3\cdot3}N_{1\cdot8}$). The liquid fraction was probably a *nitrodihydromyrcene* [Found: C, 65·6; H, 9·4; N, 7·7; I.V., 156. $C_{10}H_{17}O_{2}N$ requires C, 65·5; H, 9·35; N, 7·65%; I.V., 277 ($\frac{1}{2}$), 139 ($\frac{1}{17}$)]. 1-Methyl*cyclo*hexene similarly yielded a liquid product, b. p. 50°/0·01 mm., probably a *nitromethyl*cyclohexene (Found: C, 59·3; H, 8·0; N, 10·0; I.V., 73. $C_{7}H_{11}O_{2}N$ requires C, 59·55; H, 7·85; N, 9·9%; I.V., 180), and some viscous

1-Methyly-Johnexnes inimilarly yielded a liquid protott, b. p. 50°/0010 mm, probably a ninomethyly-cholexene (Found. C, 693; H, 80; N, 100; I.V., 73. C,H₁₁O_N requires C, 59·55; H, 785; N, 94%; I.V., 180), and some viscous min a dull light, with formation of 0·44 mol. of nitrogen; a white, crystalline solid separated continuously from the commencement of the reaction. This (2·44 g) was filtered off and the filtrate yielded on distillation a pale yellow liquid, b. p. 33—46°/0·01 mm. (ca. 6 g.), and considerable residue (1·44 g).

Crystalline solid. This was cyclobexene φ-introsite, m. p. 153° (decomp.) from benzene, mixed m. p. with authentic specimen 133° (decomp.) (Found: C, 45·55; H, 6·35; N, 17·5; I.V., 0. Calc. for C,H₁₀N,O₂: C, 45·55; H, 40·5; N, 17·5; I.V., 0. Calc. for C,H₁₀N,O₂: C,A₁C,O₂: C,A₂C,O₃: C,A₂C,O₃: C,A₃C,O₃: C,

* A high order of reactivity of groups substituted at the 3-position of cyclohexene has already been observed in halogen-substitution products (preceding paper).

Examination of other specimens of nitrocyclohexene. 1-Nitrocyclohexene, b. p. 45—50°/0·1 mm. (Found: C, 56·0; H, 7·53; N, 10·8%; I.V., 29), prepared according to Wieland (loc. cit.), was insoluble in N-sodium hydroxide; 1·9 g. gave 1·1 g. of adipic acid on oxidation, and the hydrogen uptake (Adams's catalyst) was 7 atoms per mol. 3(?)-Nitrocyclohexene, b. p. 34°/0·01 mm. (Found: C, 56·25; H, 7·1; N, 10·9%; I.V., 155), prepared by the action of alkali on cyclohexene ψ-nitrosite (Wieland) was soluble (with partial hydrolysis *) in N-sodium hydroxide; 0·8 g. gave 0·15 g. of adipic acid, corresponding to the presence of 16% 1-nitrocyclohexene, and the hydrogen uptake (Adams's catalyst) was 7.5° atoms per mol

adipic acid, corresponding to the presence of 16% 1-mitrocyclonexene, and the hydrogen uptake (Radian 1977) (Residue. This slowly decomposed at room temperature and could not be distilled without severe decomposition. An effective separation from cyclohexene ψ-nitrosite was achieved by dissolution in methanol at 0°, in which the nitrosite was insoluble; removal of the alcohol gave a brown viscous oil. The products of four separate reactions were remarkably constant in composition [Found: C, 44·6±0·5; H, 6·1±0·1; N, 13·9±0·1%; I.V., 4; M (f. p. depression in benzene), 164], which corresponded to the formula C₆H₁₀O_{3·6}N_{1·6}. Oxidation of 4·6 g. with alkaline permanganate (3%, 270 c.c.) proceeded only slowly and yielded an unidentified neutral oil (0·55 g.) and adipic acid (1·7 g.). Oil (5·4 g.) in alcoholic potassium hydroxide (100 c.c., 0·82N) at the b. p. for 15 mins. yielded a large and a small portion respectively soluble and insoluble in alkali. Distillation of the former yielded a little impure nitrocyclohexene, b. p. 40—47° [0·1 mm. (Found: C, 57·1; H, 7·4; N, 10·6. Calc. for mixture of 96·75% (C₆H₉·NO₂ and 3·25% (C₆H₉·OH: C, 57·15; H, 7·25; N, 10·5%), and a considerable residue from which an unidentified crystalline solid separated, m. p. 107—108° from benzene (Found: C, 46·0; H, 5·15; N, 17·7. C₆H₈O₃N₂ requires C, 46·1; H, 5·15; N, 17·95%).

The reaction of nitric oxide with cyclohexene did not proceed in alcoholic solution, or in a medium containing acetic acid. Nitric oxide did not react with nitrocyclohexene in carbon tetrachloride.

acid. Nitric oxide did not react with nitrocyclohexene in carbon tetrachloride.

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