

OXIDATIVE CYCLIZATION OF N,N -DIMETHYLTRYPTAMINE, 3-INDOLEPROPANOL,
AND 3-INDOLEPROPANETHIOL WITH N-BROMO- OR N-CHLOROSUCCINIMIDE

Tohru Hino^{*}, Hidetoshi Miura, Toru Nakamura, Ryoichi Murata, and Masako Nakagawa

Faculty of Pharmaceutical Sciences, Chiba University, Yayoi-cho, Chiba-shi 280

The reaction of N,N -dimethyltryptamine(5b) with NBS in carbon tetrachloride yielded a quaternary salt of pyrrolo[2,3-b]indole(7b). The similar oxidative cyclization of 3-indolepropanol(5c) and 3-indolepropanethiol(5e) with NBS in methylene chloride gave pyrano- and thiopyranoindoles(7c and 7e). Further halogenation of 7c with NCS yielded a spirooxindole(12, Y=O) and a dichloride(13), whereas 7e with NBS gave 12(Y=S) and bromothiopyranoindole(14).

Ionic bromination of 3-alkylindole(1) with N-bromosuccinimide(NBS) has been known to proceed via 3-bromoindolenine intermediate(2) which collapsed to 2-bromoindoles(3) or oxindoles(4) depending on the reaction media¹. When a nucleophilic group is present at an appropriate position in the 3-substituent, the nucleophile may attack to 2-position of the indolenine intermediate(2) to form a cyclic product. Witkop and his coworkers² have been reported the oxidative cyclization of N-acetyltryptamine(5a) and N-acetyltryptophan with NBS to give pyrroloindole(7a). We now report oxidative cyclization of

N,N -dimethyltryptamine(5b)³, 3-indolepropanol(5c)⁴, and 3-indolepropanethiol(5e)⁵ which contain a nucleophilic center such as NMe_2 , OH, or SH group in the 3-substituent.

The reaction of 5b, 5c and 5e with NBS or N -chlorosuccinimide(NCS) were carried out under various conditions and the results are summarized in Table 1, indicating the superiority of NBS over NCS in oxidative cyclization.

Table 1 Oxidative Cyclization of 5

Compd	Condition	Products(% yield)
5b	CCl_4 -NBS, rm. temp	7b(65%)
5c	CH_2Cl_2 -NBS, -14 - -18°	7c(57%)
	CH_2Cl_2 -NCS, -14 - -16°	12(30%)
5e	CH_2Cl_2 -NBS, -13- -10°	7e(80%)
	CH_2Cl_2 -NCS, -50- -48°	7e(32%)

The structures of these cyclized products were confirmed by elemental analyses and spectral data which are summarized in Table 2. Pyranoindole(7c) was easily hydrolyzed by an acid to 3-(3-hydroxypropyl)oxindole, mp 105-105.5°⁶.

On the other hand the expected cyclized products such as 7d or 7f have not been obtained by the reaction of 3-indoleethanol(5d) and 3-indoleethanethiol(5f) with NBS or NCS under various conditions. The reaction of 5d gave complicated products which included oxindolic products⁷, while 5f yielded the corresponding disulfide(8) as major product. These results may suggest that a thiol group is more reactive than an indole ring towards NBS(NCS) and an intermediate of the cyclization of 5e to 7e would not be an indolenine(6e), but a spiroindolenine(10) which derived from a sulfonyl halide(9). The disulfide(8) may be

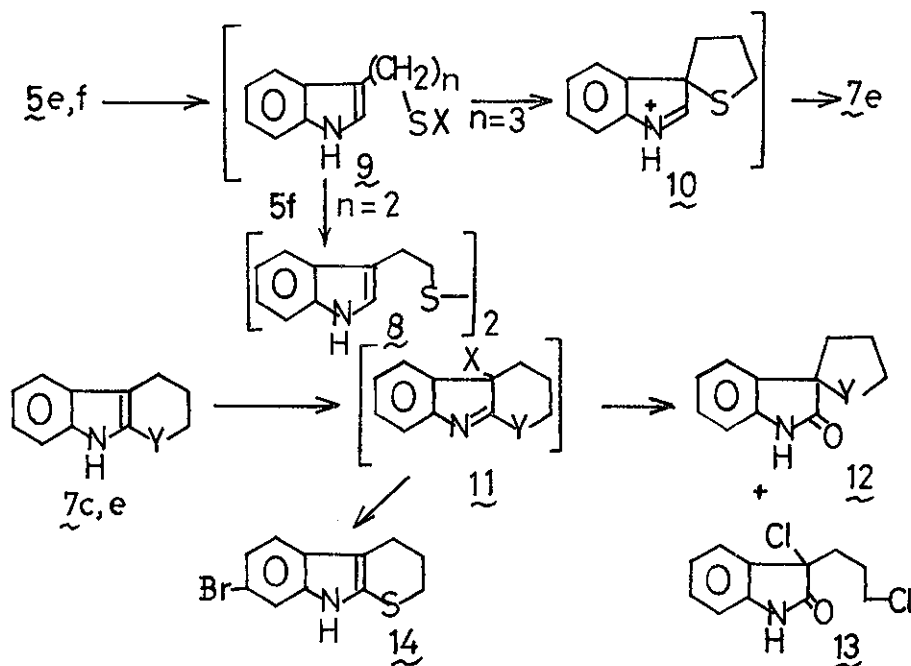
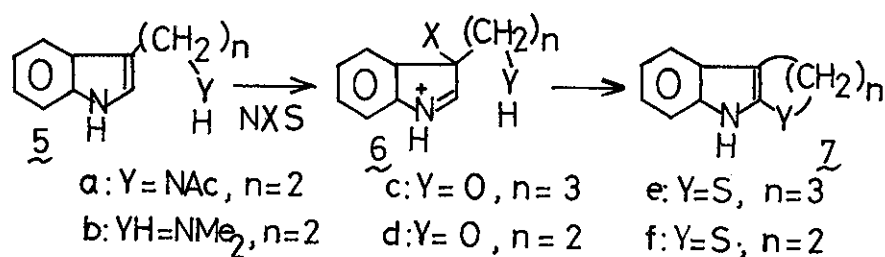
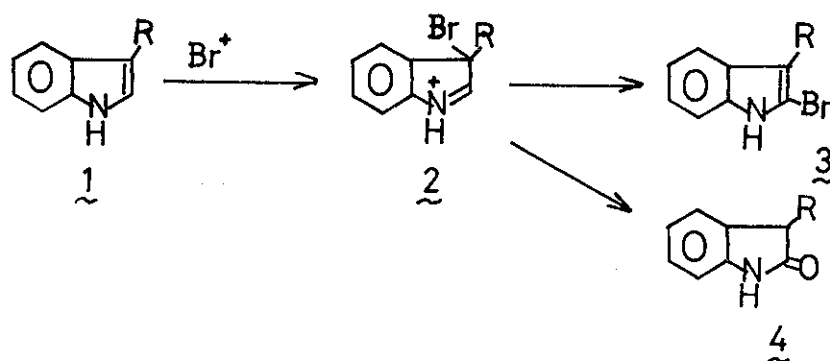
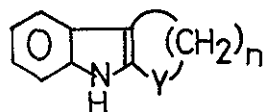


Table 2 Spectral Data of 7



Compd No	mp	$\lambda_{\text{max}}^{\text{EtOH}}$ nm(ϵ)	Nmr (δ -value)	Mass (m/e (rel.intens.))
7b Y=N ⁺ Me ₂ , Br ⁻ n=2	275°(decomp)	263(6,900) 275 ^s (5,900) 278 ^s (5,400) 286(4,400)	in D ₂ O 3.24(f, 2H, 3-CH ₂) 3.54(s, 6H, NMe ₂) 4.61(t, 2H, 2-CH ₂)	186(100, M ⁺ - HBr)
7c Y=O, n=3	91.5-92.5°	219(22,200) 230(28,700) 275(7,700) 293(5,000)	in CDCl ₃ 1.90-2.20(m, 2H, 3-CH ₂) 2.63(t, 2H, 4-CH ₂) 4.30(t, 2H, 2-CH ₂) 7.50(br.s. NH)	173(M ⁺ , 100) 145(M-CH ₂ =CH ₂ , 75)
7e Y=S, n=3	147-148°	224(21,000) 238(28,000) 275 ^s (6,800) 286(9,300) 302(10,900)	in CDCl ₃ 2.10-2.32(m, 2H, 3-CH ₂) 2.7-2.9(m, 2H, 2-CH ₂) 3.05-3.15(m, 2H, 4-CH ₂) 8.50(br.s, NH)	189(M ⁺ , 99) 161(M-CH ₂ =CH ₂ , 100)

formed by the reaction of 5f and 9(n=2) due to the difficulty in intramolecular cyclization of 9 to form a four membered spiroindolenine as an intermediate to thienindole⁸.

Further halogenation of pyranoindole(7c) and thiopyranoindole(7e) was investigated in order to compare with our previous results obtained by bromination of 2-ethylthioindoles⁹. Treatment of 7c with NCS in methylene chloride at $-5 - -10^{\circ}$ ¹⁰, followed with an acid gave spirooxindole(12, Y=O), mp 97-98°, in 80% yield. On the other hand followed treatment of the reaction mixture with ammonium chloride in aqueous ethanol provided 12(Y=O, 36%) and dichloride(13, 7%), mp 111-112°, which were also obtained by the reaction of 5c with two moles of NCS in methylene chloride at $-5 - 0^{\circ}$. Reaction of 7e with NBS in methylene chloride at room temperature gave unstable bromoindolenine(11, Y=S, X=Br)¹¹ which was readily converted to the spirooxindole(12(Y=S), mp 126-127°, by addition of an acid and to 7-bromo derivative(14), mp 152°, on heating. While the chloroindolenine(11, Y=S, X=Cl)¹² obtained by the reaction of 7e with NCS in carbon tetrachloride, was converted to the spirooxindole(12, Y=S) by addition of an acid, but 7-chloro derivative corresponding to 14 was not obtained on heating in carbon tetrachloride, suggesting that the reactivity of chloroindolenine and bromoindolenine may be different.

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REFERENCES AND FOOTNOTES

- 1 W.J.Houlihan(Ed), *Indoles*. Part 2, pp 127 Wiley-Interscience, 1972, New York;
- T.Hino, M.Tonozuka, and M.Nakagawa, *Tetrahedron*, **30**, 2123(1974).
- 2 M.Ohno, T.F.Spande, and B.Witkop, *J.Am.Chem.Soc.*, **92**, 343(1970).

- 3 H.Kondo, H.Kataoka, Y.Hayashi, T.Dodo, Itsuu Kenkyusho Nempo, 10, 1 (1959).
- 4 F.Lingens and K.H.Weiler, Ann., 662, 139(1963).
- 5 V.N.Buyanow and N.N.Suvorov, Tr.Mosk.Khim.Tekhol.Inst. No 52,114(1967)(Chem. Abst., 68, 114344z(1968)); Khim.-Farm. Zh., 1, 4 (1967)(Chem.Abst., 67, 7347a(1967)).
- 6 All new compounds shown with mps gave correct elemental analyses and reasonable spectral data.
- 7 Oxidation of 3-indoleethanol with bromine in acetic acid was reported to give 3-(2-hydroxyethyl)oxindole. E.Wenkert, J.S.Bindra, C-J.Chang, D.W.Cochran, and D.E.Pearick, J.Org.Chem., 39, 1662(1974).
- 8 Solvolysis of *w*-(3-indolyl)alkyl tosylates has been reported to proceed via spiroindolenine as the intermediate and the solvolysis via a four membered spiroindolenine has been known to be very slow. A.H.Jackson and B.Naido, Tetrahedron, 25, 4843(1969).
- 9 T.Hino, M.Endo, M.Tonozuka, and M.Nakagawa, Heterocycles, 2, 565(1974).
- 10 An intermediate, probably an chloroindolenine(11, X=Cl, Y=O) was recognized in the reaction mixture by UV spectra(λ_{max} 298nm) and tlc, but was not isolated.
- 11 The bromoindolenine could not be isolated, but its presence was recognized by UV (λ_{max} 325 nm) which is similar to that of 3-bromo-2-ethylthio-3-methylindolenine⁹.
- 12 The chloroindolenine was isolated as crystals, but decomposed on recrystallization. Mass m/e 223(M^+ , 42%). IR(KBr) 1520 cm^{-1} (C=N). $\lambda_{\text{max}}^{\text{EtOH}}$ 228, 238, 325 nm.

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