## EFFICIENT SYNTHESIS OF BENZ[c,d]INDOL-3(1H)-ONE DERIVATIVES BY INTRAMOLECULAR CYCLIZATION OF 3-(4'-METHYLCINNAMOYL)INDOLES AND SUBSEQUENT ELIMINATION OF TOLUENE.

## Taichi Komoda and Shin-ichi Nakatsuka\*

The United Graduate School of Agricultural Science, Gifu University
1-1 Yanagido, Gifu 501-1193, Japan

**Abstract**: Benz[c,d]indol-3(1H)-one 10 and 11, a major part of indole alkaloids 0231A 2 and 0231B 3 produced by *Streptomyces sp.* HKI0231, new enzyme inhibitors of  $3\alpha$ -hydroxysteroid dehydrogenase, were efficiently synthesized by cyclization of 3-(4'-methylcinnamoyl)indole derivatives 6c and 7c and elimination of toluene.

In the biosynthesis of ergot alkaloids 1, a phenyl group is first introduced at 4-position of tryptophan.<sup>1)</sup> Although many attempts introducing such substituent at the position of indole nucleolus have been studied for a long time, no successful method was reported<sup>2)</sup> except a few cases.<sup>3-5)</sup>

R Me Me Me OMe OMe OMe OMe 
$$^{1}$$
 Ergot alkaloids  $^{2}$ : R = OMe  $^{0231}$ A  $^{3}$ : R = H  $^{0231}$ B

Figure 1.

In our synthetic studies on indole alkaloids, we have found intra- and intermolecular cyclization toward the 4-position of indole nucleus in cases of dehydrotryptophan and N-acylindole derivatives.<sup>5)</sup> Recently, we found a novel synthetic method of benz[c,d]indol-3(1H)-one derivatives 5, whose common structure was found in enzyme inhibitor 0231A 2 and 0231B 3 from Streptomyces sp. HKI0231,<sup>6)</sup> via intramolecular cyclization of substituted 3-cinnamoylindoles 4 and subsequent elimination of benzene (Fig. 2).<sup>7)</sup>

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Figure 2. Cyclo-elimination of 3-cinnamoylindole derivatives.

In this reaction, a phenyl group was used as a promoter of the cyclization and also as a protective group of a double bond because its elimination gave aromatized benz[c,d]indol-3(1H)-one derivatives. Although the benz[c,d]indol-3(1H)-one derivatives were expected to be useful key intermediates for indole alkaloids containing substituent at the 4-position, the reaction yields were insufficient (45-79%). In this paper, we report much more efficient synthetic method for the substituted benz[c,d]indol-3(1H)-one 10 and 11 by studies on the efficiency of several substituted benzene as a promoter and eliminator.

The starting materials **6a-c** and **7a-d** were prepared from 1-pivaloylindole by Friedel-Crafts acylation<sup>8)</sup> with corresponding 3-cinnamoyl chlorides and subsequent de-protection of the pivaloyl group. Reaction conditions and results of AlCl<sub>3</sub> catalyzed cyclo-elimination of substituted 3-cinnamoylindole derivatives **6a-c** and **7a-d** are shown in the Table 1.

An electron withdrawing nitro group at 4'-position of benzene ring completely obstructed the cyclization (entry 7 and 8). Although a stronger electron donating methoxy group is expected to be a suitable function for cyclo-elimination, cyclization and also elimination were delayed to afford the intermediates **8b**<sup>9)</sup> or **9b**<sup>10)</sup> (entry 2 and 5). The elimination of 4'-methoxyphenyl group of **9b** was completely impeded and no eliminated product **11** was detected in the reaction mixture (entry 5). House *et al.* reported that the excess AlCl<sub>3</sub> converted methoxy group to complex compound with positive charge and it inactivates phenyl group. (11) Our results agree with their theory.

On the other hands, an electron donating methyl group accelerated cyclization and also elimination of the phenyl group (entry 3 and 6). Typical reaction in entry 3 is as follows: 5 eq. AlCl<sub>3</sub> was added to a stirred solution of **6c** in CHCl<sub>2</sub>CHCl<sub>2</sub> at 80°C and stirred for 10 min. The cooled reaction mixture was poured into saturated aq. NaCl, and extracted with CH<sub>2</sub>Cl<sub>2</sub>. The organic layer was dried over Na<sub>2</sub>SO<sub>4</sub>, and the solvent was removed in *vacuo*. Purification of the residue by silica gel chromatography (100% ethyl acetate) gave benz[c.d]indol-3(1H)-one **10** (85%). [**10**: m.p. 157-162°C (dec.); H-NMR  $\delta$  ppm(500MHz, 10%CD<sub>3</sub>OD in CDCl<sub>3</sub>) 6.71(1H, d, J=9.4Hz), 7.40(1H, dd, J=7.3 and 8.0Hz), 7.58(1H, d, J=7.3Hz), 7.66(1H, d, J=8.0Hz), 7.79(1H, d, J=9.4Hz), 8.31(1H, s);  $^{13}$ C-NMR  $\delta$  ppm(100MHz, 10%CD<sub>3</sub>OD in CDCl<sub>3</sub>) 115.5, 115.8, 123.6, 123.7, 124.3, 125.8, 131.8, 133.4, 133.9, 139.1, 182.3; UV  $\lambda_{max}$  (MeOH) 232( $\epsilon$ =20000), 340( $\epsilon$ =7600), 381( $\epsilon$ =7100), 398( $\epsilon$ =9000), 419( $\epsilon$ =6500)nm; 1R  $\nu_{max}$  (KBr) 3147, 3094, 3054, 2931, 2850, 2795, 1640, 1590, 1565, 1500, 1321, 1186, 1137, 821, 740cm<sup>-1</sup>; EIMS (70eV. m/z) 169(M<sup>+</sup>, base peak), 141, 114; HREIMS Calcd for C<sub>11</sub>H<sub>7</sub>NO: 169.0528. Found 169.0536].

Table 1. Cyclo-elimination of substituted 3-cinnamolylindole derivatives 6a-c and 7a-d.

Entry	S. M.	Conditions	Time (min)	Products (Yie	Products (Yield%)	
1	6a	CHCl <sub>2</sub> CHCl <sub>2</sub> , 80°C	20	8a (-)	<b>10</b> (45) <sup>7)</sup>	
2	6b	CHCl <sub>2</sub> CHCl <sub>2</sub> , 80°C	20	<b>8b</b> (16)	10 (11)	
3	6c	CHCl <sub>2</sub> CHCl <sub>2</sub> , 80°C	10	8c (-)	10 (85)	
4	7a	CH <sub>2</sub> Cl <sub>2</sub> , r. t.	30	9a (-)	-11 (79) <sup>7)</sup>	
5	7b	CH <sub>2</sub> Cl <sub>2</sub> , r. t.	120	9b (84)	11 (-)	
6	7c	CH <sub>2</sub> Cl <sub>2</sub> , r. t.	10	9c (-)	11 (93)	
7	7 <b>d</b>	CH <sub>2</sub> Cl <sub>2</sub> , r. t.	30	No Reaction		
8	7d	CHCl <sub>2</sub> CHCl <sub>2</sub> , 120°C	30	No Reaction		

Figure 3. Cyclo-elimination mechanism of 6a-c and 7a-d.

Thus, we could develop an efficient synthesis of benz[c,d]indol-3(1H)-one derivatives 10 and 11<sup>12)</sup> by intramolecular cyclization of 3-(4'-methylcinnamoyl)indoles 6c and 7c and subsequent elimination of toluene. Application of this cyclo-elimination to total synthesis of ergot alkaloids 1 and enzyme inhibitor 0231A 2 and B 3 are now in progress.

## References and Notes

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- 7) Very recently, our preliminary results on cyclo-elimination of 3-cinnamoylindole derivatives 4 were submitted for publication.
- 8) (a) D. M. Ketcha and G. W. Gribble, J. Org. Chem., 50, 5451(1985); (b) S. Nakatsuka, K. Teranishi and T. Goto, Tetrahedron Lett., 35, 2699(1994).
- 9) **8b**: <sup>1</sup>H-NMR δ ppm (200MHz, 10%CD<sub>3</sub>OD in CDCl<sub>3</sub>) 3.08(2H, d, *J*=8.1Hz), 3.81(3H, s), 4.70(1H, t, *J*=8.1Hz), 6.80-6.89(3H, m), 7.15-7.23(3H, m), 7.33(1H, d, *J*=6.8Hz), 7.83(1H, s).
- 10) **9b**: <sup>1</sup>H-NMR δ ppm(400MHz, CDCl<sub>3</sub>) 1.80(3H, s), 2.94(1H, d, *J*=15.9Hz), 3.27(1H, d, *J*=15.9Hz), 3.76(3H, s), 6.79(2H, br.d, *J*=8.8Hz), 7.02(1H, d, *J*=7.0Hz), 7.20-7.32(4H, m), 7.73(1H, d, *J*=2.9Hz), 9.03(1H, br.d, *J*=2.9Hz).
- (a) H. O. House and J. K. Larson, J. Org. Chem., 33, 448(1968); (b) H. O. House and C. B. Hudson, ibid, 35, 647(1970); (c) W. S. Jhonson and H. J. Glenn, J. Am. Chem. Soc., 71, 1092(1949).
- 12) 11: m.p. 159-164°C (dec.);  ${}^{1}$ H-NMR  $\delta$  ppm(500MHz, 10%CD<sub>3</sub>OD in CDCl<sub>3</sub>) 2.54(3H, d, J=1.1Hz), 6.52(1H, q, J=1.1Hz), 7.42(1H, dd, J=7.6 and 8.0Hz), 7.63(1H, d, J=8.0Hz), 7.64(1H, d, J=7.6Hz), 8.23(1H, s);  ${}^{13}$ C-NMR  $\delta$  ppm(100MHz, 10%CD<sub>3</sub>OD in CDCl<sub>3</sub>) 18.3, 115.4, 115.7, 121.0, 124.0, 124.9, 126.1, 130.2, 133.2, 133.5, 149.5, 182.6; UV  $\lambda_{max}$  (MeOH) 233( $\epsilon$ =18000), 321( $\epsilon$ =7300), 378( $\epsilon$ =6000), 395( $\epsilon$ =7600), 415( $\epsilon$ =5600)nm; IR  $\nu_{max}$  (KBr) 3386, 3081, 3040, 2936, 2846, 1633, 1565, 1502, 1440, 1362, 1307, 1130, 950, 848, 755cm<sup>-1</sup>; EIMS (70eV, m/z) 183(M<sup>+</sup>), 154(base peak), 127; HREIMS Calcd for C<sub>12</sub>H<sub>9</sub>NO: 183.0684. Found 183.0680.

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