FIVE NEW DITERPENOIDS FROM ACONITUM DOLICHORHYNCHUM

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Abstract - From the root extracts of Aconitum dolichorhynchum Wang var. subglabratum T. L. Ming, we have isolated five new minor alkaloids: dolichotine A (I), dolichotine B (II), dolichotine C (III), dolichotine D (IV) and dolichotine E (V), besides yunaconitine (VI), 8-deacetylyunaconitine (VII), crassicauline A (VIII), talatisamine (IX), columbidine (X) and cammaconine (XI). The structures of these alkaloids were determined with the aid of spectral data and correlation with compounds of established structures. The structure of dolichotine C was confirmed by synthesis from talatisamine and partial hydrolysis. All the new alkaloids are characteristic of aromatic acid ester or palmitic acid ester substituted at C8 of C19-diterpenoid alkaloids.

The root of Aconitum dolichorhynchum Wang var. subglabratum T. L. Ming was collected in Zhongdian, the northwest of Yunnan. Being strong poisonous, it has been used to smear on an arrowhead to kill animal. After our investigating this plant material, it showed that the major component is yunaconitine in 0.13 % yield. The toricity of yunaconotine was reported. From the root of A. dolichorhynchum, five new minor alkaloids - dolichotine A (I), dolichotine B (II), dolichotine C (III), dolichotine D (IV) and dolichotine E (V), as well as six known alkaloids - yunaconitine (VI), 8-deacetylyunaconitine (VII), crassicauline A (VIII), talatisamine (IX), columbidine (X) and cammaconine (XI), have been isolated. In this report we wish to describe the separation and structure determination of these compounds. The crude alkaloid mixture was obtained by acidification with 28 H₂SO₄, basification with 30% NH₄OH to pH 8-9 and extraction with CHCl₃. Column chromatography and preparative tlc afforded eleven alkaloids.

- (I) Dolichotine A R1=As; R2=Ac
- (II) Dolichotine B R1-Vr; R2-Ac
- (III) Dolichotine C R1=Cn; R2=Ac
- (IX) Talatisamine R1=R2=H
- (IXa) 14-Acetyltalatisamine R1-H; R2-Ac
 - (I) Clumbidine R1=C2H5; R2=H

(XI) Cammaconine

Table 1. ¹³C nmr chemical shifts and assignments for talatisamine(IX), 14-acetyltalatisamine(IXa), cammaconine(XI), columbidine(X), dolichotine A(I), dolichotine B(II), dolichotine D(IV), dolichotine E(V), crassicauline A(VIII) and vilmorrianine C(XII). (CDCl₃)

Carbons	IX	IXa	π	X	I	11	IA	XII	٧	AIII
C(1)	86.3	85.5	86.0	85.6	85.1	85.2	85.1	85.1	84.2	84.8
C(2)	25.9	26.0	25.6	26.0	26.1	26.3	26.4	26.4	25.8	25.9
C(3)	32.8	34.2	32.2	32.0	32.2	32+3	34.8	34•9	35•3	35•4
C(4)	38.6	38.5	38.7	38.5	38.1	38.4	39•1	39.1	39•6	39•1
C(5)	38.9	40.9	38.7	38.5	41.4	41.8	49.1	49.2	48.8	48.9
C(6)	24.8	24.9	24.7	23.9	24.7	25.1	83.4	82.6	83.8	83.3
c(7)	46.0	46.3	45•9	40.0	45.6	45 .6	44•9	44.9	49.1	50•4
c(8)	72.8	73.6	73•4	78.2	85.9	86.5	85.8	85.9	85.4	85.3
C(9)	47.1	45.3	47.3	45•4	42.1	42.1	49.2	49•3	40.7	40.9
C(10)	46.0	40.6	45•7	45.7	38.6	39.2	43.9	43.9	44.6	44•9
C(11)	48.8	48.8	48.9	49•1	48.5	49.0	50.6	50.3	50.0	49.8
0(12)	27.9	28.4	28.0	28.9	28.4	28.8	29.2	29.0	34•5	34.5
C(13)	45•1	45.8	37.8	39.1	44.8	45.2	39.8	39.1	75.8	75.1
C(14)	75.6	76.9	75•7	75•1	75•3	75.8	75•3	75•4	79.0	78.9
C(15)	38.8	37.8	39.1	35•2	37.5	37.8	38 .1	37 •9	38.4	38.9
C(16)	82.4	81.7	82.4	82.6	82.7	83.1	82.9	83 .5	84.0	83.9
C(17)	62.9	62.1	62.9	62.4	61.6	61.7	61.3	61.7	62.3	61.6
0(18)	79•5	80.0	68.9	79•2	79-1	79•4	80.4	80.6	80.4	80.4
C(19)	53•3	53.2	53•5	53.2	52.7	53•2	54.0	53.8	54.2	53.8
N-CH ₂	49•5	49•3	49•5	49•4	49.0	49•3	49•3	49.0	49•4	49.1
ĊH3	13.7	13.4	13.4	13.6	13.1	13.3	13.0	13.4	13.6	13.1
C(1)'	56.3	56.0	56.1	56.1	55•1	56.0	56.6	56.6	55•7	56.1
c(6)'	-	-	-	-	-	-	58.1	57.8	58 .0	5 7. 8
0(16)1	56.4	56.0	55.8	56•4	55•8	56.0	55•9	56.0	58 .9	58.7
0(18)1	59•5	59•4		59•5	59•1	59•4	59•1	59.1	59•4	59•1
C(8)-OCH2		-		55•9	-	-	-	-	-	-
çн3		-		16.2	-	-	-	-	-	-
C=0		170.6			171.1	171.5	-	169.8	-	169.8
ĊH ₃		21.2			21.1	21.6	~	21.8	-	21.3
C=0					-	-	172.5	-	172.6	-
CH ₂					-	-	22.6	-	22. 8	-
(CH ₂) ₁₃					-	-	24.0-30.	0 –	24.0-30.	0 -
CH ₃					-	-	11.4		11.6	-
Ç=0					164.3	164.7	166.0	166.2	166.3	166.3
6// 72	2				1 123.8	124.0	123.0	123.0	123.2	123.1
5 人 人	NO STE				2 131.1	112.0	131.8	131.8	132.1	131.9
400H2	осн3				3 113.2	149.8	113.8	113.7	114.2	114.1
5					4 162.9	153.0	163.5	163.5	163.9	163.8
					5 113.2	110.4	113.8	113.7	114.2	114.1
					6 131.1	123.4	131.8	131.8	132.1	131.9
					3' -	56.1	-	-	-	-
					4' 56.2	56.6	55•4	55•4	55•5	55•4

Dolichotine A (I) was obtained as an amorphous compound, [cl] +15.2° (CHCl3), C3MH47NO8 (deduced from ms, 1H- and 13C-nmr spectral data). The 13C-nmr spectrum exhibited 34 lines corresponding to 34 carbon atoms of the molecule (see Table 1). The ir spectrum showed no absorption above 3200 cm-1, indicating the absence of hydroxyl groups in delichotine A. The 1H-nmr spectrum gave signals at 6 1.09 (3H, t, J = 7 Hz, NCH2CH3), 1.79 (3H, s, OCOCH3), 3.24, 3.30, 3.33 (each 3H, s, OCH_3), 3.85 (3H, g, Ar- OCH_3), 4.83 (1H, t, J = 4.5 Hz, C_14 - β -H), 6.91, 7.94 (each 2H, d, J = 9 Hz, A2B2 type, Ar-H), showing that dolichotine A is a C19-diterpenoid alkaloid having an Nethyl, three methoxyls, an acetyl and one anisoyl group. Comparison of 1H-nmr spectral data of dolichotine A (I) with those of anisoezochasmaconitine² which appeared a signal at δ 4.10 (18, dd, $J_1 = 6$ Hz, $J_2 = 1$ Hz, $C_6 - B + H$) in its spectrum, showed that dolichotine A has no methoxyl group at C6. The loss of 31 mass unit from the molecular ion to give an intense peak suggested a methoxyl group at C1 of a C19-diterpenoid alkaloid and the methoxyl group at C1 being oforient, 3 which is also supported by the chemical shifts of the 13C-nmr spectrum at \$85.1 (d), 26.1 (t) and 32.3 (t) corresponding to C1, C2 and C3 of dolichotine A, respectively.4 The presence of C18-OCH3 in I was supported by the carbon signals at \$ 59.3 (q) and 79.7 (t) of the 13C-nmr spectrum. 4 According to the biogenesis of C19-diterpenoid alkaloids, 5,6 there is usually β-OCH_λ substituent at C₁₆. Alkaloids without an oxygen substituent at C₁₃, but bearing C8-OBz or C8-OAs and C14-OAc, show a 3H singlet for the acetate methyl group between 6 1.76-1.79 and a 1H triplet for C14-H between & 4.80-4.82.2 However, alkaloids with the reverse arrangement, viz. C8-OAc and C14-OBz or C14-OAs, such as 8-acetyl-14-benzoylneoline, show a 3H singlet for the acetate methyl between & 1.34-1.46 and a 1H triplet for C14-H between \$ 5.00-5.11.7 Because dolichotine A revealed a 3H singlet at 6 1.79 and a 1H triplet at 6 4.83, it was proposed for structure I. Hydrolysis of dolichotine A with 2% KOH in MeOH gave talatisamine and anisic acid. So dolichotine A is 8-anisoyl-14-acetyltalatisamine.

Dolichotine B (II) was obtained as an amorphous compound, $C_{35}H_{49}NO_{9}$ (deduced from ms, ¹H- and ¹³C-nmr spectral data), $[\alpha]_{\mathbf{D}}$ 0° (CHCl₃). The ¹H- and ¹³C-nmr spectral data of dolichotine B (II) are very similar to those of dolichotine A (I), except for differences of signals in the aromatic region. Comparison of aromatic signals of II with those of I revealed that dolichotine B possesses a veratroyl group instead of an anisoyl group as in dolichotine A. In addition II is 30 mass unit more than I, showing that II has one methoxyl group more than I. The ¹³C-nmr data for dolichotine B are given in Table 1. Hydrolysis of dolichotine B with 26 KOH in MeOH gave talatisamine and veratric acid. So dolichotine B is 8-veratroyl-14-acetyltalatisamine.

Dolichotine C (III) was obtained as an amorphous compound, $C_{35}H_{47}NO_{7}$ (derived from ms, $^{1}H_{-}$ and $^{1}3C_{-}$ nmr spectral data). The ir and $^{1}H_{-}$ nmr spectral data showed that dolichotine C possesses a cinnamoyl group⁸ instead of an anisoyl group as in dolichotine A and a veratroyl group as in dolichotine B. The signal for the acetoxyl protons in the $^{1}H_{-}$ nmr spectrum of dolichotine C occurs at δ 1.95, which is at a little higher field than the normal signal⁹ but is at lower field than the corresponding signal in dolichotine A (δ 1.79) and dolichotine B (δ 1.74). In dolichotine C which was esterified with cinnamic acid, the plane of the benzene ring is further away from the acetate methyl group which, however, is close to the double bond. Hence the smaller shift (δ 1.95) in the acetoxyl protons signal must be due to the anisotropic action of the double bond, which is not as strong as that of the aromatic ring. In addition, chasmanthinmine⁸ manifests a δ H singlet for the acetoxyl protons at δ 1.77 and a 1H signal for C_{14} —H at δ 4.80. But dolichotine C shows a

3H singlet (\$\delta\$ 1.95) and a 1H signal at \$\delta\$ 4.82. So dolichotine C was designated as 8-trans-cinnamoyl-14-acetyltalatisamine. It was confirmed by synthesis from talatisamine (IX). Acetylation of IX with Ac20 and pyridine gave 14-acetyltalatisamine (IXa). The \$^{1}\$C-mmr spectral data for IXa are given in Table 1. Cinnamoylation of IXa with cinnamic anhydride and p-TsOH in toluene afforded 8-trans-cinnamoyl-14-acetyltalatisamine (IXb), which is identical in every respect with III. Furthermore, partial hydrolysis of IXb and III with dioxane-H20 (1:1)11 gave IXa showing one spot on tlc. Therefore, dolichotine C has structure III.

- (IV) Dolichotine D R1=COC₁₅H₃₁; R2=As
- (XII) Vilmorrianine C R1-Ac; R2-As
- (XIII) Chasmanine R1=R2=H

- (V) Dolichotine E R1=H; R2=COC15H31; R3=As
- (VI) Yunaconitine R1=OH; R2=Ac; R3=As
- (VII) 8-Deacetylyunaconitine R1=OH; R2=H; R3=As
- (VIII) Crassicauline A R1=H; R2=Ao; R3=As
- (XIV) Bikhaconine R1=R2=R3=H

Dolichotine D (IV) was an amorphous compound, C49H77NOg (derived from the ms, 1H- and 13C-nmr spectral data). The 13c-nmr spectral data are given in Table 1. The 1H-nmr spectrum showed that dolichotine D has the functional formula of C19-diterpenoid alkaloids - C19H22(NCH2CH3)(OCH3)4 (CH30C6H4COO)(C15H31COO).12 The mass spectrum exhibited two fragments at m/z 256 and m/z 135 corresponding to palmitic acid and anisoyl group, respectively. According to the ms fragmentation patterm^{3,13} of C19-diterpenoid alkaloids substituted with ester, molecular ion preferred losing C8-ester to losing C1-methoxy when the ester group attaching at C8 is large; whether first losing Cg-ester or C1-methoxy the intense characteristic peak is always corresponding to the fragment ion which has just lost C1-OCH2. The mass spectrum of dolichotine D exhibited an intense peak at m/z 536 (M+- C15H31COOH - OCH3, 90) showing a x-methoxyl group at C1, and a fragment peak at m/z 567 $(M^+- C_{15}H_{31}COOH, 27)$ but no fragment peak at m/z 671 $(M^+- 152)$ or m/z 688 $(M^+- 135)$, so this revealed that dolichotine D possesses a palmityl group at Cg. Comparison of 130-nmr spectral data of dolichotine D with those of vilmorrianine C (XII) indicated that dolichotine D possesses a palmityl group instead of an acetyl group as in vilmorrianine C. Hydrolysis of delichotine D with 2% KOH in MeOH gave chasmanine, palmitic acid and anisic acid. Dolichotine D was designated as structure IV.

Dolichotine E (V) was obtained as an amorphous compound, $C_{49}E_{77}NO_{10}$ (deduced from the ms, $^{1}H_{-}$ and $^{1}3C_{-}$ nmr spectral data). The $^{1}3C_{-}$ nmr spectral data for dolichotine E are given in Table 1. Compari-

son of ir, ¹H- and ¹³C-nmr spectra of dolichotine E with those of dolichotine D revealed that there is a hydroxyl substituent in dolichotine E. Alkaloids with a hydroxyl group at C₁₃, bearing C₁₄-OAs and C₁₆-OCH₃, show a 1H doublet for the C₁₄-H between \$5.00-5.11 and a 3H singlet for the C₁₆-OCH₃ between \$3.50-3.65. Examples are crassicalline A, crassicalsine and forestine. ¹⁴ But alkaloids without a hydroxyl group at C₁₃, still bearing C₁₄-OAs and C₁₆-OCH₃, show a 1H triplet for the C₁₄-H between \$5.00-5.10 and a 3H singlet for the C₁₆-OCH₃ between \$3.30-3.40, such as dolichotine D, fotesaconitine and crassicaldine. ¹⁴ Because dolichotine E shows a 1H doublet at \$5.10 and a 3H singlet at \$3.54, it indicated that V possesses a hydroxyl group at C₁₃. Hydrolysis of V with 2\$ KOH in MeOH gave bikhaconine, palmitic acid and anisic acid. So dolichotine E was assigned as structure V.

EXPERIMENTAL

Melting point was determined on a Thomas-Kofler hot stage equipped with a microscope. Ir spectra were taken on Perkin-Elmer model 577 spectrophotometer. ¹H-Nmr spectra were run on BRUCKER WH-90 spectrometer with TMS as an internal reference. ¹³C-Nmr spectra were operated on BRUCKER AM-400 spectrometer in CDCl₃; chemical shifts are reported in ppm downfield from TMS. Mass spectra were measured on a Finnigan-4510 instrument.

<u>Plant material</u>. The root of <u>Acconitum</u> <u>dolichorhynchum</u> Wang var. <u>subglabratum</u> T. L. Ming was ∞ llected in Zhongdian, Yunnan, China. This plant was identified by Professor Ming Tianlu, Kunming
Institute of Botany.

Extraction and fractionation. Powdered roots of A. dolichorhynchum (10.3 Kg) were extracted with 85% ethanol (4 x 4 1) at room temperature for one week. After evaporation of the solvent, the residue (304 g) was acidified with 2% $\rm H_2SO_4$ (2 1) and extracted with CHCl₃ (5 x 1.5 1) to give a crude alkaloid mixture - base A (160 g), which is due to unperfect acidification. The acidic water phase was basified with 30% NH₄0H and extracted with CHCl₃ (5 x 1.5 1) to give a crude alkaloid mixture - base B (100 g).

Isolation of dolichotine A (I), dolichotine B (II), dolichotine C (III), dolichotine D (IV), dolichotine E (V), crassicauline A (VIII) and yunaconitine (VI). A solution of base A (160 g) in CHCl₃ (250 ml) was evaporated with 400 g of silica gel. The mixture was placed on the top of a column filled with 4.5 Kg of silica gel and eluted with petroleum ether-acetone (9:5) to afford fraction A₁ (91 g), and with acetone to afford fraction A₂ (4.5 g). A₁ was chromatographed on a column containing 2 Kg of Al₂O₃ (neutral, activity II) and eluted with petroleum ether, petroleum ether-EtOAc (98:2) to afford dolichotine D (IV, 31 mg), petroleum ether-EtOAc (98:6) to give fraction A₃ (1.5 g), petroleum ether-EtOAc (98:10) to afford fraction A₄ (2.1 g), petroleum ether-EtOAc (98:20) to give dolichotine E (V, 30 mg), petroleum ether-EtOAc (98:30) to give orassicauline A (VIII, 1.5 g) and EtOAc. By preparative the over silica gel (GF₂S₄, cyclohexane-20% Et₂NH), A₃ and A₄ afforded dolichotine A (I, 120 mg), dolichotine B (II, 140 mg) and dolichotine C (III, 20 mg). Crystallization of A₂ from ether gave yunaconitine (VI, 3.5 g).

Isolation of yunaconitine (VI), 8-deacetylyunaconitine (VII), talatisamine (IX), columbidine (X) and cammaconine (XI). A solution of base B (100 g) in CHCl₃ (200 ml) was evaporated with 300 g of Al₂O₃ (neutral, activity II). The mixture was placed on the top of a column filled with 2.5 Kg of Al₂O₃ and eluted with ether to give columbidine (X, 30 mg), hexane-EtOAc (90:1) to afford talatisamine (IX, 2.4 g), hexane-EtOAc (1:2) to give yunaconitine (VI, 10 g), hexane-EtOAc (1:50) to give 8-deacetylyunaconitine (VII, 15 mg) and EtOAc to afford cammaconine (XI, 32 mg).

Identification of dolichotine A (I). Dolichotine A is an amorphous compound, [\$\times 7_p + 15.2\circ\$ (CHCl3);

14 nmr (CDCl3) \$\int 1.09 (3H, \(\frac{t}\), J = 7 Hz, NCH2CH3), 1.79 (3H, \(\frac{s}\), 0CCH3), 3.24, 3.30, 3.33 (each

3H, \(\frac{s}\), 0CH3), 3.85 (3H, \(\frac{s}\), 0CH3), 4.83 (1H, \(\frac{t}\), J = 4.5 Hz, C14-\(\beta\)-H), 6.91, 7.94 (each 2H, \(\frac{d}\), J =

9 Hz, A2B2 type, Ar-\(\frac{H}\)); ir (KBr) 1735, 1700 (ester), 1600, 1510, 850, 770 (Ar); ms m/z 597 (M+,

1.6), 566 (M+- OCH3, 47.1), 445 (M+- CH3OC6H4COOH, 68.1), 414 (M+- OCH3 - CH3OC6H4COOH, 80), 152

(CH3OC6H4COOH, 100); the \(^{13}C-NMR spectral data are given in Table 1.

Hydrolysis of dolichotine A. Dolichotine A (50 mg) was dissolved in 5 ml of 2% KOH in MeOH and allowed to stand at room temperature for 6 h. Removal of solvent under reduced pressure gave a residue which was mixed with a small amount of H₂O and extracted with CHCl₃. The CHCl₃ extract was dried over anhydrous Na₂SO₄ and evaporated to give a pale yellow residue which was crystalized from acetone to give colorless needles (25 mg) being identical with those of talatisamine in its co-tlc, ir spectrum and ¹H-NMR spectrum. The water phase was acidified with 2% H₂SO₄ and extracted with CHCl₃ to give anisic acid (10 mg), mp 183-184 °C; ms m/z 152 (M+, 100), 135 (M+-OH, 95); ir (KBr) 2720 (br, COOH), 1116, 830, 770 (Ar), which was identical with those reported.15,16

Identification of dolichotine B (II). Dolichotine B is an amorphous compound, (cd)_b 0° (CHCl₃);

1H nmr (CDCl₃) 6 1.10 (3H, t, J = 7 Hz, NCH₂CH₃), 1.74 (3H, g, OCOCH₃), 3.26, 3.30, 3.38 (each 3H, g, OCH₃), 3.90, 3.98 (each 3H, g, OCH₃), 4.79 (1H, t, J = 4.5 Hz, C₁₄-H), 7.02 (1H, d, J = 9 Hz, Ar-5'-H), 7.60 (1H, dd, J₁ = 9 Hz, J₂ = 3 Hz, Ar-6'-H), 7.73 (1H, d, J = 3 Hz, Ar-2'-H);

ir (KBr) 1730, 1700 (ester), 1650, 1510, 910, 765 (Ar); ms m/z 627 (M+, 1), 596 (M+- OCH₃, 41),

445 (M+- C₂H₆O₂C₆H₃COOH, 40), 414 (M+- C₂H₆O₂C₆H₃COOH - OCH₃, 100), 386 (M+- C₂H₆O₂C₆H₃COOH - CH₃COO, 20), 182 (C₂H₆O₂C₆H₃COOH, 76); the ¹³C-NMR spectral data are given in Table 1.

Hydrolysis of dolichotine B. Dolichotine B (50 mg) was hydrolized by the same method to dolichotine A as colorless needles (26 mg) which was identical as talatisamine (IX) by comparison of the mp 145-146 °C, co-tlc, ir spectrum and ¹H-NMR spectrum with those of talatisamine, and veratric acid, mp 180-181 °C; ms m/z 182 (M⁺, 60), 135 (100); ir (KBr) 2710 (br, COOH), 910, 765 (Ar), which were identical with those reported. 17,18

Identification of dolichotine C (III). Dolichotine C is an amorphous compound, ¹H nmr (CDCl₃) **6** 1.08 (3H, <u>t</u>, J = 7 Hz, NCH₂CH₃), 1.95 (3H, <u>s</u>, OCOCH₃), 3.22, 3.25, 3.36 (each 3H, <u>s</u>, OCH₃), 4.82 (1H, <u>t</u>, J = 4.5 Hz, C₁₄-\(\beta\) H, 6.32, 7.60 (each 1H, <u>d</u>, J = 16 Hz, trans-\(\beta\)CCH₃, 6.90 - 7.50 (5H, \(\beta\), Ar-\(\beta\)); ir (KBr) 1730, 1705 (ester), 1680, 970 (trans-double bond), 1600, 1460, 770, 710 (Ar); ms m/z 593 (M⁺, 0.1), 562 (M⁺- OCH₃, 31), 445 (M⁺- C6H₅C₂H₂COOH, 100), 414 (M⁺- C6H₅C₂H₂COOH - OCH₃, 82), 386 (M⁺- C6H₅C₂H₂COOH - OCCH₃, 42), 148 (C6H₅C₂H₂COOH, 70).

Preparation of 14-acetyltalatisamine (IXa). A solution of talatisamine (100 mg) in 5 ml of acetic anhydride and 5 ml of pyridine was allowed to stand at room temperature for 2 days. To the residue obtained on evaporation of solvent was added 10 ml of H_2O ; the mixture was basified with 30% NH_4OH to pH 8 and then extracted with CHCl₃. The CHCl₃ extract was dried over Na_2SO_4 and evaporated to give a brownish foam (100 mg) containing 14-acetyltalatisamine (IXa) as major component in 91 % yield. ¹H Nmr (CDCl₃) 6 1.09 (3H, \pm , J = 7 Hz, NCH₂CH₃), 2.09 (3H, \pm , 0COCH₃), 3.28, 3.30, 3.31 (each 3H, \pm , 0CH₃), 5.09 (1H, \pm , J = 4.5 Hz, C_{14} - β -H); ms m/z 463 (M+, 1), 432 (M+- OCH₃, 100); the ¹³C-NMR spectral data are given in Table 1. Structure IXa was assigned on the basis of ¹H- and ¹³C-nmr spectra, and was identical with an authentic sample. ¹⁹

Synthesis of dolichotine C. IXa (100 mg) and cinnamic anhydride (1.5 g) were heated at 110 °C with toluene (50 ml) and p-TsOH (10 mg) for 12 h. This reaction gave 8-cinnamyl-14-acetyltalatisamine (IXb, 20 mg) in 12.8 % yield. IXb was identical with dolichotine C on the basis of their 1H-nmr spectrum, ir spectrum, ms and co-tlc.

Partial hydrolysis of dolichotine C and IXb. 11 Dolichotine C (10 mg) and IXb (15 mg), dissolved in 10 ml of dioxane-H₂O (1:1), were heated on an oil bath at 120° C with stirring for 1 h and extracted with CHCl₃. The CHCl₃ extract was purified by preparative tlc over silica gel (GF₂₅₄, cyclohexane-20% Et₂NH) and gave IXa (15 mg, in 78 % yield) which was identical with the IXa prepared above on the basis of their co-tlc, ir spectra, ms and ¹H-nmr spectra.

Identification of dolichotine D (IV). Dolichotine D is an amorphous compound, ${}^{1}H$ nmr (CDCl₃) δ 1.05 (3H, \pm , J = 7 Hz, NCH₂CH₃), 3.17, 3.28, 3.39 (each 3H, \pm , OCH₃), 3.84 (3H, \pm , Ar-OCH₃), 4.05 (1H, \pm d, J₁ = 6 Hz, J₂ = 1 Hz, C₆- β - \pm), 5.03 (1H, \pm , J = 4.5 Hz, C₁₄- β - \pm), 6.89, 7.99 (each 2H, \pm d, J = 9 Hz, A₂B₂ type, Ar- \pm); ir (KBr) 2920, 2850, 2820 (-CH₃, -CH₂-), 1715, 1710 (ester), 1600, 1510, 1460, 850, 770 (Ar); ms m/z 823 (M⁺, 0.01), 792 (M⁺- OCH₃, 16), 567 (M⁺- C₁₅H₃₁COOH, 27), 536 (M⁺- OCH₃ - C₁₅H₃₁COOH, 90), 416 (M⁺- C₁₅H₃₁COOH - OCH₃C₆H₄COO, 100), 384 (M⁺- OCH₃ - C₁₅H₃₁COOH - OCH₃C₆H₄COOH, 66), 256 (C₁₅H₃₁COOH, 25), 238 (C₁₅H₃₀COO, 60), 135 (OCH₃C₆H₄COO, 78); the 13C-nmr spectral data are given in Table 1.

Eydrolysis of dolichotine D. With the same method to dolichotine A, dolichotine D (15 mg) was hydrolized and gave a compound (6.5 mg) which was identical with chasmanine (XIII) on the basis of their co-tlc, ir spectra and mass spectra. On the water phase was acidified with 2% H₂SO₄ and extracted with CHCl₃. The CHCl₃ extract was dried over anhydrous Na₂SO₄ and evaporated to give a residue which was further esterified with 1% H₂SO₄ in MeOH (10 ml) and then operated on go-ms to show molecular ion peaks at m/z 166 and m/z 270 for methyl anisic ester and methyl palmitic ester, respectively.

Identification of dolichotine E (V). Dolichotine E is an amorphous compound, ir (KBr) 3500 (OH), 2920, 2850, 2820 (-CH₃, -CH₂-), 1715, 1710 (ester), 1604, 1576, 1455, 850, 770 (Ar); ¹H nmr (CDCl₃) \$ 1.05 (3H, t, J = 7 Hz, NCH₂CH₃), 3.18, 3.25, 3.33, 3.54 (each 3H, s, OCH₃), 3.77 (3H, s, Ar-OCH₃), 4.09 (1H, dd, J₁ = 6 Hz, J₂ = 1 Hz, C₆-β-H), 5.10 (1H, d, J = 4.5 Hz, C₁4-β-H), 5.70 - 5.90 (1H, br, disappearing after exchanging with D₂O, OH), 6.90, 8.10 (each 2H, d, J = 9 Hz, Ar-H); ms m/z 839 (M+, 0.02), 808 (M+- OCH₃, 6), 583 (M+- C₁5H₃1COOH, 35), 552 (M+- OCH₃ -

 $C_{15}H_{31}COOH$, 50), 432 (M+- $C_{15}H_{31}COOH$ - $CH_{30}C_{6}H_{4}COO$, 20), 256 ($C_{15}H_{31}COOH$, 20), 238 ($C_{15}H_{30}CO$, 60), 135 ($CH_{30}C_{6}H_{4}CO$, 100); the 13C-nur spectral data are given in Table 1.

Hydrolysis of dolichotine E. With the same method to dolichotine D, dolichotine E (15 mg) was hydrolized and gave a compound (6 mg) which was identical with bikhaconine (XIV) on the basis of their co-tlo, ir spectra and mass spectra. 21 The water layer was acidified with 25 H₂SO₄ and extracted with CHCl₃. The CHCl₃ extract was dried over anhydrous Na₂SO₄ and evaporated to give a residue which was esterified with 15 H₂SO₄ in MeOH (10 ml) and then operated on go-ms to show molecular ion peaks at m/z 166 and m/z 270 for methyl anisic ester and methyl palmitic ester, respectively.

Identification of yunaconitine (VI). VI was crystallized as rhombus from ether, mp 140-141 °C; ms m/z 659 (M+, 2), 628 (M+- OCH3, 50), 135 (CH30C6H4CO, 65), 43 (CH3CO, 75); ir (KBr) 3450 (OH), 1715, 1708, 1251 (ester), 1608, 1510, 1480, 850 (Ar); [†]H nmr (CDCl₃) δ 1.11 (3H, \pm , J = 7 Hz, NCH2CH3), 1.34 (3H, \pm , 0COCH3), 3.16, 3.26, 3.30, 3.55 (each 3H, \pm , 0CH3), 3.87 (3H, \pm , Ar-OCH3), 4.06 (1H, \pm dd, \pm

Identification of 8-deacetylyunaconitine (VII). VII was obtained as an amorphous compound, ms m/z 617 (M+, 0.1), 586 (M+- OCH3, 45), 135 (CH30C6H4CO, 50); ir (KBr) 3530, 3420 (OH), 1705, 1250 (ester), 1602, 1520, 840, 770 (Ar); ¹H nmr \bullet 1.12 (3H, \bullet , J = 7 Hz, NCH2CH3), 3.25, 3.29, 3.31, 3.41 (each 3H, \bullet , OCH3), 3.86 (3H, \bullet , Ar-OCH3), 4.08 (1H, \bullet , J₁ = 6 Hz, J₂ = 1 Hz, C6- \bullet -H), 5.02 (1H, \bullet , J = 4.5 Hz, C₁₄- \bullet -H), 6.80, 8.03 (each 2H, \bullet , J = 9 Hz, AB q, Ar-H). It was identified as 8-deacetylyunaconitine by comparison of its spectral data with those reported.²²

Identification of crassicauline A (VIII). VIII was crystallized as prism from acetone, mp 162-163 °C; ms m/z 643 (M+, 1), 612 (M+-0CH₃, 100), 583 (M+-CH₃COOH, 50), 552 (M+-0CH₃-CH₃COOH, 80), 135 (CH₃OC₆H₄CO, 50); ir (KBr) 3500 (OH), 1725, 1710, 1256 (ester), 1600, 1508, 849, 770 (Ar); ¹H nmr (CDCl₃) & 1.06 (3H, \pm , J = 7 Hz, NCH₂CH₃), 1.33 (3H, \pm , OCCH₃), 3.20, 3.27, 3.50 (each 3H, \pm , OCH₃), 3.91 (3H, \pm , Ar-OCH₃), 4.04 (1H, \pm d, J = 6 Hz, J = 1 Hz, C₆- \pm H), 4.84 (1H, \pm d, J = 4.5 Hz, C₁4- \pm H), 7.07, 8.07 (each 2H, \pm d, J = 9 Hz, AB q, Ar-H). The ¹³C-nmr spectral data are given in Table 1. Hydrolysis of VIII (20 mg) with 25 KOH in MeOH (15 ml) gave bikhaconine (9 mg) which was identified as crassicauline A by comparison of its spectral data and co-tlc with those of an authentic sample.²³

Identification of talatisamine (IX). IX was crystallized as needles from acetone, mp 141-142 °C; ms m/z 421 (M+, 2), 390 (M+- OCH3, 100); ir (KBr) 3520, 3410 (OH), 1100, 1080 (C-0); ¹H nmr (CDCl3) δ 1.07 (3H, t, J = 7 Hz, NCH2CH3), 3.26, 3.31, 3.40 (each 3H, s, OCH3), 4.20 (1H, t, J = 4.5 Hz, C14- β -H3; ¹³C-nmr spectral data (see Table 1). It was proved to be talatisamine by comparison of its spectral data and co-tlc with those reported.²⁴

Identification of columbidine (X). X was isolated as an amorphous compound, $C_{26}H_{4}3NO_{5}$ (deduced from the ms, $^{1}H_{-}$ and $^{1}3C_{-}$ nmr spectral data); ms m/z 449 (M+, 2), 418 (M+- 0CH3, 100), 404 (M+- 0C2H5, 5), 386 (M+- 0C2H5 - H20, 2), 373 (M+- 0CH3 - 0C2H5, 4); ir (KBr) 3540 (OH), 2965, 2910, 2870, 2810, 1493, 1382, 1260 (-CH3, -CH2-); $^{1}H_{-}$ nmr (CDC13) & 1.04, 1.07 (each 3H, $_{\pm}$, J = 7 Hz, NCH2CH3), 3.20, 3.23, 3.29 (each 3H, $_{\pm}$, 0CH3); $^{1}3C_{-}$ nmr spectral data (see Table 1). It was identified as columbidine by comparison of its spectral data with those reported.4

Identification of cammaconine (XI). XI was crystallized as prism from CHCl₃-MeOH (1:1), mp 135-136 °C; ms m/z 407 (M+, 1), 376 (M+- OCH₃, 100); ir (KBr) 3530, 3460 (OH), 2930, 2860, 1382, 1100 (-CH₃, -CH₂-); ¹H nmr (CDCl₃) δ 1.06 (3H, \pm , J = 7 Hz, NCH₂CH₃), 3.21, 3.29 (each 3H, \pm , OCH₃), 4.20 (1H, \pm , J = 4.5 Hz, C₁₄- β - \pm); ¹3C-nmr spectral data (see Table 1). It was proved to be cammaconine by direct comparison of its ir spectrum, ¹H- and ¹3C-nmr spectra with those reported.⁴

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