Two New Diterpenoids from Isodon rubescens

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Abstract: Two new diterpenoids taibairubescensin A (1) and B (2) have been isolated from *Isodon rubescens*. The structures of compound 1 and 2 were elucidated as $2\beta.3\beta$ -diacetoxy- $11\beta.13\alpha$ -dihydroxy-ent-kaur-16-en-15-one (1) and $3\beta.11\beta$ -diacetoxy- $2\beta.6\alpha$ -dihydroxy-ent-kaur-16-en-15-one (2) on the basis of spectroscopic analysis.

Keywords: Isodon rubescens, taibairubescensin A, taibairubescensin B, ent-kaurene diterpenoids.

In order to further study on minor diterpenoid constituents of *Isodon rubescens*, we reinvestigated this species, which was collected in Taibai mountain, Shaanxi Province. Two new diterpenoids, taibairubescensins A (1) and B (2), were isolated. In this paper, we present the structure elucidation of these two new diterpenoids.



1.R₁=R₂=OAc, R₃=H, R₄=R₅=OH 2.R₁=OH, R₂=OAc, R₃=OH R₄=OAc, R₅=H 3.R₁=H, R₂=OH, R₃=H, R₄=R₅=OH 4.R₁=OH, R₂=OAc, R₃=R₄=OH, R₅=H

Taibairubescensin A (1), $C_{24}H_{34}O_7$ (FABMS m/z 435[M+1]⁺), an amorphous powder, showed UV and IR absorption bands for the existence of hydroxyl, acetoxyl and a five-membered ring ketone conjugated with an *exo*-methylene functions (240.5nm; 3468, 1740, 1732 and 1649 cm⁻¹). The ¹³C-NMR (**Table 1**) and DEPT spectra of **1** showed signals for this compound with $5 \times \text{CH}_3$, $5 \times \text{CH}_2$, $5 \times \text{CH}$, $4 \times \text{C}$, two olefinic carbons, one ketonic carbon and two ester carbonyl carbons. These data suggested that **1** possessed a basic skeleton of *ent*-kaur-16-en-15-one with two acetoxyls and two hydroxyls. The ¹H-, ¹³C-NMR data of **1** were very similar to those of deacetylisodopharicin A (**3**)¹ except for one more acetyl groups. Comparison of their ¹³C-NMR data indicated that the difference between **1** and **3** was only in A ring. This ment

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that two hydroxyls were at C-11 β and C-13 α , and two acetoxyls were in A ring, respectively, in compound 1. In the ${}^{1}H^{-1}H$ COSY spectrum of 1, the signal at δ 4.98 (1H, d, J = 2.6 Hz, H-3 α) showed correlation with the signal at δ 5.24 (1H, ddd, J = 12.3, 2.6, 3.9 Hz, H-2 α), the latter showed correlation with both signals at δ 1.59 (1H, dd, J = 11.9, 3.9 Hz, H-1 α) and δ 1.91 (1H, dd, J = 11.9, 12.3 Hz, H-1 β). Thus two acetoxyl groups should be located at the C-3 and C-2 positions, respectively. The relative configurations were established as 2 β -OAc and 3 β -OAc by considering the coupling constants of H-2 and H-3. These cases were further confirmed by NOESY spectrum of 1. Therefore, compound 1 should be elucidated as 2 β -3 β -diacetoxy-11 β ,13 α -dihydroxy-ent-kaur-16-en-15-one.

Taibairubescensin B (2), $C_{24}H_{34}O_7$ [HRFABMS(pos.) m/z: 435.2364[M+1], calc. 435.2382], an amorphous powder, showed UV and IR absorption bands for the existence of hydroxyl and a five-membered ring ketone conjugated with an *exo*-methylene functions (243.5nm; 3473, 1738, 1648 cm⁻¹). The ¹³C-NMR (**Table 1**) and DEPT spectra of **2** clearly indicated that the compound **2** was an *ent*-kaurene diterpenoid derivative with two acetoxyl groups and two hydroxyl groups. The ¹³C-NMR spectrum of **2**, compared with that of Lusanrubescensin D (4)², differed from that of **4** only in chemical shift at C-11. The chemical shift of the C-11 is δ 65.1 in **4**, but it is δ 68.5 in **2**. This fact indicated that the acetoxyl group at the C-11β position in **2** had replaced the hydroxyl group in **4**. These assignments were further confirmed by ¹H-¹H COSY and HMBC spectra of **2**. Thus, taibairubescensin B (**2**) was established as 3β.11β-diacetoxy-2β,6α-dihydroxy-*ent*-kaur-16-en-15-one.

Carbon 2 Carbon 2 2 Carbon Carbon 379t 1 43.2t 7 32.5 t 41.1t 13 75.3 s 37 0d 19 21.3q 22 6q 2 67.5d 8 65.0d 52.8 s 48.4s 14 18.3q 19.9q 44.9 t 37.7 t 20 3 9 76.6d 80 9d 61.9d 62 9d 15 207.3 s208.7s OAc 170.4s 171 9s 4 10 38.1s 38.48 39 6 s 39 4s 16 151.9 s 149.1s 170 6s 169.8s 5 48.6d 48 4d 11 66.7d 68.5d 17 113.7 t 113.7t 20 9q 21.1q 6 17.6 t 65.8d 12 48.2 t 37..3t 18 27.8q 28.2q 21.0q

Table 1. 13C NMR data* for 1 and 2

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^{*}Recorded in CDCl₃; chemical shift values reported as δ values (ppm) from TMS at 100.6MHz.