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Synthesis of licochalcone analogues with increased *anti*-inflammatory activity

Si-Jun Kim^a, Cheol Gi Kim^a, So-Ra Yun^a, Jin-Kyung Kim^b, Jong-Gab Jun^{a,*}

^b Department of Chemistry and institute of Applied Chemistry, Hallym University, Chancheon 200-702, Republic of Korea

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ABSTRACT

Licohalcones have been reported to have various biological activities. However, most of licochalcones also showed cytotoxicity even though their versitile utilities. Licochalcones B and D, which have common substituents at aromatic ring B, are targeted to modify the structure at aromatic ring A for inflammatory studies. Licochalcone derivatives (**1–6**) thus prepared are compared for their suppression ability of nitric oxide (NO) production and showed 9.94, 4.72, 10.1, 4.85, 2.37 and 4.95 μ M of IC₅₀ values, respectively. © 2013 Elsevier Ltd. All rights reserved.

Licochalcone A (LicoA), licochalcone B (LicoB), licochalcone C (LicoC), licochalcone D (LicoD), echinatin and Isoliquiritigenin are major active components in licorice which is a traditional medicine used in the Northeast Asia for the treatment of ulcer, asthma, inflammation and other diseases.¹ Licochalcones have been isolated and characterized from the root of *Glycyrrhiza inflata*, and have been reported to show various biological properties, including antibacterial,² antitumor,³ *anti*-inflammatory,⁴ and antioxidative⁵ activities. *G. inflata* is the main species in licorice and contains about 40 kinds of flavonoids of which having no hydroxyl at the position 2' (or 6') which is different from usual flavonoid have been found to show the major contribution on biological activities.⁶ These unusual chalcones are called retrochalcone or 'reversely constructed chalcone' in which the ring A would be derived from shikimate and the ring B from polyketide of malonate orgin.⁷

The *anti*-inflammatory activities of licochalcones as shown in Scheme 1 were compared by the inhibition effect of the mast cell degranulation which play a key role in allergic inflammation in RBL (rat basophilic leukemia)-2H3 cells.⁸ Also, the *anti*-inflammatory activities were compared by the inhibition effect of NO production in inflammatory regions.⁹ The 50% inhibitory concentration (IC₅₀) against degranulation, 30% cytotoxicity (CC₃₀), and IC₅₀ of LPS-induced NO production of each licochalcones are listed in Scheme 1. Since LicoA, LicoC and LicoD exhibited similar inhibitory effects on the degranulation with the IC₅₀ at 17, 24 and

21 μ M. Also, LicoB and LicoD showed IC₅₀ of LPS-induced NO production at 2.3 and 2.2 μ M, respectively. From these inhibitory results, we found that LicoB and LicoD having common substituents at aromatic ring B showed higher activity with relatively lower cytotoxicity than the others.

The structure of LicoB is similar to echinatin except the presence or absence of 3-hydroxy substituent at ring B, however, the *anti*-inflammatory activities are quite different and only LicoB showed the inhibition activity of 2.3 μ M for IC₅₀ of NO production, but both compounds showed lower cytotoxicity. Also the structure of LicoB is exactly same with LicoD on ring B which has 3,4-dihydroxy-2-methoxy substituents, and both showed the higher inhibition activities for IC₅₀ of NO production. In order to find a highly active *anti*-inflammatory licochalcone derivative having lower cytotoxicity, we designed the structures same as LicoB and LicoD at ring B, which have 3,4-dihydroxy-2-methoxy substituents, but modified at ring A to resorcinol type (**2**), catechol type (**3**) and rearranged isopropenyl analogues **5** and **6** (Scheme 2).

The natural LicoB (1) was first identified in 1975 from the roots of *Glycyrrhiza glabra* Linn. (licorice from Sinkiang, China)⁷ and has been reported many biological activities,^{8–10} however, only single report for synthesis has been known.¹¹ Recently, we reported the first total synthesis of LicoD¹² and we used the similar route for the synthesis of LicoB as shown in Scheme 3 since the ring B of LicoB is exactly same as LicoD. The aldehyde portion (9) of protected ring B is condensed with THP protected acetophenone (11) using 3 M NaOH in MeOH to produce the chalcone (12), and following deprotection using Dowex 50X2 resin afforded the LicoB

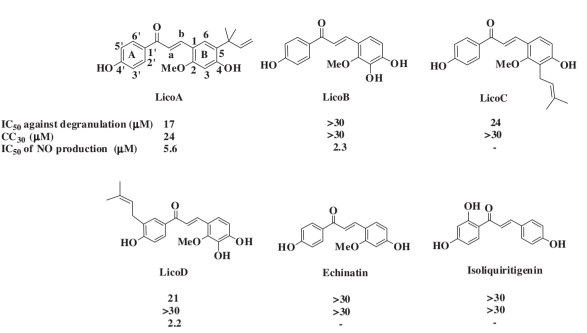




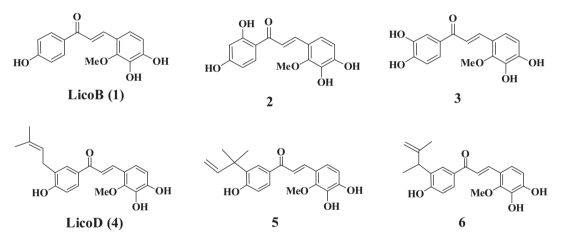


^{*} Corresponding author. Tel.: +82 33 248 2075; fax: +82 33 256 3421. *E-mail address:* jgjun@hallym.ac.kr (J.-G. Jun).

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Scheme 1. anti-Inflammatory activity of licochalcones.



Scheme 2. The designed structures of licochalcone B and D derivatives.

in total 25% yield of 5 steps. The spectral data for LicoB agreed well with the literature values.^{7,10,11}

Resorcinol (2) analogue of LicoB, which is a new compound, was prepared by conventional Claisen-Schmidt condensation of 2,4diethoxymethoxyacetophenone (14) with the benzaldehyde (9) using basic condition followed by deprotection in moderate yields (Scheme 4).

3,4-Diethoxymethoxyacetophenone (19) was prepared from commercially available 3,4-dihydroxybenzaldehyde (16) by protection using diisopropylethylamine with EOMCl followed by CH₃₋ MgCl reaction and PDC oxidation (Scheme 5). It is noteworthy that the deprotonation of 2-hydroxyacetophenone (13) requires stronger base than 3-hydroxy analogue (16) because of intramolecular hydrogen bonding. The acetophenone 19 afforded the catechol analogue (3), which is a known natural licochalcone as tetrahydroxymethoxychalcone,¹³ by condensation and deprotection.

Compound 5, a new compound, is an anologue of LicoD, which has a similar isopropenyl substituent to LicoA. [3,3]-Sigmatropic rearrangement in LicoD synthesis has been applied for this synthesis.¹² 4-Hydroxyacetophenone (**10**) is O-prenylated using

1-bromo-3-methyl-2-butene in K₂CO₃ to give **21** in 98% yield, which is then condensed with the aldehyde 9 to give chalcone 22 in 86% yield, and following water-accelerated Claisen rearrangement¹⁴ produced the analogue **5** in 52% yield (Scheme 6).

Another isopropenyl analogue 6, which is also a new compound, is prepared from **10** using same methododlogy via O-prenylated acetophenone 23 and chalcone 24 as shown in Scheme 7.

The synthetic licochalcones 1-6 are subjected to comparison of their anti-inflammatory activities. The suppression results of nitric oxide (NO) production in lipopolysaccharide (LPS)-stimulated RAW264.7 machrophages for the licochalcones 1-6 are compared in Table 1. Isopropenyl analogue 2 and resorcinol analogue 5 showed 75.9% and 71.1% suppression of NO production at 2 µM, respectively. LicoD (4) and its analogue 6 also showed moderate% inhibition at same concentration, however, LicoB (1) and its catechol analogue 3 showed very low or no inhibition activities. Also, Resorcinol 2, isopropenyl 5, LicoD (4) and its analogue 6 showed 96.5%, 96.1%, 95.4% and 94.3% inhibition, respectively, at 20 µM. Interestingly, catechol analogue showed 92.5% inhibition, however, LicoB (1) still showed only 69.6% inhibition at this concentration.

CC₃₀ (µM)

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