

form a tetrahydropyran system, and a 10-membered ring lactone bearing a side chain with a doubly unsaturated acylenamine moiety. Here we report a formal synthesis of apicularen core unit and its synthetic derivatives. In retrosynthetic analysis, we equated the introduction of an acetate unit to a two-step procedure involving allylation followed by ozonolysis. The reiterations of the allylation-ozonolysis sequence were to be performed in the designated order. These steps not only would "mimic" nature's polyacetate biosynthetic pathway to apicularen A, but also would have the potential of yielding the correct stereochemistry at each chiral center of the target molecule through the judicious choice of appropriate reagents and conditions. After fifth allylation, exposure of intermediate to NaH for 2h followed by addition of water furnished the macrolide lactone. Then terminal homoallyl group was ozonolyzed followed by semicarbazonation and derivatizations. Another derivatizations was carried out as follows, the terminal homoallyl group was homologated by hydroboration-oxidation, then the alcohol obtained was oxidized by TPAP to give the corresponding aldehyde, which was also semicarbazonated and derivatized.

[PD1-3] [2003-10-10 14:00 - 17:30 / Grand Ballroom Pre-function]

Synthesis and COX-2 Inhibitory Properties of Luotonin A Homologues

Park Jae Gyu^o, Kim Dong Hyun, Rahaman A. F. M. Motiur, Chang Hyeun Wook, Lee Eung Seok, Jahng Yurngdong
Yeungnam University

Luotonin A was isolated from *Peganum nigellastrum* Bunge (Zygophyllaceae) which was named Luo-Tuo-Hao in China and used as a Chinese traditional medicine for the treatment of rheumatism, abscess, and inflammation. The basic fractions of *P. nigellastrum* showed antitumor activity, and the origin of such an activity was recently revealed by identifying its constituent luotonin A which inhibited the growth of leukemia P-388 cells ($IC_{50} = 1.8 \mu\text{g/mL}$). Such an intriguing properties of luotonin A led developments of efficient methods for total synthesis. Although Luo-Tuo-Hao has been used more likely for the treatment of inflammation-related symptoms, no efforts towards to explore antiinflammatory activity of *P. nigellastrum* itself as well as its components have been pursued as yet. Our continuing interests in the conformational effects on biological activity as well as search for anti-inflammatory agents spurred us to design a series of luotonin A related compounds in which the dihedral angles between planar 4(3H)-quinazolinone and quinoline rings could be controlled in a regular fashion by a methylene bridge connecting N3 of 4(3H)-quinazolinone and C2 of quinoline. We herein described the synthesis and properties of homologous series of luotonin A.

[PD1-4] [2003-10-10 14:00 - 17:30 / Grand Ballroom Pre-function]

Protective effects of synthetic of 3-Alkoxy-6-allylthiopyridazine against aflatoxin B₁-induced hepatotoxicity

Shin Hea-Soon^o, Kang Joo-Yeon, Park Myung-Sook, Kwon Soon-Kyoung
College of Pharmacy, Duksung Women's University

3-Alkoxy-6-allylthiopyridazine derivatives showed the strongest protective effect against oxidative stress and their anticancer effect determined on the growth of SK-Hep-1 hepatocellular carcinomar cells. The allylthio group as a pharmacologically active group was introduced into pyridazine nucleus and a substituent such as halogen or alkoxy was also introduced into paraposition of allylthio group. Five kinds of 3-alkoxy-6-allylthiopyridazine derivatives were synthesized and their chemoprotective activities examined in rats exposed to aflatoxin B₁-toxicant. Rats were pretreated with five 3-alkoxy-6-allylthiopyridazine derivatives at daily oral doses of 50 mg/kg for 10consecutive days, and during this period with one or three repeated doses of the potent hepatotoxin, aflatoxin B₁. The hepatoprotective effects of the 3-alkoxy-6-allylthiopyridazine derivatives against aflatoxin B₁ administration were showed the significantly normal as compared with control in body and liver weights. Aspartate aminotransferase and alanine aminotransferase levels were markedly elevated after aflatoxin B₁ administration, and pretreatment with 3-alkoxy-6-allylthiopyridazine derivatives, before aflatoxin B₁ administration, resulted in decreased levels of these enzymes. In addition, the 3-alkoxy-6-allylthiopyridazine derivatives,