[PD2-58] [10/17/2002 (Thr) 09:30 - 12:30 / Hall C]

Astragali Radix extract as a therapeutics on osteoporosis

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Aging and estrogen deficiency after menopause induce bone loss and result in osteoporosis. This study was investigated effects of n-hexane fraction (Hx) extracted from Astragali Radix on osteoporosis with osteoblast-like cell line (MG-63 and Saos-2) and an ovariectomized (OVX) rat model. Proliferation of osteoblast-like cells, MG-63 and Saos-2, was tested with MTT and alkaline phosphatase (ALP) assays. Young adult SD rats (10 weeks old) and senile SD rats (52 weeks old) were divided into four groups consisted with sham-operated and placebo dose, OVX and placebo dose as control, OVX and 17β-estradiol at 1 µg/kg/day (E2), and OVX and Hx at 1 mg/kg/day (Hx), respectively. Animals in each group were given i.p. daily for 9 weeks. Trabecular bone areas (TBAs) of tibia and lumbar were measured by bone histomorphometry. In results, Hx increased osteoblast proliferation (approximately 130% of control) but did not increase ALP activities on osteoblast. The TBAs of tibia in young Hx group were increased 174% of control. However, in senile Hx group, the TBAs of lumbar were 126% of control (P<0.017) and those of tibia were only 122% of control. Effects of Astragali Radix extract on osteoporosis were not the same in both old and young rats. (Supported partially by a grant. #HMP-98-D-4-0043, from HPEB, Korea)

[PD2-59] [10/17/2002 (Thr) 09:30 - 12:30 / Hall C]

Analgesic Effect of the Essential Oil from the Rhizomes of Cnidium officinale

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The rhizomes of Cnidium officinale Makino (Umbelliferae) has been used as sedatives and analgesics in the Traditional Medicine in Korea. The essential oil of this rhizomes has been reported to possess CNS depressant activity, however, its analgesic activity has not yet been investigated. We evaluated the analgesic action of the essential oil by administration in oral, on inhalation and after inunction with the phenylquinone- and acetonic acid-induced Writhing test and hot plate test. The essential oil showed the most potent analgesic effect after inunction in the phenylquinone-induced Writhing test, but exhibited almost no analgesic activity in the acetonic acid-induced Writhing test. Moreover, the analgesic action of the essential oil (100mg/kg, p.o.) was more effective than that of acetaminophen (80mg/kg, p.o.), a positive control, in the hot plate test, which suggests that this essential oil may affect central nerve system, revealing an analgesic activity.

[PD2-60] [10/17/2002 (Thr) 09:30 - 12:30 / Hall C]

Antinociceptive effects of 3,4-Dicaffeoyl Quinic Acid of Ligularia fischeri var. spiciformis, caffeic acid and its methyl ester

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The plant Ligularia fischeri var. spiciformis(Compositae) is a candidate for available functional foods. It has been used to treat diabetes mellitus and rheumatoid arthritis. We have reported the isolation of a new eremophilone from the plant with the isolation of hydrophilic constituents, chlorogenic acid, 3,4-di-O-cafeoylquinic acid (1), and 5-O-[1-butyl]-3,4-di-O-cafeoylquinic acid. Compound 1 was again isolated by combination of silica gel- and ODS column chromatography for the anti-nociceptive action. Compound 1 along with caffeic acid(2) and its methyl ester (3) were assayed in hot plate-
and writing tests in the rat. Although the three derivatives of caffeic acid exhibited significant anti-nociceptive effects at 10 mg/kg dose (i.p.), compound 3 was the most potent (activity potency: 3 > 2 > 1). These results suggest that compound 1 is responsible for at least rheumatoid arthritis, and chemical modification of active moiety, caffeoyl group, may increase the activity potency.

[PD2–61] [10/17/2002 (Thr) 09:30 – 12:30 / Hall C]

**Two new acylated neoline derivatives from Aconiti Tuber**

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Aconiti Tuber (Aconitum spp. tuber, Ranunculaceae) which contains bioactive but toxic alkaloids has been used as analgesic, cardiotoxic, diuretic, and stimulant. We have previously reported two new C-19 nortriterpenoid alkaloids and five known nortriterpenoid alkaloids. Further study has now led to the isolation of two new nortriterpenoid alkaloids, 14-O-anisoyleneoline and 14-O-veratroyleneoline. The structures of these compounds were characterized by spectroscopic methods.

[PD2–62] [10/17/2002 (Thr) 09:30 – 12:30 / Hall C]

**Anti-Oxidative compounds from Quercus salicina bark**

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College of pharmacy Chung-Ang University

Quercus species have been used for diarrhea, dysentery, dermatitis, haemoptoe, and haemorrhagia in Korean folk medicine. Specially Quercus salicina have been used for diuretic, anti-inflammatory, antiedemic, and litholytic agent.

In order to investigate the efficacy of antioxidative activity, the activity guided fraction and isolation of physiologically active substance were performed. Its 30%, 60%, 100% MeOH, H2O, and CHCl3 fractions were examined antioxidative activity by DPPH method. It was revealed that H2O, 30% MeOH fractions have significant antioxidative activity.

From 30% MeOH fraction, four phenolic compounds were isolated and elucidated gallic acid, 6'-galloyl salidroside, 2,4-(4-hydroxyphenyl)-ethyl-(6-O-cafeoyl)-gh-D-glucopyranoside, and 4',6'-hexahydroxydiphenoyl salidroside through their physicohemical data and spectroscopic methods.

To investigate the antioxidative activities of each compound, we were measured radical scavenging activity with DPPH method. Gallic acid, 6-galloyl salidroside, and 4',6'-hexahydroxydiphenoyl salidroside showed significant radical scavenging activity against DPPH radical.

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**Poster Presentations - Field D3. Oriental Medicine**

[PD3–1] [10/18/2002 (Fri) 13:30 – 16:30 / Hall C]

**Development of Quantitative Extraction Method of Amygdalin without Enzymatic Hydrolysis from Kyonin(Armeniacae Semen) by High Performance Liquid Chromatography**

Kim DongMin, Hong SeonPyo