protein level. In cells transiently transfected with nuclear facto-kB (NF- κ B) promoter-luciferase reporter construct, this compound clearly inhibited the LPS-stimulated NF- κ B activation. Moreover, this compound inhibited I κ B- κ B-adegradation in a concentration and time-dependent manner. These results indicate that FPP-3 inhibit NO production via inhibition of degradation of I κ B- κ B activation.

[PC1-39] [2003-10-10 09:00 - 13:00 / Grand Ballroom Pre-function]

Anti-Inflammatory action and Cellular Toxicity of Resina Pini on Human Gingival Fibroblast

Suk Kui-Duk, <u>Suh Young-Ah</u>°, Chang Su-Jin College of Pharmacy, *Catholic University of Daegu*

This study was carried out to evaluate the cytotoxicity and anti-inflammatory effects of Resina Pini on cultured human gingival fibloblasts. We carried out a study of cytotoxic effects of Resina Pini on cultured cells by MTT assay. Various treatments on Resina Pini reduced its toxicity on cultured cells in order of natural Resina Pini, water extracted mixture of Resina Pini and Ramus Mori Albae and recrystalized Resina Pini. However, Resina Pini showed harmless levels of cytotoxicity to cultured human gingival fibroblast. Anti-oxidative activity was evaluated by DPPH radical scavenging test, and PGE₂ by PGE₂ EIA system. Resina Pini suppressed productions of free radicals and PGE₂, which causes tissue inflammation and clinical pain. Interestingly, Resina Pini extract samples displayed superior inhibitory activity upon PGE₂ synthesis, compared to contrast group aspirin. This fact may suggest safe and efficient periodontal hygienic and therapeutic uses of Resina Pini.

[PC1-40] [2003-10-10 09:00 - 13:00 / Grand Ballroom Pre-function]

Differentiation and authentication of Panax ginseng (Korea and China), Panax quinquefolius, and development of genetic marker by AFLP analysis.

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Panax ginseng is one of the most important medicinal plant in the Orient. The international trade of ginseng is increasing yearly. The disguise of Chinese and American ginseng into Korean ginseng became a problem in recent years in Korea and an abroad. Obviously, an effective method of authentication of Korean ginseng from others at a DNA level, is necessary for the healthy development of the ginseng market. In order to develop convenient and reproducible methods for the identification of Korean ginseng, amplified fragment length polymorphism (AFLP) analysis was applied within Panax species (Korean cultivatied and wild ginseng, Chinese wild ginseng, American cultivatied and wild ginseng). The genetic distance coefficients between the P. ginseng and P. quinquefolius were high, ranging from 0.573 to 0.692, whereas samples of P. ginseng (cultivatied and wild type) from the different area in Korea and China were very low, ranging from 0.056 to 0.164. By detailed AFLP analysis, some important different bands between wild type of P. ginseng from Korea and China were obtained. These results support that this approach could be applied to distinguish Korean ginseng (Panax ginseng) from others (Chinese and American ginseng) and to authenticate cultivatied and wild ginseng at the molecular level.

[PC1-41] [2003-10-10 09:00 - 13:00 / Grand Ballroom Pre-function]

Antioxidative effect and anti-apoptosis effect of extract from Betula platyphylla var. japonica

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The antioxidant and anticancer properties of a medicinal plant, Betula platyphylla var. japonica were

investigated. The total methanol extract of B. platyphylla var. japonica had protective effects against hydrogen peroxide (H₂O₂) in the Chinese hamster lung fibroblast (V79-4) cell line and induced apoptotic cell death in human promyelocytic leukemia (HL-60) cells, a cancer cell line. B. platyphylla var. japonica extract significantly increased cell viability against H₂O₂. The extract also showed high 1,1-diphenyl-2-picrylhydrazyl (DPPH) radical scavenging activity (IC₅₀ 2.4 mg/ml) and lipid peroxidation inhibitory activity (IC₅₀ below 4.0 mg/ml). Furthermore, B. platyphylla var. japonica extract reduced the number of V79-4 cells arrested in G₂/M in response to H₂O₂ treatment and increased the activities of several cellular antioxidant enzymes, including superoxide dismutase, catalase and glutathione peroxidase. Treatment with B. platyphylla var. japonica extract induced cytotoxicity and apoptosis in HL-60 cells, as shown by nucleosomal DNA fragmentation, increases in the subdiploid cell population, and fluorescence microscopy. B. platyphylla var. japonica extract gradually increased the expression of pro-apoptotic Bax and led to the activation of caspase-3 and cleavage of PARP. These findings suggest that B. platyphylla var. japonica exhibits potential antioxidant and anticancer properties.

[PC1-42] [2003-10-10 09:00 - 13:00 / Grand Ballroom Pre-function]

Conformational Study of Pseudo-Proline Dipeptide in the Gas Phase and Solutions Park Hae Sook^o, Kang Young Kee^o

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We report here the results on N-acetyl-N"-methylamide of oxazolidine (Ac-Oxa-NHMe) calculated using the ab initio molecular orbital method with the self-consistent reaction field (SCRF) theory at the HF level of theory with the 6-31+G(d) basis set. The displacement of the γ -CH₂ group in proline ring by oxygen atom has affected the structure of proline, cis-trans equilibrium, and rotational barrier. The up-puckered structure is found to be prevalent for the trans conformers of the Oxa amide. The higher cis populations of the Oxa amide can be interpreted due to the longer distance between the acetyl methyl group and the 5-methylene group of the ring for the trans conformer of the Oxa amide than that of the Pro amide. The changes in charge of the prolyl nitrogen and the decrease in electron overlap of the C-N bond for TS structures seem to play a role in lowering rotational barriers of the Oxa amide compared to that of the Pro amide. The calculated preferences for cis conformers in the order of Oxa > Pro amides and for trans-to-cis rotational barriers in the order of Pro > Oxa amide in water are consistent with experimental results on Oxa-containing peptides. The pertinent distance between the prolyl nitrogen and the N-H amide group to form a hydrogen bond might indicate that this intramolecular hydrogen bond could contribute in stabilizing the TS structures of Oxa and Pro amides and play a role in prolyl isomerization

[PC1-43] [2003-10-10 09:00 - 13:00 / Grand Ballroom Pre-function]

Involvement of Akt in mitochondria-dependent apoptosis induced by a naphthoquinone analog

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Vitamin K-related analogs induce growth inhibition in various cancer cell lines. We report that 2,3-dichloro-5,8-dihydroxy-1,4-naphthoquinone (DDN), a naphthoquinone analog, induces mitochondria-dependent apoptosis in human promyeloid leukemic HL-60 cells. DDN induced cytochrome c release, cleavage of Bid, and activation of caspases -8, -9 and -3. Cleavage of Bid, the caspase-8 substrate, was inhibited by the broad caspase inhibitor zVAD-fmk, whereas cytochrome c release was not affected by zVAD-fmk. These results indicate that DDN induces activation of caspas-8 and subsequent processing of Bid downstream of cytochrome c release. DDN inhibited the activation of Akt detected by decreasing levels of phosphorylation. Overexpression of constitutively active Akt protected cells from DDN-induced apoptosis. Furthermore, Akt prevented release of cytochrome c in DDN-treated HL-60 cells. In conclusion, DDN-induced apoptosis in HL-60 cells is associated with mitochondrial signaling which involves cytochrome c release through the inhibition of Akt pathway.