Lee Sang-Soo, Hong Seon-Pyo, Yoon Hye-Ran
College of Pharmacy, Kyung Hee University, Department of Oriental Pharmaceutical Science, Kyung Hee University, Seoul Medical Science Institute

In galactose metabolic pathway: there are three inborn metabolic disorders galactokinase deficiency (galactosemia type II), galactose-1-phosphate uridyl transferase (GALT) deficiency (galactosemia type I), uridine diphosphate galactose-4-epimerase deficiency (galactosemia type III). Among these disorders GALT deficiency is the most severe and common. Infants with GALT deficiency fail to metabolize galactose-1-phosphate. As a consequence, galactose-1-phosphate and galactose are accumulated in blood in which GALT enzyme plays the role of a pathognomonic marker. In the previous paper, we reported a reversed-phase HPLC method using 8-Amino-2-naphthalenesulfonic acid as derivatization reagent for the determination of galactosemia. But, this method has the defects such as a relatively longer pretreatment, the reduction of sensitivity. We developed an advanced diagnostic method for galactosemia by shortening pretreatment and increasing the sensitivity.

[PD4-8] [2003-10-10 14:00 - 17:30 / Grand Ballroom Pre-function]

Indirect chiral separation of α-arylmethylpropionic acids by liquid chromatography
Min Chung Sik, Jang Seung Jae, Choi Bo Kyung, Kim Young Lim, Jung Hae Yun, Bae Kyung Min, Lee Kyung Hee, Jo Keang In, Gu You Ni
KFDA

A various α-arylmethylpropionic acids(profen) have been widely used as non-steroidal anti-inflammatory drugs for the relief of acute and chronic rheumatoid arthritis and osteoarthritis, as well as for other connective tissue disorders and pains. Example is fenoprofen, ibuprofen, ketoprofen, and naproxen. All are chiral and, except for naproxen and ibuprofen, are marketed in racemic form. Enantioseparations of profens have been of considerable interest because their anti-inflammatory and analgesic effects have been attributed almost exclusively to their (S)-enantiomer. A simple method for determination of optical purity of (+) and (-)-α-arylmethylpropionic acids has been developed. By means of EEDQ, α-arylmethylpropionic acids were coupled to (S)-naphthylethylamide. The diastereoisomeric derivatives was then separated by normal-phase liquid chromatography. And separation process of diastereomeric isomer was interpreted by molecular mechanics and quantum mechanics calculation of diasteromeric conformation.

[PD4-9] [2003-10-10 14:00 - 17:30 / Grand Ballroom Pre-function]

Studies on the tyrosinase inhibitory comound of Potentilla bifurca L. var. glabrata Lehm
Piao Xiang Lan, Lim Geun Sil, Lee Youn Ju, Kim Bak Kwang, Park Man Ki, Park Jeong Hill
College of Pharmacy, Seol National University, Seoul 151-742, Korea

Tyrosinase is an important enzyme involved in the transition steps from tyrosine to melanin. Inhibition of the tyrosinase activity could block melanin formation from tyrosine and thus prevent melanin pigmentation on skin. This may contribute to the development of new whitening agent that would be useful in the prevention of pigmentation. In this study, we isolated tyrosinase inhibitory compound from BuOH fraction of Potentilla bifurca L. var. glabrata Lehm by activity guided fractionation method. Based on spectroscopic data, the active compound was identified as a quercetin 4'-O-glucopyranoside.

[PD4-10] [2003-10-10 14:00 - 17:30 / Grand Ballroom Pre-function]

Physical properties and determination of eupatilin, a new antigastritic agent, by high performance liquid chromatography
Jang Ji Myun, Shim Hyun Joo, Ahn Byung Ok, Kim Soon Hoe, Kim Won Bae
Dong-A Pharmaceutical Co. LTD., 47-5, Sanggal-ri, Kiheung-up, Yongin-si, Kyunggi-do, 449-905, Korea
Eupatilin is a major active component of Stillen?(Artemisia Herba Extract) having a potent antiagastriotic effect. We investigated the physical properties of eupatilin using high performance liquid chromatography. Solubility, stability & partition coefficient of eupatilin were investigated. pH-stability of eupatilin was examined over the broad range through pH1-9 at 37°C & it has good stability above the broad range pH. The solubility of eupatilin was extremely low but the value of logP was more than 2. Also, a high performance liquid chromatographic method was developed for the determination of eupatilin in rat plasma. The method involved deproteinization of biological sample with the same volume of acetonitrile, 0.2M zinc sulphate, and 0.15M barium hydroxide. The mobile phase employed was ammonium acetate buffer(1% ammonium acetate and 0.5% acetic acid) – acetonitrile (58:42,v/v) and the flow rate was 1.0 ml/min. The quantitation limit of eupatilin in rat plasma was 10 ng/ml. No interferences from endogenous substances were found.

[Cycloxygenase Inhibitory Activity of Ginsenosides from Panax ginseng]
Yoo Hye Hyun, Kang Ki Sung, Lee Yang Beom, Kim Bak Kwang, Park Man Ki, Park Jeong Hill
College of Pharmacy, Seoul National University

P. ginseng C.A. Meyer is one of the most widely used herbal medicine in Asia. It has been used for the treatment of many disorders. Its major constituent is known to be ginsenosides, and there are many documents about bioactivities of ginsenosides such as anti-oxidant, anti-tumorogenic, anti-fatigue, and anti-inflammatory activities. Some of these activities are supposed to have some correlation with inhibitory action of cyclooxygenase (COX). Ginsenosides from P. ginseng and sapogenins were evaluated for their inhibitory effects against both cyclooxygenase-1 and -2 (COX-1 and -2). Inhibitory activity was evaluated by measuring prostaglandin E2 (PGE2) production from arachidonic acid with an ELISA reader. As a results, Rg1(S), and Rg3 and Rk1 showed COX-2 inhibitory activity in a selective manner (COX-1: IC50 = >100, 77.01 µg/mL, COX-2: IC50 = 35.47, 18.6 µg/mL). Protopanaxatriol (PPT) showed moderate activity on COX-1 and -2 (COX-1: IC50 = 39.16, COX-2: IC50 = 35.56 µg/mL), while Re, Rg3 (R), and protopanaxadiol (PPD) showed little activity.

[HPLC Analysis of Phytosphingosine and Its Metabolites in Mammalian Cells with TCPO- H2O2 Chemiluminescence Reaction]
JIN YouXun, Choi Chang-Hwan, Yoo Hwan-Soo, Lee Yong-Moon
College of Pharmacy. Chungbuk National University

Sphingolipids has been known to induce apoptosis, cell proliferation, differentiation and migration in a variety of cell types. Recently, its phosphate form was suggested that they may act both as an agonist ligand to S1PRs and a second messenger in intracellular action. Phytosphingosine(PHS) is not easily detected due to trace component of cellular lipids in mammalian and human tissues while this is a major sphingolipid in yeast and plants. We therefore developed highly sensitive and reproducible analytical method for PHS and its phosphate by oxalic acid bis(2,4,6-tri-chlorophenyl) ester(TCPO)-hydrogen peroxide(H2O2) chemiluminescence. The NDA derivatives of PHS exhibited stable fluorencesces & was enhanced their detectability at low concentrations by post-column chemiluminescence detection with TCPO- H2O2. The dried lipid extracts or sphingoid base standards for the calibration curve were dissolved in 40 ul of ethanol. NDA derivatization was accomplished by adding the following stock solutions: 40 ul 0.05M NaHCO3 / 0.1M NaOH buffer(pH 10.5), 20 ul 13% (w/v) NaCN, and 20 ul 0.5%(w/v) NDA. The tube was tightly sealed with PTFE film and heated at 67°C in a water bath for 90 min, glycine was added to stop the derivatization reaction. We successfully measured the amount of PHS and PHS-1-P in LLC-PK1 cells. Collectively, this method can be thus used to detect and distinguish PHS and PHS-1-P with high sensitivity from other sphingolipids in mammalian cells.