and decreased the plasma glucose levels and FFA in HF diet-induced obesity of C57BL/6 mice.

[PA1-10] [04/17/2003 (Thr) 14:00 - 17:00 / Hall P]

Metabolism of YH3945, a novel anticancer drug, in rats using 14C-labeled compound

<u>Lee Jaeick</u>^o, Son Junghyun, Ahn Byung-Nak, Lee Bongyong, Kim Dong-Hyun Kim

Korea Institute of Science and Technology, Yuhan Research Institute

The metabolism of a novel anticancer agent 1–{3-[3-(4-Cyano -benzyl)-3H-imidazol-4-yl] - propyl}-3-(6-methoxy-pyridin-3-yl)-1-(2-trifluoromethyl-benzyl)-thiourea (YH3945) were investigated in the Sprague-Dawley rat after single oral and i.v. administration of [14C]-YH3945. Bile, feces, urine and plasma were collected and analyzed by an HPLC system equipped with multiple detectors. The present analysis system includes the simultaneous detection technique of three different detectors (diod array detector-radioactivity flow detector-tandem mass spectrometry) in single run. The structures of each metabolite were characterized based on UV, tandem mass (MS2 and MS3) and NMR (1H and TOCSY) spectral analyses. YH3945 was metabolized to seventeen different metabolites including glucuronide conjugate. The four major metabolic pathways of YH3945 in rat were identified as O-demethylation of pyridine moiety, N-debenzylation of imidazol moiety, hydroxylation of aromatic ring and recyclization between pyridine and benzylic carbon. Especially, nonenzymatic reaction mechanism of metabolite generated by the recyclization was theoretically postulated.

[PA1-11] [04/17/2003 (Thr) 14:00 - 17:00 / Hall P]

Attenuation of nicotine-induced locomotor sensitization in μ -opioid receptor knockout mice

Yoo JiHoon^o, Yang EunMi, Kim KyungIn, Lee SeokYong, Jang ChoonGon

Department of Pharmacology, College of Pharmacy, Sungkyunkwan University, Suwon. Korea

The present study was undertaken to examine the hypothesis that μ -opioid receptors play a crucial role in behavioral sensitization to nicotine using μ -opioid receptor knockout mice. All mice were treated acutely or repeatedly with nicotine 0.05 mg/kg twice daily for 7 consecutive days. The mice were challenged with nicotine on day 11. And locomotor activity was measured for 30min. Locomotor activity challenged by acute nicotine was no difference between μ -opioid receptor knockout and wild-type mice. Repeated treatment with nicotine induced behavioral sensitization in wild-type mice on days 7 and 11. In contrast, nicotine exposure failed to develop behavioral sensitization in μ -opioid knockout mice. This behavioral sensitization was accompanied by an incerese in D(2) receptor binding in striatum of the μ -opioid knockout mice compared with the wild-type mice on day 7. However, D(1) receptor expression was not changed in the striatum and nucleus accumbens.

These results suggest that abolishment of nicotine-induced behavioral sensitization in the μ -opioid receptor knockout mice may be related to the increase of D(2) receptor binding in the striatum.

[PA1-12] [04/17/2003 (Thr) 14:00 - 17:00 / Hall P]

The effects of the novel IDPc inhibitor, DA-11004, on NADPH generation, insulin secretion, and glucose level in zucker rats

Shin ChangYell^o, Jeong MiYoung, Sohn JinBup, Lee InKi, Son Miwon, Bae CheolJun, Byun JongSoo, Kim DongSung, Kim SoonHae, Yoo Moohi, Huh TaeLin∗, , Kim WonBae

Research Laboratories, Dong-A Pharm. Co. Ltd., 47-5, Sanggal, Kiheung, Yongin, Kyunggi 449-900, Korea. *Department of Genetic Engineering, College of Natural Sciences, Kyung Pook National University

The biological effects of NADPH-dependent isocitrate dehydrogenase (IDPc) inhibitor, DA-11004, was examined in obese zucker rats or streptozotocin-induced diabetic SD rats. Diabetes was induced by injection of streptozotocin (50mg/kg) dissolved in citrate buffer (pH 4.8) into the tail vein and induction of diabetes was confirmed by the measurement of the tail blood glucose level at 48h. DA-11004 (30mg/kg, po) was injected for successive 7days and significantly reduced the plasma glucose in streptozotocin-induced diabetic rats (P<0.05). In obese zucker rats, DA-11004 (10mg/kg, ip) inhibited the increases of body weight and diet consumption when compared to control, but oxalomalate (30mg/kg, ip) had no effects. DA-11004 significantly reduced the weight of epididymal and retroperitoneal fats and NADPH generation in plasma (P<0.01). We measured the levels of free fatty acids (FFA) and insulin in plasma of zucker rats. DA-11004 reduced the level of FFA from 821.4 \pm 229.6 to 587.2 \pm 47.1 (µEq/L) and significantly decreased the plasma insulin responses when compared to control (P<0.05). Administration of DA-11004 into zucker rats significantly decreased the levels of glucose in liver and the levels of triglycerides, GOT, GPT, and LDH in plasma, but oxalomalate had no effects (P<0.05).

In summary, DA-11004 reduced the fat synthesis via inhibition of NADPH generation and inhibited the insulin secretion, FFA generation and declined the glucose and triglycerides levels in plasma and liver of zucker rats.

[PA1-13] [04/17/2003 (Thr) 14:00 - 17:00 / Hall P]

KR-32158 protects heart-derived H9c2 cells from oxidative stress-induced cell death

Kim Mi Jeong^o, Jung Yi-Sook, Kim Sun-Ok, Lee Dong Ha, Lim Hong, Yi Kyu Yang, Yoo Sung Eun, Lee Soo Hwan, Baik Eun Joo, Moon Chang-Hyun

1 Department of Physiology, Ajou university School of Medicine, Suwon, South Korea 2Bioorganic Division, Korea Research of Chemical Technology, Daejon, South Korea

A benzopyranyl derivative, KR32158, synthesized as a plausible KATP opener, has been shown to exert cardioprotective effect in vivo myocardial infarction model. Myocardial ischemia, induced by oxidative stress, mental stress and fever, result in artheroscleosis, myocardial infarction and hypertrophy. In this study, we investigated in vitro effect of KR32158 by determining whether KR32158 produce cardioprotective effect against oxidative stress—induced death in heart—derived H9c2 cells. Oxidative stress was induced by buthionine sulfoximine (BSO) which inhibit GSH level and subsequently increase ROS level in H9c2. Cell death was determined by measuring lactate dehydrogenase(LDH) release. KR32158 significantly decreased LDH release from H9c2 induced by oxidative stress, in dose—dependent manner. And also, we measured BSO—induced ROS generation to confirm whether KR32158 had protective effect from ROS—induced cardiac injury. As the result, KR32158 significantly decreased BSO—induced ROS generation. We also observed cardioprotevtive effect of KR32158 morphologycally by using microscope. These results suggest that KR32158 has protective effects through inhibiting the production of ROS.