

in JNK1(-)-transfected cells. Conclusions: DA-125 potently inhibited topoisomerase II activity and induced apoptosis with the high rate of prooxidant production. DA-125 exhibited high-affinity DNA binding with improved cellular drug accumulation. Apoptosis induced by DA-125 involved the pathway of JNK1, but not ERK1/2 or p38 kinase

[OA-6] [04/20/2001 (Fri) 14:45 - 15:00 / Room 1]

Altered Expression of Ferritin Light Chain (FLC) by Sulfur Amino Acid Deprivation in Hepa1c1c7 and Raw264.7 Cells: The Role of Cellular Ca²⁺ and Free Iron for Prooxidant Production

Kim HJ^o, Kim SG

College of Pharmacy and Research Institute of Pharmaceutical Sciences, Seoul National University (*
A doctoral candidate from Duksung Women's University)

Ferritin expression is induced by oxidative stress, which confers resistance to oxidative insults. Sulfur amino acid deprivation (SAAD) induces oxidative stress through a decrease in a GSH content. The molecular mechanisms of cell-type specific ferritin light chain (FLC) expression in association with increases in intracellular Ca²⁺ and free iron pools were investigated in Hepa1c1c7 and Raw264.7 cells exposed to SAAD. Intracellular Ca²⁺ level was rapidly increased by SAAD, followed by returning to control at later times. Sulfhydryl-containing compounds prevented the increase in intracellular Ca²⁺ by SAAD, supporting the role of redox-state in the regulation of Ca²⁺. Thapsigargin or Ca²⁺-free medium inhibited the increase in intracellular Ca²⁺, showing that Ca²⁺ originated from endoplasmic reticulum as well as from extracellular source. Inhibition of Ca²⁺ mobilization decreased fluorescence of free iron pool inside cells and inhibited dichlorofluorescein oxidation. Deferoxamine also inhibited dichlorofluorescein oxidation. Hence, the increase in cellular Ca²⁺ content coupled with elevation in intracellular free iron pool and subsequent prooxidant production. FLC protein level was detected by Western blotting in Raw264.7 cells, but not in Hepa1c1c7 cells. SAAD alone or in combination with FeSO₄, however, down-regulated FLC expression. On the contrary, the FLC mRNA level was increased by SAAD in both Hepa1c1c7 and Raw264.7 cells. Calcium or iron chelators prevented increases in the FLC mRNA. These results provided evidence that oxidative stress by SAAD inhibited FLC protein expression but increased the mRNA level through intracellular Ca²⁺ and subsequent release of iron.

[OA-7] [04/20/2001 (Fri) 15:00 - 15:15 / Room 1]

DNA Adduct Formation, Induction of Apoptosis and Cell Cycle Arrest by N-Nitroso Metabolite of Carbofuran

Yoon JY¹, Oh SH¹, You SM¹, Lee SJ¹, Ko WG¹, Park MR¹, Lee HS¹, Choi SJ¹, Moon CK², Lee BH¹

¹ College of Pharmacy and Medicinal Resources Research Center, Wonkwang University, Iksan, Jeonbuk, Korea ²College of Pharmacy, Seoul National University, Seoul, Korea

Carbofuran (CF) is one of the most widely used carbamate pesticide in the world applied to insect and nematode control. Due to its widespread use in agriculture and households, contamination of food, water, and air has become imminent, and consequently adverse health effects are inevitable in humans, animals, wildlife and fish. It has been reported that CF alone or in combination with other carbamate insecticides influences the level of reproductive and metabolic hormones such as thyroxine and corticosterone, and results in impairment of endocrine, immune and behavioral functions. In this study, we evaluated the effects of CF and its N-nitroso derivative N-nitrosocarbofuran (NOCF) on DNA adduct formation, genotoxicity, cell growth and apoptosis of CHL cells. NOCF, but not CF, induced the formation of O⁶- and N⁷-methylguanine DNA adducts in calf thymus DNA and induced genotoxicity determined by Ames test. NOCF inhibited the growth of Chinese hamster lung fibroblast