

THF, and $N^{5,10}\text{-CH}_2$ THF. Folic acid did not detected by both determinations. Two peaks which were detected with radiometry were likely to the degradation products of $^3\text{H-FA}$.

When rats were treated with 10 mg of amethopterin, incorporation of radioactivity into these derivatives were extremely small and peaks due to degradation were not detected. But it was observed that small amount of folic acid was remained unchanged in the liver.

An aliquot of human urine collected in a brown bottle containing potassium ascorbate

for 3 hours following an intravenous injection of $20\ \mu\text{g}$ of $^3\text{H-FA}$ was applied on a column. Most of folic acid derivatives assessed by microbiological assay was folic acid and relatively large amount $N^5\text{-CH}_3$ THF, and small amount $N^{10}\text{-CHO}$ THF, whereas the radioactivity was mostly in folic acid and relatively small in $N^{10}\text{-CHO}$ THF and extremely small in $N^5\text{-CH}_3$ THF. Peaks of radioactivity due to, probably, degradation products were also observed. Thus, $N^5\text{-CH}_3$ THF appearing in the urine following intravenously injected folic acid was mainly not derived from the injected folic acid.

Intravenous Pentose Solution Injection in the Surgical Patients

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In the 4th and 6th Annual Meeting of the Japanese Society of Nuclear Medicine, we had reported that, both P-incorporation into the hepatic RNA and S-incorporation into the hepatic RNA and C-incorporation into the hepatic ribosomal protein and serum albumin of mice were markedly increased in response to surgical stress. To evaluate the effects of xylitol injection in the surgical patients, we have studied the changes of these hepatic RNA metabolism following intraperitoneal injection of xylitol, both in the laparotomized mice and 70% hepatectomized rats.

H-incorporation in vivo of hepatic RNA during the periods from 3 to 12 hours after the 31 ml 10% xylitol and $10\ \mu\text{Ci}$ H-uracil administration per 20 g body weight, was as similar as the 10% glucose or saline solution injected control mice. However, in the laparotomized mice, the H-specific activity of hepa-

tic RNA after xylitol injection was somewhat more remarkable than the glucose or saline solution injected groups. In the same time, TPN and DPN activity in the xylitol administrated mice liver was more rapidly decreased following surgical stress. These findings about RNA and pyridine nucleotides metabolism suggest that the metabolic flow via uronic acid pathway is stimulated first of all in order to metabolize xylitol, especially under the surgical stress.

In the 70% hepatectomized rats, H-incorporation of DNA in the regenerating liver following $30\ \mu\text{Ci}$ H-thymidine injection was also stimulated by xylitol administration in the early peak of cell division. However, it should be solved by the further investigation why fatty acid deposition early after hepatectomy was most prominent in the xylitol injected rats.