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FATTY ACID COMPOSITION OF PHOSPHOINOSITIDES IN RAT LIVER NODULES

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Rat liver nodules produced by treatment with carcinogens exhibit elevated proliferation rate and differs from normal liver in several biochemical properties.

The composition of individual phospholipids in nodular tissue is changed with a two-fold increase in phosphatidylinositol (PI) compared to normal liver. Since PIs play a critical role in cell regulatory mechanisms, it is of great importance to understand the action of PIs in nodules. Earlier studies show a connection between cell proliferation, PI-metabolism and arachidonic acid release.

The fatty acid composition of the phosphoinositides in nodules was studied, with special interest focused on arachidonic acid (20:4) and its precursor linoleic acid (18:2). Preliminary results indicate no difference in fatty acid composition in phosphoinositides between nodular and normal liver.

INHIBITION OF COLONIC NEOPLASIA AND CRYPT CELL PRODUCTION RATES BY INTRALUMINAL CALCIUM

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Small bowel resection and intrarectal administration of sodium deoxycholate each stimulate cell proliferation and promote carcinogenesis in the large intestine; oral supplements of calcium reduce the mitogenic effect of bile acids on colorectal mucosa. Potential suppression of intestinal and carcinogenesis by adaptation intraluminal calcium was tested in 120 male Sprague-Dawley rats weighing 186±9 g. Rats were randomised to receive azoxymethane s/c 15 mg/kg/week for 6 weeks or vehicle, followed by 80% mid small bowel resection or transection with reanastomosis. Half the animals in each group received supplemental calcium in the drinking water (calcium lactate 24g/1). Crypt cell production rate (CCPR) in descending colon was determined 7 weeks postoperatively in vehicle-treated rats; in the remainder colonic tumour yield was assessed at 26 weeks. Among rats with transection calcium supplements reduced colonic CCPR by 26% from 4.49±0.33 to 3.32+0.40 cells/crypt/hr (p<0.05) and more than halved tumour yield from 4.3 to 1.8 tumours/rat (p=0.0007). Jejunoileal resection increased both CCPR (by 51 to 61%; p<0.001) and tumour yield (by 65 to 105%; p<0.005), but again calcium lowered CCPR by 31% (7.23+0.44 vs 4.98±0.70; p<0.02) and tumour yield by 46% (6.9 vs 3.7; p=0.0006). Increased dietary levels of calcium diminish both adaptive and neoplastic growth in the colon, and calcium also blunts the co-carcinogenic stimulus of massive enterectomy.

ASSESSMENT OF RAT COLONIC TUMOURS BY FLOW CYTOMETRY: METASTASES ARE COMMONLY DNA ANEUPLOID

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The study of azoxymethane-induced colonic carcinogenesis in rats helps in understanding human colorectal cancer. We have used flow cytometry to investigate the presence of DNA aneuploidy in rat intestinal tumours, and to evaluate the rat model as an experimental system. Fifty male Sprague-Dawley rats weighing 185.6+9.2 g were given azoxymethane 15 mg/kg/week s/c for 6 weeks and then underwent 80% small bowel resection (n=25) or jejunal transection (n=25). Half the animals in each group had calcium lactate 24g/1 added to the drinking water. Ten further non-operated rats (NOP) received azoxymethane 10 days later than the others. Forty-three rats survived 26 weeks and yielded 149 colonic and duodenal tumours of which 140 were measurable by flow cytometry. The incidence of DNA aneuploidy was 43% in NOP which was higher than in rats with resection (9%; p<0.0005) or transection (24%; p<0.0005). There was no significant difference in the prevalence of DNA aneuploidy between adenomas (32%) and carcinomas (17%) or between calcium treated (11%) and non-calcium groups (12%). However metastases were more commonly DNA aneuploid than the primary tumours (62% vs 20%; p<0.005). DNA aneuploidy is present in rat intestinal tumours and levels can vary widely with manipulation of the model. Metastases are associated with a high incidence of DNA aneuploidy.

ASSESSMENT OF COLONIC ADAPTATION BY CRYPT CELL PRODUCTION RATES IN ORGAN CULTURE: AN