Short Communication

REJECTION OF MURINE MAMMARY TUMOURS IN BALB/c MICE

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The presence of circulating antibodies to murine mammary tumour virus (MuMTV) related antigens in mice as well as in humans has been reported by a number of investigators (Muller et al., 1976; Priori et al., 1972; Hoshino & Dnochowski 1973). Tumour growth, according to a hypotheses of Nordquist et al. (1977) is facilitated by antibody-induced redistribution and shedding of mammary-tumour antigens into the circulatory system. The antigen-denuded cells are not then recognized by effector agents. Efforts to inhibit mammary-tumour growth and reduce its incidence in susceptible strains of mice by immunization with MuMTV were not successful (Charney et al., 1976). Sarkar & Moore (1978), however, have recently shown that C57BL mice that were immunized with purified formalin-inactivated MuMTV showed increased resistance to exogenous MuMTV infection, and to subsequent development of mammary tumours. They concluded that successful immunization required the use of relatively high doses of killed virus.

The recorded incidence of spontaneous-tumour rejection by inbred mice is very rare. Hewitt et al. (1976) found that, in ~20,000 transplantsations of different tumours in WHT/Ht and CBA/Ht lines of mice, only 2 cases of tumour rejection occurred.

The purpose of this study was to explain the repeated rejection of mouse mammary-tumour implants by 2 mice in our laboratory, and to determine whether these mice produced antibody reactive against human breast-tumour tissue.

In the course of 2 years, murine mammary tumours derived from BALB/cf C3H mice were implanted into 290 mice. Tumours were allowed to grow to a size of about 1 cm³ before s.c. transplantation of 2 pieces of tumour tissue (1 mm³) in the lateral abdominal area of recipient mice. The tumours were propagated in the animals for 10 generations. Only 2 mice, one in the 4th and the other in the 7th generation, rejected the tumours. Three separate attempts to implant tumours were made in each mouse, without success. At each attempt the same tumour was successfully implanted into several other mice.

Sera obtained by retro-orbital sinus bleeding from the mice that rejected the tumour, 8 healthy mice, and 3 tumour-bearing mice from the same colony, were absorbed with lyophilized human benign breast-tumour tissue and examined for the presence of antibodies to antigens in human breast carcinomas. Sera from the 2 mice that rejected the tumors reacted positively when tested by indirect immunofluorescence for reactivity against frozen sections of human breast adenocarcinomas. FITC-labelled anti-mouse IgG serum from Meloy Laboratories was used as the secondary antibody in the indirect immunofluorescence assay. Sera from all other mice were negative. All sera were

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negative when tested against sections of human benign breast tumours. Positive immunofluorescence was also seen when sections of human breast adenocarcinomas were examined for reactivity with rabbit anti-MuMTV serum. The results are shown in the Table.

| Origin of serum | No of animals | Benign | Carcinoma |
|-----------------|--------------|--------|-----------|
| Mice rejecting  | Cancer       |        |           |
| tumour          | Healthy      | 2      | +         |
| Healthy mice    | Tumour-bearing mice | 8 | - | - |
| Rabbit anti-    | Rabbit       | 3      | -         |
| MuMTV           |              |        | +         |

The results of this investigation indicate that sera from the mice that rejected the mammary tumours contained antibodies directed against antigens in human breast-carcinoma tissue. The immunological reactivity of sera from these mice was similar to the reactivity of rabbit anti-MuMTV sera when tested against sections of human breast tumour. The sera of control and tumour-bearing mice did not possess this reactivity.

It has been previously shown (Muller et al., 1976; Priori et al., 1972; Hoshino & Dmochowski, 1973; Tomana et al., 1979) that an immunological relationship exists between mouse and human breast cancer. Although the existence of a human breast-tumour virus is still questionable, components were detected in human breast-carcinoma patients that are antigenically related to a component of MuMTV. Although our results suggest that the mice rejecting the tumour had natural or spontaneously acquired antibodies to MuMTV, it is possible that the piece of tumour used for the first implantation consisted of non-viable tissue which resulted in immunization of the mouse. Rejection of the 2nd and 3rd implants that were done at about one-month intervals could thus be explained. This explanation is consistent with previous findings of Sarkar & Moore (1978) who demonstrated successful immunization of mice with formalin-treated MuMTV. The results of this investigation support the hypothesis that there is an aetiologic relationship between mouse and human breast carcinomas.

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