Short Communication

Pre-existing anti-mouse immunoglobulin in a patient receiving $^{131}$I-murine monoclonal antibody for radioimmunolocalisation

A.G. Davies¹, S.P. Bourne¹, R.B. Richardson², R. Czudek⁴, T.B. Wallington⁴, J.T. Kemshead⁵ & H.B. Coakham¹,³

¹Brain Tumour Research Laboratory; Departments of ²Medical Physics, and ³Neurosurgery, Frenchay Hospital, Bristol, BS16 1LE; ⁴South Western Regional Transfusion Centre, Southmead Road, Bristol, BS10 5ND; ⁵Imperial Cancer Research Fund Oncology Laboratory, Institute of Child Health, London, UK.

Radiolabelled monoclonal antibodies (MCAs) which recognise tumour-associated antigens are being used increasingly for radioimmunolocalisation (RIL) and trials of treatment for various malignancies, (Miller et al., 1982, 1983; Foon et al., 1984; Dillman & Royston, 1984). The majority of these MCAs are murine.

We have previously reported on the use of the mouse monoclonal antibody UJ13A for in vivo radioimmunolocalisation of gliomas in humans, (Richardson et al., 1985), and there has been a recent report of the treatment of a glioma using MCA-linked $^{131}$I -irradiation (Epenetos et al., 1985).

We report here pre-existing anti-mouse immunoglobulin activity in the serum of a patient undergoing RIL. A 38-year old man with a parietal gliosarcoma and no significant exposure to mice was given an i.v. injection of radiolabelled control and specific MCAs: 300 µg of mouse MCA HMFG2 (Taylor-Papadimitriou et al., 1981) labelled with 2.06 mCi $^{131}$I and 60 µg of UJ13A (Allan et al., 1983) labelled with 0.41 mCi $^{125}$I.

Throughout the study the patient suffered no ill effects and urinary and haematological investigations remained normal. Base line investigations were normal. These included serum IgM and IgG levels, tests for rheumatoid factor, an auto-immune profile, and liver function tests. The patient had no cutaneous reaction following an i.d. injection of 10 µg (100 µl) of UJ13A prior to RIL.

A plasma sample of the two radiolabelled antibodies taken 10 minutes post-injection was passed through a Sephacryl S300 superfine gel (Pharmacia). Both the $^{131}$I and $^{125}$I conjugates eluted significantly earlier than the IgG peak, (Figure 1), indicating the label to be in a higher molecular weight form than the monomeric radiolabelled IgG antibody. Figure 2 shows the elution of radiolabelled MCA in a different patient who had no anti-mouse response.

Incubation in vitro of the specific $^{131}$I-labelled MCA with a pre-injection plasma sample and subsequent passage through a Sephacryl S300 gel column showed the majority of the radioactivity to elute significantly in advance of the IgG peak, thus paralleling the in vivo behaviour of the injected antibodies (Figure 3).

Plasma taken before injection of the antibodies was pre-cleared in vitro by incubation with mouse IgG (Sigma) prior to adding $^{131}$I-specific MCA, the elution of the mouse radiolabelled MCA then coincided with the IgG peak, indicating that the murine MCA now remained as the monomer (Figure 4).

Haemagglutination tests on the pre-injection serum using appropriately sensitised red cells confirmed the presence of a pre-existing anti-mouse

Correspondence: H.B. Coakham.

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immunoglobulin. A six-times washed Group O, RhD, WrA red cell suspension was sensitised with a predetermined sub-agglutinating dose of one of the following antibody preparations: (i) Human IgG: (incomplete anti-D); (ii) Bric 14.2/1AB: a mouse monoclonal anti-WrA, IgG2a subclass, (Ridgwell et al., 1983); (iii) Bric 20.3: a mouse monoclonal anti-WrA, IgM subclass, (D. Anstee, personal communication).

A sensitised red blood cell suspension was diluted to 5% in PBS. One drop was added to 2 drops of the patient’s serum, centrifuged for 10 sec, and agglutination observed macroscopically.

The patient’s serum reacted very strongly with Bric 14 sensitised cells (mouse IgG), unconvincingly with Bric 20 (mouse IgM) and not at all with incomplete anti-D. All appropriate controls behaved as anticipated. Thirteen other patients with gliomas were studied and two strongly positive reactions were seen. No such reaction was observed in seventeen controls. Furthermore, a positive response did not correlate with blood group.

The pharmacokinetics of the injected MCAs in this patient (RM), did not differ significantly from 10 other patients studied. The clearance from the blood was biphasic. The fast component of the half-life in serum was 3.6 h for both antibodies and the slow component was 58 and 63 h for the specific and non-specific antibodies respectively. This is not significantly different from a mean of 4.5 ± 4.9 and 62 ± 15 h for the fast and slow half-life components of the specific MCA in 7 other patients studied.

Comments It has been known for some years that individuals may possess antibodies to various animal proteins (Rotheberg & Farr, 1965), but the introduction of clinical studies with MCAs now means that specific anti-mouse reactivity becomes relevant (Hunter et al., 1984).

Other workers have described human anti-mouse serological responses in patients following injection of murine MCAs (Miller et al., 1982). These immune responses have been reported in 50% of patients with melanoma that were given repeatedly large doses of a MCA. These responses consisted of both IgG and IgM (Oldham et al., 1984).

There are few reports in the literature concerning the presence of pre-existing anti-mouse activity in
the serum of patients undergoing RIL, and this phenomenon has not been related to skin test positivity (Schroff et al., 1985). Primus et al. (1980), using goat heteroantisera to carcinoembryonic antigen, have demonstrated pre-existing anti-goat IgG in several patients.

The high molecular weight eluates seen in the serum of patient RM suggest the formation of immune complexes. Similar elution profiles were seen by James et al. (1971) in renal transplant recipients given equine anti-lymphocyte globulin. It has previously been observed in studies of RIL that immune complexes (ICs) may form when there is circulating free antigen (Goldenberg et al., 1978). These ICs did not seem to interfere with the success of the RIL in the circumstances studied (Primus & Goldenberg, 1980). Of more importance, however, would be the consequence of administering a large quantity of animal antibody to an individual possessing pre-existing anti-species immunoglobulin, as might occur in attempted therapy. This might result in serum sickness in such a susceptible patient (Oldham et al., 1984), together with altered pharmacokinetics of potentially therapeutic conjugates.

The reported effects of anti-mouse activity on MCA pharmacokinetics are variable. In the treatment of certain leukaemias and lymphomas, the production of anti-mouse antibodies is said to effectively neutralise any clinical effects of the therapy (Dillman & Royston, 1984). The in vivo handling of radiolabelled MCAs in patients with melanoma who developed an anti-mouse response following repeated injections is not so clear. Oldham et al. (1984), reported unaltered pharmacokinetics but Larson et al. (1983), using a different anti-

melanoma MCA, observed an increase in the rate of clearance from the circulation with an increase in liver uptake and a decrease in tumour uptake. An immune response in a sensitised individual to a radiolabelled MCA may result in its accelerated catabolism and dehalogenation, with rapid excretion of most of the unbound radionucleide within 24 h (J. Kemshead unpublished observation).

Patients with various malignancies, including gliomas, are known to produce a range of heterophile antibodies, (Pfreundschuh et al. 1978), which may explain the present observations. The incidence of naturally occurring anti-mouse activity in the serum of cancer patients remains to be established. Our observation of this activity in 20% of a small series of glioma patients sera may suggest that this phenomenon is more common than previously suggested. The consequence of complex formation, should it occur, are likely to differ with each patient, the type of malignancy and with the MCA used. In view of these facts, and our present findings, we would advise testing for such antibodies against the particular immunoglobulin which is to be used for in vivo immunotargeting, especially if repeat injections or large doses are contemplated. The usual intradermal skin test (for hypersensitivity) is probably inappropriate, and direct serological testing is suggested.

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