Preparation, characterisation and tumour targeting of cross-linked divalent and trivalent anti-tumour Fab' fragments

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Summary The monoclonal anti-CEA antibody, ASB7, has previously been administered to patients for radioimmunotherapy (RIT). Long circulation time and the formation of an immune response have limited therapeutic success in the clinic. Antibody fragments can be used to reduce the in vivo circulation time, but the best combination of fragment and radionuclide to use for therapy is far from clear. In this study we have compared the biodistribution of ASB7 IgG and F(ab')₂ with chemically cross-linked divalent (DFM) and trivalent (TFM) ASB7 Fab' fragments in nude mice bearing human colorectal tumour xenografts. The cross-linkers were designed to allow site-specific labelling using yttrium 90 (³⁰Y), a high-energy β-emitter. We have also compared the above antibody forms conjugated to both ¹³¹I and ⁹⁰Y. Both DFM and TFM were fully immunoreactive and remained intact after radiolabelling and incubation in serum at 37°C for 24 h. Biodistribution results showed similar tumour uptake levels and an identical blood clearance pattern for F(ab')₂ and DFM with high tumour–blood ratios generated in each case. However, unacceptably high kidney accumulation for both F(ab')₂ and DFM and elevated splenic uptake of DFM labelled with ²³⁵T were observed. Kinetic analysis of antigen binding revealed that DFM had the fast association rate (kₐ = 1.6 x 10¹³ M⁻¹ s⁻¹) of the antibody forms, perhaps owing to increased flexibility of the cross-linker. This advantage implies that DFM may be more suitable than F(ab')₂ radiolabelled with ¹³¹I for RIT. TFM cleared from the blood significantly faster than ASB7 IgG when labelled with both ¹³¹I and ⁹⁰Y, producing an improved therapeutic tumour–blood ratio. Kidney accumulation was not observed for ⁹⁰YT/TFM, but a slightly higher splenic uptake was observed that may indicate reticuloendothelial system (RES) uptake. Overall, tumour uptake was higher for ⁹⁰Y-labelled antibodies than for ¹³¹I-labelled antibodies. Because of the faster clearance, it should be possible to administer a higher total dose of ⁹⁰Y-labelled TFM than IgG, which is attractive for RIT. Both ASB7 DFM and TFM, therefore, show favourable properties compared with their parent antibody forms.

Keywords: maleimide cross-linking; radioimmunotherapy; Di-Fab; Tri-Fab

The use of antibodies to deliver radiation selectively to tumours for therapeutic and diagnostic purposes is now well established (Waldmann, 1991; Juric and Scheinberg, 1994; Larson et al., 1994). The murine ASB7 antibody raised against carcinoembryonic antigen (CEA) has been used for both imaging and therapy in nude mice bearing human colorectal tumour xenografts (Pedley et al., 1993) and in patients (Lane et al., 1994). However, results from clinical trials using this antibody, and many other studies on the treatment of solid tumours, have been disappointing, considering the success in mice (Begent and Pedley, 1990; Delaloye and Delaloye, 1995). In spite of this, some patient responses to radioimmunotherapy (RIT) have been reported (Begent and Pedley, 1990; Lane et al., 1994), suggesting that improvements to current methodology may be clinically beneficial.

There are a number of factors thought to influence antibody localisation to the tumour, which may account for the wide variation in tumour uptake levels between patients (Boxer et al., 1992). These include heterogeneous expression of antigen, variable levels of antibody achieved in the blood, the presence of circulating antigen, tumour vascularisation and the penetration of antibody into tumour tissue. In addition, RIT is often limited by the toxicity of circulating activity, particularly to the bone marrow. In attempts to overcome these problems, several strategies have been attempted to improve the delivery to tumours, while removing circulating radiolabelled antibody more rapidly, in order to reduce the dose to the bone marrow. These include the use of specific clearing regimens either in vivo or ex vivo (Begent et al., 1987; Norrgrnen et al., 1993), the use of two-step targeting regimens (Goodwin et al., 1994) and the use of antibody fragments.

F(ab')₂ fragments have been widely investigated for RIT, both in animal models and in the clinic (Lane et al., 1994). It has been shown that, in some situations, F(ab') fragments may be more suitable for RIT than intact IgG labelled with ¹³¹I (Buchegger et al., 1990; Yorke et al., 1991). However, comparatively few studies have attempted to identify the most suitable fragments for RIT with different isotopes. Small antibody fragments such as Fv and single-chain Fv (scFv) fragments showed improved penetration into tumour tissue (Yokota et al., 1992). scFv fragments also show favourable tumour–blood ratios and are potentially excellent reagents for tumour imaging (Chester et al., 1994). However, the dose of antibody delivered to the tumour by these rapidly cleared fragments is relatively low and further development is required before a therapeutic tumour dose could be delivered.

We have been evaluating alternative reagents comprising multivalent Fab' produced by chemical cross-linking. Chemically cross-linked di-Fab (DFM) and tri-Fab (TFM) have been prepared from the mouse–human chimeric Fab' fragment of B7.2.3 (King et al., 1994) and an engineered human form of the antibody (King et al., 1995). Biodistribution studies with these antibodies have revealed higher tumour accumulation than seen with scFvs, although clearance from the blood is still rapid. In addition, the choice of antibody fragment for RIT may be different for different isotopes. It is well known that the biodistribution of antibody fragments labelled with radiiodine is very different from those labelled with metallic radionuclides such as ¹¹¹In, ⁹⁰Y, ¹⁷⁷Lu, ⁶⁷Cu and ⁹⁹ᵐTc (Brown et al., 1987; Sharkey et al., 1990; Schott et al., 1992). The purpose of this study was, therefore, to determine which was the optimal form of the antibody ASB7 for RIT with the isotopes ¹³¹I and ⁹⁰Y.
In particular, we have developed methods for the production of murine versions of DFM and TFN of ASB7 to allow biodistribution studies of the murine antibody fragments in the nude mouse xenograft system. Although models of this type have been shown to have many limitations, particularly with regard to relative tumour size and accessibility, they have been shown to generate tumour–normal tissue ratios in mice similar to those observed in humans administered with the same antibody (Begent and Pedley, 1990). The biodistribution of the murine reagents produced here may be more representative of the behaviour of engineered human antibodies in patients than direct mouse studies with the recombinant humanised reagents.

Material and methods

Preparation of F(ab')2 fragments

ASB7, (IgG 1) was dialysed into 0.1 M sodium acetate, pH 5.5, containing 3 mM EDTA and concentrated by ultrafiltration using a pressurised stirred cell fitted with a YM10 membrane (Amicon) to 10 mg ml⁻¹. Bromelain (5 mg) (Sigma) was incubated with freshly prepared 50 mM cysteine (4 ml) in the same buffer for 30 min at 37°C. Activated bromelain was buffer exchanged on a Sephadex G-25 prepacked PD-10 column (Pharmacia) and the concentration of enzyme determined by absorbance at 280 nm. Digestion was performed by incubation of antibody (approximately 100 mg) with enzyme at a ratio of 50:1 (w/w) with gentle mixing at 37°C. Digestion was periodically monitored by high-performance liquid chromatography (HPLC) gel filtration analysis on a Zorbax GF-250 analytical column (DuPont), at a flow rate of 1 ml min⁻¹ with a 0.2 M sodium phosphate, pH 7.0, mobile phase, and was usually complete in 1 h. The reaction mix was adjusted to pH 6.0 with 0.1 M sodium hydroxide and bromelain was rapidly removed at 4°C by loading onto an SP-Sepharose (Pharmacia) column (approximately 30 ml) equilibrated with 0.1 M sodium acetate, pH 6.0. F(ab')₂ was eluted with 0.5 M sodium chloride and further purified by gel filtration on a 2 x 2.6 cm diameter Sephacryl S-200 (Pharmacia) column equilibrated in 0.1 M sodium acetate/0.1 M potassium chloride buffer, pH 6.0, containing 3 mM diethylenetriamine penta-acetic acid (DTPA), at a flow rate of 0.3 ml min⁻¹.

Cross-linking

Purified F(ab')₂ was concentrated by amicon ultrafiltration to approximately 5 mg ml⁻¹ and buffer exchanged to 0.1 M sodium acetate, pH 8.0, containing 2 mM DTPA. F(ab')₂ was partially reduced to form Fab' fragments to produce a free hinge thiol for cross-linking, using 2-mercaptoethanolamine (Fluka) at a final concentration of 5 mM for 30 min at 37°C. The reducing agent was then removed by desalting on a PD-10 column, and the presence of free thiols was measured by titration with 4,4'-dithiodipyridine (Sigma) and measurement of the thiopyridine released at 324 nm. Cross-linking to DFM was performed by incubating desalted reduced Fab' with 12-N4-macrocycle containing dimaleimide linker CT52, for 2 h at 37°C at a molar ratio of 2.1. TFN was prepared using Fab' in the same way by cross-linking with a 12-N4-macrocycle containing trimaleimide linker CT998, at a 3-fold molar excess of Fab'–linker. Detailed synthesis and chemical structure of the cross-linkers, CT52 and CT998, has been previously described by King et al. (1994).

Preparation of antibodies for ⁹⁹ᵐY labelling was performed under metal-free conditions. This was achieved by use of Milli-Q SP deionised water (Millipore), high-quality reagent buffers and metal-free containers. Purification of DFM and TFN after cross-linking was achieved by HPLC gel filtration using a Zorbax GF-250XL column run at a flow rate of 3 ml min⁻¹ in 0.2 M sodium phosphate, pH 7.0, containing 2 mM DTPA. To confirm purity, SDS–PAGE under non-reducing conditions was performed. To check for the presence of reoxidised F(ab')₂, in the DFM, a small sample of DFM was reduced as above and alkylated with excess N-ethylmaleimide. HPLC analysis was used to calculate the percentage of F(ab')₂ in the sample relative to DFM.

Preparation of IgG and F(ab')₂ macrocycle conjugates

ASB7 IgG and F(ab')₂ were dialysed into 0.1 M sodium phosphate, pH 8.0, containing 2 mM DTPA and incubated with a 10-fold molar excess of 2-iminothiolane hydrochloride (Sigma) for 30 min at room temperature. Unreacted reagent was removed by desalting on a PD-10 column equilibrated with the same buffer at pH 6.0, and a thiol assay was performed. A 3-fold molar excess of CT52 containing the 12-N4-macrocycle group for ⁹⁹ᵐY labelling (Harrison et al., 1991) was added to thiolated antibody or F(ab')₂, and incubated for 2 h at room temperature. The number of macrocycles per antibody was determined by measuring the number of thiol groups remaining after addition of macrocycle. Suitable controls revealed that the loss of thiol groups by oxidation over time was negligible.

Antigen binding and kinetic analysis

To determine the immunoreactivity of DFM, TFN and macrocycle conjugates compared with unmodified IgG and F(ab')₂, enzyme-linked immunosorbent assays (ELISAs) were performed. Microtitre plates were coated with 100 µl of CEA antigen (perchloric acid extracted from a patient and affinity purified in the department of Clinical Oncology) at 2 µg ml⁻¹ in phosphate-buffered saline (PBS, Sigma) for 1 h at room temperature, then blocked with 3% bovine serum albumin (BSA, Sigma) in PBS overnight at 4°C. Serial doubling dilutions (100 µl) of 10 µg ml⁻¹ DFM, TFN, F(ab')₂, IgG and macrocycle conjugates of IgG and F(ab')₂ in PBS/0.05% Tween 20, were applied to washed coated plates, for 1 h with gentle mixing. Plates were washed four times with PBS/0.05% Tween 20 and four times with distilled water, and 100 µl of anti-mouse peroxidase 1:1000 dilution (Amersham) was added and incubated for 1 h with gentle mixing. After washing, 100 µl of the substrate O-phenylenediamine dihydrochloride (OPD, Sigma, 10 mg tablet dissolved in 40 ml of 50 mM citrate buffer, pH 5.0, with 8 µl of hydrogen peroxide) was applied to each well. After approximately 5 min, the reaction was stopped by the addition of 4 M hydrochloric acid. The optical density at 490 nm was measured using a plate reader (Becton-Celltech Therapeutics). This assay was repeated to ensure consistent results.

Kinetic analysis was carried out by surface plasmon resonance using the BIAcore system (Pharmacia Biosensor) to measure on and off rates of the above antibodies. CEA antigen was immobilised to biosensor chips using surface thiol chemistry or, alternatively, by aldehyde chemistry, and the immobilised antigen density was optimised in a similar way to that described previously (Abrahám et al., 1995). Kinetic binding parameters were calculated using BIA evaluation software after correcting concentrations to nm binding sites (assuming two binding sites per antibody for IgG, F(ab')₂, and TFN, three for TFN and one binding site for F(ab')₂), such that antibody forms with different numbers of binding sites could be compared.

Radiolabelling and animal studies

Labelling with ¹²⁵I was performed using the chloramine T method. Free iodine was removed using a PD-10 column blocked with 3% BSA and equilibrated in PBS, and percentage incorporation of radiolabel was analysed by thin-layer chromatography (TLC) analysis in 80% methanol. Antibodies for ⁹⁹ᵐY labelling were first desalted into 0.1 M MES buffer, pH 6.0, at concentrations >1 mg ml⁻¹. ⁹⁹ᵐYtrium chloride (⁹⁹ᵐYCl₃, Amersham) at 50 mCi ml⁻¹ was added to achieve a specific activity of 2 mCi µg⁻¹ and...
incubated for 20 min at room temperature. The reaction was quenched by addition of 10 mM DTPA for 10 min at room temperature. Incorporation was measured by TLC in a mobile phase of 0.1 M citrate buffer, pH 5.0, and HPLC gel filtration was used to remove any unreacted \(^{90}\)Y. Characterisation of antibodies after labelling was performed to ensure full immunoreactivity by applying a dilution of the radiolabelled antibody to a 1 ml CEA affinity column and measuring the percentage bound, as described previously by Casey et al. (1995). Stability of radiolabelled antibody was analysed by application of an aliquot of the sample to a Sepharyl S-300 column (110 x 1 cm). Fractions (1.3 ml) were monitored for \(^{111}\)I or \(^{90}\)Y levels. Stability at 37°C in human serum for 24 h was also assessed.

Comparative biodistribution experiments were performed in the nude mouse colorectal xenograft model, LS174T (Pedley et al., 1993).

**Results**

**Preparation of F(ab')\(_2\), DFM and TFM**

Digestion of A5B7 with bromelain enzyme routinely produced approximately 70% fully immunoreactive F(ab')\(_2\) in 1 h. Purification of this F(ab')\(_2\) resulted in material which was >90% pure as assessed by SDS-PAGE (Figure 1). Cross-linking of Fab\(^\prime\) to DFM was monitored by HPLC gel filtration after a 2 h incubation period and resulted in yields of approximately 40% cross-linked dimer with less than 5% reoxidised F(ab')\(_2\). For TFM cross-linking, yields of approximately 25% cross-linked trimer were obtained. These yields are relatively low compared with those obtained previously with recombinant antibody fragments, where yields of 70% (DFM) and 60% (TFM) have been observed (King et al., 1994). This is probably a result of using murine A5B7 Fab\(^\prime\), which contains three hinge cysteine residues compared with recombinant Fab\(^\prime\), which has a single hinge thiol residue. Optimisation studies have shown that higher cross-linking levels may be achieved on a larger scale at higher antibody concentrations. DFM and TFM were purified by HPLC gel filtration to >90% purity as illustrated by SDS-PAGE analysis in Figure 1.

**Antigen binding and kinetic analysis**

Antigen binding analysis by ELISA demonstrated full reactivity of DFM and F(ab')\(_2\)-macrocycle conjugate compared with that of native F(ab')\(_2\) (Figure 2a). DFM also showed slightly higher binding than F(ab')\(_2\), and the F(ab')\(_2\)-macrocycle conjugate. TFM and IgG-macrocycle retained full reactivity (Figure 2b) compared with unmodified IgG, and TFM also demonstrated a slightly higher binding than IgG and the macrocycle conjugate. It was important to analyse the immunoreactivity of antibody after modification, as high levels of reducing agent or 2-iminothiolane have been reported to cause loss of immunoreactivity or aggregation (Turner et al., 1994).

Kinetic analysis of antigen binding was performed using surface plasmon resonance with a BIAscore instrument (Pharmacia Biosensor). The amount of CEA coupled to the sensor chip was optimised in preliminary experiments using either PDEA surface thiol or aldehyde immobilisation. The optimal antigen binding density for kinetic analysis was determined at which the antibody–antigen interaction was minimal for mass transport (768 RU for aldehyde coupling and 2544 RU for surface thiol immobilisation of CEA). Table 1 shows a comparison of association and dissociation rate constants (\(k_{on}\) and \(k_{off}\)) for the antibodies studied.
comparing both immobilised CEA surfaces, the values being a mean of four serial dilutions of antibody (333 nM, 167 nM, 83 nM and 42 nM). To compare forms of antibody with different numbers of binding sites, the values have been converted to nM binding sites for each entity. Monovalent Fab' was included in the evaluation and showed a considerably slower on rate (\(k_{\text{on}} 3.55 \times 10^4\) mean of values for both surfaces) and faster off rate (\(k_{\text{off}} 1.25 \times 10^{-4}\) mean) than the multivalent species, presumably owing to the lower avidity of a single binding site compared with the divalent or trivalent antibody forms. The on rates were significantly superior for DFM (\(k_{\text{on}} 1.6 \times 10^6\) mean) when compared with mean values for both surfaces of F(ab')\(_2\) (\(k_{\text{on}} 7.34 \times 10^6\)), IgG (\(k_{\text{on}} 5.3 \times 10^6\)) and TFM (\(k_{\text{on}} 7.33 \times 10^6\)). However, there was little difference in off rate between the divalent and trivalent antibodies, F(ab')\(_2\) and IgG: (\(k_{\text{off}} 0.17 \times 10^{-4}-0.47 \times 10^{-4}\)). Surprisingly, the increase in avidity of TFM did not produce higher kinetic binding, although there was a clear advantage between mono- and divalent constructs. The dissociation rate constants measured were close to the lower limit of detection possible with the BLAcore system. In general, the results obtained with the CEA immobilised via surface thiol

Table 1 BLAcore kinetic parameters for antibodies determined by binding to CEA immobilised by either aldehyde coupling or surface thiol chemistry

|        | \(k_{\text{on}} (1 \times 10^4 \text{ Ms}^{-1})\) | \(k_{\text{off}} (1 \times 10^{-4} \text{ s}^{-1})\) |
|--------|-----------------------------------------------|-----------------------------------------------|
|        | Aldehyde | Thiol | Mean | Aldehyde | Thiol | Mean |
| Fab'   | 3.25 (±1.3) | 3.84 (±0.6) | 3.55 | 1.44 (±4.0) | 1.06 (±1.4) | 1.25 |
| F(ab')\(_2\) | 5.17 (±1.0) | 9.50 (±1.2) | 7.34 | 0.45 (±0.04) | 0.18 (±0.04) | 0.32 |
| DFM    | 12.5 (±2.6) | 19.5 (±2.0) | 16.0 | 0.47 (±2.5) | 0.17 (±0.3) | 0.32 |
| TFM    | 5.46 (±1.7) | 9.2 (±2.1) | 7.33 | 0.43 (±2.1) | 0.17 (±1.3) | 0.30 |
| IgG    | 3.64 (±1.2) | 6.96 (±1.6) | 5.30 | 0.34 (±0.2) | 0.20 (±0.03) | 0.27 |

Values are corrected for number of binding sites and molecular weight and presented as association and dissociation constants. The values are average (mean) values calculated from analysis of four concentrations of antibody, standard deviations are in brackets. The overall mean values of both coupling methods for \(k_{\text{on}}\) and \(k_{\text{off}}\) are shown in the ‘mean’ columns.

Figure 3 Gel filtration profile of radiolabelled antibodies incubated in human serum for 24 h at 37°C. (a) DFM (■) and F(ab')\(_2\) (▲) labelled with \(^{131}\)I. (b) DFM (■) and F(ab')\(_2\) (▲) labelled with \(^{90}\)Y. (c) TFM (■) and IgG (▲) labelled with \(^{131}\)I. (d) TFM (■) and IgG (▲) labelled with \(^{90}\)Y. Molecular weight markers for 200kDa and 60kDa are marked with an arrow. C.p.m is a measure of radioactivity in counts per minute.
chemistry showed faster on rates and slower off rates than for aldehyde immobilisation chemistry shown in Table I. This probably reflects the fact that the antigen density was significantly higher for the thiol-immobilised surface allowing more rebinding events to occur during the dissociation phase than with the alternative surface.

**TLC** analysis of radiolabelled antibodies routinely revealed 96—99% incorporation of 131I and 90Y after purification. Antigen binding by application of a small sample of radiolabelled antibody to a 1 ml CEA affinity column revealed retention of 90—96% binding. A non-CEA antibody was radiolabelled as a control and 2—10% total activity bound to the column.

Antibodies after labelling and incubation at 37°C in normal human serum remained intact and non-aggregated, as shown in Figure 3 by S-300 gel filtration chromatography.

**Biodistribution study**

The biodistribution of A5B7 F(ab')2 and DFM, and also A5B7 IgG and TFM were compared in tumour-bearing mice over a 6 day period after radiolabelling with both 131I and 90Y.

**F(ab')2 and DFM**

The tissue distributions of 131I- and 90Y-labelled F(ab')2 and DFM were compared at 3, 24, 48 and 144 h time points. The biodistribution of A5B7 F(ab')2 and DFM labelled with 131I (Figure 4a and b) proved to be very similar, suggesting there is no significant difference in the stability of these fragments in vivo. Rapid clearance from the blood and normal tissues by 24 h produced high therapeutic tumour–blood ratios illustrated in Table II (38:1 F(ab')2; 26:1 DFM), which were

| Table II Tumour to blood ratios of percentage injected dose at various time intervals after injection of radiolabelled antibody |
|---------------------------------------------------|
| 3 h | 24 h | 48 h | 144 h |
| 131I[F(ab')2] | 0.8 | 38 | 50 | 71 |
| 131I[DFM] | 1.0 | 26 | 43 | 61 |
| 90Y[F(ab')2] | 0.9 | 23 | 22 | 5.0 |
| 90Y[DFM] | 0.9 | 40 | 27 | 7.0 |
| 125I[IgG] | 0.3 | 2.0 | 4.3 | 23 |
| 131I[TFM] | 0.5 | 6.0 | 12 | 44 |
| 90Y[IgG] | 0.4 | 4.0 | 5.8 | 17 |
| 90Y[TFM] | 0.4 | 15 | 15 | 14 |

Data are expressed as a mean of four mice.

**Figure 4** Biodistribution of (a) 131I[F(ab')2], (b) 131I[DFM], (c) 90Y[F(ab')2] and (d) 90Y[DFM] in nude mice bearing LS174T human tumour xenografts. Time points at 3 h (first column), 24 h (second column), 48 h (third column) and 144 h (fourth column) after i.v. injection. Results are expressed as percentage injected dose per gram of tissue; columns are a mean of four mice and bars represent standard deviations.
retained over the 6 day period. Labelling with $^{90}$Y dramatically altered the biodistribution (Figure 4c and d) leading to very high accumulation in the kidney at early time points and throughout the 6 days, producing a very different clearance pattern to the iodinated fragments. High splenic uptake was also observed, which accumulated over time, especially for DFM (144 h: F(ab’)$_2$: 8.5%; DFM 20%). Despite unfavourable uptake in the kidney and spleen, tumour uptake levels were similar to the iodinated fragments (48 h: $^{90}$YDFM 10%, $^{131}$I DF 6% injected dose per gram).

**IgG and TFM**

The blood clearance of TFM, despite its similar size (150 kDa), was significantly faster than IgG (P<0.02, using the Mann–Whitney U-test non-parametric statistical analysis) as illustrated in Figure 5. This produced superior therapeutic tumour–blood ratios (Table II) at 24 h $^{131}$I (IgG 2:1; TFM 6:1), $^{90}$Y (IgG 4:1; TFM 15:1) and 48 h $^{131}$I (IgG 4:1; TFM 12:1), $^{90}$Y (IgG 5.8:1; TFM 15:1). The levels of activity accumulated at the tumour were increased by labelling with $^{90}$Y (Figure 5c and d); this persisted over time and was most apparent at later time points, e.g. at 48 h: $^{131}$I (IgG 15% ID g$^{-1}$; TFM 10% ID g$^{-1}$), $^{90}$Y (IgG 25% ID g$^{-1}$; TFM 17% ID g$^{-1}$). This may be caused by higher retention of yttrium-labelled conjugates in cells compared with iodinated conjugates that are dehalogenated. The levels of radiolabel in the tumour for IgG and TFM were also greater than for F(ab’)$_2$, and DFM fragments, probably as a result of slower clearance from the circulation. However, the lower molecular weight fragments produced high therapeutic ratios earlier, potentially reducing toxicity and, thus, allowing larger doses to be given. In contrast to $^{90}$Y-labelled DFM and F(ab’)$_2$, kidney uptake levels of both $^{90}$YIGG and $^{90}$YTFM were relatively low. However, referring to the 144 h time point, it does appear that a greater proportion of $^{90}$YTFM clears through the kidney than $^{90}$YIgG (TFM 9%, IgG 3.4%), and this is also reflected by the higher values in the liver for $^{90}$YIgG at this time point (TFM 4.4%, IgG 12.8%). Slightly higher splenic and femur uptake was observed for the $^{90}$YTFM compared with $^{90}$YIgG, which may indicate non-specific uptake by the reticuloendothelial system (RES).

**Discussion**

It has been widely documented that antibody fragments injected into tumour-bearing nude mice give higher tumour to normal tissue ratios than the parent IgG (Buchegger et al., 1988; Pedley et al., 1993). Removal of the Fc portion may also be beneficial owing to the removal of binding sites for Fc receptors, thus lowering toxicity (Buchegger et al., 1992).
However, despite these potential benefits of using F(ab')$_2$ fragments, the lack of the Fc portion exposes the hinge region, making it more susceptible to enzymatic and/or reductive breakdown. In addition, different F(ab')$_2$ fragments have different in vivo stabilities. For example, in a study comparing murine F(ab')$_2$ from an IgG$_1$ antibody with chimeric F(ab')$_2$ of human subclasses 1, 2, and 4, it was reported that human IgG, F(ab')$_2$ fragments were relatively unstable in vivo, human IgG$_2$ the most stable and murine IgG and human IgG$_1$, F(ab')$_2$ of intermediate stability (Buchegger et al., 1992). Chemically cross-linked F(ab')$_2$ fragments and trispecific F(ab')$_2$, derivatives have been produced through use of bis-maleimide linkers using techniques developed by Glennie et al. (1987) and Tutt et al. (1991). Recombinant Fab's have also been synthesised with a single hinge thiol to facilitate cross-linking reactions (Carter et al., 1992; King et al., 1994) and scFv molecules with a C-terminal cysteine expressed, which allow cross-linking (Cumber et al., 1992). In some reports, increased in vivo stability of the cross-linked fragments has been demonstrated (Quadi et al., 1993; King et al., 1994).

However, as F(ab')$_2$ fragments vary in their in vivo stability themselves, the nature of the F(ab')$_2$ fragments must be taken into account and direct comparisons in the same system are necessary. In this study, we have shown that ASB7 chemically cross-linked DFM and native F(ab')$_2$, fragments have similar in vitro and in vivo stability. Although higher splenic uptake was observed for [9Y]DFM compared with [9Y]F(ab')$_2$, there was no significant difference in levels of activity in tissues, including blood, kidney and tumour when labelled with either $^{131}$I or $^{90}$Y.

The major site of antibody fragment catabolism in mice is the kidneys (Covell et al., 1986) and this was the normal organ with the highest level of $^{131}$I and $^{90}$Y[F(ab')$_2$] and DFMs at early time points. Radioiodinated fragments undergo metabolism to release low molecular weight fragments, which are rapidly released from cells and cleared (Press et al., 1990), whereas radiometals are retained longer owing to cellular trapping of the chelate (Pimm et al., 1989). Biodistribution of antibody fragments labelled with $^{131}$I and $^{90}$Y have been compared previously (Andrew et al., 1988). For $^{131}$I labelled F(ab')$_2$, >60% of administered activity was retained in kidneys compared with <10% for $^{90}$Y[F(ab')$_2$]. This is similar to the difference we observe here for $^{90}$Y-labelled F(ab')$_2$ and DFM; kidney: blood ratios are increased 3-fold at 3 h and over 100-fold at 24 h compared with $^{131}$I-labelled fragments.

The stability of the attachment of radiochelate to antibody is an important determinant of therapeutic ratio, since some labelling methods, involving derivatives of the chelating ligand DOTA, FPA, mononuclearised while conjugates in vivo (Harrison et al., 1991). Weak ligands will lead to both a reduction in the amount of $^{90}$Y-conjugate in the tumour and irradiation of the normal tissues with free $^{90}$Y, especially proliferating tissues of the bone marrow, since $^{90}$Y is a bone-seeking isotope (Hnatiowich et al., 1983). The development of new chelating agents, such as the macroyclic chelating agent DOTA (Harrison et al., 1988; Cox et al., 1989) may be more stable in vivo (Hird et al., 1991; Harrison et al., 1991).

The cross-linkers used in this study contain the 12N4 DOTA macrocycle, which allows the site-specific attachment of $^{90}$Y and this has the added advantage that there is no loss of antigen binding after radiolabelling. Attachment of radiolabel in random positions, as we describe here for IgG and F(ab')$_2$, leads to non-homogenous preparations owing to variation in the distribution of thiol groups generated, and may reduce immunoreactivity, if residues which contribute to antigen binding are modified.

Engineering an additional C-terminal cysteine residue on scFv or Fab' subunits and thus providing a free thiol group, may also be useful for site-specific radiolabelling with technetium-99m (Verhaar et al., 1996), again allowing retention of immunoreactivity as a result of conjugation distant from the antigen binding sites.

The increase in valency and, therefore, avidity of divalent fragments over monovalent fragments produced a significant increase in kinetic association and dissociation rate-binding parameters. However, the increase in valency from divalent to trivalent molecules did not further increase kinetic binding. Three Fab' arms may not all be accessible for binding at the same time to a solid surface. Therefore, steric hindrance to antigen sites on solid surface may explain why an increase in association and dissociation rate was not achieved. A cell binding assay may be more representative of binding parameters in vivo; in a previous study improved binding of trivalent molecules over divalent F(ab')$_2$ was demonstrated using this type of assay (Werlen et al., 1996).

The biodistribution of IgG and TFM in the xenograft model showed surprisingly faster clearance for TFM compared with whole antibody, despite having the same molecular weight. The same phenomenon has been observed with trivalent bis-maleimide-linked Fab' fragments described previously (King et al., 1994; Schott et al., 1993), and may be partially caused by the lack of the Fc portion thought to be responsible for the long circulation time of the intact immunoglobulin. Tumour accumulation of TFM was greater than for the divalent fragments, and similar levels to IgG were observed up to 24 h. The major advantage of faster clearance leading to lower blood toxicity is that for therapy a lower dose may be given. Again, by labelling with $^{90}$Y, tumour retention is superior than for iodinated IgG and TFM owing to the absence of dehalogenation and/or $^{131}$I metabolism. Several publications (Schott et al., 1992; Sharkey et al., 1990) have also reported this finding, which could also explain the higher percentage injected dose of the $^{90}$Y-labelled antibodies in normal tissues, such as kidney spleen and liver. The splenic uptake of $^{90}$Y-labelled DFM and TFM was notably high and IgG$_1$, which indicates that there may be some RES uptake of the cross-linked fragments. This will require further investigation and dosimetry evaluation with higher doses of radioconjugate.

In contrast to the divalent fragments, kidney accumulation was greatly reduced. This is presumably mainly due to the increase in molecular weight, although the slight increase in kidney levels of TFM compared with IgG suggest that there may be other factors, such as shape and charge, which also influence the filtration process (Sumpio and Hayslett, 1985). Catabolism of TFM to form F(ab')$_2$, or Fab' fragments could also occur, which may also contribute to increased kidney uptake. However, there is no evidence to suggest reabsorption of fragmented TFM or DFM, or breakdown occurring in the serum, by HPLC analysis of serum from mice injected with labelled DFM and TFM (unpublished findings).

In the present study, DFM showed faster blood clearance than TFM and IgG and similar biodistribution to F(ab')$_2$ in the xenograft model. High kidney accumulation of both $^{90}$YDFM and $^{90}$Y[F(ab')$_2$], and increased splenic uptake of $^{90}$YDFM is clearly unacceptable for RIT. However, this high kidney and splenic uptake was not seen on labelling with $^{131}$I. DFM has a faster $k_{mm}$ rate than F(ab')$_2$ and all the other antibody forms, which may be a consequence of the increased spacer or flexibility of the chelating cross-linker. Therefore, although the biodistribution data demonstrate equivalent tumour uptake levels, we conclude that ASB7 DFM, owing to its faster $k_{mm}$ rate, may be more suitable than F(ab')$_2$ when radiolabelled with $^{131}$I for RIT.

TFM clears faster from the blood than IgG and produces higher tumour uptake and lower kidney accumulation than DFM and F(ab')$_2$, radiolabelled with $^{90}$Y. Although there is a slight elevation in splenic uptake, this combination avoids the long blood circulation time, which is dose limiting, and spares the kidneys from damage by radiation accumulation. Therefore, from these results we conclude that $^{90}$Y-labelled ASB7 TFM would be the most suitable antibody species for RIT.

The superior pharmacokinetics of murine versions of DFM and TFM compared with the parent IgG and F(ab')$_2$, in vivo, coupled with the ability to re-treat patients by using
humanised forms, should provide improved clinical therapy in the future. A humanised Fab' version of ASB7 has been constructed (Adair et al., 1992) and preliminary data involving hDFM and hTFM are promising.

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