The effect of an anti-membrane antibody–methotrexate conjugate on the human prostatic tumour line PC3

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Summary Methotrexate (MTX) was linked, via an active ester intermediate, to a purified IgG fraction of rabbit polyclonal antiserum raised against a cell membrane preparation from the human prostatic cell line PC3. The resulting conjugates contained an average of 0.044 mg of MTX per mg of antibody with acceptable losses in both the binding activity of the immunoglobulin (27.5%) and the enzyme inhibitory activity of the drug (32% at a MTX concentration of 3 x 10⁻⁷M). Using cultures of PC3 cells the antibody–MTX (Ab–MTX) conjugates were observed to be as effective as free drug in causing cell death and more effective than non-immune IgG–MTX (NilG–MTX) conjugates. When androgen nude mice bearing PC3 tumours were administered with Ab–MTX conjugates, significant reductions in tumour growth rates were observed compared to animals given saline, MTX alone or NilG–MTX conjugates (P<0.01 in all cases). Furthermore, the accumulation of radioactive MTX in the tumour tissue of animals injected with these Ab–MTX conjugates was 16-fold greater than those given free drug and 8.6-fold greater than those administered with NilG–MTX conjugates. Uptake by the reticuloendothelial system, however, was not significantly different when animals from each treatment group were compared.

Following androgen ablation for the treatment of prostatic cancer, any initial positive response is almost universally followed by a relapse to a hormone independent state (Menom & Walsh, 1979). Treatment of such relapsed tumours by chemotherapy is limited by the toxicity of the agents used, and thus the targeting of such drugs to the tumour site may provide a potentially more effective therapy for hormone independent tumours, with fewer side-effects.

Antibodies have been suggested as potential carriers for drugs (Ghose & Blair, 1978) and methods of linking the drug methotrexate to antisera while retaining both antibody and drug activity have been documented (Kulkarni et al., 1981). In cultures of the prostate cell line LNCaP, incubation with a monoclonal antiserum to human prostatic acid phosphatase (PAP) linked to methotrexate has been shown to be more effective than a similar conjugate in which the antiserum was substituted by non-immune IgG (Deguchi et al., 1986).

Using this same conjugate in vivo on LNCaP tumours in athymic nude mice, however, demonstrated that although MTX from the conjugate accumulated more in the tumour than free drug, accumulation was also higher in the spleen, kidney and liver (Deguchi et al., 1986), which are known sites of MTX toxicity. As PAP from the prostate tumour can be released into the circulation, it is probable that immune complexes form between the antibody in the conjugate and circulating PAP and are taken up by the reticuloendothelial system. Antisera raised against an antigen such as a tumour cell membrane should provide a preferential localisation at the tumour site and thus reduce complex formation within the circulation. This study was therefore undertaken to assess the effectiveness of a cytotoxic drug, methotrexate, conjugated to an antiserum raised against cell membranes isolated from a human prostate cancer cell line PC3.

Materials and methods

Summary Methotrexate (sodium salt) was purchased from Lederle Laboratories (Hampshire, UK) and MTX (free carboxylic acid), NADPH, dihydrofolate reductase, dihydrofolate acid and N,N-dicyclohexylcarbodiimide (DCC) were purchased from Sigma Chemical Co. Ltd (Dorset, UK). ³¹H-MTX (250 mCi (9.25 GBq) mmol⁻¹) and ⁵¹Cr (350–600 mCi (12.95–22.2 GBq) mg⁻¹) were products of Amersham International plc (Buckinghamshire, UK). N-Hydroxysuccinimide (NHS) was supplied by Aldrich Chemical Company Ltd (Dorset, UK) and AH Sepharose 4B was obtained from PharmaChem Ltd (Uppsala, Sweden). Diethylaminoethyl cellulose (DE-52) was purchased from Whatman Ltd (Maidstone, Kent, UK) and dust-free Isoton (Pharmacia Ltd, Upton, UK), both Soluene-100 and Dimilume-30 were purchased from United Technologies (Packard, Berks., UK) and microtitre strips obtained from Costar Corporation Ltd (Cambidge, UK). All tissue culture dishes and plates were supplied by Becton-Dickinson UK Ltd (London).

PC3 cell line in vitro and in vivo

Athymic nude mice were routinely bred from Nu/Nu males and heterozygous Nu/+ females and housed in polycarbonate filter-top cages. Irradiated breeding diet (Pilsbury’s Ltd, Birmingham, UK) and acidified water (pH 2.8) were given ad libitum. Controlled light and temperature conditions were maintained with a 12 h light/dark cycle and a temperature of 20–22°C.

The human prostate cell line PC3 (Kainth et al., 1979) was maintained in Dulbecco’s modified Eagles medium (DMEM) supplemented with fetal calf serum (7% v/v), penicillin (200 IU ml⁻¹), streptomycin (100 μg ml⁻¹), fungizone (5 μg ml⁻¹) and glutamine (1.46 mg ml⁻¹). Cultured PC3 cells required for implantation were trypsinised (trypsin 0.025%; versene 0.02%, v/v) and resuspended in phosphate buffered saline (PBS). The cells were injected subcutaneously into the flanks of 6–8-week-old mice (10⁵ cells per site) and tumours appeared 2–3 weeks later.

Isolation of PC3 cell membranes, immunisation of animals and preparation of an immunoglobulin fraction

Cell membranes were partially purified according to the method of Thom et al. (1977). PC3 cell monolayers were grown in 75 cm² tissue culture flasks, harvested using a cell scraper and lysed in hypo-osmotic borate/EDTA (boric acid 0.02M, EDTA 0.2mM, pH 10.2). The cytoplasmic contents were expelled and gelatinised and the empty membrane sacs collected by differential centrifugation (450 g for 10 min; 24,000 g for 1 h). Samples were routinely prepared for electron microscopy to check for purity. An aliquot of the membrane protein, equivalent to 5 mg of soluble protein

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(as assessed by the Bio-Rad protein assay) was injected intradermally with Freund's complete adjuvant into New Zealand white rabbits according to the multiple site injection technique described by Vaitukaitis et al. (1971). The rabbits were boosted with the same amount of immunogen at monthly intervals and blood samples (20 ml) obtained at 14-day intervals by incision of the marginal ear vein.

IgG fractions were obtained from both immune and non-immune sera by ammonium sulphate precipitation (45% saturation) followed by ion-exchange chromatography on DE-52 by the method described by Fahey and Terry (1973), in which the IgG fraction was eluted from the column by 0.005 M phosphate buffer (pH 7.4). The specificity of the antiserum was assessed by immunocytochemical techniques on 5 μm sections of various rat, mouse and human tissues using the unlabelled antibody enzyme method developed by Sternberger et al. (1970) and an adapted protocol reported previously (Sibley et al., 1984).

Preparation of immunoglobulin–methotrexate conjugates

Trinitiated MTX was mixed with unlabelled MTX to give a final specific activity of 4.2 μCi (0.155 MBq) mg⁻¹ and conjugated to both anti-membrane IgG and normal rabbit IgG via the active ester intermediate method described by Kulkarni et al. (1981). MTX (30 mg) was dissolved in dimethylformamide (DMF, 0.1 ml) and added to N-hydroxysuccinimide (15.21 mg per 0.1 ml DMF) and N,N'-dicyclohexylcarbodiimide (27.26 mg per 0.1 ml DMF). The reaction mixture was allowed to stand at room temperature for 1 h followed by 18 h at 4°C in the dark. The resulting precipitate was removed by centrifugation (10,000 g for 10 min) and the supernatant containing the active ester was stored at 4°C in the dark. Determination of the amount of active ester produced was carried out using the method described by Kulkarni et al. (1981), but employing AH Sepharose 4B columns instead of AFFI-Gel 102. The amount of active ester was calculated by subtracting the amount of 3H-MTX passing through the column after binding from the amount of 3H-MTX in the eluate following hydrolysis, and gave an average yield of 77.6% (five separate experiments).

The active ester of MTX (1 mg in DMF) was incubated with anti-PC3 membrane IgG or normal rabbit IgG (10 mg per 2 ml) on a roller-bed for 2 h at 4°C in the dark. Precipitated protein was removed by centrifugation (10,000 g for 10 min) and the supernatant containing the IgG–MTX conjugates dialysed against PBS (pH 7.4) for 18 h at 4°C, to remove free drug. Further separation from free MTX was carried out by gel filtration on Sephadex G-50 with PBS (pH 7.4). The concentration of MTX in the conjugates was estimated by measurement of the radioactivity present, and calculation with reference to the original standard of 4.2 μCi (0.155 MBq) mg⁻¹ MTX employed in the conjugation procedure. Protein concentration in the Ig–MTX conjugates was estimated using the Bio-Rad protein assay kit. The conjugates contained an average of 25 μg of MTX ml⁻¹ and 560 μg of protein ml⁻¹ of PBS.

Inhibition of dihydrofolate reductase activity by Ab–MTX conjugates

Pharmacological activity of MTX in the conjugate was assayed by its ability to inhibit the enzymatic activity of dihydrofolate reductase (DHFR) and measured spectrophotometrically using the method described by Balk et al. (1976). MTX standards and Ab–MTX conjugates (2 × 10⁻⁸ to 3 × 10⁻⁴ M) were incubated separately with the enzyme (0.1 units ml⁻¹) at room temperature for 15 min to allow the MTX to bind to the enzyme. Dihydrofolinic acid (50 μl of 0.25 mg ml⁻¹) was added to each tube and the absorbance measured spectrophotometrically, at 340 nm, after 6 min.

Inhibition of DHFR activity was calculated using the following formula:

\[
\% \text{ inhibition} = \left( \frac{\text{DHFR activity (control)} - \text{DHFR activity (MTX or conjugates)}}{\text{DHFR activity (control)}} \right) \times 100
\]

Competitive binding of Ab–MTX conjugates to PC3 membrane fraction

Assessment of the binding of the Ab–MTX conjugates to PC3 membranes was determined by a solid-phase competitive binding assay using anti-membrane IgG labelled with ¹²⁵I by the chloramine-T method (Hunter & Greenwood, 1962). Membrane fractions prepared from PC3 cells at a concentration of 10 μg ml⁻¹ in 0.1 M sodium carbonate/bicarbonate buffer (pH 9.5) were adsorbed onto plastic microtitre plates by incubation (100 μl per well) overnight at 4°C. Known amounts (1–250 μg protein ml⁻¹ PBS) of anti-membrane antibody or Ab–MTX conjugates were incubated (100 μl per well) with 0.2 μCi (0.74 MBq) of ¹²⁵I-labelled anti-membrane antibody (2 μCi mg⁻¹; 74 MBq) at 37°C for 3 h. The wells were washed (10 × 100 μl PBS) and the radioactivity counted on a gamma counter. The percentage inhibition of membrane IgG or Ab–MTX conjugates was calculated using the following formula:

\[
\% \text{ inhibition} = \left( \frac{\text{c.p.m. (control)} - \text{c.p.m. (anti-membrane IgG or conjugates)}}{\text{c.p.m. (control)}} \right) \times 100
\]

The loss of binding activity displayed by the Ab–MTX conjugates was then calculated from the following formula:

\[
\% \text{ loss of inhibition} = \left( \frac{\% \text{ inhibition (anti-membrane IgG)} - \% \text{ inhibition (conjugates)}}{\% \text{ inhibition (anti-membrane IgG)}} \right) \times 100
\]

Assessment of the cytotoxicity of the Ab–MTX conjugates in vitro

Estimation of cell number and cell size distribution

The cytotoxicity of the Ab–MTX conjugates was assessed by measuring the number of cells remaining in the cultures after treatment in comparison to controls. Monolayers of PC3 cells were subcultured with trypsin/versene and seeded onto 24-well plates at a density of 5 × 10⁵ cells ml⁻¹ DME. The cells were allowed to recover from trypsinisation for 24 h at 37°C before incubation with Ab–MTX conjugates (90 μg Ab ml⁻¹, 4 μg MTX ml⁻¹), Ab alone (90 μg ml⁻¹), MTX alone (0.4–4 μg ml⁻¹), Ab plus MTX combined but not conjugated (90 μg Ab ml⁻¹, 4 μg MTX ml⁻¹), and IgG–MTX conjugates (65 μg IgG ml⁻¹, 4 μg MTX ml⁻¹) in DME, or DME alone (control) for 2 h at 37°C, using quadruplicate cultures per experimental group. After this time medium was aspirated from the wells, the cultures washed with PBS (5 × 2 ml), fresh medium (1 ml) added to each well and the cultures incubated for 72 h at 37°C. Cells were counted at the end of this period by disrupting the monolayers with trypsin/EDTA and diluting the cells with isoton (15 ml). The suspended cells were counted on a Coulter counter (model ZBI) equipped with a multi-channel analyser facility and an x–y plotter.

Total cell population per well was determined by taking account of machine coincidence error, sample volume and dilution of the suspension. Any accumulation of cells within a certain phase of the cell cycle was determined by analysis of the cell size distribution of each experimental group using the multichannel analyser.

Release of ¹⁵⁷Cr from pre-loaded cells

Release of ¹⁵⁷Cr from pre-loaded PC3 cells was measured by a modification of the method described by Wigzell (1965), and gives an estimation of cell death. PC3 cells were seeded onto 24-well plates, at a density of 5 × 10⁵ cells ml⁻¹ and allowed to recover from trypsinisation for 24 h at 37°C. Pre-labeling of the cells was carried out by the addition of ¹⁵⁷Cr in DME (20 μCi
(0.74 MBq) x 10^6 cells) for 30 min and each well washed individually with PBS (6 x 2 ml). Monolayers of PC3 cells were incubated with Ab – MTX conjugates (90 μg Ab ml⁻¹, 4 μg MTX ml⁻¹) pure Ab alone (90 μg ml⁻¹) MTX alone (0.4 – 4 μg ml⁻¹), Ab plus MTX combined but not conjugated (90 μg Ab ml⁻¹, 4 μg MTX ml⁻¹), NlgG – MTX conjugate (83 μg IgG ml⁻¹; 4 μg MTX ml⁻¹) in DME, or DME alone (control) for 2 h at 37°C. After this time the media was aspirated from the cells, each well washed with PBS (5 x 2 ml) and fresh medium added (1 ml) to each culture. The ³¹Cr released from the cells was measured 24 h later by removal of the media, washing each well individually (2 x 2 ml DME) and counting the total radioactivity in the medium from each well on a gamma counter.

Assessment of the cytotoxicity of Ab – MTX conjugates in vivo

The anti-tumour activity of the Ab – MTX conjugates was assessed in vivo by their action on the growth of PC3 tumours in athymic nude mice. Mice bearing subcutaneous tumours (0.3 – 1.1 cm diameter) were divided into six groups (16 animals per group) and injected with Ab – MTX conjugates, Ab alone, MTX alone, Ab + MTX combined but not conjugated, NlgG – MTX conjugates or saline. Drug and antibodies were administered i.m. every 48 h for 12 days at a dose of 1 mg kg⁻¹ H-MTX (22 μg per animal) in saline (100 μl) and an antibody concentration equivalent to that found in the Ab – MTX conjugate (approximately 498 μg per animal). Tumour size was recorded every 2 days as the mean of two perpendicular diameters, one across the greatest width, and the tumour volume estimated using the formula 4/3πr³, where r is the mean radius. Tumour volume calculated in this way was found to have a linear relationship with both tumour weight and tumour volume as measured by water displacement. The errors involved in using this formula were found to be negligible.

Tissue distribution of H in animals treated with Ab – MTX conjugates

Twenty-four hours after the sixth injection of the animals under study, the animals were anaesthetised, bled by cardiac puncture and killed by cervical dislocation. The tumour and various body tissues were excised and known amounts (up to 100 mg wet weight) were individually placed in counting vials. Tissues were solubilised by the addition of Soluene (1 ml) and incubated at room temperature for up to 48 h before the addition of Dimilume scintillation fluid (5 ml). The radioactivity in each sample preparation was determined by scintillation counting and quenching corrected for by internal standardisation. The concentration of MTX in the tissues was estimated per g wet weight of tissue by extrapolation from the specific activity of the starting material of H-MTX (4.2 μCi (0.155 MBq) mg⁻¹).

Statistical evaluation

A Mann – Whitney U test was used to compare the treatment data derived from ³¹Cr release and cell population studies in vitro and also from the tissue distribution of H-MTX from the various treatment groups studied in vivo. Tumour volume data were analysed by two-way analysis of variance (ANOVA) providing values for an F statistic and hence a P value, reflecting the level of statistical significance.

Results

Conjugation procedures and assessment of antibody and drug activity

Methotrexate was conjugated to anti-membrane IgG and non-immune IgG via an active ester intermediate. The extent of substitution was 14.6 mol of MTX per mol of IgG with a 64% recovery and a 27.5% loss in immunological activity, as assessed by a competitive binding assay (Table I).

The immunoglobulin fraction used in this study was not completely specific for prostatic tissue as assessed by immunocytochemical techniques. At an antibody dilution of 1/500 (PBS, pH 7.5), human breast, bladder, kidney and intestinal epithelium also exhibited some staining although with a much lower intensity than that observed with benign and carcinomatous prostatic tissue. Staining was always associated with the epithelial cell membrane and no cross-reactivity was observed with any mouse or rat tissues. When equivalent concentrations of Ab and Ab – MTX conjugates were compared in the immunocytochemical procedure, no diminution in membrane staining or change in tissue specificity of the antibody was observed after conjugation (data not shown).

Comparison of free MTX and Ab – MTX conjugates to inhibit the enzyme activity of DHFR indicated that the MTX within the conjugate retained approximately 68% of its ability to bind to the enzyme at a drug concentration of 3 x 10⁻² M (Figure 1).

Cytotoxicity testing in vitro

Cytotoxicity testing was carried out using cultured PC3 cells, each experiment being carried out at least three times. Conjugate and drug concentrations were adjusted to 4 μg MTX ml⁻¹ unless otherwise stated, and antibody concentrations to 90 μg protein ml⁻¹ (equivalent to the amount of Ab in the Ab – MTX conjugates at 4 μg ml⁻¹). Incubations with drugs and conjugates were carried out for 2 h at 37°C and the results expressed as the means from quadruplicate cultures.

Estimation of cell number

The differences in cell population with the various treatments are shown in Figure 2 and are expressed as number of cells per ml of medium. Incubations with increasing concentrations of MTX (0.4 – 4 μg ml⁻¹) produced a progressive decrease in the number of cells, at 4 μg ml⁻¹ MTX the value being 14.6% of the control population. The Ab – MTX conjugates reduced the cell population to 14.9% of the control, and no significant difference was observed between the values obtained for Ab – MTX conjugates and free drug (4 μg ml⁻¹).

Significantly larger cell numbers were observed (P < 0.05) when the PC3 cells incubated with NlgG – MTX conjugates were compared to cells treated with Ab – MTX conjugates. The addition of membrane Ab alone to cultured cells led to a reduction in cell number compared to controls (P < 0.05). Cultures incubated with Ab plus MTX, combined but not conjugated, however, displayed a larger cell population than incubation with MTX alone (P < 0.05).

Cell size distribution

As the inhibition of DHFR leads to inhibition of DNA, RNA and protein synthesis (Bertino, 1963), cells treated with MTX would be expected to be

Table I

| Concentration of MTX-antibody conjugates (μg) | % inhibition of ¹²⁵I-antibody binding | % loss of binding activity in the conjugates |
|---------------------------------------------|-------------------------------------|-------------------------------------------|
| 250                                         | 48.7                                | 66.3                                      |
| 100                                         | 40.2                                | 56.8                                      |
| 50                                          | 39.3                                | 52.5                                      |
| 25                                          | 31.2                                | 45.7                                      |
| 10                                          | 28.2                                | 37.6                                      |

PC3 membrane fractions were adsorbed onto microtitre wells (100 μg protein per well) and known amounts (1 – 250 μg) of Ab – MTX conjugates or Ab were incubated with ¹²⁵I labelled Ab (2 μCi, 7.4 MBq mg⁻¹; 10μg per well) for 3 h at 37°C. After washing, (10 x 100 μl PBS) total radioactivity was assessed per well. Data are expressed as the mean percentage inhibition from 16 experimental wells.
expressed 3). Methotrexate and Ab-MTX conjugates (2 x 10⁻⁸ to 3 x 10⁻⁷ M, 100 μl per tube) were incubated with the enzyme (0.1 units per tube) in the presence of NADPH (16.6 μg) for 15 min at 25°C. Dihydrofolinic acid (12.5 μg in 50 μl) was dispensed into each tube and the change in absorbance measured spectrophotometrically at 340 nm after 6 min.

Figure 1 The inhibition of dihydrofolate reductase (DHFR) by MTX (---) and Ab-MTX conjugates (-----). Methotrexate and Ab-MTX conjugates (2 x 10⁻⁸ to 3 x 10⁻⁷ M, 100 μl per tube) were incubated with the enzyme (0.1 units per tube) in the presence of NADPH (16.6 μg) for 15 min at 25°C. Dihydrofolinic acid (12.5 μg in 50 μl) was dispensed into each tube and the change in absorbance measured spectrophotometrically at 340 nm after 6 min.

Figure 2 The effect of the Ab-MTX conjugates on the number of PC3 cells remaining in culture after treatment. PC3 cells (5 x 10⁵ per well) were incubated with DME (1 ml) containing MTX (0.4-4 μg ml⁻¹), Ab-MTX conjugates (90 μg Ab ml⁻¹; 4 μg MTX ml⁻¹), Ab-MTX conjugates (83 μg IgG ml⁻¹; 4 μg MTX ml⁻¹), Ab-MTX conjugates (90 μg Ab ml⁻¹; 4 μg MTX ml⁻¹ or DME alone. Treatments were carried out for 2 h at 37°C, the media removed from each well and replaced with fresh DME (1 ml). After a further incubation of 72 h cells were trypsinised, diluted with Isoton and total cell population per well measured on a Coulter counter. Data represent mean and + s.d. of quadruplicate cultures.

arrested in the S phase of the cell cycle. The distribution of cell size within control and drug treated cultures was expressed as the number of cells against cell volume (Figure 3). Control cultures displayed a distribution pattern as seen in Figure 3a and cells treated with Ab alone had an identical profile (Figure 3b). Incubation of the cells with Ab-MTX conjugates showed a slight shift towards a larger cell volume (Figure 3b), whereas cultures treated with either MTX alone (4 μg ml⁻¹, Figure 3a) or Ab-MTX conjugates (Figure 3b) displayed a pronounced shift towards larger cell volumes indicating an accumulation of cells in the S phase of the cell cycle for these latter two treatments.

The release of ⁵¹Cr from pre-loaded cells The release of ⁵¹Cr into the medium from pre-loaded PC3 cells incubated with the various drug concentrations and conjugate are shown in Figure 4 and expressed as d.p.m. ml⁻¹. With increasing concentrations of MTX there was a progressive rise in the amount of ⁵¹Cr liberated from the cells, reaching a value 5.6 times higher than the control of 4 μg ml⁻¹ of drug. Essentially similar results were observed after incubation of the cells with Ab-MTX conjugates as obtained with free drug. The IgG-MTX conjugates were not as effective in causing the release of ⁵¹Cr from the cells as the Ab-MTX conjugates (P < 0.05), but did display some cell killing activity as the ⁵¹Cr content of the media was significantly larger than that seen in control cultures (P < 0.01). Cells treated with membrane Ab plus MTX displayed a significantly lower liberation of ⁵¹Cr than that observed with free MTX alone (P < 0.05). Incubation of the cultured cells with membrane Ab alone demonstrated a higher release of ⁵¹Cr into the medium than determined in medium from control cultures (P < 0.05).
Assessment of the cytotoxicity of Ab–MTX conjugates in vivo

Measurement of the size of individual PC3 tumours growing in mice receiving the various drugs and conjugates was carried out over a 12-day period and is expressed as the percentage change in tumour volume against time (Figure 5). Control animals, treated with saline only, showed a 15-fold increase (mean from 16 animals) in tumour volume over the 12-day period studied. Although NlgG–MTX conjugates appeared to suppress tumour growth, analysis of variance on the tumour data showed no significant difference from controls. Injections of MTX alone, Ab alone or Ab plus MTX resulted in a significant inhibition of tumour growth when compared to the control group ($P<0.002$). Animals treated with Ab–MTX conjugates, however, displayed an inhibition of tumour growth that was significantly greater than all other treatment groups ($P<0.001$).

Distribution of $^3$H-MTX in vivo

The Ab–MTX conjugates, NlgG–MTX conjugates and free MTX given to animals all contained $^3$H-MTX (4.2 μCi (0.155 MBq) mg$^{-1}$ MTX). Measurement of the label was undertaken in five animals from each treatment group in order to assess the uptake and distribution of the tritium label within the animals. The results for each tissue and treatment are expressed as ng MTX g$^{-1}$ wet weight of tissue and also as tissue/blood ratios (Table II). The highest accumulation of radioactivity was observed in the tumours of animals treated with Ab–MTX conjugates, the tumour: blood ratio being 59.7, compared with 7.4 for animals treated with MTX alone, 6.6 for those given Ab plus MTX and 6.3 for those receiving NlgG–MTX conjugates. The results show (Table II) that the distribution of label in the lung, heart, kidney, liver, spleen and skin vary slightly, but not significantly between the treatment groups.

Significantly higher levels of radioactivity ($P<0.05$) were observed within the blood of animals given Ab–MTX conjugates or NlgG–MTX conjugates compared to those receiving free drug indicating a difference in the rate of clearance.

Discussion

To maintain therapeutic effectiveness of cytotoxic drug linked to antibodies, conjugation should not lead to a substantial loss in either drug activity or in antibody binding and specificity. Therefore, in this present study the resulting conjugation of methotrexate with a polyclonal antiserum raised against a prostatic tumour cell line PC3 was assessed in several systems which examined both the drug cytotoxic ability in vitro and in vivo and the localisation in vivo.

The retention of antibody activity after conjugation with

Table II Tumour and tissue distribution of $^3$H-MTX and conjugated

| Tissue | $^3$H-MTX, ng g$^{-1}$ wet weight | $^3$H-MTX | $^3$H-MTX–Ab conjugates | $^3$H-MTX–NlgG conjugates |
|--------|----------------------------------|----------|-------------------------|--------------------------|
| Lung   | 290.2 (6.9) 244.9 (5.0) 185.9 (2.1) 201.7 (2.1) |
| Heart  | 394.4 (9.4) 234.1 (4.8) 156.8 (1.8) 234.4 (2.4) |
| Kidney | 718.5 (17.1) 474.6 (9.7) 1064.0 (12.2) 972.7 (10.1) |
| Liver  | 906.5 (21.6) 777.9 (15.9) 1063.7 (12.2) 1674.4 (17.4) |
| Spleen | 185.4 (4.4) 353.8 (7.2) 293.7 (3.4) 471.3 (4.9) |
| Skin   | 230.8 (5.5) 293.0 (6.0) 306.2 (3.5) 619.4 (6.4) |
| Tumour | 312.1 (7.4) 323.8 (6.6) 5214.8 (59.7) 604.2 (6.3) |
| Blood  | 41.9 48.8 87.4 96.4 |

Athymic nude mice bearing palpable PC3 tumours (0.3–1.1 cm diameter) were injected i.m. with MTX, Ab–MTX conjugates, NlgG conjugates, Ab alone, Ab + MTX or saline every 48 h over a period of 12 days. Each dose of MTX was adjusted to 1 mg MTX kg$^{-1}$ body weight (22 μg per animal in 100 μl saline) and the amount of Ab given to the control groups was equivalent to the concentration of Ab present per 22 μg of MTX in the Ab–MTX conjugates. Tumour size was measured with calipers before each injection and tumour volume calculated from the formula $4/3πr^3$. Data are expressed as a percentage increase in tumour volume from day 0, each value representing the mean from 16 experimental animals, and the standard errors of the mean.
the least interruption in structure and activity is clearly of major importance, except drug linkage or adverse reaction could then lead to denaturation or aggregation of the antibody. In the studies described in this paper, conjugation by an active ester intermediate method produced a molar incorporation ratio of drug to antibody of 14.6 with a 72.5% retention of antibody binding, as estimated by quantitative membrane binding analysis and no detectable change in membrane specificity as assessed by immunocytochemical techniques. A similar molar ratio was reported for conjugation of a methotrexate conjugate with a monoclonal antibody against prostatic acid phosphatase which retained over 90% of its antibody activity (Deguchi et al., 1986). Such differences may be explained by the availability of amino groups on the particular immunoglobulins under study, an idea supported by the results of Kanellos et al. (1985), who noted that although conjugation of methotrexate to two separate monoclonal antibodies produced the same molar ratio of drug and antibody their retention of antibody activity was markedly different.

Assessment of drug activity by the inhibition of dihydrofolate reductase showed some loss in the capacity of methotrexate to inhibit the enzyme when conjugated compared to an equimolar amount of free drug (Figure 1). Such losses of dihydrofolate reductase inhibition have been reported previously when methotrexate was conjugated to several different macromolecules (Deguchi et al., 1986; Fung et al., 1979; Garrett et al., 1983; Kanellos et al., 1985; Kulkarni et al., 1985; Tsudo et al., 1980) and is probably due to the stearic interference of the macromolecule with the binding of the pteridine moiety of the methotrexate to the active site of the enzyme (Baker, 1969). Studies suggest that a reduced ability to inhibit dihydrofolate reductase activity also occurs within the cell. Deguchi et al. (1986) observed that methotrexate conjugated to immunoglobulin was less effective than free drug at inhibiting the incorporation of 3H-deoxyuridine by LNCaP cells in culture, even though methotrexate in the conjugated form was taken up to a greater extent than free methotrexate. This finding is supported by Uadia et al. (1985), who showed that when human melanoma cells were treated with 3H-methotrexate–antibody conjugates, 35% of the radioactivity was found in the low molecular weight fraction and the methotrexate derivatives in this fraction were no more potent inhibitors of dihydrofolate reductase than the original conjugate. Their study suggests that the methotrexate derivatives digested within the cell are biologically less potent than free drug.

In our tissue culture experiments, determination of the cell volume distribution patterns (Figure 3) revealed that the antibody methotrexate conjugate displayed the same profile as methotrexate alone. As the inhibition of dihydrofolate reductase by methotrexate ultimately results in the cessation of purine and amino acid synthesis (Bertino, 1963), the altered distribution pattern displayed by the cultures incubated with free drug compared to control cultures would be expected, reflecting the accumulation of arrested cells in the S phase of the cell cycle. The similar profiles obtained with cells treated with both the antibody–methotrexate conjugates and with free drug would therefore indicate that the methotrexate moiety of the conjugate was still able to inhibit dihydrofolate reductase activity within the cell and hence arrest the cells in S phase, even though potency of the drug may be reduced.

Conjugation of cytotoxic drugs to non-immune IgG have been shown to produce a greater inhibition of tumour cell growth than free drug alone (Hurwitz et al., 1979; Johnson et al., 1981; Kulkarni et al., 1985). In the experiments reported in this study, both methotrexate–antibody conjugates were significantly less effective than either methotrexate alone or antibody–methotrexate conjugates in reducing the cell number (Figure 2) and causing in the release of 31Cr from cells in vitro (Figure 4). Furthermore, these non-immune conjugates did not significantly reduce tumour growth in vivo (Figure 5). Analysis of cell size distribution after incubation of the PC3 cells with the various drugs and conjugates demonstrated that the non-immune methotrexate conjugates failed to display the same distribution profile observed with free drug or antibody–methotrexate conjugates (Figure 3). Assuming that both immunoglobulin conjugates are broken within the cell in the same manner, the results would suggest that the non-immune conjugates are unable to bind to or enter the cell to the same extent. Such non-immune immunoglobulin conjugates appear to rely upon non-specific uptake by the cell and discrepancies in data from different laboratories may be due to the amount of non-specific binding regions on the cell surface or the pinocytic activity of the cell line under study. An altered clearance rate may also contribute to differences found in vivo.

Specific anti-body–methotrexate conjugates have been reported to enter the cell as a consequence of antibody binding and subsequent endocytosis (Smyth et al., 1987) rather than via the active carrier-mediated transport system used for methotrexate or by the non-specific uptake of non-immune conjugates. Assessments of the cytotoxic effectiveness of the antibody–methotrexate conjugates in vitro, using measurements of both 31Cr release from pre-loaded cells (Figure 4) and cell population parameters (Figure 2) demonstrated that these conjugates were not significantly different in action from an equimolar amount of free drug, even though the dihydrofolate reductase inhibitory activity of the conjugates was reduced. It would appear therefore that the membrane antibody–methotrexate conjugates are efficiently incorporated into the cells after interaction of the antibody with the cell surface and that the methotrexate from the conjugates is sufficient to cause inhibition of dihydrofolate reductase.

When a drug and tumour specific antibodies conjugates have been administered in combination, but not conjugated, several workers have observed a more potent cytotoxic action than free drug given alone (Kulkarni et al., 1981; Lee & Hwang, 1979; Tai et al., 1979) and in some cases an enhanced effect has been observed over the conjugate (Hurwitz et al., 1978). No such synergistic effect between drug and antibody could be demonstrated in our study and may be explained by the high affinity of the polyclonal antiseraum utilised for the cell membrane. As a carrier mediated active transport system has been reported for methotrexate (Goldman et al., 1968) it is possible that enhanced binding of the antibody to the cell membrane in our study may block this carrier system, thus hampering drug transport. High affinity membrane binding might also explain the cytotoxicity of the antibody when given alone, both in vitro (Figures 2 and 4) and in vivo (Figure 5). Antibody cytotoxicity is well documented (Yang & Vas, 1971) and antibodies against tumour associated cell surface antigens have been demonstrated to produce membrane mediated cell changes either directly (Yang & Vas, 1970) or through complement and other intermediaries both in vitro and in vivo (Ghose et al., 1977).

When anti-membrane antibody–methotrexate conjugates were given to nude mice bearing PC3 tumours, a highly significant reduction in tumour growth was observed when compared to all other treatment groups (Figure 5). The concentration of 3H-methotrexate in the tumours of animals given these conjugates were significantly higher than controls (P<0.01 in all cases, Table II). Animals treated with both specific anti-membrane antibody and non-immune antibody conjugates displayed significantly higher blood levels of 3H-methotrexate than animals receiving drug in its free form thus indicating a slower clearance rate. Few studies have involved measurement of the in vivo distribution of cytotoxic drugs linked to antibodies. Deguchi et al. (1986) reported an increase of 3H in the liver and kidney of animals given a conjugate of methotrexate linked to a monoclonal antibody against prostatic acid phosphatase (PAP). As PAP is a secretory product of prostate cells it is conceivable that its release into the blood stream is sufficient to elicit conformational changes of antibody complexes, which would then be taken up by the reticuloendothelial system. This may explain the elevated levels of radioactivity labelled conjugate within the liver and kidneys of their animals in comparison to controls. Selection of an antigen against prostatic cell membranes for conjugation with a cytotoxic drug, in preference to one against
secretory products of the cells, should reduce the possibility of antibody complexes forming within the blood. In our study treatment with the anti-membrane antibody–methotrexate conjugates significantly increased drug accumulation at the tumour site without increasing the level of drug in the other body tissues, suggesting that uptake by the reticuloendothelial system was not pronounced.

Although it would appear that the anti-membrane antibody–methotrexate conjugates described in this study provided an efficient means for specificity targeting the drug in vivo, other anti-membrane antibody conjugates do not appear to produce such significant tumour reduction (Kanellos et al., 1987; Smyth et al., 1986). As the antibody preparation employed in this study consisted of a purified rabbit polyclonal antiserum to prostate membranes, its effectiveness in vivo may probably relate to the existence of numerous antibodies to various cell surface antigens. A cocktail of three monoclonal antibodies against the surface of T lymphocytes, linked to the toxin ricin, was found to be more effective in inhibiting T cell proliferation than any single monoclonal antibody (Vallers et al., 1983). The approach of using a battery of antibodies against various antigenic determinants on the cell membrane may therefore provide a more efficient carrier system for drug targeting.

The polyclonal antibody used in this study was, as stated, not specific for its prostate cell target and we are fully aware of the need for a battery of anti-membrane monoclonals preferential to human prostatic epithelial cells for future targeting and clinical studies.

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