Targeted photodestruction of human colon cancer cells using charged 17.1A chlorin<sub>e6</sub> immunoconjugates

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Summary

The goal of this study was to develop a strategy for the selective destruction of colorectal cancer cells. Towards this end, photodynamic therapy (PDT) is an experimental approach for cancer treatment in which the local or systemic delivery of a photosensitizer (PS) is followed by tissue illumination with light of an appropriate wavelength (usually red light delivered by a laser) (Hasan and Parrish, 1996; Dougherty et al, 1998). Conventional PS may have some selective accumulation in tumours (Henderson and Dougherty, 1992; Hamblin and Newman, 1994), but increased tumour targeting may be obtained by the use of macromolecular carriers which form complexes or covalent conjugates with PS (Hasan, 1992). The targeting capability of these carriers may rely on physical properties of the carrier (liposomes and microspheres (Speiser, 1991)), molecular properties (molecular weight and charge (Kornkuth et al, 1989)) or the specific recognition of molecules associated with tumours (monoclonal antibodies (mAbs), lipopolysaccharides and growth factor receptor ligands). They have been used to deliver cytotoxic drugs (Johnson et al, 1995), radioisotopes (Buchsbaum et al, 1993a), protein toxins (Houston, 1993) and PS to tumours, but the latter has the distinct advantage of not being toxic until illuminated with activating light, thus reducing toxicity due to non-specific uptake of the conjugate which can be a problem with conjugates formed from toxins and radioisotopes. Conjugates between mAbs and PS have shown promise both in vitro (Pogrebniak et al, 1993; Vrouenraets et al, 1999) and in experimental animal models of cancer (Mew et al, 1983; Goff et al, 1996), but have received only minimal clinical testing (Schmidt et al, 1992). The charge borne by the immunoconjugate may markedly influence the pharmacokinetics and biodistribution (Slinkin et al, 1993), and manipulation of the overall charge may increase the therapeutic ratio.

Colorectal cancer is in need of novel and effective therapies, and certain aspects of it may be appropriate for treatment by PDT. The ability to selectively target colorectal cancer cells may have applications to the tumour bed after surgical resection of the primary tumour (Barr et al, 1990), to disseminated intraperitoneal carcinomatosis (Veenhuizen et al, 1996), and to the liver metastases (Van Hillegersberg et al, 1992a) which are a frequent cause of death (Benotti and Steele, 1992). Because of the sensitive tissues in the peritoneal cavity (Veenhuizen et al, 1997), and the high accumulation of conventional PS in normal liver (Van Hillegersberg et al, 1992b), it is attractive to explore the use of photodynamic therapy (PDT) as radioimmunoconjugates to target radioisotopes to residual tumour (Buchsbaum et al, 1993b). As part of our on-going effort to optimize intraperitoneal photodynamic therapy (PDT) (Goff et al, 1991, 1994, 1996) we have recently described the preparation of charged PICs between the F(ab')<sub>2</sub> fragment of anti-ovarian cancer Mab OC125 and the PS chlorin<sub>e6</sub> (c<sub>e6</sub>) (Hamblin et al, 1996), and have investigated their biodistribution in vivo (Duska et al, 1997). Toward the long-term goal of applying PIT to intraperitoneal and hepatic metastases of colorectal cancer, we report here our initial studies on the preparation of charged PICs.
17.1A photoimmunoconjugates

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with the intact mAb 17.1A and their in vitro interactions with a human colorectal cancer cell line recognized by the mAb (HT29) and a cell line which showed no binding to the mAb (OVCAR-5).

MATERIALS AND METHODS

Cell line and monoclonal antibody

Two tumour cells lines were employed. HT29 cells derived from a human colorectal adenocarcinoma were a generous gift from Dr K Tanabe (Massachusetts General Hospital, Boston, MA, USA). NIH:OVCAR-5 human ovarian cancer cells were purchased from Dr T Hamilton (Fox Chase Cancer Center, Philadelphia, PA, USA). The growth medium for HT29 was DMEM/F12 (50/50 mixture) and that for OVCAR-5 was RPMI-1640. Both media contained 15 mM HEPES and L-glutamine and were supplemented with 10% heat-inactivated fetal calf serum (FCS) (Whittaker Bioproduct, Walkersville, MD, USA), 100 U ml
\(^{-1}\) penicillin and 100 \(\mu\)g ml
\(^{-1}\) streptomycin, and maintained in an incubator at 37\(^{\circ}\)C in an atmosphere of 5% carbon dioxide. 17.1A murine mAb was a kind gift from Centocor (Malvern, PA, USA). Rabbit IgG and mouse IgG was obtained from Sigma (St Louis, MO, USA).

Conjugation procedure

The procedure has been described in detail elsewhere (Hamblin et al, 1996). Briefly poly-L-lysine (average MW 25 000) was treated in DMSO with the N-hydroxysuccinimide ester of \(\text{c}_{\text{e6}}\) to give \(\text{pl-c}_{\text{e6}}\). This was then reacted with pyridyldithiopropionic acid N-hydroxysuccinimide ester (SPDP) to form the functionalized derivative \(\text{pl-c}_{\text{e6}}\)-SPDP. This was then split into two parts and one part was treated with an excess of succinic anhydride to give the negatively charged functionalized \(\text{pl-c}_{\text{e6}}\)-succ-SPDP. 17.1A mAb was reduced for 1 h with 5 mM mercaptoethylamine hydrochloride, dialysed (1 mM EDTA) and then reacted with either \(\text{pl-c}_{\text{e6}}\)-SPDP or \(\text{pl-c}_{\text{e6}}\)-succ-SPDP to form the cationic and anionic PICs respectively. The procedure was repeated with rabbit IgG. The conjugates were purified by chromatography on Sephadex G200 columns, and characterized by absorption and fluorescence spectrophotometry, and polyacrylamide gel electrophoresis. The structures of the PICs are shown in Figure 1.

Enzyme-linked immunosorbent assay

Cells were grown to 100% confluence in 96-well plates for 24 h with medium containing FCS, than washed 3 times with PBS and fixed with 0.25% glutaraldehyde. After 1 h cells were washed with PBS and plates blocked with PBS containing 5% FCS for 1 h, then 0.1 ml PBS containing the appropriate dilution of PIC or mAb was added to each well. After 2 h incubation at room temperature in the dark, wells were washed 3 times with PBS containing 0.05% Tween 20 and had added 0.1 ml of horseradish peroxidase conjugated-F(ab’)
\(_{2}\) fragment rabbit anti mouse IgG, IgA, IgM (h+l) (Zymed Laboratories, South San Francisco, CA, USA) diluted 1:200 in PBS containing 0.5 mg ml
\(^{-1}\) bovine serum albumin (BSA) and 0.05% Tween-20. Cells were incubated for 2 h at room temperature in the dark then washed 3 times with PBS containing 0.05% Tween-20 and added 0.1 ml of 0.4 mg ml
\(^{-1}\) of freshly prepared o-phenylenediamine dissolved in 0.05 M sodium citrate, 0.15 M sodium phosphate, pH 6, containing 32% vol/vol of 30% hydrogen peroxide and incubated for a further hour at room temperature in the dark. Absorbance was read at 492 nm with an enzyme-linked immunosorbent assay (ELISA) reader (500 EIA, Bio-Rad Laboratories, Hercules, CA, USA).

Figure 1  Structures of the PICs. 17.1A-pl-ce6 has primary amino groups which give it a polycationic charge, while 17.1A-pl-ce6-succ has carboxylic groups which give it polyanionic charge.
Two-colour direct/indirect immunofluorescence

Approximately 3 × 10^5 cells were plated in 35-mm tissue culture dishes for 24 h with 2 ml of medium, containing coverslips that were previously washed with 95% ethanol and flamed. After 24 h of incubation in 2 ml of medium, cells were washed with PBS (2×) and fixed with 2% formaldehyde at room temperature for 5 min. Cells were washed with PBS/1% BSA (3×) and incubated for 1 h at room temperature with the mAb or conjugate, the amount of mAb in both cases was 2 μg ml⁻¹. After this time they were gently washed with PBS/1% BSA (3×) and incubated for 1 h with or without fluorescein isothiocyanate-conjugated goat antimouse IgG (FITC-GAM, Sigma) diluted 1:128 with PBS/1% BSA. Cells were washed with PBS/1% BSA (3×) and mounted on glass microscope slides using GelMount (Biomeda Corp., Foster City, CA, USA). An epi-illumination microscope (Model WL, Zeiss, Oberkochen, Germany) equipped with a CCD camera (TM 745, Pulnix, Sunnyvale, CA, USA), video monitor and computer was used to capture digital images. Two different combinations of filters were used. The first set used an excitation bandpass filter at 450–490 nm and emission 514–530 nm bandpass filter designed for visualizing fluorescein fluorescence. The second set used an excitation bandpass filter at 402–447 nm and emission 580 nm longpass filter designed for visualizing cₐ fluorescence. The images obtained with the cₐ filter combination viewing cells incubated with unconjugated 17.1A and FITC-GAM were totally negative, while those obtained with cells incubated with a cₐ conjugate without FITC-GAM and the fluorescein filter set gave a faint image.

Cellular uptake

Twelve-well plates containing 90% confluent cells had 1 ml medium containing 10% FCS and conjugates added. After the completion of the incubation time cells were washed with PBS (3×) and incubated with trypsin (0.25%)/EDTA (0.02%) (1 ml) at 37°C for 20 min. The suspension was centrifuged and the resulting cell pellet dissolved for 48 h in 1.5 ml of 0.1 M sodium hydroxide (NaOH); 1% sodium dodecyl sulphate (SDS) to give a homogeneous solution. Fluorescence was measured with a fluorometer (Fluorolog 2, Spex Industries, Edison, NJ, USA) (excitation at 400 nm, emission scanned from 580 to 720 nm). The trypsin supernatant was checked for the presence of fluorescence, which was always less than 10% of the cell extract. The cell digest was then assayed for the amount of cell protein by a modified Lowry procedure (Larson et al., 1986). Quantitation of cₐ concentration in the cell extracts was obtained by comparing the fluorescence of standard solutions of the same conjugate of known concentrations in 0.1 M NaOH 1% SDS. The fluorescence of the cell extracts was always within the linear part of the standard curve. Uptake experiments at 4°C were carried out by adding pre-cooled medium containing conjugates to the wells in 12-well plates, which were then wrapped in aluminium foil and incubated in crushed ice for 6 h.

Competition

Studies were conducted in which the cells were preincubated with a saturating concentration of unmodified 17.1A, to see if the binding of the PICs was blocked. Since 1 ml of 1 μM cₐ equivalent PIC contains approximately 7.5 μg mAb, a fivefold excess of competing 17.1A or rabbit IgG (37.5 μg) was employed. The conjugate was added at a concentration of 1 μM cₐ equivalent and incubated for 2 h. The uncompeteted uptake was compared to the uptake found by the addition of 17.1A or rabbit IgG (37.5 μg ml⁻¹) for 1 h, followed by washing and addition of PIC (1 μM cₐ equivalent) for 1 h.

Phototoxicity

Phototoxicity was measured by a colony forming assay. In P35 dishes cells were grown to 70% confluence in 2 ml medium containing 10% FBS. Cells were washed with PBS and 1 μM cₐ equivalent of PICs and P5 in serum containing medium was added. After 6 h incubation at 37°C, the medium was removed, cells were washed twice with PBS, and 2 ml of fresh serum containing medium was added. The dishes were illuminated with 666 nm light for cₐ conjugates or 654 nm light for cₐ at a power density of 48 mW cm⁻², measured with a power meter (Model 210, Coherent Inc., Palo Alto, CA, USA). An argon-pumped dye laser (Innova 100 and 599 Dye, Coherent) was focused through a ×10 microscope lens onto the end of a 1 mm diameter optical fibre which delivered light through an inverted ×4 microscope lens (No 774317, Olympus, Tokyo, Japan) to give a 35 nm diameter spot for irradiation. Controls were as follows: no conjugate and no light remaining in incubator throughout, no conjugate and no light but plates wrapped in foil for duration of the irradiation time out of the incubator, conjugate given and no light wrapped in foil, no conjugate and irradiated. At the completion of irradiation the cells were given fresh medium containing FCS and returned to the incubator for 24 h. The cells were then washed with medium and any detached cells aspirated off. The remaining cells were detached with trypsin/EDTA (0.5 ml) and an aliquot counted for viable cells using the trypan blue exclusion assay and a haemocytometer. The cell suspension was diluted with medium and plated in P60 dishes containing 4 ml of medium at densities of 50, 100, 150 and 200 cells per plate. When the colonies had formed (9 days later) the cells were fixed with 0.2% formalin (vol/vol) in MeOH and stained with crystal violet. The number of colonies, which contained 50 or more cells, was then counted. The survival fraction was calculated by multiplying the fraction of viable cells at the counting stage compared to controls (sensitizer, dark, out of incubator), together with the fraction of colonies formed by treated cells compared to controls.

RESULTS

Absorbance spectroscopy

The absorption spectra of the cationic and anionic PICs together with unconjugated cₐ are shown in Figure 2. The conjugates have a distinct absorbance at 280 nm due to the protein in the IgG, the intensity of the Soret band is somewhat reduced, and the long wavelength Q band is red shifted to 666 nm as opposed to 654 nm. It can be calculated assuming extinction coefficients of 1.5 × 10⁵ lmol⁻¹ cm⁻¹ for cₐ at the Soret band and 2.4 × 10⁴ lmol⁻¹ cm⁻¹ for IgG at 280 nm, that the anionic 17.1A-pl-cₐ-succ had 8–9 cₐ molecules attached to each IgG (two polylysine chains per IgG), while the cationic 17.1A-pl-cₐ had a lower loading of 4–5 cₐ molecules per IgG (one polylysine chain per IgG).
The uptake of c-e6 was strikingly similar (Figure 4 C, D).

ELISA

Results of ELISA binding assays using fixed cells and assaying binding of 17.1A mAb, 17.1A-pl-c-e6, 17.1A-pl-c-e6-succ and non-specific mouse IgG are shown in Figure 3. The unmodified mAb exhibited a typical binding curve with reduction in binding over the concentration range 10–0.1 μM c-e6. The anionic 17.1A-pl-c-e6-succ showed a somewhat reduced affinity, while the cationic 17.1A-pl-c-e6 gave a slightly higher affinity compared to native 17.1A. As expected, non-specific mouse IgG showed no binding.

Two-colour direct/indirect immunofluorescence microscopy

Two-colour direct/indirect immunofluorescence microscopy was carried out by treating fixed cells with a PIC, which binds to membrane antigens, and then adding a FITC-conjugated goat anti-mouse second antibody that recognizes the murine IgG of the first mAb. Fluorescent images were obtained with a dual filter system capable of isolating green fluorescence (510 nm) emitted by FITC, and red emission filter and FITC alone gave negative images (data not shown). Images with similar appearance were obtained from 17.1A-pl-c-e6-bound to fixed HT29 cells (Figure 4 A, B). In a similar fashion images obtained from 17.1A-pl-c-e6-succ with both c-e6 and FITC emissions were strikingly similar (Figure 4 C, D).

Cellular binding and uptake

The uptake of c-e6 per mg cell protein from the 17.1A-pl-c-e6 and 17.1A-pl-c-e6-succ in a range of concentrations of PICs (measured as c-e6 equivalent in the medium) is shown in Figure 5. The uptake of c-e6 equivalent from the cationic PIC is up to 4 times higher than that obtained from the anionic PIC after 6 h incubation, and both show linear relationships with increasing concentration. This linear increase is consistent with internalization of the conjugates. A concentration of 3 μM c-e6 equivalent is equivalent to approximately 300–500 nm 17.1A, which is much higher than the binding constant of the typical mAb. Thus if there was no internalization some saturation of uptake should have been observed. The cellular uptake of c-e6 obtained with the 17.1A PICs were compared to that obtained with non-specific rabbit IgG PICs, and free c-e6 (Table 1) under the same incubation conditions (1 μM c-e6 equivalent, 6 h incubation at 37°C in serum containing medium). Free c-e6 gave the lowest uptake followed by rabbit IgG-pl-c-e6-succ and 17.1A-pl-c-e6-succ. The cationic rabbit IgG-pl-c-e6 gave roughly twice the uptake of its anionic counterpart. The 17.1A conjugates gave the highest uptakes among conjugates bearing the same charge, and the cationic 17.1A-pl-c-e6 gave 3 times the uptake of the anionic 17.1A-pl-c-e6-succ.

Table 1 Comparison of uptake after incubation at 37°C and 4°C

| Conjugate                  | Uptake at 37°C | Uptake at 4°C | Ratio  |
|----------------------------|----------------|---------------|--------|
| 17.1A-pl-c-e6              | 2.4 ± 0.17     | 1.4 ± 0.053   | 1.71   |
| 17.1A-pl-c-e6-succ         | 0.73 ± 0.037   | 0.23 ± 0.006  | 3.17   |
| Rabbit IgG-pl-c-e6         | 1.1 ± 0.064    | 0.65 ± 0.035  | 1.53   |
| Rabbit IgG-pl-c-e6-succ    | 0.32 ± 0.022   | 0.071 ± 0.003 | 4.5    |
| c-e6                       | 0.063 ± 0.02   | 0.025 ± 0.002 | 2.47   |

Uptake was determined after incubation with 1 μM c-e6 equivalent concentration for 6 h. Concentrations are expressed as nmol c-e6 equivalent/mg cell protein. Each value is the mean of values from two separate experiments each containing three wells ± s.e.m.
In order to gain additional information on the degree to which the PIC which is bound to the cells is internalized, the uptake after incubation at 37°C which is bound to the cells is internalized, the uptake after incubation at 37°C was compared to that found after incubation at 4°C. The concentration was 1 μM cα equivalent and the incubation time was 6 h. The results are given in Table 1. It can be seen that the cationic species have about 65% of the uptake remaining at 4°C, while the anionic conjugates have only 30% remaining. However, the absolute amount of the conjugate internalized (difference between uptakes at 37°C and 4°C) is greater for the cationic species than the anionic species. Note that the uptake of free cα is also significantly greater at 37°C than 4°C.

### Comparison of uptake with non-target cell line

OVCAR-5 human ovarian carcinoma cell line was used as a non-target control compared to HT29 cells. Indirect immunofluorescence using 17.1A mAb and FITC-goat anti-mouse IgG on fixed OVCAR-5 cells gave a negative fluorescence image (data not shown) indicating that OVCAR-5 cells expressed very low levels of EpCAM, but the two-colour direct/indirect immunofluorescence procedure with both the cationic and anionic 17.1A PICs gave weak images for both cα and FITC fluorescence (data not shown) indicating a small amount of non-specific binding of the PICs to OVCAR-5 cells.

The uptake of cα by live OVCAR-5 and HT29 cells from the cationic and anionic 17.1A PICs and from cationic and anionic non-specific rabbit IgG PICs was compared and the results are shown in Table 2. Almost no difference was found between target and non-target cell line using either the cationic and anionic non-specific rabbit IgG PICs as expected, whereas the uptake from both the cationic and anionic 17.1A PIC showed twofold selectivity for the target HT29 cells compared to the non-target OVCAR-5 cells.

### Competition of uptake by unmodified proteins

In order to further understand to what extent the observed uptake was due to antigen binding and what extent to non-specific charge interaction, uptake of cα from cationic and anionic 17.1A PICs by HT29 cells was compared with and without saturation of the antigens by preincubation with unmodified 17.1A (Figure 6). The uptake of cα from the cationic 17.1A-pl-cα was reduced to 30% of the control level by preincubation with 17.1A, while that of the anionic 17.1A-pl-cα-succ was reduced to 10% of the control level. That this reduction was due to blocking of the antigenic binding sites by 17.1A is confirmed by the observation that preincubation with rabbit IgG had no effect on subsequent binding of 17.1A PICs. As expected the uptake of the cationic PIC had a greater contribution from non-antigen mediated charge interactions than the anionic PIC.

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**Table 2** Selectivity of conjugates for HT29 cells compared to OVCAR-5 cells

| Conjugate          | HT29 cells | OVCAR-5 cells | Selectivity for HT29 |
|--------------------|------------|---------------|----------------------|
| 17.1A-pl-cα        | 2.45 ± 0.17| 1.3 ± 0.06    | 1.9 : 1 (P < 0.01)   |
| 17.1A-pl-cα-succ   | 0.74 ± 0.04| 0.34 ± 0.02   | 2.2 : 1 (P < 0.01)   |
| Rabbit IgG-pl-cα   | 1.0 ± 0.06 | 0.9 ± 0.01    | 1.1 : 1 (n.s.)       |
| Rabbit IgG-pl-cα-succ | 0.29 ± 0.01| 0.34 ± 0.02   | 0.9 : 1 (n.s.)       |

Uptake was determined after incubation at 1 μM cα equivalent concentration for 6 h at 37°C using non-specific rabbit IgG anionic and cationic PICs and 17.1A cationic and anionic PICs. Cellular fluorescence was measured after extraction into 0.1 M NaOH/1% SDS and expressed in nmol cα per mg cell protein. Each value is the mean of values from two separate experiments each containing three wells ± s.e.m. Significance was assessed by two-tailed unpaired Student’s t-test.

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**Effect of incubation temperature on uptake**

In order to gain additional information on the degree to which the PIC which is bound to the cells is internalized, the uptake after incubation at 37°C was compared to that found after incubation at 4°C. The concentration was 1 μM cα equivalent and the incubation time was 6 h. The results are given in Table 1. It can be seen that the cationic species have about 65% of the uptake remaining at 4°C, while the anionic conjugates have only 30% remaining. However, the absolute amount of the conjugate internalized (difference between uptakes at 37°C and 4°C) is greater for the cationic species than the anionic species. Note that the uptake of free cα is also significantly greater at 37°C than 4°C.

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**Figure 5** Concentration dependence of cellular uptake of cα from cationic 17.1A-pl-cα and anionic 17.1A-pl-cα-succ. Cells were incubated for 6 h at 37°C for each PIC and cellular fluorescence was measured after extraction into 0.1 M NaOH/1% SDS and expressed in nmol cα per mg cell protein. Each point is the mean of values from two separate experiments each containing three wells and bars are s.e.m.
Phototoxicity

Cells were illuminated after 6 h incubation with 1 μM cₐₐ equivalent of all the compounds. The phototoxicity was measured by using a colony forming assay that combines short-term and long-term damage to the cells. The 17.1A cationic and anionic PICs (with HT29 target cells) showed fluence-dependent phototoxicity with greater than 99.9% killing after 10 J cm⁻² 666 nm red light (Figure 7A). To compare the phototoxicities of different pairings of cells and PICs we carried out experiments at a constant fluence of 3 J cm⁻² and the results are shown in Figure 7B. Under these conditions 10% of HT29 cells survived after treatment with 17.1A-pl-cₐₐ and 27% survived after PIT with 17.1A-pl-cₐₐ-succ. Comparing these results with the killing produced by the non-specific rabbit IgG PICs of the same overall charges the difference is seen to be significant; four times the number of surviving of cells with the rabbit IgG cationic PIC compared to the 17.1A cationic PIC, and three times more cells surviving after PIT with the anionic rabbit IgG PIC compared to the 17.1A anionic PIC. The free cₐₐ had a very low phototoxicity compared to both the 17.1A PICs. Cell type selectivity of the 17.1A PICs was shown by using OVCAR-5 cells as a non target cell line; the survival was 3–4 times higher than the target HT29 cell line gave. Interestingly OVCAR-5 cells were killed by PIT with the 17.1A cationic and anionic conjugates to almost exactly the same extent as the non-specific rabbit IgG PICs killed HT29 cells as shown in Figure 7B.

DISCUSSION

Additional therapies to target colorectal cancer metastases in the liver are urgently needed (Van Cutsem, 1996). Existing treatments include surgery (Steele and Ravikumar, 1989), chemoembolization (Sanz-Altamira et al, 1997), regional chemotherapy (McMurrick and Nelson, 1997) and cryotherapy (Yeh et al, 1997). Among investigative treatments some workers are exploring the role of PDT (Van Hillegersberg et al, 1992a). A drawback of PDT in the liver is the high accumulation of most free PS in normal liver (Van Hillegersberg et al, 1992b). The preparation of PICs that could target tumour-associated antigens on the colorectal cancer metastasis as opposed to normal liver tissue may increase the specificity of PDT for liver metastases. In addition the disseminated intraperitoneal spread of colorectal cancer may also be treated by PDT (Veenhuizen et al, 1996), but again additional selectivity for the tumour is necessary to prevent unwanted...
damage to intestines and other intraperitoneal organs (Veenhuizen et al, 1997). There are many reports in the literature which confirm the selectivity of the 17.1A murine mAb toward gastrointestinal tumours and in particular to colorectal cancer (Martin et al, 1986; Pierce et al, 1990; Buchsbaum et al, 1993b; Meredith et al, 1995). In order for PIT to be effective each PIC molecule must deliver as much PS as possible to the tumour, without sacrificing unduly the specificity and affinity of the PIC for its antigen. One way of accomplishing this is to use a polymeric linker to attach several PS molecules in a site-specific manner to the mAb, and this linker may bear positive, negative or neutral charge (Hamblin et al, 1997). There are many reports in the literature which confirm the selectivity of the 17.1A murine mAb toward gastrointestinal tumours and in particular to colorectal cancer (Martin et al, 1986; Pierce et al, 1990; Buchsbaum et al, 1993b; Meredith et al, 1995). Variation in the overall charge may affect the binding of the PIC to its target antigen, its intracellular location and its phototoxicity. A previous publication reported the effect of charge on the selectivity, uptake and phototoxicity of OC125 F(ab')2 PICs constructed in a similar fashion to the present 17.1A PICs (Hamblin et al, 1996). Alterations in the charge borne by mAb conjugates can also lead to wide variations in the biodistribution (Slinkin et al, 1993; Duska et al, 1997). The object of this study was to investigate the binding (specificity and affinity) of cationic and anionic 17.1A PICs to HT29 target human colorectal cancer cells, and their consequent phototoxicity. The results from the ELISA and two-colour direct/indirect immunofluorescence studies on fixed cell, showed that the capacity of the two differently charged PICs to recognize the membrane antigen expressed on HT29 cells, compared reasonably with that of unmodified 17.1A. Since these cells were fixed, the subsequent endocytosis of bound PICs was not an issue. Although the affinity of the anionic PIC was somewhat reduced as measured by the ELISA it was still relatively high. The fluorescence microscopy showed that with fixed cells the 17.1A and the cationic PIC delivered by the PIC had very similar localizations making it likely that the great majority of cationic mAb recognition at least doubles the uptake compared to other conjugates of the same charge. The data comparing uptake at 37°C and 4°C show that the cationic conjugates bind much better than the anionic ones to the membrane at 4°C, and the additional internalized uptake at 37°C is only another 50%; while for the anionic conjugates the additional uptake at 37°C was 200–300% of that at 4°C. This suggests that a higher proportion of the anionic 17.1A PIC was internalized than for the cationic 17.1A PIC, although the overall uptake was less. Uptake of cationic PIC from both the cationic and anionic 17.1A PICs was also approximately twofold higher by target HT29 colorectal cancer cells than by non-target OVCAR-5 ovarian cancer cells, while the non-specific rabbit IgG PICs showed no difference in cationic uptake between cell lines. However, since the EpCAM antigen is a common antigen over-expressed on cancer cells it is possible that OVCAR-5 cells also expressed this antigen, although the indirect immunofluorescence was negative. Additional evidence of the retention of antigen recognition in the PICs was provided by the experiments in which the antigen was saturated by preincubation with unmodified mAb. It should be noted that it is difficult to get any selectivity between two epithelial human cancer cell lines using unconjugated PS in vitro.

One of the critical issues to be considered in immunoconjugate therapy is the penetration of the immunoconjugate into the tumour tissue (Jain, 1990). It has been shown (Saga et al, 1995) that high affinity mAbs penetrate less well than those of lower affinity, and therefore a conjugation process which reduces the affinity of the mAb for target antigen expressing cells, may be quite acceptable. The question then arises, to what extent do the values for cationic uptake delivered by the various conjugates correlate with the phototoxicity? Under the conditions where the uptake of cationic PICs from the cationic conjugates relative to unconjugated mAb was 38:16:1 for 17.1A and rabbit IgG, the relative phototoxicity (1/survival fraction) compared to unconjugated cationic mAb with 3 J cm−2 666 nm light was 6.5:1.4:1. Similarly for the anionic series where the relative uptakes were 11.5:5.5:5:1, the relative phototoxieties were 2.4:0.8:1:1. To determine which of these conjugates is inherently the most effective photosensitizer, we can divide the relative phototoxicity by the uptake in nmol cationic mAb equivalent per mg cell protein and the resulting numbers are shown in Table 2. Free cationic mAb would appear to have more phototoxic potential than any of the conjugates per unit cationic mAb uptake, but since the uptake is exceptionally low this is not of much relevance. This is in agreement with other reports (Bachor et al, 1991) that free mAb has such low cellular uptake that it is difficult to get any phototoxicity in vitro. The remaining conjugates have roughly similar values but the values obtained when mAb conjugates interact with target cells are consistently higher than non-matching pairs. These results can be compared with those obtained (Hamblin et al, 1996) with polycationic and polyanionic OC125F(ab')2 conjugates prepared in a similar manner and directed towards target OVCAR-5 cells. In this previous study the charge effect was found to be of larger magnitude but in the same direction as the present study, with the polycationic PIC having 6 times the uptake and 10 times the phototoxicity of the polyanionic PIC. This difference in magnitude between charge effects may be attributed to differences in the magnitude of the negative charge expressed on the outside of the cells (Bischoff et al, 1981), or to differences in the extent to which polycations stimulate endocytosis between cell lines (Duncan et al, 1979). It is accepted that imparting cationic charge to a protein

### Table 3

| Conjugate | HT29 cells | OVCAR-5 cells |
|-----------|------------|--------------|
| 17.1A-pl-cα | 4.17 ± 0.56 | 2.17 ± 0.06 |
| 17.1A-pl-cα-succ | 5.07 ± 0.11 | 4.79 ± 0.07 |
| Rabbit IgG-pl-cα | 2.17 ± 0.07 | |
| Rabbit IgG-pl-cα-succ | 3.59 ± 0.07 | |
| cα | 24.5 ± 1.31 | |

The phototoxicity (1/survival fraction) after incubation at 37°C for 6 h at 1 μM cα equivalent concentration and 3 J cm−2 666 or 654 nm light, was divided by the cellular uptake in nmol cα equivalent per mg cell protein. Errors are the s.e.m. of the ratio of the means calculated in quadrature.
(Shen and Ryser, 1978), receptor ligand (Cotten et al, 1990), mAb (Pardridge et al, 1994) or immunoconjugate (Hamblin et al, 1996) increases the absolute uptake and the degree to which it is internalized in vitro. It is thought that the mechanism of increased uptake is due to non-clathrin-coated pit-mediated endocytosis leading to endosomal processing (Hansen et al, 1993), and to accumulation in lysosomes where proteolysis may take place. In vivo, however, polycationic moieties have high and fast uptake in the liver and kidney when administered i.v. (Clegg et al, 1990), which suggest that polycationic PICs would be better suited to intracavitary administration (Hamblin et al, 1996). This hypothesis was confirmed with a series of biodistribution experiments conducted with OC125F(ab\')2 PICS prepared in a similar manner to the present PICS and injected i.p. in nude mice bearing i.p. OVCAR-5 tumours (Duska et al, 1997). However, in order to effectively target colorectal tumour cells growing in the liver where the PIC must be administered i.v., it is likely that a polycationic PIC will outperform a polycationic species. This hypothesis will be tested in a forthcoming report.

In conclusion we have demonstrated advantages of conjugating \( c_{\text{e}} \) to mAb 17.1A by a site-specific synthetic route. The immunoreactivity is preserved, the PICS show selectivity to target colorectal cancer cells over non-target ovarian cancer cells, and the absolute uptake by tumour cells is very much higher for both charges than that given by the free \( c_{\text{e}} \). In vitro there is little difference in the amount of killing per molecule of \( c_{\text{e}} \) delivered by polycationic and polyanionic PICS, thus leaving the choice of charge borne by the PIC for in vivo PIT to be made on the basis of biodistribution and pharmacokinetic data. While these initial data on the preferential photodestruction of target cells are encouraging, there remain many questions which will only be answered by in vivo experiments. Will the PICS be able to penetrate solid tumours after i.v. administration? In addition to binding to tumour cells, will the PICS be taken up by cells of the monocyte/macrophage lineage which are especially prevalent in liver? Will the pharmacokinetics and biodistribution of the PICS be suitable for effective photodestruction of tumours in vivo? These questions will be addressed in forthcoming reports.

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