Site-specific Prodrug Activation by Antibody-β-Lactamase Conjugates: Regression and Long-Term Growth Inhibition of Human Colon Carcinoma Xenograft Models


ABSTRACT

Antibody-directed catalysis (ADC) is a two-step method for the delivery of chemotherapeutic agents in which enzyme-antibody conjugate, prelocalized to antigen-bearing tumor cells, catalyzes the site-specific conversion of prodrug to drug. An ADC system consisting of F(ab')-β-lactamase conjugates and a cephalosporin derivative of the oncolytic agent 4-desacetylvinblastine-3-carboxhydrazide was investigated.

The ability of the system to mediate antitumor activity was compared with that of free drug given alone and with covalent drug-antibody conjugates in LS174T and T380 colon carcinoma xenografts in nude mice. Efficacy increased from moderate tumor growth inhibition by using free 4-desacetylvinblastine-3-carboxhydrazide to tumor regression and long-term stabilization with the ADC system. Labile covalent drug-antibody conjugates prepared from the same antibodies were less effective than ADC and required much higher antibody doses.

The antigens KSI/4, carcinoembryonic antigen, and tumor-associated glycoprotein-72, TAG-72, present on the model cell lines, were chosen to investigate the effect of differences in subcellular location and expression heterogeneity on the efficacy of ADC delivery. Response was equivalent with the three tumor antigens. Hence, heterogeneous expression and membrane shedding of carcinoembryonic antigen and TAG-72, did not diminish the suitability of these antigens as targets for ADC therapy. In contrast, drug-antibody conjugate efficacy was more sensitive to subcellular location and heterogeneity.

Thus, ADC is a highly effective form of immunochemotherapy in preclinical models, with applicability toward a variety of antigen targets.

INTRODUCTION

The fundamental rationale for targeted delivery systems is that efficacy can be maximized and toxicity minimized by biasing the biodistribution of the chemotherapeutic agent toward the disease site and away from sites of organ toxicity. To achieve advantage over more traditional forms of treatment, a targeted delivery system must also produce a higher concentration of drug at the tumor than can be produced by nontargeted administration of the same drug. Several recent reports (1–3) have described ADC systems which attempt to provide, simultaneously, a biodistribution bias favoring the tumor, and a mechanism for obtaining higher tumor concentrations of drug than could be obtained by direct administration of active agent. This is achieved by using an enzyme covalently conjugated to a tumor-selective antibody. Such a conjugate catalytically converts an independently administered prodrug into the active oncolytic agent, potentially reaching a concentration far greater than that of the conjugate itself.

The current study was undertaken to determine whether ADC is capable of overcoming two of the most important theoretical limitations of targeted delivery with drug-antibody covalent conjugates: the requirement for large antibody doses and the paucity of antigens that are both sufficiently specific for effective targeting and sufficiently concentrated at the tumor to mediate delivery of therapeutic concentrations of drug (4). To this end experiments were performed to determine: (a) the pharmacokinetic behavior of an enzyme-antibody conjugate; (b) the efficacy of the ADC system directed toward three different antigens in two different cell lines, relative to nontargeted drug treatment and to covalent drug-antibody conjugate treatment with the same drug; and (c) the minimum conjugate dose required to obtain maximal efficacy.

The primary goals in selecting the enzyme and substrate components of this ADC system were (a) that the enzyme and substrate be absent from mammalian tissues; (b) that the system offer potential for release of a variety of drugs with drug release mediated via several different functional groups; (c) that the drug released (at least in initial studies) not require further activation or metabolism; and (d) that the molecular weight of the conjugate be minimized in order to facilitate rapid clearance of the conjugate from circulation. The enzyme chosen to meet these criteria was the β-lactamase from the 265A strain of Enterobacter cloacae, and the prodrug was a cephalosporin sulfoxide derivative of 4-desacetylvinblastine-3-carboxhydrazide. Similar considerations have also led others to adopt P99 or other β-lactamas tos for site-specific prodrug activation (5–7).

The E. cloacae 265A β-lactamase, designated P99, was selected because of its ability to hydrolyze a large number of cephalosporins.3 The P99 enzyme has been well studied (8, 9). Its amino acid and gene sequences have been determined and the active site residues are known. Several of its properties were considered to be advantageous for ADC: it is of moderate molecular weight (40,000), is readily available via one-step affinity chromatography (10), and requires no cofactors. The enzyme catalyzes the hydrolysis of cephalosporins resulting in expulsion of substituents, if they are sufficiently good leaving groups (11, 12), from the cephalosporin 3'-position.

To facilitate rapid clearance from the circulation, conjugates were designed with a single F(ab')-binding arm. Use of this fragment for the antibody moiety allowed regiospecific conjugation through the hinge region sulfhydryls and facilitated purification by gel permeation chromatography (13). Hence, the studies reported here also address the question of whether single binding arm conjugates are capable of mediating this form of targeted delivery.

MATERIALS AND METHODS

Prodrug Synthesis. Synthetic methodology was developed to attach the acyl hydrazide functionality of the potent vinblastine derivative DAVLBHYD to the cephalosporin 3'-position via an azacarbamate linkage (Fig. 1) (14). The synthesis began with the commercially available cephalosporin, cephalothin, which was converted to the 3-(hydroxymethyl)-2-cephem derivative in two steps (15). The carboxylic acid function was protected as a benzhydryl ester, and the 3'-hydroxyl moiety was acylated with p-nitrophenyl chloroformate to form the carbonate. Oxidation of the sulfide, with concomitant olefin isomerization, provided the 1-β-sulfoxide-3-cephem.

3 T. Parr, private communication.
Selective nucleophilic displacement of the carbonate with DAVLBHYD (16, 17) formed the azacarbamate linkage, thus coupling the oncolytic agent to the cephalosporin. Acid-catalyzed hydrolysis of the ester provided the desired produg. Previous experiment (14) had shown that the sulfoxide form of the cephalosporin would be a better substrate with hydrolytic stability at least comparable to the unoxidized cephalosporin. The azacarbamate functional group is relatively resistant to uncatalyzed hydrolysis at physiological pH, but behaves as a leaving group after enzyme catalyzed hydrolysis of the ß-lactam.

Hydrolysis of the ß-lactam ring of the cephalosporin sulfoxide results in expulsion of the 3' substituent leading to generation of DAVLBHYD.

**Construction of F(ab')-ß-Lactamase Conjugates.** Antibody-enzyme conjugates were constructed from F(ab') fragments (M, ~50,000) for targeting. Fragments were obtained by pepsin digestion of antibodies directed to three human tumor-associated antigens: KS1/4, CEA, and TAG-72 [antibodies: P99 ß-lactamase was treated with sulfosuccinimidyl 4-(A'-maleimidomethyl)-cyclohexane-1-carboxylate (Pierce) to produce the desired size before dosing was initiated. Animals (10 in the saline control group; 5 in the treatment groups) were treated with specific (α-KS1/4-, α-CEA-, or α-TAG-72-ß-lactamase) or nonspecific conjugate (35 μg), or phosphate-buffered saline (10 mm NaPO₄, 150 mm NaCl, pH 7.4) on the days indicated in the figure. These injections were followed after 96 h by a single injection of prodru (4 mg/kg), drug (2.4 mg/kg), or phosphate-buffered saline. Tumor masses were calculated from tumor volume (assuming a tissue density of 1 g/cm³), which was determined by caliper measurements, using the formula mass = 1 g/cm³ × 0.5 × L × W², where L and W are the longest dimension and its perpendicular in cm, respectively. All injections were made in the tail vein.

T380 tumors (Figs. 4 and 5) were implanted on day 0 by s.c. inoculation of a 0.1 ml tumor slurry, prepared by suspending 1 g of fresh tumor from a "passage" mouse in 1 ml buffer, in the flank of female athymic nude mice. Tumors reached the desired size in 2 weeks. Treatment groups of 8 mice each were given injections of specific (α-CEA-ß-lactamase) or nonspecific conjugate (35 μg, except in the experiment of Fig. 5 wherein conjugate was administered at 3.5, 14, or 35 μg/course of therapy, or 35 μg in the first course of therapy only; see Fig. 5 legend), or saline (50 mm sodium borate, 100 mm NaCl, pH 8.0) on days 14, 21, and 28. These injections were followed by four daily injections of 1 mg/kg/day of prodru LY266070 beginning 72 h after conjugate injection. In control arms, prodru injections were substituted with either saline (150 mm NaCl) or DAVLBHYD at the molar equivalent of the prodru dose (0.6 mg/kg/day).

All conjugate and prodru injections were given i.v. All animals tolerated the multiple injections in fractionated dosing protocols without apparent toxicity.

**Covalent Conjugate Tumor Therapy Studies.** Drug-antibody conjugates were studied in nude mice bearing LS174T tumors in the "ADC protocol" (Fig. 7) and in a "covalent conjugate protocol" (Fig. 8). Under the ADC protocol mice were dosed starting on day 16 after tumor implantation, 96 h after saline injections. Drug-antibody conjugate doses were sized to keep the level of DAVLBHYD constant at 2.4 mg/kg/injection between experiment arms. Under the covalent conjugate protocol nude mice bearing established LS174T tumors were given injections twice per week for 2 weeks with sufficient conjugate to make the DAVLBHYD dose 2 mg/kg/injection, or with DAVLBHYD or saline. Doses of drug-antibody conjugate (in mg) administered to provide the desired level of drug are shown in Table 1.

**Data Analysis.** Rate of tumor growth was the indicator of activity used to evaluate the ADC delivery system. Efficacy was inferred from either regression or 1 mg/kg/day of prodru LY266070 beginning 72 h after conjugate injection. In control arms, prodru injections were substituted with either saline (150 mm NaCl) or DAVLBHYD at the molar equivalent of the prodru dose (0.6 mg/kg/day).

All conjugate and prodru injections were given i.v. All animals tolerated the multiple injections in fractionated dosing protocols without apparent toxicity.

**RESULTS**

The two cell lines used in this study, LS174T and T380, have served as models for localization and treatment with various antibody and drug conjugates. The initial tumor therapy experiments were designed to determine whether ADC delivery could immunospecifically enhance the efficacy of DAVLBHYD in two cell lines. Subsequent experiments were designed to determine which antigen best mediated the enhanced effect, what dose of antibody-enzyme conjugate was required to obtain the maximum effect, and how ADC compared with labile covalent drug-antibody conjugate-targeted delivery in terms of overall efficacy, dependence on a particular antigen, and antibody dose requirement. Comparison of ADC delivery with covalent drug-antibody conjugate delivery was performed on the basis of an ADC-dosing protocol and a drug-antibody conjugate-dosing protocol.

**ADC Tumor Therapy Studies.** The cell lines used for in vivo studies are colon carcinoma lines designated T380 and LS174T. LS174T cells have been shown to express the KS1/4 (24), TAG-72 (25, 26), and CEA (27–29) antigens. CEA has also been detected on T380 cells (30). Antigen expression on the LS174T cells is described in "Results."

Tumor growth experiments were performed somewhat differently for the two tumor types. LS174T tumors (Figs. 3 and 6) were implanted as in the biodistribution study (s.c. inoculation of 1 × 10⁷ cells) and allowed to reach the desired size before dosing was initiated. Animals (10 in the saline control group; 5 in the treatment groups) were treated with specific (α-KS1/4-, α-CEA-, or α-TAG-72-ß-lactamase) or nonspecific conjugate (35 μg), or phosphate-buffered saline (10 mm NaPO₄, 150 mm NaCl, pH 7.4) on the days indicated in the figure. These injections were followed after 96 h by a single injection of prodru (4 mg/kg), drug (2.4 mg/kg), or phosphate-buffered saline. Tumor masses were calculated from tumor volume (assuming a tissue density of 1 g/cm³), which was determined by caliper measurements, using the formula mass = 1 g/cm³ × 0.5 × L × W², where L and W are the longest dimension and its perpendicular in cm, respectively. All injections were made in the tail vein.

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All conjugate and prodru injections were given i.v. All animals tolerated the multiple injections in fractionated dosing protocols without apparent toxicity.

**Table 1 Doses of drug-antibody conjugate (in mg) administered to provide desired level of drug**

<table>
<thead>
<tr>
<th>Drug/antibody conjugate</th>
<th>in mgi administered to provide desired level of drug</th>
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<tbody>
<tr>
<td>α-KS1/4</td>
<td>1.9</td>
</tr>
<tr>
<td>α-CEA</td>
<td>4.3</td>
</tr>
<tr>
<td>α-TAG-72</td>
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</tr>
<tr>
<td>IgG</td>
<td>2.2</td>
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<tr>
<td>α-KS1/4</td>
<td>1.6</td>
</tr>
<tr>
<td>α-CEA</td>
<td>3.6</td>
</tr>
<tr>
<td>α-TAG-72</td>
<td>0.8</td>
</tr>
<tr>
<td>IgG</td>
<td>1.8</td>
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conjugate constructs in the past. Previous studies have shown that neither KS1/4 (31) nor TAG-72 internalize efficiently (25, 26). Other studies have shown that TAG-72 and CEA antigen expression can be markedly affected by whether the cells are grown in vitro or in vivo as human tumor xenografts (32). We therefore determined the relative staining intensity and heterogeneity of all three antigen systems in LS174T tumor xenografts. Frozen sections of s.c. implanted LS174T xenografts were subjected to standard immunoperoxidase staining techniques by using biotinylated MoAbs directed against the antigen epitopes utilized in the therapy experiments described throughout this paper. An irrelevant biotinylated MoAb was also included as a negative control. This study demonstrated that the antigen expression detected by the MoAbs used to construct the β-lactamase conjugates was similar to that in previously published reports. Thus, the KS1/4 antigen was characterized by homogeneous, intense staining which was similar to that observed against human colon tumor tissue (24). CEA antigen expression was observed to be intense cytoplasmic and membrane staining in 70–80% of the tumor cells, in good agreement with the findings of Esteban et al. (29) who examined the reactivity of the ZCE025 anti-CEA MoAb in LS174T tumor xenografts. ZCE025 binds to the same epitope as the CEM-231 MoAb used in the present study (33). Similar staining was observed with the anti-CEA antibody CEM231 on T380 tumors. The anti-TAG-72 MoAb, CC-49 (20, 34), bound to approximately 50–70% of the LS174T tumor section and demonstrated reactivity which was associated with tumor cells as well as extracellular mucin. The binding characteristics of CC-49 against LS174T xenografts was very similar to that reported for another anti-TAG-72 MoAb, B72.3, against this same tumor (25).

Before testing the effects of ADC treatment on tumor growth it was necessary to show that tumor localization of the antibody-enzyme conjugate occurred, and to determine an appropriate prelocalization time. Tissue distribution of radiolabeled, α-KS1/4-β-lactamase conjugate (Fig. 2) showed substantial localization to LS174T tumors, with favorable, (≥10), tumor: blood ratios beyond the 24-h point. Within 72 h radiolabeled conjugate was barely detectable (0.1% of the injected dose) in the blood. Consequently, 72 or 96 h was considered to be a sufficient interval between conjugate and prodrug injections. The apparent decrease in tumor conjugate concentration between 72 and 120 h (Fig. 2) was shown to be due to increase in tumor mass during this period, not conjugate egress (see “Discussion”). Conjugate concentration in serum has also been followed by enzyme assay (35) and was shown to drop to an acceptably low level (200 ng/ml) within 72 h after injection.

The prodrug dose and schedule used for the ADC experiments shown in Figs. 3 and 6, i.e., 4-mg/kg bolus i.v. injection once weekly for 3 consecutive weeks, was chosen because this was shown in preliminary experiments (not shown) to be the maximum amount of prodrug that could be administered by itself which did not result in significant antitumor efficacy or animal toxicity. The equivalent amount of free DAVLBHYD (2.4 mg/kg), used in Figs. 3 and 7, was also near its maximum tolerated dose since free drug treatment caused the mice to lose between 13 and 15% of their pretreatment body weight. In contrast, the weight loss due to the 4-mg/kg prodrug treatment ranged from 7 to 10%. The prodrug dose was fractionated in the experiments of Figs. 4 and 5 to determine whether fractionation could be used to decrease treatment toxicity, thereby permitting higher prodrug doses. The results supported the hypothesis, in that neither the mice treated with free drug nor the mice treated with conjugate/ prodrug exhibited any weight loss under this protocol, while efficacy appeared to have been similar to that seen with bolus dosing (see below).

ADC treatment of LS174T tumors with either CEA- or KS1/4-targeted conjugates and prodrug caused the tumors to regress. Using α-KS1/4 conjugate, tumors regressed from 0.37 ± 0.17 (SE) g on day 16 to 0.01 ± 0.01 g on day 44, while with α-CEA conjugate the tumors regressed from 0.30 ± 0.15 to 0.05 ± 0.03 g (Fig. 3) over the same period. These growth curves show that while antigen-specific ADC treatment caused regression of tumors, treatment with free drug at the same dose caused only a delay of about 23 days in the time required for the tumors to reach 2 g. Growth curves for CEA- and KS1/4-directed treatments were indistinguishable. Prodrug alone had little effect on tumor growth. Irrelevant conjugate/prodrug had activity at least as great as free drug alone, but significantly less than antigen-targeted treatments, indicating that antigen specificity contributes to treatment efficacy.
TUMOR-SPECIFIC ACTIVATION OF A Vinca PRODRUG

Fig. 4. T380 tumor growth curves following ADC treatment directed at CEA. Treatments were similar to those in Fig. 3, except that prodruk administration was begun 72 h after conjugate instead of 96, and was fractionated over 4 days. Thus, conjugate administration, 35 µg of α-CEA, ⊳, or irrelevant, ○, open arrows, was followed after 72 h by 1.0 mg/kg/day of LY266070 for 4 days. The filled arrows denote the first day of prodruk dosing. Additional control groups were treated with saline followed by LY266070, ⊳, or by saline, ○, on the same schedule. Tumor masses averaged 0.3 g at initiation of treatment.

Fig. 5. Response of T380 tumors to ADC treatment with different doses of α-CEA-ß-lactamase conjugate. Treatment schedule was identical to that in Fig. 4, e.g., conjugate followed after 72 h by 4.0 mg/kg prodruk fractionated over 4 days. Conjugate was administered at 35, 14, ⊳, and 3.5, ⊳, µg in each course of therapy. A fourth group of mice, ⊳, were treated with 35 µg conjugate in the first therapeutic course only, e.g., on day 14 only, but with prodruk in all three courses. Control groups were treated with saline followed by DAVLBHYD, ⊳, or saline, ○, on the same schedule. Tumor masses averaged 0.3 g at initiation of treatment.

T380 xenografts also regressed, from 0.34 ± 0.03 g on day 18 to 0.22 ± 0.07 g on day 60, on treatment with α-CEA-ß-lactamase conjugate and LY266070 (Fig. 4; note that the solid arrows in the figure denote the first of 4 days of prodruk dosing in each course of therapy as described in “Materials and Methods”). As in the LS174T model free DAVLBHYD produced only a minor growth delay (9 days in the time required for tumors to reach 2 g), while ADC treatment produced regression and long-term stabilization of tumor mass with an equivalent drug dose. Irrelevant conjugate/LY266070 had an intermediate effect in which growth to reach 2 g was delayed 26 days without regression.

The relationship between antibody-enzyme conjugate dose and response was examined by holding the prodruk dose constant at 4 mg/kg/course of therapy fractioned over 4 days as in the experiment of Fig. 4, and decreasing the α-CEA-ß-lactamase conjugate dose. An additional treatment group received conjugate in the first treatment course only, but received prodruk in all three courses, and a control group received free drug at a molar equivalent dose. Growth curves are compared in Fig. 5. All ADC treatment groups showed initial tumor regression, but the duration of the response varied with conjugate dose. Responses duplicating those observed in Fig. 4 were observed in response to ADC treatment with 35 and 14 µg conjugate/course, while the duration of response to 3.5 µg/course was shorter. Thus, maximal efficacy was obtained at between 3.5 and 14 µg α-CEA-ß-lactamase conjugate/course.

Administration of conjugate with only the first course caused a diminished effect in which tumor growth was delayed 39 days, compared to saline-treated controls, in the time required to reach 2 g. Histological examination of the small tumor areas present at various stages of treatment in additional sets of mice showed that CEA was available to bind the second and third courses of conjugate (data not shown).

Coexpression of KS1/4 (24, 36), CEA (30, 37), and TAG-72 (25) antigens (see above) by LS174T tumor cells enabled the direct comparison of two modes of targeted chemotherapy on a single cell line: ADC (Fig. 6) and covalent drug-antibody conjugates (Figs. 7 and 8). For the ADC study doses and schedule were identical to those in the experiment of Fig. 3. Large initial tumor volumes averaging 0.6 to 0.7 g were used in these experiments to provide a significant challenge to treatment. Tumors in animals treated with α-CEA-ß-lactamase/prodrug regressed from 0.83 ± 0.24 g on day 20 to 0.19 ± 0.07 g on day 41; α-KS1/4-ß-lactamase/prodrug-treated tumors changed from 0.96 ± 0.23 g to 0.49 ± 0.26 g over the same period; while tumors treated with α-TAG-72-ß-lactamase/prodrug changed from a maxi-
curves are compared in Fig. 6. While no significant difference between the targeted treatment groups was observed, only the α-CEA-β-lactamase/prodrug treatment strictly met the criterion for regression, as described in “Materials and Methods.” Irrelevant antibody-β-lactamase conjugate/prodrug treatment resulted in an intermediate effect in which growth was delayed (18 days to reach 2 g) but no regression was observed.

Treatment with DAVLBHYD delivered as a covalent drug-antibody conjugate was measured for comparison with ADC delivery by using α-KS1/4-, α-CEA-, and α-TAG-72-DAVLBHYD conjugates. Doses were calculated to provide the same molar quantity of DAVLBHYD as the ADC treatments (see “Materials and Methods”). The growth curves (Fig. 7) show that no tumor regression occurred in any treatment arm and that only KS1/4-DAVLBHYD may have been more effective than irrelevant conjugate, although the difference did not meet the criterion for significance.

An additional experiment was performed to test the efficacy of the covalent drug-antibody conjugates in a more intense dose schedule, typical of previous experiments with these agents (25). Mice with initial tumor burden averaging 0.5 g were given injections twice per week for 2 weeks of sufficient α-KS1/4 and α-TAG-72 drug-antibody conjugate to give 2 mg/kg DAVLBHYD per injection. Mice with initial tumor burdens of 0.15 g were treated with α-CEA-DAVLBHYD conjugate or the identical control materials. In one set of animals treated with free drug the dose was 60% lethal dose, in the other it was 40% lethal dose. Growth curves (Fig. 8) show no tumor regressions. KS1/4- and TAG-72-DAVLBHYD conjugates appeared to suppress LS174T tumor growth relative to the saline-treated control group (Fig. 8A). Neither of these conjugates, however, was significantly more effective than an irrelevant IgG-DAVLBHYD conjugate (Fig. 8A). The anti-CEA-DAVLBHYD conjugate, on the other hand, was more effective than either free DAVLBHYD or an irrelevant IgG-DAVLBHYD conjugate when used in a lower tumor burden model (Fig. 8B).

DISCUSSION

Tumor Growth Inhibition. The data in Figs. 3–6 demonstrate that the activity of the oncolytic agent DAVLBHYD can be qualitatively increased from moderate tumor growth inhibition to tumor regression and long-term stabilization by utilizing the antibody-directed catalysis delivery system. Since this enhancement of activity was observed in both the T380 and LS174T colon carcinoma models it is unlikely that the result is strongly cell line dependent.

The data in Figs. 3 and 6 show that the ADC delivery system mediates tumor regression with the three antigen targets, CEA, KS1/4, and TAG-72, which have different patterns of expression. The choice of antigen to be utilized for antibody-targeted therapy is generally critical to its success. Many targeted delivery systems require cellular internalization for anti-tumor activity (38, 39), but only a few of the known tumor-selective antigens internalize efficiently (40). For those antigens that do internalize, each cell that the antibody fails to target, whether because of antigen heterogeneity, slow antibody diffusion rate, or poor vascularization, represents a locus from which the tumor can regenerate without inhibition.

Internalization is not required for chemoimmunoconjugates constructed with certain suitable linkers, like those used in the experiments of Figs. 7 and 8. These “labile-linked” conjugates have demonstrated preclinical activity against a variety of target antigens (31). But even for drug-antibody conjugates that do not require internalization, antitumor efficacy has been shown to correlate with target antigen expression level (41). The results in Figs. 7 and 8 showing that KS1/4 targeting is more effective than CEA or TAG-72 targeting for DAVLBHYD-antibody conjugates are in agreement with this conclusion. It would be clearly advantageous, therefore, to use a site-specific delivery system which is relatively unaffected by antigen characteristics such as copy number and heterogeneous expression, and which does not require internalization.

Anti-CEA and anti-TAG-72 conjugates do not internalize sufficiently (25, 26, 30, 37). Both bind antigen in the interstitial fluid as well as at cell membranes and both antigens are heterogeneously distributed in LS174T tumors, TAG-72 being more heterogeneous than CEA. Conjugates targeting KS1/4, in contrast, bind exclusively at the cell membrane and may bind virtually every LS174T cell (see “Results”). These three antigens, therefore, represent a diverse panel of molecular targets with which to evaluate the therapeutic potential of the ADC system. The data in Figs. 3 and 6 indicate that ADC therapy was quite successful for all three antigens with no significant difference in activity among them, and is, therefore, relatively unaffected by the differing properties of these antigens.

Within this set of experiments two slightly different dosing schedules were also compared. While the total prodrug doses were the same in all experiments reported, the weekly 4-mg/kg prodrug dose was fractionated over 4 days instead of being given in a single bolus in the T380 experiments (Figs. 4 and 5). Fractionation should lead to greater specificity, because conjugate not bound in the tumor has more time to clear before the full prodrug dose is administered. Furthermore, it was reasoned that if the prodrug elimination was rapid, fractionation might result in a higher percentage of the total dose reaching the tumor, while a lower peak concentration of the drug might reduce its overall toxicity. The only clearly discernible difference between the fractionated and bolus dosing was that mice in the free drug control...
groups showed no weight loss with fractionated dosing but significant weight loss with bolus dosing.

The data also show that single binding arm antibody-enzyme conjugates used in the experiments of Figs. 3–6 mediate drug delivery. Since comparable results were obtained with prodrug fractionated over days 4–7 (Figs. 4 and 5) to results with prodrug given in a bolus day 5 (Figs. 3 and 6), retention of the single binding arm conjugate in tumor appears to be adequate. However, the result obtained with injection of conjugate in the first therapeutic course only (Fig. 5, ) suggests that the tumor residence time of the conjugate during treatment is less than 3 weeks. Egress of conjugate from the tumor compartment during treatment could be due to the antigen-antibody-binding equilibrium, in which case a dual binding arm conjugate [e.g., an antibody- or F(ab')2-enzyme conjugate] would be advantageous; but it could also be due in part to metabolic processes at the tumor. Physiological changes, resulting from, for instance, cell lysis, could also decrease retention of conjugate at the tumor.

It has been noted that one potential disadvantage to the use of covalent drug-antibody delivery is that high doses of antibody are likely to be required in humans (4). The doses of DAVLBHYD-antibody conjugate given in the experiments of Figs. 7 and 8 ranged from 800 to 4300 μg. Doses of 270 to 4800 μg/course of therapy were required to effectively treat LS174T (25, 33) and other (41) tumor xenografts with labeled-linked chemiomunocoujugates in previous studies. The experiment of Fig. 5 was performed to determine the relationship of response to antibody-enzyme conjugate dose for the ADC system. The results showed that response increased with dose up to 14 μg, but not above this level, so that in the T380 model maximum efficacy was achieved with between 3.5 and 14 μg conjugate. Thus, the quantity of conjugate localized at the tumor following a 14-μg dose apparently hydrolyzed all LY266070 that reached the tumor and additional catalytic activity had no additional benefit at this prodrug dose. (In a separate test of prodrug dose response, tumor growth inhibition decreased when prodrug dose was decreased at constant conjugate level. Data not shown.)

The ratio of antibody doses used to give equivalent quantities of drug in the different delivery formats (drug-antibody conjugate/ADC) are summarized in Table 2. They vary from 16 to 69 on a molar basis depending on the number of drug residues per antibody for the covalent drug-antibody conjugates. These ratios are calculated from the doses actually used in the experiments of Figs. 6 and 7, and are artificial in the sense that drug dose for ADC can be increased without increasing the antibody-enzyme conjugate dose. Since the data in Fig. 5 show that considerably less than 35 μg of F(ab')2-β-lactamase may be required to achieve the full effect, it is estimated that drug-antibody conjugates require doses ranging upward from 100 times greater than required by ADC to deliver equivalent drug. Furthermore, even larger amounts of drug-antibody conjugate administered in a more intense schedule (Fig. 8) failed to produce the regressions observed with ADC treatment. Hence, these data suggest that ADC has the advantage over drug-antibody conjugate delivery that, in addition to improved efficacy, it requires much less antibody.

Table 2: Ratio of antibody doses: DAVLBHYD antibody:β-lactamase antibody

<table>
<thead>
<tr>
<th>Antibody</th>
<th>wt/wt</th>
<th>mol/mol</th>
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<tbody>
<tr>
<td>α-KS1/4</td>
<td>53</td>
<td>30</td>
</tr>
<tr>
<td>α-CEA</td>
<td>123</td>
<td>69</td>
</tr>
<tr>
<td>α-TAG-72</td>
<td>28</td>
<td>16</td>
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</table>

The antibody-enzyme conjugate dose required to reach maximal efficacy in ADC is expected to be a function of the kinetic constants for the enzyme substrate pair, and the potency of the drug. If enzyme-catalyzed hydrolysis of prodrug is slow, then a large concentration of enzyme would be required to activate sufficient drug to reach a therapeutic threshold. Likewise, at a given enzyme turnover rate, a lower potency drug should require a larger amount of enzyme to produce the therapeutic effect. The β-lactamase-LY266070 system of the current study involves both a relatively rapid turnover rate (kcat/KM ~11 (s μM)-1, (13)), and a relatively potent drug (50% inhibiting concentration 10^-8 M). Hence, the conjugate dose requirements for other systems could be considerably higher (or lower) than the one determined in this study, depending on the potency of the drug and the rate at which it is produced by the enzyme. We note for comparison that antibody doses in preclinical radioimmunoimaging and radioimmunotherapy experiments performed in our laboratories typically use 10–20 μg conjugate/mouse (42, 43). The radioimmunoimaging studies have been successfully translated into human patients by using doses between 1 and 50 mg of antibody. We predict that dose requirements would translate similarly from mice to humans for antibody-enzyme conjugates if the drug potency and enzyme kinetic parameters were similar to those of this study. If the prodrug were a significantly worse substrate (lower kcat/KM), or significantly less potent, then higher conjugate dose requirements would be expected.

In the experiments depicted in Figs. 3 and 4, irrelevant antibody-enzyme conjugate/prodrug was more effective than a molar equivalent dose of free drug. This efficacy enhancement may be due to conjugate which localizes in tumor independent of antigen binding ["preferential accumulation" (4)]. Theoretical considerations suggest that some non-antigen-mediated localization of proteins to tumors is expected and can be attributed to tumor physiology (44).

Pharmacokinetics. The pharmacokinetic behavior of the α-KS1/4-β-lactamase conjugate (Fig. 2) is similar to that reported previously for F(ab')2 fragments (45–47). Rapid clearance from the blood and good tumor/blood ratios indicate favorable pharmacokinetics of the conjugate. Since the behavior of the α-KS1/4 conjugate resembled closely that of other constructs of similar molecular weight, and since tumor targeting had previously been demonstrated with each of the antibodies from which the conjugates were constructed, it was assumed that the α-CEA and α-TAG-72 conjugates would also localize and would clear from the serum within 72 h. The data showing that the efficacy and toxicity resulting from treatment with these conjugates were no different than those resulting from treatment with α-KS1/4 conjugates support this assumption.

The data reported in Fig. 2 suggest that conjugate level in the tumor decreased continuously from 24 to 120 h after injection. However, examination of the organ masses showed that the tumor mass change between 72 and 120 h accounted for essentially all of the conjugate concentration decrease during this period. Egress of conjugate from the tumor after 72 h was found to be very slow.

Reports on the pharmacokinetics of antibody-enzyme conjugates with different molecular properties have appeared previously (48). Bagshawe (3), investigating F(ab')2-carboxypeptidase G2 conjugates, noted a slow rate of serum clearance of anti-CEA conjugate relative to anti-hCG conjugate. This difference was attributed to accelerated clearance of the hCG conjugate due to complex formation between conjugate and the hCG antigen (which is secreted into the serum at high level), and subsequent removal of the immune complex by the reticuloendothelial system. Since the KS1/4 antigen is a transmembrane glycoprotein (36), it is unlikely that shedding and immune complex formation were involved with the pharmacokinetics of the α-KS1/4-F(ab') conjugate of Fig. 2. The relatively rapid clearance of this conjugate suggests that immune complexes are not required for
rapid serum clearance, and that other molecular properties are responsible for the slow serum clearance of the F(ab′)_2-carboxypeptidase G2 conjugate.

Tumor Growth Inhibition in Comparable Systems. Antibody-directed catalysis systems based on carboxypeptidase G2 (referred to by the authors as Antibody-Directed Enzyme Prodrug Therapy or ADEPT) showed efficacy against LS174T and other tumor xenografts (3). In the LS174T studies targeting the CEA antigen, tumor growth inhibition, but not regression, resulted from a three-step regimen: (a) localization of anti-CEA F(ab′)_2-carboxypeptidase G2 conjugate; (b) galactosylated enzyme-blocking antibody; and (c) nitrogen mustard-derogrded prodrg. The galactosylated enzyme-blocking antibody had the effect of clearing conjugate from the serum to allow prodrg administration while tumor level of conjugate was still high (49). This study was conducted with a relatively small initial tumor volume, used a prodrg dose higher than the maximum tolerated free drug dose, and used only a single course of therapy (3). In studies targeting the hCG antigen in a different cell line, cures were reported (3).

Antibody-directed catalysis systems based on alkaline phosphatase and other enzymes have also been reported to be active in tumor xenograft models (2). Whole antibody-alkaline phosphatase conjugates were used in combination with prodrgs that are phosphorylated derivatives of etoposide and mitomycin C to obtain tumor growth inhibition, but if it is based on toxicity, diminished toxicity must lead to better treatment. Labile-linked drug-antibody conjugates in this system (a nitrogen mustard) was more bioavailable than the free drug. The nonspecificity was attributed to substantial conjugate re-formation free drug were used. However, efficacy was relatively nonspecific: 4.3 versus 3.7 log cell kill for tumor-specific versus irrelevant conjugate, respectively, in one experiment using mitomycin phosphate for prodrg. The nonspecificity was attributed to substantial conjugate remaining in the serum when the prodrg was administered, 24 h after injection (2). In all in vivo studies reported to date (2, 3), however, ADC delivery has proven more effective than direct injection of the free chemotherapeutic agent.

As noted, the previous studies on the efficacy of ADC delivery systems (2, 3) have used prodrg doses greater than the molar equivalent free drug dose, taking advantage of the difference in toxicity between drug and prodrg in these systems (2, 50). The remarkable enhancement for molar equivalent doses observed here with ADC delivery (Figs. 3–6) may result from an improvement in the bioavailability of the prodrg over the free drug. Pharmacokinetic studies to investigate this question are in progress. A study by Antoniw et al. (51) demonstrated that prodrg can have pharmacokinetic properties different from those of the parent drug. It was shown that the parent drug in that system (a nitrogen mustard) was more bioavailable than its prodrg derivative (glutamic acid amide of the nitrogen mustard).

The ultimate intent of this series of experiments is to assess the practical and theoretical advantages and disadvantages of different methods of administering toxic agents to humans with the use of preclinical models. The comparison could be based on either efficacy or toxicity, but if it is based on toxicity, diminished toxicity must lead to better treatment. Labile-linked drug-antibody conjugates in this study showed efficacy comparable to, or slightly better than direct injection of drug, but the activity was strongly antigen dependent, being most pronounced for the antigen present in highest concentration, and least pronounced for a heterogeneously expressed antigen. Drug-antibody conjugates were substantially less toxic than free drug, but efficacy comparable to or better than free drug required a very high antibody dose. ADC delivery, in contrast, showed substantial improvement in efficacy over free drug, in addition to toxicity reduction. For ADC the improvement was obtained at a much lower antibody dose, and the effect was equivalent for three dissimilar antigens. Thus, ADC improved the efficacy of the oncolytic agent and overcame two of the most troubling aspects associated with antigen-directed delivery of chemotherapeutic agents: antibody dose and antigen suitability.

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Site-specific Prodrug Activation by Antibody-β-Lactamase Conjugates: Regression and Long-Term Growth Inhibition of Human Colon Carcinoma Xenograft Models


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