**Short Communication**

A VINDESINE-ANTI-CEA CONJUGATE CYTOTOXIC FOR HUMAN CANCER CELLS IN VITRO

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Eighty to ninety per cent of patients with lung cancer are potential candidates for chemotherapy (Selawry, 1977). At present a major limitation to the use of chemotherapeutic agents is their lack of specific anti-tumour activity. Doses required for adequate cancer treatment are invariably accompanied by unwanted systemic side effects, myelosuppression, alopecia, neurotoxicity and damage to gastrointestinal epithelium being among the more prominent complications. The concept of using anti-tumour antibodies as carriers is an attractive one, in that the potency of the cytotoxic drugs may be combined with the specificity provided by the antibodies. Since Ehrlich (1956) first suggested this possibility, various drugs have been linked to anti-tumour antibodies. The multiplicity of methods used for conjugation and the variety of test systems investigated have been well reviewed recently (Ghose & Blair, 1978; Lee & Hwang, 1979; De Weger & Dullens, 1980; Rowland, 1981). We have previously reported the ability of an anti-CEA Ig to enhance the cytotoxicity of vincristine in a $^{51}$Cr-release assay, using cultured human lung-tumour cells (Johnson et al., 1980). In this communication we report the cytotoxic effects of a vindesine-anti-CEA Ig conjugate on cells from the same line, using a terminal $^3$H-uridine uptake assay.

The sheep anti-CEA Ig was donated by Dr A. R. Bradwell from the Immuno-Diagnostics Research Laboratory, University of Birmingham. Its preparation and characterization have previously been described (Dykes et al., 1980). In these experiments the initial protein concentration of the preparation was adjusted to 1.28 mg/ml with medium. Vindesine–Ig conjugates were prepared at Lilly Research Centre Ltd from desacetylvincaleucoblastine acid hydrazide by a modification of the procedure described for vindesine-BSA (Conrad et al., 1979) and purified by gel filtration. A solution of the anti-CEA conjugate had a protein concentration of 1.28 mg/ml as assayed by the Lowry method, and a drug concentration of 27.5 μg/ml, determined by difference spectroscopy, giving a molar conjugation ratio of 4.6. A conjugate with normal sheep Ig had a protein concentration of 2.3 mg/ml and a drug concentration of 43 μg/ml, a molar ratio of 3.7. For comparison with the anti-CEA conjugate, this and unconjugated vindesine were diluted to an initial drug concentration of 27.5 μg/ml in medium. Calu-6, the lung-tumour line used in this study, was supplied by Dr Jørgen Fogh, Sloan Kettering Institute, New York. Calu-6 cells were originally obtained from an anaplastic lung tumour and maintained as a monolayer in Eagle's
minimal essential medium, supplemented with 15% foetal calf serum, non-essential amino acids, L-glutamine and antibiotics.

We used a terminal labelling method based on the ability of surviving cells to incorporate 3H-uridine into their RNA-precursor pools. This assay, described by Smith & Nicklin (1979), has shown a direct correlation between counts incorporated and the number of viable cells, indicating that it is a reliable method of assessing cell survival. Calu-6 cells in the exponential phase of growth were trypsinized, washed and resuspended in medium. One hundred μl containing either $5 \times 10^3$ or $10^4$ cells were dispensed into each well of a microtitre plate (Sterilin: M29ARTL) and incubated at 37°C in a humidified atmosphere of air and 5% CO₂. After 24 h, doubling dilutions of drug, antibody or conjugate were prepared and 50 μl added to the appropriate wells with 50 μl of medium. Tests and medium-only controls were performed in triplicate. After a further incubation of 72 h, the supernatants were discarded, the monolayers washed ×3 with warm (37°C) phosphate-buffered saline (PBS) and 100 μl of medium added to each well. After recovery for 24 h, the medium was again decanted and replenished with 50 μl of medium containing 1 μCi 3H-uridine (Radiochemical Centre, Amersham). After 3 h, when we have shown that the transportation mechanisms but not the RNA-precursor pools are still saturated by uridine, the medium was removed, the cells washed with PBS, and 150 μl of ice-cold 5% trichloracetic acid added to each well for 20 min. One hundred and twenty-five μl of this was then removed for scintillation counting.

The surviving fraction was calculated as:

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\text{Mean radioactive count (ct/min) in test wells} \times 100\% \text{ Mean radioactive count (ct/min) in medium control wells}
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The interrupted line in Fig. 1 shows the dose–response curve for vindesine in this system. The anti-CEA immunoglobulin on its own had a low cytotoxicity, and when added to the drug at the relevant concentrations the effect closely paralleled that of drug alone. However, with the vindesine–anti-CEA conjugate there was a marked dose-dependent decrease in surviving fraction.

Fig. 2 shows the results of a second experiment, performed to compare the effects of the drug conjugated to anti-CEA Ig and to the normal sheep Ig using $10^4$ cells per well. Whereas the conjugate of vindesine with normal sheep Ig had little effect on cell survival, the anti-CEA conjugate produced a dose-dependent decrease in surviving fraction. In a 2-stage immunoperoxidase test, the anti-CEA Ig localized on Calu-6 cells and on CEA-secreting colonic adenocarcinoma cells, both before and after conjugation. Similarly in a competitive-inhibition test with
peroxidase and ELISA results demonstrate that the anti-CEA Ig after conjugation retains its ability to recognize CEA determinants. However, preliminary evidence with these tests suggests that there is a loss of anti-CEA activity with storage at 4°C. This may explain the difference in surviving fraction of the anti-CEA conjugate seen in Fig. 1 (31 days after conjugation) and Fig. 2 (52 days).

Although CEA is not tumour-specific, its concentration may be greatly increased in gastro-intestinal, lung and other malignancies. This difference has been exploited by investigators who have reported in vivo tumour localization using radiolabelled antibodies against CEA, in some cases despite high circulating levels of the antigen (Goldenberg et al., 1978; Mach et al., 1980). Antibodies to CEA may, therefore, provide useful carriers for cytotoxic compounds, thereby achieving a selective accumulation of the drug at its desired site of action. The specificity and potency of such therapy may be further improved by using monoclonal antibodies from a hybridoma source. In this context, xenogeneic (Ghose et al., 1975; Newman et al., 1977; Goldenberg et al., 1978; Mach et al., 1980) and monoclonal (Nadler et al., 1980) antibodies have already been administered safely to cancer patients.

In the system reported here it would appear that the potency of a vindesine alkaloid is remarkably increased by the ability of the antibody to deliver it to tumour cells. We are currently extending our investigations into the stability, affinity and specificity of the vindesine–anti-CEA conjugate, with a view to its clinical applications in the treatment of cancer.

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