Pre-clinical evaluation of a novel chloroethylating agent, Clomesone

A.M. Matthew, R.M. Phillips, P.M. Loadman & M.C. Bibby

Clinical Oncology Unit, University of Bradford, West Yorkshire, BD7 1DP, UK.

Summary The in vitro activity of the novel chloroethylating agent, Clomesone, was investigated in a panel of established murine and human tumour cell lines. In vivo anti-tumour activity was examined against three transplantable adenocarcinomas of the mouse colon and in vivo bone marrow toxicity was assessed using a spleen colony forming unit assay. The pharmacokinetic behaviour of the drug in vivo and drug stability in vitro was analysed by gas chromatography with electron capture detection. Clomesone exhibited no activity in vitro against the majority of cell lines derived from solid human colorectal carcinomas. Anti-tumour activity against the murine tumours in vivo was not impressive and was accompanied by myelosuppression. Pharmacokinetic data suggested that the lack of in vivo activity was due to the failure to achieve effective anti-neoplastic drug concentrations at the tumour site. It was concluded that this study found no evidence to suggest that Clomesone was toxicologically more selective than the chloroethylnitrosoureas.

Clinical responses to cytotoxic agents have been largely restricted to the haematological malignancies while the majority of human solid tumours have remained refractory to chemotherapy (Marsoni et al., 1987). Thus the search for more effective compounds has continued. Chloroethylnitrosoureas are highly active anti-neoplastic agents with a broad spectrum of anti-tumour activity in experimental systems. However, their clinical usefulness has been limited by non-selective host toxicity, particularly myelosuppression.

The chloroethylnitrosoureas decompose spontaneously in aqueous solution to generate reactive intermediates that are capable of alkylating and carbamoylating nucleophilic targets (Cheng et al., 1972). Alkylation reactions consist of chloroethylations and hydroxyethylations of DNA (Tong et al., 1982). Chloroethylolation of O'-guanine with subsequent cross-link formation is considered to be the most important cytotoxic reaction (Lown et al., 1978) while hydroxyethylation reactions are thought to be responsible for their mutagenic and carcinogenic effects (Pelfrene et al., 1976; Swenson et al., 1979). Carbamoylation activity is not required for cytotoxicity but does contribute to the overall effect, probably by inhibiting DNA repair (Erickson et al., 1980) and it has been suggested that the carbamoylating activity of some chloroethylnitrosoureas could interfere with the ability of normal cells to recover from the action of these drugs (Sarban et al., 1984).

In the search for other types of compound which would chloroethylate DNA, a series of 2-haloethyl sulphphonates was synthesised (Shealy et al., 1983). 2-Chloroethyl(methylsulphonyl)methanesulphonate (Clomesone), the most active analogue tested, was shown to be highly effective against P388 leukaemia in vivo (Shealy et al., 1984). The chemical structure of Clomesone is shown in Figure 1. Clomesone was further evaluated against a spectrum of animal tumour models and was found to be generally as effective as the chloroethylnitrosoureas (Dykes et al., 1989).

The results of an investigation of the action of Clomesone in vitro indicated that it affected cellular DNA in a manner similar to the chloroethyl nitrosoureas with cross-link formation via chloroethylolation of O'-guanine residues (Gibson et al., 1985). However, it was found to be more specific in its reaction with DNA in that it produced less variety of products than the chloroethyl nitrosoureas with no apparent generation of hydroxyethyl adducts (Gibson et al., 1986). In addition, the chemical nature of Clomesone precludes the formation of isocyanate breakdown products (Shealy et al., 1984). It was suggested that this lack of carbamoylating potential of Clomesone, together with the lack of hydroxyethylating activity, might result in fewer unwanted side reactions and that Clomesone would be a more toxicologically selective agent. On this basis Clomesone was selected for clinical development and is presently undergoing Phase 1 clinical trials in the UK.

Recently, the NCI has introduced a new disease-orientated screening programme based on the assessment of the in vitro cytotoxicity of a panel of cell lines representing the common clinical forms of human cancers using a colorimetric assay (Boyd, 1989). While in vitro colorimetric assays have the advantage that they can rapidly evaluate novel anti-tumour agents against a large number of cell lines, a recent review of the literature has revealed that marked differences in the response of cells in vitro and tumours in vivo exist (Phillips et al., 1990). One reason for this discrepancy is that the in vitro tests ignore the potential role played by the pharmacokinetic behaviour and bioavailability of a drug in determining tumour responses in vivo. This has led some workers to advocate that appropriate transplanta-ble mouse tumour models, similar in sensitivity to solid cancers in man where therapeutic indices are low, also have a role in the pre-clinical evaluation of novel compounds which should include toxicological and pharmacokinetic studies (Corbett et al., 1987; Double & Bibby, 1989).

A series of transplantable murine adenocarcinomas of the colon (MAC tumours) has been shown to exhibit a similar spectrum of histology and chemosensitivity to human large bowel cancer with responses to standard agents only normally seen close to the maximum tolerated dose (Double & Ball, 1975). It has been used extensively as part of the pre-clinical evaluation of new anti-cancer agents within the Screening and Pharmacology Group of the EORTC (Bibby et al., 1988).

The feasibility of performing minimal toxicity studies in conjunction with anti-tumour studies has been described (Bibby et al., 1988b). Measurement of bone marrow damage, the major dose-limiting toxicity of the chloroethyl nitrosoureas, will aid the identification of alternative chloro-

\[
\begin{align*}
\text{CH}_3 & \quad \text{S} \quad \text{CH}_2 \quad \text{S} \quad \text{O} \quad \text{CH}_2 \text{CH}_2 \text{Cl} \\
\end{align*}
\]

2-chloroethyl(methylsulphonyl)methanesulphonate

Figure 1 Structure of Clomesone.
ethylating agents with improved therapeutic indices. The assessment of stem cell survival has been recommended to study irreversible cytotoxic bone marrow injury (Schofield, 1986) and this can be performed readily in mice using a spleen colony forming unit assay.

The purpose of this present study was to further evaluate the novel chloroethylating agent, Clomesone. In vitro activity was assessed against a panel of established murine and human tumour cell lines while in vivo anti-tumour responses were evaluated against an ascitic tumour, MAC 15A, and two solid subcutaneous (sc) tumours, MAC 13 and MAC 26. MAC 13 is relatively nitrosourea-sensitive due to a low level of the repair enzyme O'alkylguanine-DNA alkyltransferase while MAC 26 is relatively nitrosourea-resistant due to a high repair enzyme level (Lunn et al., 1989). In vivo bone marrow toxicity and pharmacokinetic studies and in vitro drug stability studies were performed in conjunction with the anti-tumour studies. A novel, clinically active nitrosourea, 1-(2-chloroethyl)-3-[2-(dimethylaminosulphonyl)ethyl]-1-nitrosourea (TCNU) and a non-carbamoylating nitrosourea, Chlorozotocin, were included in the study as reference compounds.

Materials and methods

Test compounds

Clinically formulated Clomesone was obtained from Dr S.M. Crawford through his involvement in the ongoing Phase I clinical trial. Chlorozotocin was received from the NCI while TCNU was a gift from Pharmacia Leo Therapeutics, Helsingborg, Sweden. Clomesone was dissolved in sterile water while TCNU was dissolved in 0.9% sterile saline. Chlorozotocin was dissolved in 10% dimethylsulphoxide (DMSO)/arachis oil. Drug solutions were prepared at an appropriate concentration for the desired in vivo dose to be administered in 0.1 ml per 10 g body weight by intraperitoneal (ip) or intravenous (iv) injection and for the desired final concentration in vitro to be achieved upon a 1 in 10 dilution.

In vitro studies

Cell culture The in vitro activity of Clomesone was assessed against a panel of established tumour cell lines (Phillips et al., 1992) which consisted of three murine colon adenocarcinomas (MAC 15A, MAC 16, MAC 26), a murine myelomonocytic leukaemia (WEHI-3B), a human chronic myelogenous leukaemia (K562), a human rectal carcinoma (HRT-18) and four human colon adenocarcinomas (DLD-1, HT-29, HCT-18, HCLO). They were routinely maintained as monolayer cultures, except K562, WEHI-3B and MAC 16 which do not adhere strongly to plastic cultures vessels, in RPMI 1640 tissue culture medium containing sodium pyruvate (1 mM), penicillin/streptomycin (50 IU/ml/150 µg/ml) and supplemented with 10% foetal calf serum.

Chemosensitivity testing In vitro chemosensitivity was assessed using a modified micro-tetratrazolium (MTT) assay (Twentyman & Lascombe, 1987). Cell suspensions were obtained from monolayer cultures in the exponential growth phase and 0.5-10 x 10⁴ tumour cells in culture medium were plated into 96-well dishes. Drug solutions were added to give final concentrations ranging from 0.1-100 µg/ml at four log increments and the dishes incubated for 4 days at 37°C in an atmosphere of 5% CO₂. Prior to the addition of 20 µl MTT (5 mg/ml-1), 150 µl old medium was removed and replaced with fresh medium. The dishes were then incubated for a further 4 h and the purple formazan crystals produced were dissolved in MSO. Optical densities of the resulting solutions were read with an ELISA spectrophotometer at a wavelength of 550 nm. Each drug concentration was assayed against each cell line eight times and in four independent experiments. Cytotoxic effects were expressed as percentage survival of treated plates compared to control plates and the initial drug concentration required to inhibit cell survival by 50% (IC₅₀) was obtained from semi-logarithmic plots of cell survival versus concentration.

In vivo studies

Animals Pure strain NMRI mice, aged 6-8 weeks, from our inbred colony were used throughout this study. They were fed with a pellet diet (CRM, Labsure, Croydon, England) and water ad libitum.

Tumour system The development of several transplantable adenocarcinomas of the colon in mice from primary tumours induced by prolonged administration of 1,2-dimethylhydrazine has been described elsewhere (Double et al., 1975). MAC 13 tumours were transplanted into female mice and MAC 26 tumours into male mice by sc implantation of tumour fragments (~1 x 2 mm) in the flank. MAC 15A ascites tumour cells were transplanted into male mice by ip inoculation of 10³-10⁶ tumour cells in 0.2 ml 0.9% saline.

Chemotherapy The differing morphology and growth characteristics of the tumour lines necessitated the use of different chemotherapy protocols. Animals bearing the more rapidly growing MAC 13 and MAC 15A tumours were allocated into groups of 5 and chemotherapy commenced 2 days after implantation. Anti-tumour responses against MAC 13 were assessed 17 days later by comparing the tumour weights of treated and control groups and expressed as percentage tumour weight inhibition while responses against MAC 15A were determined by comparison of median survival times (MST) of treated and control groups as described by Geran et al. (1972). MAC 26 tumour bearers were allocated into groups of 10. The administration of cytotoxic drugs did not commence until these slower growing tumours could be reliably measured (>4 x 4 mm), approximately 17 days after transplantation. Anti-tumour effects against MAC 26 were assessed by twice weekly caliper measurements. Tumour volumes were calculated from the formula a² x b/2, where a is the smaller diameter and b is the larger (Geran et al., 1972) normalised with respect to the starting volume and semi-logarithmic graphs of relative tumour volume against time were plotted. Therapeutic effects were expressed as the regrowth delay which was obtained by comparing the times taken by treated and control tumours to reach a tumour volume ten times that of the starting volume.

Bone marrow toxicity Acute bone marrow toxicity was assessed using a modified version of the spleen colony forming unit assay of Till and McCulloch (1961). The drugs were administered by a single ip injection to pairs of male mice and bone marrow toxicity was assayed 24 h later. Marrow cells were obtained from both femora of each pair of treated mice and pairs of untreated control mice and suspended in RPMI 1640 tissue culture medium. Cell suspensions were diluted so that a 0.2 ml aliquot contained an appropriate number of cells for each experimental group. A number of 5 x 10⁶-7.5 x 10⁶ were initially obtained from the marrows of control mice to investigate the relationship between the number of spleen colonies formed and the number of marrow cells inoculated. Subsequently, a cell inoculum of 1.0 x 10⁶-2.5 x 10⁶ was prepared from both treated and control marrows. The marrow cells were then injected iv via the tail vein into mice which had previously been exposed to irradiation from a Newton Victor Superficial Therapy Unit (GX x 10) at a dose of 11.7 Gy. Groups of six mice were used for each experimental point. After 8 days the mice were killed, the spleens removed and fixed in Bouin’s fluid and the nodules on the spleen surface were counted. The survival fraction was determined by comparing the number of colonies observed with the number of colonies expected for a given cell inoculum of untreated marrow cells.
Pharmacokinetic studies

Sample collection Clomesone was administered to female non-tumour bearing mice by a single ip injection at a dose of 50 mg kg\(^{-1}\). Blood samples were obtained at time intervals ranging from 2 min to 2 h by cardiac puncture under diethyl ether anaesthesia. The samples were placed in heparinised tubes, centrifuged at 4°C (1000 \(g\) for 5 min) and the plasma stored at –20°C until analysed. Each time point was represented by two mice and three independent experiments were performed.

Sample extraction Silanised glassware was used throughout the extraction procedure to minimise drug binding to vessel walls and all samples and reagents were kept on ice whenever possible. A 200 \(\mu\)l aliquot of mouse plasma was mixed with 50 \(\mu\)l 2-chloroethyl-p-toluenesulphonate (Sigma Chemical Company Limited, Poole, UK) as an internal standard, diluted with 0.9% saline and extracted with 6 ml diethyl ether. Following centrifugation (1000 \(g\) for 5 min at 4°C), the organic layer was decanted and 1 g anhydrous sodium sulphate was added to remove any excess water. After a further centrifugation step to remove the sodium sulphate, the diethyl ether layer was evaporated in a stream of nitrogen in a water bath at 25°C and the dried sample was reconstituted in 100 \(\mu\)l ethyl acetate.

Sample analysis Plasma Clomesone levels were analysed by gas chromatography. The chromatographic system consisted of a Shimadzu GC-14A chromatograph fitted with a \(^{60}\)Ni electron capture detector (Dyson Instruments Limited, Tyne & Wear, UK) and linked to a Varian 4290 integrator (Varian Instrument Group, California, USA). The analytical column was a quartz capillary column (25 m \(x\) 0.25 mm id) coated with 0.2 \(\mu\)m SE-30 (Philips Analytical, Cambridge, UK). Operating temperatures for injector, column and detector were 225, 195 and 295°C respectively. High purity CP grade nitrogen (BOC Limited, London, UK) was used as the carrier gas at a flow rate of 1.0 ml min\(^{-1}\) and as the detector make up gas at a flow rate of 30 ml min\(^{-1}\). A sample injection volume of 1 \(\mu\)l was used and the split ratio was set at 1:40. Plasma drug concentrations were determined using an internal standard method based on peak areas and plotted as a function of time. Linear regression analysis of the terminal log-linear phase of the curve was used to determine the 1st order elimination rate constant \((k_d)\) and the terminal half-life \((t_{1/2})\) was calculated from the equation:

\[ t_{1/2} = \ln 2/k_d \]

The area under the plasma concentration versus time curve (AUC) was calculated from \(t = 0\) to the last measured time point \((t_e)\) using the trapezoidal rule. The remaining area from \(t_1\) to \(t_e\) was calculated using the equation \(C_t/k_d\) where \(C_t\) is the concentration at \(t\).

In vitro stability studies

The stability of Clomesone was assayed in complete RPMI 1640 tissue culture medium at 37°C at an initial drug concentration of 10 \(\mu\)g ml\(^{-1}\). Sample aliquots were removed at time intervals up to 7 h and extracted and analysed by gas chromatography as described for mouse plasma samples. The 1st order rate constant \((k)\) was obtained from a semi-logarithmic plot of concentration versus time and used to calculate the drug half-life. In this case the rate constant was used to construct the drug decay curves for the time course of the MTT assay of the initial concentrations required to achieve a 50% cell kill (IC\(_{50}\) values) for each tumour cell line using the equation:

\[ C_t = C_0e^{-kt} \]

Concentration-time products \((c \times t)\), a measure of the total drug exposure of the tumour cells in vitro, were then determined by calculating the areas under the decay curves using the trapezoidal rule.

Results

Table I shows the in vitro chemosensitivity of the panel of tumour cell lines to Clomesone. In general, the murine cell lines were more sensitive than the human cell lines. MAC 15A was the most sensitive with an IC\(_{50}\) value of 10.1 \(\mu\)g ml\(^{-1}\). The human leukaemia K562 and a colon cell line HCLO showed some sensitivity to Clomesone, but it was relatively inactive against the majority of human cell lines derived from solid tumours with IC\(_{50}\) values in excess of 100 \(\mu\)g ml\(^{-1}\).

The in vivo activity of Clomesone against the MAC tumour lines was compared with that of TCNU and Chlorozotocin (Tables II–IV). Full dose range data to the point of toxicity were obtained to assure that maximum tolerated doses were achieved for each compound and are presented in Table IV. Activity of Clomesone against each tumour line was reproduced in a separate experiment as was the activity of TCNU and Chlorozotocin against MAC 26. The values obtained for TCNU and Chlorozotocin against MAC 13 and MAC 15A were similar to those in previously reported studies (Bibby et al., 1988a; McElhinney et al., 1989).

Table II shows that Clomesone was active against the ascitic MAC 15A tumour at doses of 50 mg kg\(^{-1}\) and

| Table I | In vitro chemosensitivity to Clomesone |
|---------|--------------------------------------|
| **Cell line** | **IC\(_{50}\) \(\mu\)g ml\(^{-1}\)** | **c \(\times\) t** |
| (mean ± SD; \(n = 4\)) | (\(\mu\)g h ml\(^{-1}\)) |
| MAC 15A | 10.1 ± 1.5 | 54 |
| MAC 16 | 33.6 ± 12.5 | 177 |
| MAC 26 | 185 | |
| WEHI-3B | 284 ± 4.8 | 150 |
| K562 | 39.2 ± 14.6 | 207 |
| HCT-18 | >100 | >528 |
| HKT-18 | >100 | >528 |
| HCL0 | 146 | |
| DLD-1 | >100 | >528 |
| HT-29 | >100 | >528 |

*One experiment only.

| Table II | Anti-tumour activity against MAC 15A |
|---------|------------------------------------|
| **Compound** | **Dose** | **Route** | **T/C%** |
| (mg kg\(^{-1}\)) | (mg kg\(^{-1}\)) | | |
| Clomesone | 100 | ip | 133, 163 | |
| | 50 | ip | 163 | |
| | 50 | iv | 125 | |
| | 25 | ip | 138 | |
| | 25 | iv | 100 | |
| | 12.5 | ip | 88 | |
| | 12.5 | iv | 100 | |
| | 30 | ip | 200 | |
| TCNU | 60 | ip | 288 | |
| Chlorozotocin | 60 | ip | 288 | |

*Maximum tolerated dose. *Two independent experiments.

| Table III | Anti-tumour activity against MAC 13 |
|---------|-----------------------------------|
| **Compound** | **Dose** | **Route** | **% Tumour weight inhibition** |
| (mg kg\(^{-1}\)) | (mg kg\(^{-1}\)) | | |
| Clomesone | 100 | ip | 48.3, 45.0 | |
| | 50 | ip | 33.8, 35.1 | |
| | 25 | ip | 47.3 | |
| | 12.5 | ip | 13.5 | |
| | 25 | iv | 21.6 | |
| | 12.5 | iv | 45.9 | |
| TCNU | 30 | ip | 91.3 | |
| Chlorozotocin | 60 | ip | 59.2 | |

*Maximum tolerated dose. *Two independent experiments.
100 mg kg⁻¹ ip. The increases in median survival times were significant at the 1% level when compared to the control group using the Mann-Whitney U test. However, at the maximum tolerated doses the activity of Clomesone was inferior to that of TCNU and Chlorozotocin. Anti-tumour activity against the relatively nitrosourea-sensitive solid sc MAC 13 tumour is presented in Table III. Although the tumour weight inhibitions achieved by Clomesone at 100 mg kg⁻¹ ip and 50 mg kg⁻¹ ip or iv were significant (P = 0.05; student’s t-test), they were less than that achieved by TCNU. Clomesone was inactive against the relatively nitrosourea-resistant solid sc MAC 26 tumour as described in Table IV. The relative tumour volumes of the group treated with the maximum tolerated dose of 100 mg kg⁻¹ ip were not significantly different (P > 0.1) from those of the control group when analysed using the student’s t-test.

The spleen colony forming unit assay calibration curve is shown in Figure 2. Linear regression analysis showed a strong positive correlation (r = 0.997) between the number of surface spleen colonies and the number of marrow cells inoculated. Linearity was observed over the range 1.2 x 10⁷–7.4 x 10⁸ cells injected. A cell inoculum below 1.2 x 10⁷ produced no visible surface colonies while a cell inoculum of 3.4 x 10⁷ resulted in the total repopulation of the spleen with no discrete colony formation.

Acute bone marrow toxicity following ip drug administration is described in Table V. The results at the maximum tolerated dose for each drug represent the survival fractions obtained in two independent experiments. Clomesone was less myelosuppressive than Chlorozotocin but still produced a 5-fold reduction in colony forming units at the maximum tolerated dose of 100 mg kg⁻¹.

The solvent extraction procedure used in the pharmacokinetic studies gave recoveries for six replicate plasma samples, at a concentration of 1 μg ml⁻¹, of 98.8% for Clomesone and 88.9% for the internal standard, 2-chloroethyl-p-toluenesulphonate with an overall intersample variation for the assay of 9.4%. Adequate separation of Clomesone and the internal standard from plasma interferents was achieved under the assay operating conditions and a typical chromatogram is presented in Figure 3(a). No underlying plasma interfering peaks co-eluted with Clomesone as shown in Figure 3(b). The limit of detection for Clomesone in plasma was 0.05 μg ml⁻¹. A drug calibration curve was constructed using least squares linear regression analysis and linearity was observed in the concentration range 0.05–10 μg ml⁻¹ (r = 0.987).

The plasma concentration of Clomesone following ip administration of 50 mg kg⁻¹ as a function of time is shown in Figure 4. Each point is the mean of duplicate samples and each curve represents an independent experiment. Peak plasma levels were reached within 2 min of drug administration and ranged from 18.1–35.9 μg ml⁻¹ with a mean value of 28.0 μg ml⁻¹ while the mean terminal half-life was 8.1 min (range 4.8–10.8 min). The areas under the plasma concentra-

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**Table IV** Anti-tumour activity against MAC 26 following ip administration

| Compound     | Dose (mg kg⁻¹) | Survivors | Tumour growth delay (days) |
|--------------|---------------|-----------|---------------------------|
| Clomesone    | 200           | 0/10      | Toxic                      |
|              | 150           | 0/10      | Toxic                      |
|              | 100           | 10/10     | 0.8, 0.8³                  |
|              | 50            | 10/10     | 0                         |
|              | 25            | 10/10     | 0                         |
| TCNU         | 50            | 0/5       | Toxic                      |
|              | 30³           | 10/10     | 11.6, 12.4³                |
| Chlorozotocin| 80            | 3/9       | Toxic                      |
|              | 60³           | 9/9       | 0, 0³                      |

*Maximum tolerated dose, bTwo independent experiments.

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**Table V** Bone marrow toxicity following ip administration

| Compound     | Dose (mg kg⁻¹) | No. of colonies observed (mean ± SD; n = 6) | Survival fraction |
|--------------|---------------|--------------------------------------------|-------------------|
| Clomesone    | 100³          | 4.8 ± 0.75                                  | 0.20              |
|              | 100³          | 6.5 ± 2.0                                   | 0.16              |
|              | 50            | 11.8 ± 1.17                                 | 0.59              |
|              | 25            | 17.8 ± 0.84                                 | 0.80              |
| TCNU         | 30³           | 0                                           | 0.20              |
|              | 30³           | 6.0 ± 0.9                                   | 0.01              |
| Chlorozotocin| 60³           | 21.4 ± 1.14                                 | 0.80              |

*Maximum tolerated dose, two independent experiments.

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Figure 2 Spleen colony forming unit assay calibration curve (points are mean ± SD; n = 6).

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Discussion

Examination of the in vitro activity of Clomesone in a panel of established murine and human tumour cell lines revealed that, in general, the human cell lines were insensitive. The colon adenocarcinoma HC10 and the leukaemia K562 were the only human cell lines that were sensitive at similar IC₅₀ values. Similarly, in the murine cell lines the concentration required for a 50% cell kill of the line derived from the relatively nitrosourea-resistant MAC 26 tumour was of the
same order of magnitude as that required by the leukaemia WEHI-3B. These findings suggested that Clomesone had no preferential specificity for cell lines derived from solid tumours.

The lack of selectivity was confirmed by the results of the in vivo anti-tumour activity and toxicity studies. The activity of Clomesone in the MAC tumour system in vivo was not impressive. Moderate activity was exhibited against the ascitic MAC 15A. The solid sc MAC 13 did respond to Clomesone over a broad dose range but the maximum anti-tumour activity exhibited was not great in this relatively nitrosourea-sensitive tumour. No significant activity was demonstrated against the relatively nitrosourea-resistant solid sc MAC 26 tumour. Thus, the anti-tumour activity of Clomesone was similar, if slightly inferior, to that of the non-carbamoylating nitrosourea, Chlorozotocin, and it was much less effective than the carbamoylating nitrosourea, TCNU, which remains the most active nitrosourea analogue in the MAC tumour system to date. Although Clomesone exhibited less bone marrow toxicity compared to both TCNU and Chlorozotocin, the reduction in colony forming units produced at the maximum tolerated dose was still marked. In addition, the dose of TCNU which produced a similar level of bone marrow toxicity has been reported to give a better response than Clomesone against the nitrosourea-sensitive MAC 13 tumour (Bibby et al., 1988a).

The development of the selective and sensitive method for the quantitative determination of Clomesone in mouse plasma allowed the pharmacokinetic behaviour of the drug to be characterised with a view to relating the activity-toxicity pattern to bioavailability. The AUC values for Clomesone following a single ip injection of 50 mg kg\(^{-1}\), when considered in conjunction with the in vitro c x t product for MAC 26, suggested that the inactivity in vivo was due to ineffective anti-neoplastic drug concentrations at the tumour site. A higher drug exposure of the tumour cells could be achieved in vitro as the drug half-life in tissue culture medium of 3.38 h (Chan & Barrientos, 1988). The dose of 50 mg kg\(^{-1}\) was chosen as the optimal dose as the highest maximum tolerated dose of 100 mg kg\(^{-1}\) resulted in no improvement in anti-tumour activity at the expense of increased.

Figure 3  a, Chromatogram of mouse plasma extract containing Clomesone and internal standard (1 = Clomesone; 2 = internal standard). b, Chromatogram of mouse plasma extract containing internal standard only (1 = position of Clomesone peak; 2 = internal standard).

Figure 4  Plasma concentration of Clomesone following administration of 50 mg kg\(^{-1}\)ip (points are mean values; n = 2; three independent experiments).
bone marrow toxicity.

Clomesone was selected for clinical trial on the basis that although it was only as effective as the chloroethylnitrosoureaes in some murine tumour models, its chemistry suggested that it would be more toxicologically selective. The broader pre-clinical evaluation performed in this study could find no evidence to support this theory. On the contrary, the findings suggested that the effectiveness of Clomesone, in common with the chloroethylnitrosoureaes, would be limited by myelosuppression. Whether the results of this study will be reflected in the clinical setting remains to be seen and the results of the recently completed Phase I clinical trial are awaited with great interest.

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