An in vitro study comparing the cytotoxicity of three platinum complexes with regard to the effect of thiol depletion

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Summary The cytotoxicity of three platinum complexes, cis-diamminedichloroplatinum(II) (cis-platin), cis-dichloro-trans-dihydroxy-cis-bis (isopropylamine) platinum(IV), (CHIPP) and diammine (1, 1-cyclobutane-dicarboxylato) platinum(II) (carboplatin) on Chinese Hamster ovary (CHO) and mouse sarcoma RIF-1 cells cultured in vitro has been compared.

The tumour cell line was much more sensitive to the cytotoxic action of the three agents compared to the CHO cell line. CHIP and carboplatin gave similar dose-response curves, both being much less toxic than cis-platin.

The effect of thiol modification on platinum toxicity was also investigated. Substantial reduction in the intracellular non-protein sulphydryl content markedly enhanced the cytotoxicity of CHIP but had much less effect on carboplatin and cis-platin. Thiol depletion by diethylmaleate had a negligible effect on cis-platin toxicity.

Cis-platin is an effective anti-tumour agent and is well established in the clinic (Gottlieb & Drewinko, 1975; Yagoda et al., 1976; Young et al., 1976), but has dose-limiting nephrotoxicity (Krackoff, 1979). In order to combat the side effects of cis-platin, second generation platinum complexes have been developed with much less toxicity to normal tissue (Harrap et al., 1980). Two such compounds, CHIP and carboplatin, have emerged for clinical studies. CHIP has reached phase II clinical trials (Wong et al., 1985), and carboplatin, phase III clinical trials (Canetta et al., 1985).

Manipulation of intracellular thiols may be useful clinically in either enhancing drug toxicity to the tumour or by protecting against normal tissue damage to increase the therapeutic gain. There are numerous reports that thiols can protect animals and cells against the toxicity of alkylating agents (Connors et al., 1964; Connors, 1966; Contractor, 1963; Walker & Smith, 1969).

Glutathione (GSH) is the major intracellular non-protein sulphydryl (NPSH) in living cells and is generally accepted to play a mechanistic role in cancer treatment (Arrick & Nathan, 1984), the remaining component of NPSH is comprised of cysteine, γ-glutamylcysteine, lipoic acid and co-enzyme A. GSH can interact with a variety of anti-neoplastic drugs, mainly through nucleophilic thioether formation or oxidation-reduction reactions. To examine the effect of thiol depletion on platinum sensitivity we have used diethylmaleate (DEM) or DL-buthionine-SR-sulphoximine (BSO) prior to platinum treatment. BSO is a specific inhibitor of glutathione biosynthesis (Griffith & Meister, 1979) and over 90% reduction in glutathione levels can be achieved in cell culture. DEM, on the other hand, is less specific and reacts with non-protein thiols to form a stable adduct (Bump et al., 1982).

Materials and methods

Cell culture

Cells were routinely cultured as a monolayer in Heps-buffered MEM plus 15% foetal calf serum supplemented with non-essential amino acids and glutamine (Gibco). One hundred units ml⁻¹ of penicillin/streptomycin were added to the RIF-1 cultures. CHO and RIF-1 cells were grown in plastic tissue culture flasks (Nunc.) at 37°C. The RIF-1 cultures were incubated in a humidified atmosphere containing 5% CO₂.

Drug treatments

Monolayers of cells were treated with fresh medium containing the required drug concentration for one hour at 37°C. The drug was then removed and the cells were made into a suspension using 0.01% trypsin in 0.02% EDTA for CHO cells, and 1 mg ml⁻¹ protease (type IX Sigma Chem. Co.) in complete medium for RIF-1 cells. Appropriate cell numbers were reseeded into 25 cm² tissue culture flasks and incubated for either 6 days for CHO cells or 8 days for RIF-1 cells. Colonies were stained with carbol fuchsin and the surviving fraction determined by comparing the colonies counted in drug-treated to untreated controls. The plating efficiencies for the CHO and RIF-1 cell lines were approximately 90% and 40% respectively.

Drugs

Cis-platin (Johnson Mathey), CHIP (Bristol Myers) and carboplatin (Bristol Myers) were all freshly dissolved in saline prior to use.

BSO was kindly supplied by Dr R.W. Middleton, Brunel University and the Gray Laboratory of the Cancer Research Campaign, and was dissolved in saline.

DEM (Sigma Chem. Co.) was initially diluted in absolute alcohol at 100 mM and subsequently diluted in saline.

NPSH determination

Ice-cold trichloroacetic acid was used to precipitate protein and the supernatant was assayed spectrophotometrically for NPSH using Ellmans reagent (1959). NPSH was measured in control CHO and RIF-1 cells and also after 1 hour's treatment at 37°C with a platinum dose sufficient to reduce cell survival to 10%.

GSH determination

This was assayed spectrophotometrically by the procedure developed by Tietze (1969).

Results

Platinum toxicity

Figures 1, 2 and 3 show the drug dose responses obtained with cis-platin, CHIP and carboplatin on CHO and RIF-1 cells cultured in vitro. CHIP and carboplatin are very similar in toxicity and are ~40 times less toxic than cis-platin towards CHO cells. The RIF-1 cells are much more sensitive to all three platinum agents than are CHO cells and the
Effect of thiol depletion on platinum toxicity

Monolayers of CHO cells were treated with 0.2 mM DEM for 2 h at 37°C prior to platinum exposure for a further 1 h at 37°C. Under these conditions the NPSH content of CHO cells was reduced to a level such that it was no longer detectable. DEM pretreatment did not alter the cellular response to cis-platin (Figure 4) and potentiated carboplatin toxicity by reduction of the shoulder. At very high carboplatin doses where the survival curve begins to plateau, the effect of thiol depletion was minimal (Figure 5). The effect of thiol depletion by DEM pretreatment was much more pronounced on CHIP sensitivity, reducing the D₀ from 200 μM to 83 μM (Figure 6). For comparison CHO cells were also treated with 100 μM BSO for 24 h prior to platinum.

shoulders of the RIF survival curves were almost absent with CHIP and carboplatin and completely absent with cis-platin.

Figure 1 Dose response survival curves for CHO (●) and RIF-1 cells (○) treated with cis-platin for 1 h at 37°C. Data points represent the mean of 3 or more experiments with the range of values indicated.

Figure 2 Dose response survival curves for CHO (●) and RIF-1 cells (○) treated with CHIP for 1 h at 37°C. Data points represent the mean of 3 or more experiments with the range of values indicated.

Figure 3 Dose response survival curves for CHO (●) and RIF-1 cells (○) treated with carboplatin for 1 h at 37°C. Data points represent the mean of 3 or more experiments with the range of values indicated.

Figure 4 Cis-platin response of CHO cells after thiol depletion either by BSO (●) or DEM (×). Data points (●) for the cis-platin control curve are as for Figure 1. Individual data points from several experiments on thiol-depleted cells are shown.
exposure. The results were similar to that obtained after DEM pretreatment except for cis-platin (Figure 4) where a modest degree of enhancement was observed (a dose modification of 1.35).

Intracellular thiol levels

The GSH and NPSH contents of CHO and RIF-1 cells are shown in Table I. The levels were not significantly altered after platinum treatment.

Discussion

Carboplatin and CHIP are much less toxic in vitro on both the normal and tumour cell lines, when compared with the parent compound, cis-platin. In agreement with Ohnuma et al. (1986) we found no effect of cell density on the cytotoxicity of cis-platin or carboplatin. However, we did find CHIP response varied significantly with cell density, i.e. CHIP became more cytotoxic with a decrease in cell density (data not shown). Despite the fact that CHIP and carboplatin are quite different in chemical structure, they exhibit similar dose-response curves. The absence of the shoulder on the RIF survival response may indicate that the tumour line is less able or unable to repair sub-lethal damage induced by platinum complexes. The greater sensitivity of RIF-1 cells to platinum agents compared to CHO cells could be due to differences in drug uptake (Eicholtz-Wirth & Hietel, 1986), increased binding of platinum to DNA in the sensitive line (Roberts & Fraval, 1978) or the ability to excise platinum adducts from DNA (Fraval & Roberts, 1979). Cis-platin (Roberts & Fraval, 1978) and CHIP (Edwards & Nias, 1986) have both been shown to be preferentially cytotoxic towards cells in the $G_1$ phase of the cell cycle and therefore cell lines with longer $G_1$ phases may be more sensitive to platinum treatment than cells with shorter $G_1$ times. This agrees with the cell cycle time of 24 hours in the RIF-1 compared to 13-14 hours in the CHO cells assuming cells with longer cell cycles have longer $G_1$ phases.

Other workers have looked at the effect of platinum toxicity on sensitive and resistant human carcinoma cell lines in vitro in relation to the intracellular glutathione content, and found in some instances that resistance to platinum agents correlated with higher levels of glutathione (Wolf et al., 1985; Hamilton et al., 1985; Behrens et al., 1985). However, Andrews et al. (1985) showed that their platinum resistant line had the same glutathione content as the sensitive cell line. Our work indicates that the RIF-1 cells have a similar glutathione content compared to CHO cells. Many investigators are now looking at intracellular thiol levels with respect to drug toxicity in different cellular systems. Care must be taken when comparing results from

Figure 5 Carboplatin response of CHO cells after thiol depletion either by BSO (■) or DEM (×). Data points represent the mean of 3 or more experiments with the range of values indicated.

Figure 6 CHIP response of CHO cells after thiol depletion either by BSO (■) or DEM (×). Data points represent the mean of 3 or more experiments with the range of values indicated.

Table I Intracellular GSH and NPSH concentrations after thiol depletion.

| Cell line | Treatment | GSH content $\times 10^{-6}$ nmol cell$^{-1}$ | NPSH content $\times 10^{-6}$ nmol cell$^{-1}$ |
|-----------|-----------|---------------------------------|-----------------------------------|
| CHO       | control   | 8.1 ± 1.8$^a$                   | 10.8 ± 1.16$^b$                   |
|           | after BSO | 0.12$^b$                        | non detectable                    |
|           | after DEM | 0.2$^b$                         | non detectable                    |
| RIF       | control   | 8.0 ± 1.1$^a$                   | 13.8 ± 0.45$^b$                   |
|           | after BSO | 0.32$^b$                        |                                   |

$^a$S.d.

$^b$Denotes the mean of 2 determinations.
different cell lines and different laboratories because Post et al. (1983) have shown that GSH content varies with time after passage and serum concentration in cell culture conditions. In our studies the RIF-1 and CHO cells were both cultured in the same serum concentration and were assayed for GSH levels 48 h after passage.

Although DEM and BSO reduce glutathione levels by more than 90% only BSO pretreatment was able to modify the cis-platin response. The reason for this could be that BSO is more efficient than DEM at removing glutathione from platinum sensitive sites.

Of the three platinum drugs only CHIP was substantially affected by thiol depletion in CHO cells (ER = 2.4). A similar result was also confirmed with the RIF-1 cells. Recent data using synchronised CHO cells have shown that the cell cycle dependence of CHIP is removed if the cells are pretreated with BSO (Edwards, 1986). The higher oxidative state of CHIP compared to cis-platin and carboplatin may be a contributory factor in its modification by thiol. It has been proposed that CHIP requires intracellular reduction to a Pt II complex for activity (Blatter et al., 1984), also that CHIP can be reduced by oxidised DNA (Butler et al., 1985). However, both these processes are less likely to occur in thiol depleted cells. To add further controversy Andrews et al. (1985) found no effect on CHIP toxicity in human ovarian carcinoma cells which had been previously treated with BSO. The carcinoma cell line was 10 times more sensitive to cis-platin than RIF-1 cells and therefore potentiation of CHIP by BSO might depend on the inherent sensitivity of the cell line to CHIP such that extremely sensitive cell lines may not be made more sensitive to CHIP by thiol depletion.

It has been shown that thiourea can reverse Pt(II) induced DNA cross-links and lethal lesions in isolated DNA (Filipski et al., 1979), and that thiourea can protect cells against the toxicity of platinum II complexes (Zwelling et al., 1979). We have also shown that thiourea can protect against the toxicity of cis-platin, CHIP and carboplatin (unpublished data). Naturally occurring thiols inside the cell may also be able to protect against platinum toxicity in a similar fashion to thiourea but to a lesser degree. Thus reduction of intracellular glutathione located near Pt-DNA adducts may be responsible for enhanced platinum toxicity.

In conclusion, RIF-1 cells did not differ appreciably in their GSH or NPSH content from CHO cells and yet they are much more sensitive to platinum toxicity. The concentration of NPSH in specific cell organelles may be more important than the total cellular NPSH with regard to platinum toxicity. Cellular sensitivity to Pt IV complexes can be increased significantly in thiol depleted cells. Thiol depletion has little effect on Pt II complexes. As with all chemotherapeutic drugs, effective therapy is marred by dose-limiting toxicities and the emergence of drug resistance by the tumour. The use of biochemical modifiers that can either sensitize tumour cells, or minimise drug resistance, may improve the efficacy of platinating agents. BSO is a relatively non-toxic compound and its use in combined modality treatments may be of some clinical benefit.

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