Radiation-induced transient cisplatin resistance in murine fibrosarcoma cells associated with elevated metallothionein content

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Summary Cisplatin resistant mouse fibrosarcoma cells were isolated after fractionated irradiation in the absence of any drug treatment. Several sublines have been established; clone SSK-rad, was used for further studies. These cells exhibit unchanged radiosensitivity and are compared to cisplatin resistant sublines, SSK-cis, previously induced by low dose cisplatin exposure. Both resistant sublines are characterised by reduced CdCl₂ sensitivity, indicating enhanced metallothionein content; loss of cisplatin resistance occurs after 10 to 25 generations and correlates with rising CdCl₂ toxicity. Increase of MT is demonstrated directly by ¹⁰⁹Cd binding to the MT containing region after FPLC. Both sublines differ in GSH level, which is increased only in SSK-rad, cells, and in cellular platinum content, which is reduced in SSK-cis₂ cells compared to the parental SSK cell line. These factors may contribute to cisplatin resistance but are not the main cause responsible for the transient nature of the drug resistance observed.

Our results indicate that transient cisplatin resistance in SSK cells can be induced not only by the drug itself but also by γ-irradiation and is based on the same mechanism of increased cellular MT content.

Resistance in cancer therapy usually means chemoresistance following drug treatment. However, resistance can also be induced by irradiation (Hill et al., 1988; 1990b; Osmak & Petrovic, 1989). Since tumour therapy often includes both sequential or concomitant radio-chemotherapy, it is important to study the mechanisms leading to radiation-induced resistance or cross-resistance.

Most of the in vitro studies on drug and radiation interaction have looked into cross-resistance after drug induction and the results are rather controversial. Findings range from no effect (Wallner & Li, 1987; Eichholtz-Wirth et al., 1993; Mitchell et al., 1988) to cross-resistance (Schwartz et al., 1988; Louie et al., 1985) and even to collateral sensitivity (Hill et al., 1988). Lehner et al. (1989 and 1990) discuss the contribution of GSH and its related enzymes to the radiation response in drug resistant human tumour cells. They show that two drug resistant sublines with different underlying mechanisms are both radioresistant and only these sublines are radiosensitised by GSH depletion.

There are only few publications concerned with radiation induced drug resistance. They demonstrate altered responsiveness to different classes of antineoplastic agents, such as the topoisomerase II inhibitor VP-16 (Lock & Hill, 1988), cisplatin (Dempke et al., 1992; Hill et al., 1990c; Osmak & Petrovic, 1989), the antimetabolic drug methotrexate (Sharma & Schimke, 1989) as well as the development of multi-drug-resistance (Mattern et al., 1991; Hill et al., 1990a).

We have previously described cisplatin resistance in a murine fibrosarcoma cell line (SSK-cis), induced by intermittent low dose cisplatin treatment (Eichholtz-Wirth et al., 1993). This drug resistance was transient and was characterised mainly by elevated metallothionein content.

In the present study we report on cisplatin resistant SSK-rad cells that were isolated after fractionated irradiation without any prior drug treatment. The following experiments were performed to identify the mechanisms underlying this radiation induced drug resistance and to compare them to those after cisplatin induction. The data could help to clarify whether the way of generating resistance determines the operating mechanism of resistance, as suggested by Hill et al. (1990b).

Materials and methods

Cell lines and induction of resistance

Mouse fibrosarcoma cells (SSK) were grown as monolayer culture in Eagle's minimal essential medium (MEM), supplemented with 10% calf serum, 0.01% neomycin, and 0.035% NaHCO₃ in a humidified CO₂ incubator at pH 7.4 and 37°C. Their mean doubling time was 12 h.

About 10⁵ cells were exposed to fractionated γ-irradiation (6 Gy per fraction), using a gamma cell 40 caesium-137 source (AECL-Industria, Canada) at a dose rate of 1.2 Gy min⁻¹. After each fraction the cells were allowed to grow until confluency at which time irradiation was repeated up to a total dose of 90 Gy. Twelve clones, denoted SSK-rad, were isolated. These clones exhibited cisplatin resistance. Clone SSK-rad, was used for all experiments. SSK-rad, cells are more elongated than the parental cells but they didn’t differ in cell size. Their colonies exhibit more fascicular growth whereas the SSK colonies rather pile up in the centre.

Induction of cisplatin resistance by drug treatment has been described elsewhere (Eichholtz-Wirth et al., 1992). Briefly, cisplatin resistance was generated by exposure of 10⁵ SSK cells to 10 μg ml⁻¹ cisplatin for 1 h. Subsequently, the cells were allowed to grow to confluency at which time the treatment was repeated. After only 3–5 treatment cycles, resistant clones, designated SSK-cis, were isolated and subcultured in drug free medium. In this study clone SSK-cis₂ was used which is identical to SSK-R2, previously characterised. The drug induced cisplatin resistant cells are denoted now SSK-cis, to differentiate from the radiation induced cisplatin resistant cells SSK-rad. They have similar growth characteristics as the SSK-rad, cells.

Measurement of drug and radiation sensitivity

Exponentially growing cells of SSK-rad and SSK-cis clones were subcultured, appropriately diluted and allowed to attach to the glass surface overnight. Exposure to cisplatin (cisplatin solution, Behring, Marburg, Germany) was carried out in culture medium for 1 h at different drug concentrations. The drug was diluted in Hanks' solution and added to the culture medium (maximum cell number of 10,000). After the allotted exposure time the medium was decanted, the cells rinsed with Hanks’ solution and fresh culture medium was added. For cadmium chloride (CdCl₂) toxicity studies, cells were
exposed for 1 h to various concentrations of CdCl₂.

For radiation treatment, cells were exposed to graded single doses of γ-rays from a gamma cell 40 caesium-137 source at a dose rate of 1.2 Gy min⁻¹.

Following any of the indicated treatments, cells were incubated for either 8 days (SSK cells) or 10 days (SSK-rad and SSK-cis sublines). The colonies were then stained with methylene blue and those containing more than 50 cells were counted. The surviving fraction (SF) was calculated from the ratio of mean colony yield of treated to untreated cells. All experiments were carried out with four replicate bottles and repeated at least three times. Experimental data were accepted if the colony forming efficiency of the untreated cells was higher than 35%.

Resistance factor (R₀) was defined as the ratio of drug doses, D₀, of resistant over parent SSK cells.

**Cellular cisplatin concentration**

The cellular concentration of platinum was determined with proton-induced characteristic X-ray emission (PIXE) as described in detail by Eichholtz-Wirth et al. (1990 and 1992). After exposure of 10⁶ cells to 0, 10, 20 and 40 µg ml⁻¹ cisplatin for 1 h, cellular platinum content was determined. Regression lines were calculated (correlation coefficient >0.99) and the cellular platinum content compared.

**GSH and protein determination**

Cells in the logarithmic growth phase were used for GSH determination according to Tietze (1969). GSH-S-transferase (GST) was assayed according to the method of Habig et al. (1974), using 1-chloro-2,4-dinitrobenzene (CDNB) as substrate. GST activity is expressed as nmol GSH-CDNB conjugate formed per min mg⁻¹ protein. For protein determination the Lowry assay was used (Lowry et al., 1951).

**Metallothionein test**

Cells were tested for metallothionein content by binding of radioactive ¹⁰⁹Cd to the cytosol followed by FPLC. An aliquot of 5 × 10⁶ exponentially growing SSK or SSK-rad cells were harvested by trypsinization and washed twice with PBS (4°C). The cells were disrupted by sonification and then centrifuged (20,000 g for 20 min). The supernatant was incubated with trace amounts of ¹⁰⁹Cd for 30 min at 37°C. Thereafter the solution was centrifuged (12,000 g, 2 min) and filtered through a 0.45 µm membrane filter. Aliquots of the filtrate were injected to FPLC (gelfiltration on a Superose-12 column (300 × 10 mm), using 0.05 m sodium-phosphate pH 7.0, 0.15 m sodium chloride as eluent at a flow rate of 0.5 ml min⁻¹. Fractions of 0.5 ml were collected and measured for radioactivity. Cadmium binding metallothionein was identified by comparison with rabbit liver metallothionein (Sigma).

**Results**

SSK-rad₁ clones have almost the same radiosensitivity as the parental SSK cells as well as the drug induced cisplatin resistant subline SSK-cis₂ (Figure 1). If survival is measured as a function of cisplatin concentration after a 1 h drug exposure, SSK-rad₁ cells exhibit a distinctly enhanced drug resistance. Resistance is only slightly less than that observed in the SSK-cis₂ cells as indicated by the R₀ values of 5.8 and 7.8 respectively (Figure 2). Twelve clones were isolated that were drug resistant; however, in this study the induction rate of drug resistance after irradiation was not determined systematically. The degree of cisplatin resistance could not be enhanced, if the resistant clones were given three additional radiation fractions (five clones tested). For all further experiments, the same SSK-rad₁, clone was used.

Growth characteristics of both resistant sublines are similar, colony forming efficiencies are between 60–90% as in the parental SSK cells. Both sublines differ from SSK cells by longer doubling times of 15–18 h as compared to 11–13 h in the SSK cells (Table I). There is no cross resistance to other cytostatic agents such as doxorubicin, vinblastin or melphalan (Table II). Protein content is not significantly different in all cell lines, whereas cellular drug content is reduced in SSK-cis₂ cells but not in SSK-rad₁ cells (Table I).

Since thiols were shown to play an important role for cisplatin detoxification in SSK-cis cells, non-protein and protein thiols were determined in SSK-rad₁ cells. GSH is increased by a factor of 1.9 in SSK-rad₁ cells compared to the parental SSK cells but is in the normal range in SSK-cis₂ cells (Table I). When resistance is lost the enhanced total GSH level is not significantly changed. GST is slightly increased in both SSK-rad₁ and SSK-cis₂ cells. For SSK-cis...
Table I Characteristics of the sensitive SSK cells and the resistant sublines SSK-rad, and SSK-cis<sub>2</sub>

| Parameter | SSK | SSK-rad<sub>1</sub> | SSK-cis<sub>2</sub> |
|-----------|-----|-------------------|-------------------|
| Doubling time (h)<sup>a</sup> | 11–13 | 15–18* | 15–18 |
| Protein content<sup>b</sup> (μg (10<sup>6</sup> cells)<sup>-1</sup>) | 161±8 | 148±12* | 153±9* |
| Total GSH<sup>c</sup> (nmol (mg protein)<sup>-1</sup>) | 14.5±1.2 | 27±2.6** | 16.5±2.2* |
| GST<sup>c</sup> (nmol (DNB min mg protein)<sup>-1</sup>) | 63±8 | 92±10** | 98±6*** |
| Cellular drug content<sup>d</sup> (ng (10<sup>6</sup> cells)<sup>-1</sup>) | 1.6 | 1.5 | 1.0 |

<sup>a</sup>Calculated from the exponential portion of the growth curve; <sup>b</sup>Means ± s.d. from three separate experiments; <sup>c</sup>Data are derived from the regression curves of cellular platinum content at 20 μg ml<sup>-1</sup> at 1 h exposure time. **P<0.01 as compared to the data of the SSK cells; *P<0.05 as compared to the data of the SSK cells. *n.s.

Table II Comparison of drug resistance of parental SSK cells and cisplatin resistant SSK-rad, and SSK-cis<sub>2</sub> cells

| Cytostatic drug | SSK | SSK-rad<sub>1</sub> | SSK-cis<sub>2</sub> |
|----------------|-----|-------------------|-------------------|
| Melphalan      | 3.6 | 3.8               | 3.9               |
| Doxorubicin    | 6.4 | 5.6               | 5.8               |
| Vinblastin     | 4.2 | 4.8               | 5.0               |

<sup>a</sup>C<sub>50</sub>, drug concentration necessary to reduce cell survival to 10% after 1 h drug exposure; data are derived from the survival curves and are expressed as μg ml<sup>-1</sup>.

cells, reduced CdCl<sub>2</sub> toxicity, an indirect measure of MT, was shown to correlate with drug resistance and to increase, when the transient drug resistance was lost. Therefore it was tested whether the same mechanism is also responsible for cisplatin resistance in SSK-rad<sub>1</sub> cells. Figure 3 demonstrates that CdCl<sub>2</sub> toxicity is reduced to almost the same extent in SSK-rad<sub>1</sub> cells as in SSK-cis<sub>2</sub> cells. The CdCl<sub>2</sub> concentration necessary to reduce survival to 10% is by a factor of 2.0 (SSK-rad<sub>1</sub> cells) and 2.2 (SSK-cis<sub>2</sub> cells) higher in the resistant cells than in the parental SSK cell lines. After 10–20 passages, cisplatin resistance decreases in SSK-rad<sub>1</sub> cells. Figure 4 demonstrates the loss of drug resistance between passages number 9 and 13. This loss of drug resistance coincides with an increase in CdCl<sub>2</sub> toxicity (Figure 5). Various SSK-rad<sub>1</sub> clones that were isolated and tested had the same characteristics as described above; only the time they retained their cisplatin resistance differed, lasting from only 10 passages (as shown in Figures 4 and 5) to 25–30, corresponding to 120–150 cell cycles. Five passages after loss of cisplatin resistance, revertant cell lines did not retain the same cisplatin sensitivity as the parental cells (R<sub>v</sub> = 1.6); doubling times remained longer. The revertants were then no longer followed up.

Increased metallothionein (MT) content is measured directly by specific binding of trace amounts of <sup>109</sup>Cd to the cytosolic fractions (Figure 6). In this semiquantitative assay, fraction numbers 31 to 33 correspond to the MT region (molecular weight of 6,000–10,000). These fractions contain 27% of the cumulative activity in the SSK-rad<sub>1</sub> cells and only 11% in the parental SSK cells. In contrast, in the GSH region (fractions 36–38), the <sup>109</sup>Cd activity is not significantly increased in the SSK-rad<sub>1</sub> cells as compared to the parental cells. As a result of these changes, SSK-rad<sub>1</sub> cells contain only 17% of the <sup>109</sup>Cd activity in the high molecular weight region but the amount is 30% in the parental SSK cells.

**Discussion**

In this study we have demonstrated that cisplatin resistance can be induced in murine fibrosarcoma cells in vitro by...
ionising irradiation without prior drug treatment. This acquired drug resistance is not associated with a decrease in radiosensitivity. The mechanisms leading to transient cisplatin resistance are similar to those described after cisplatin induced resistance (Eichholtz-Wirth et al., 1993). Both in subclones SSK-rad, and SSK-cis, several mechanisms may contribute to cisplatin resistance: however, the main factor correlating with the development and loss of cisplatin resistance is the cellular content of metallothioneins.

These enhanced MTs do not confer radiation resistance, although they are suggested to play a role in radiation protection by scavenging hydroxyl and superoxide radicals (Thornalley & Vasak, 1985). Matsubara et al. (1987) showed that the induction of MTs in mouse liver is a significant factor for radiation protection. Also, Hodgkiess (1990) described higher endogenous levels of protein and non-protein thiols in irradiated cells which might reduce the efficacy of radiation itself. Similar to our findings, he observed that the resistant phenotypes can persist through many cell generations in the absence of selection pressure but eventually revert to the same phenotype as the unirradiated population. This suggests that there is no classical gene mutation. However, he did not correlate the increased thiols to the cytotoxic effect of chemotherapeutic agents. In both of our resistant SSK-rad, and SSK-cis, clones, increased MT content also lasts for a limited number of generations and it is associated with transient cisplatin resistance, but not with radioresistance. This is independent of the selection procedure. As hypothesised by Kaina et al. (1990), MTs are required to be in close proximity to the DNA in order to neutralise free radicals in the nucleus and to be an efficient radical scavenger. These authors conclude, - which may be confirmed by our data --, that MT concentrations in the nucleus are probably insufficient for radiation protection. Our results also correspond to those of Miura and Sasaki (1990), using mouse squamous carcinoma cells, demonstrating that the MT level does not determine intrinsic radiosensitivity. These authors also confirm that the cytotoxic effect of Cd is correlated to the MT content; however, they did not compare the MT level and the cellular sensitivity to cisplatin. Data of Kelley et al. (1988) also indicate that cells with acquired resistance to cisplatin frequently have an increased MT level and overexpress MT mRNA. There is no evidence for gene amplification, suggesting enhanced rate of gene transcription or increased mRNA stability. Reversal of cisplatin resistance is also accompanied by a decrease in MT content.
Comparison of the two differently derived sublines SSK-rad and SSK-cis shows that only SSK-rad cells exhibit elevated levels of total GSH content which remain elevated also after loss of drug resistance. Moreover, the importance of MTS relative to GSH content for cisplatin resistance is demonstrated by FPLC: the 106Cd activity was unchanged in the GSH fraction but it was 2.5 times higher in the MT region in SSK-rad cells compared to the parental SSK cells. SSK-cis cells derived from SSK-rad cells by reduced cellular platinum content, which was also unchanged upon loss of cisplatin resistance (Eichholtz-Wirth et al., 1993). These factors – increased GSH in SSK-rad cells and reduced cellular platinum content in SSK-cis cells – may contribute to cisplatin resistance in the two sublines, showing that the mechanisms involved in cisplatin resistance are multifactorial. However, the dominating factor that correlates with the transient nature of cisplatin resistance is the elevation of the MTS and this is demonstrated for both sublines.

This is also stressed by our data on cross resistance to doxorubicin, melphanal and radiation. Cross-resistance to these agents and radiation is reported mainly in cells with altered Pt-DNA binding, reduced cross-links and elevated GSH levels or GSH-dependent enzymes (Hamilton et al., 1985; Hospers et al., 1988). Since there is no cross resistance in SSK-cis and SSK-rad cells, this would also suggest other mechanisms responsible for the acquisition of cisplatin resistance in SSK cells.

Resistance has been demonstrated to be multifactorial not only for drug induced resistance but also for radiation induced cisplatin resistance, as also reported by Dempke et al. (1992). In their human ovarian cells, resistance was associated mainly with enhanced repair and increased tolerance of DNA damage; cisplatin uptake was decreased and cytotoxicity could be enhanced by verapamil, but not by inhibition of GSH with BSO. In our SSK cells, verapamil has no effect on SSK and SSK-cis2 cells (Eichholtz-Wirth, 1993).

whereas BSO treatment may be used to increase cisplatin toxicity for the sensitive and resistant cells (Eichholtz-Wirth, 1993; data for SSK-rad, cells not shown). DNA repair was not studied in our SSK cells.

Drug resistance after fractionated irradiation was described in a series of publications by the group of Hill et al. (1988–1990b) for various cellular systems. These authors propose that the patterns of response to anti-tumour drugs and the associated mechanisms differ depending on the agent employed to induce resistance. According to our results a similar pattern of resistance may rather develop in one model system, independent of the way how drug resistance was induced. This agrees with data on MTX resistance after radiation in different cells by Sharma and Schimke (1989). They suggest that different tumour cell types may have different propensities for developing MTX resistance by different mechanisms. Moreover, the degree of resistance, which is also supposed to correlate to the way of induction and which is usually enhanced upon increasing treatment doses, is only slightly lower in SSK-rad cells compared to SSK-cis2 cells and cannot be enhanced either by additional irradiation (SSK-rad cells) or by further drug exposure (SSK-cis2 cells).

In our study as well as in most of the published data cited above cisplatin resistance was generated after high dose fractionated radiation. It remains to be determined now whether this induction of cisplatin resistance is a general phenomenon also after a low total radiation dose as well as lower dose per fraction in vitro and in vivo. These data may have implications for combined modality therapy using sequential or simultaneous drug and radiation treatment.

Abbreviations: BSO, buthionine sulfoximine; CDNB, 1-chloro-2,4-dinitrobenzene; Cisplatin, cis-diaminedichloroplatinum (II); FPLC, fast protein liquid chromatography; GSH, glutathione; GST, glutathione S-transferase; MT, metallothionein; Rf, resistance factor; SF, surviving fraction.

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