Radiosensitization with Chemotherapeutic Agents and Hyperthermia: Effects on Linear-quadratic Parameters of Radiation Cell Survival Curves

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Abstract

The radio-sensitizing effects of several chemotherapeutic agents and hyperthermia have been investigated in several animal and human cell culture systems. Cells are first treated with Cisplatin, Gemcitabine, Halogenated pyrimidines or hyperthermia and thereafter irradiated with different dose of radiation up to 8 Gy. After treatment the clonogenic survival was determined and from the survival curves the values of the linear and quadratic parameters were determined using the formula S(D)/S(O) = exp(-(αD+βD²)).

An increase in the value of the linear parameter, α, was observed in most cases, which corresponds to an enhanced (potentially) direct lethal damage (PLD) at low doses. The quadratic parameter β, which is assumed to depend on the interaction of sublethal lesions (SLD), was rarely affected. Furthermore, it appeared that more radioresistant cell lines were more sensitised than the radiosensitive lines. Furthermore it can be concluded that radiosensitization is also dependent on cell cycle stage like plateau or exponentially growing phase or post treatment plating conditions.

Introduction

The radiosensitisation effects of chemotherapeutic agents and hyperthermia have been investigated in mammalian normal and cancer cell systems growing in vitro with respect to the linear quadratic parameters of dose-survival curve [1]. These treatment modalities are currently exploited in the clinic. The studies on the different human tumour cell lines show that a synergistic interaction can be obtained between chemotherapy, hyperthermia and radiation and that this interaction is more likely to occur in cell lines which are relatively sensitive to chemotherapy. The influence of modifying agents on radiation dose survival curves can adequately be analysed with the use of the linear-quadratic model: S(D)/S(O) = exp-(αD+βD²). The linear parameter, a, represents lethal damage from single particle events and describes the low dose area while the quadratic parameter, β, indicating sub lethal damage (SLD) dominates the effectiveness in the high dose region [2-5]. The radiation dose survival curves have been obtained by carrying out clonogenic assays and values of the linear and quadratic parameters have been calculated [6].

The linear-quadratic model is based on well accepted biophysical concepts, involving the assumption that lethal damage can be induced by single-particle tracks and by interaction of damage from multiple particles. The LQ-model does not have a current biological basis. However, it has been found to describe the low-dose region of the survival curves up to 6 Gy rather accurately. Furthermore the LQ-model has been shown to describe adequately dose fractionation effects for normal tissue tolerance and for experimental tumours. The LQ-model has also the advantage that it requires only two parameters to describe radiation dose-survival curves. It allows the separate analysis of changes in effectiveness in the low dose range, mainly determined by the linear term and in the high dose range determined mainly by the quadratic term [7,8]. An additional advantage of the LQ model is that its parameters can be discussed in terms of specific mechanisms of cell inactivation by radiation [2,3].

Materials and Methods

Cell cultures

Several different cell lines have been used in the studies. The human squamous lung carcinoma cell line SW-1573 was grown as a monolayer in L-15 medium (Invitrogen, Breda, The Netherlands) supplemented with 10% heat-inactivated fetal bovine serum (FBS) and penicillin, streptomycin and glutamine at 37°C without additional CO₂. The human ovarian carcinoma cell lines A2780 and its dFdC-resistant variant AG6000 were grown as monolayers in DMEM (Invitrogen) supplemented with 5% heat-inactivated FBS and with penicillin, streptomycin and glutamine at 37°C at 10% CO₂. The doubling time of the human cell lines in exponential growth is 22-24 h.

V79 (hamster fibroblast cells), RUCHI (Rat urether carcinoma) and R1 (Rat rhabdomyosarcoma) were grown as monolayers in minimal essential medium (Invitrogen) supplemented with 10% foetal bovine serum, glutamine and penicillin at 37°C at 2% CO₂ in exponentially growing - and in plateau-phase. The doubling time of these cells is about 14 h.

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Chemicals

Platinon® (cis-diaminedichloroplatinum II, cDDP, cisplatin) was provided by Pharmachemie (Haarlem, The Netherlands). Gemzar® (Gemcitabine, 2',2'-difluorodeoxycytidine, dFdC) was provided by Eli Lilly Inc. (Indianapolis, IN, USA). All other chemicals were of analytical grade and commercially available. For chemo-therapeutically induced radiosensitizing experiments cells were incubated with: 1) Medium containing cisplatin at a concentration of 1 or 5 µM for 1 h before irradiation or continuously (in this case cisplatin was present during the clonogenic assay). 2) Gemcitabine at a concentration of 10, 50 or 100 nM, which was added 24 hours before irradiation. Before the start of irradiation medium containing gemcitabine was removed, cells were washed twice with PBS and fresh medium was added. 3) Medium with Iodo-deoxyUridine (IdUrd). For these experiments cells were incubated for 72 h with 0 or 4 µM of IdUrd and 2.5 µM thymidine was added to mimic the average level of thymidine in rodent plasma.

Hyperthermia

Hyperthermia was carried out in thermostatically regulated waterbath at 41 or 43°C for 60 min.

The atmosphere of the waterbath was adjustable by a connection with air and CO2 supplies. Hyperthermia treatment was performed directly before the irradiation.

Irradiation

Irradiations were performed with single doses of γ-rays from a 137Cs source at a dose rate of 0.7 Gy/min.

Clonogenic assay for radiosensitivity

After treatment, the cells were trypsinized directly (ip) or 24 h after irradiation (dp) and replated in appropriate dilutions in 6-well macroplates (6). Eight to ten days after inoculation colonies were fixed in 6% glutaraldehyde and stained with 0.05% crystalviolet. Colonies of 50 cells or more were scored as originating from a single clonogenic cell. Surviving fractions (S(D)/S(0)) after dose (D) were corrected for the toxicity of dFdC alone (S(0)) and survival curves were analyzed using SPSS (Chicago, IL, USA) statistical software by means of a fit of the data by a weighted, stratified, linear regression, according to the linear-quadratic formula: S(D)/S(0)= exp-(αD+βD²).

Cisplatin

Cisplatin is a widely used anti-cancer drug, often combined with radiotherapy [9]. Chemo-radiation application based on cisplatin has now become the standard treatment for, among others, locally advanced cervical carcinoma [10] and locally advanced non-small cell lung cancer (NSCLC) [11]. There have been many studies on the radiation sensitizing effect of cisplatin, but results vary from a clear cisplatin-induced radiosensitization [12-15] to only an additive effect on cell survival [16]. Cisplatin and radiation have in common that their cellular target is DNA [17].

Cisplatin causes DNA damage by the formation of inter- and intrastrand adducts [18]. The cisplatin-DNA adducts can cause cell cycle arrest, inhibition of DNA replication and transcription, and eventually apoptosis [19]. Repair inhibition of DNA has also been implicated [20]. The most important repair pathways reported to be involved in cisplatin-induced DNA damage repair are nucleotide excision repair (NER) and/or homologous recombination (HR) [21, 22]. An additional route for the repair of cisplatin-DNA interstrand adducts is the post-replication/translation repair pathway which helps the cell to tolerate or bypass the lesion [23]. Irradiation causes repairable (potentially lethal) and non-repairable (lethal) lesions to the DNA which are induced independently. The ultimate effect of the repairable lesions depends on competing processes of repair and misrepair. The repair of the potentially lethal damage (PLDR) is reflected by the difference in survival between immediately and delayed plated cells. Inhibition of PLDR is implicated to play a role in cisplatin-induced radiation sensitization [14]. More specifically, cisplatin-induced radiation sensitization has been shown to occur through inhibition of the non-homologous end joining (NHEJ) pathway and recombination repair [18, 22, 24].

The radiation sensitization of cisplatin on the lung tumour cell line SW1573 and the cervical tumour cell line Siha is described as changes in linear and quadratic parameters of radiation dose survival curves. In Figure 1 the survival curves are shown for SW1573 lung tumour cells after radiation alone and after radiation combined with cisplatin treatment (1 µM for 1 h). Cisplatin was added to the cultures just before radiation. The survival curves are obtained directly (ip=immediately plated) and 24 h after (dp=delayed plated) treatment to determine potentially lethal damage repair. A slight, but statistically significant effect of cisplatin on the radiosensitivity was only observed in delayed plated cells (p = 0.02). This was also described by an increase in the α-viscosity parameter.

Figure 1: Radiation survival curves of confluent cultures of SW-1573 cells plated immediately after irradiation, ip (left) or 24 h after irradiation, dp (right) with or without 1µM cisplatin treatment for one hour. Means with standard errors of at least three experiments are shown.
and β-value (Table 1). Only for the delayed plated cells an increase with a factor of 2.5 for the value of a was obtained by cisplatin treatment. For both plating conditions an increase with a factor of 1.2 was obtained for the value of β. In the Table 1 also the effects on the linear and quadratic parameters of different plating conditions are presented as well as a 1 h incubation with 1 or 5 μM cisplatin and a continuous incubation with cisplatin during the complete duration of the clonogenic assay. It is obvious that the cervical tumour cells SiHa are more radiosensitized with 1 μM continuous cisplatin incubation than the SW1573 lung tumour cells.

Gemcitabine

Gemcitabine (dFdC, Difluorodeoxycytidine) is a deoxycytidine analogue with clinical activity in non-small cell lung cancer (NSCLC) and pancreatic cancer [25-27]. It requires phosphorylation to its active metabolites, gemcitabine-diphosphate (dF-dCTP) and gemcitabine-triphosphate (dF-dCTP), with the initial phosphorylation by deoxycytidine kinase (dCK) being the rate limiting step [28,29]. The dF-dCTP inhibits ribonucleotide reductase which regulates the production of deoxynucleotides necessary for DNA synthesis and repair [30]. The depletion of the deoxynucleotides leads to an increased production of deoxynucleotides necessary for DNA synthesis and repair [30].

Table 1: Values of the linear and quadratic parameters of the different cell lines obtained after analyses of the radiation dose survival curves for radiation alone and after combined radiation and gemcitabine treatment are summarized. SWp and SWg were almost equally sensitive to ionizing radiation alone with respect to the low dose region described by the α-value of the linear quadratic formula (Table 2). A slight increase in survival was observed in SWg cells in the high dose region which was reflected by a slightly lower β-value of the linear-quadratic formula (0.040 ± 0.006 vs 0.055 ± 0.008). The human ovarian carcinoma cell line A2780 and its gemcitabine-resistant variant AG6000 were equally sensitive to ionizing radiation. The surviving fractions of the different cell lines after incubation with gemcitabine alone are: SWp

![Table 1](image)

Significant difference with *P<0.01, †P<0.05, na is not applicable

Table 2: Values of the linear-quadratic parameters α and β and enhancement factors from cells treated with ionizing radiation only and after combined radiation cisplatin (1 μM for 1h; 1 μM continuously; 5 μM continuously) treatment.
10 nM: 0.52 ± 0.06; SWg 10 μM: 0.95 ± 0.03, 100 μM: 0.24 ± 0.11; A2780 2 nM: 0.82 ± 0.08, 10 nM: 0.21 ± 0.08; AG6000 20 μM: 0.62 ± 0.07, 50 μM: 0.22 ± 0.04.

As can be observed in Figure 2 and Table 2 radiosensitization is observed with gemcitabine-sensitive as well as in gemcitabine resistant cells. For the resistant cells much higher gemcitabine doses are needed for the radiation sensitization to result in similar cytotoxicity. Both gemcitabine-sensitive cell lines SWp and A2780 are sensitized by incubation with 10 nM of gemcitabine for 24 h before irradiation while the SWg and AG6000 are not sensitized with this dose of gemcitabine. The sensitization is described by an increase in the α-values with factors of 3 and 1.4 respectively, whereas the β-values are not significantly altered. Higher concentrations of Gemcitabine (50 and 100 nM resp.) are required to sensitise gemcitabine-resistant AG6000 and SWg cells to irradiation. For the SWg cells, the radiosensitization was reflected by an increase by a factor of 2.25 in the value of β, whereas in the AG6000 only the α-value was increased by factor of 1.3.

Iodo-deoxyUridine (IdUrd)

Incorporation of halogenated pyrimidines (HPs), chloro-, bromo- and iodo-deoxyuridine (CldUrd, BrdUrd, IdUrd) into DNA is known to sensitise cells to ionizing radiation [43-51]. Halogenated pyrimidines are incorporated into the DNA replacing the thymidine. The induced radiosensitisation increases with the degree of thymidine-replacement. The mechanism of radiation sensitisation by the HPs has been suggested to be either an increase in the amount of DNA damage induced by radiation, an influence on repair of sublethal damage (SLD), and/or an enhanced expression of potentially lethal damage (PLD) [43,52]. Since different processes are involved in these phenomena several mechanisms might contribute to the radiosensitisation.

HPs have been suggested to provide an advantage in radiotherapy as radiosensitisers of cells in rapidly growing tumours, in particular in clinical conditions in which critical normal tissues show limited proliferation and as a consequence take up less HP. Labelling depends on the growth fraction, cell loss, cell cycle time and potential doubling time. Of special importance for sensitisation is the rate at which non-cycling cells are recruited into the proliferative compartment during exposure to HPs and a course of radiotherapy. However, even in rapidly growing tumours, cells may, after proliferative cycles, move into a non-proliferative stage. This might compromise the degree of radiation sensitisation if resting cells are less affected by HPs, or are better able to cope with additional damage by repair of PLD.

Here the results of radiosensitization after incubation with 4 μM IdUrd for 72 h are presented. IdUrd-induced radio sensitisation was obtained in all studied cell lines, SW-1573 (human lung cancer), RUCII (rat urether carcinoma), R1 (rat rhabdomyosarcoma) and V79 (hamster fibroblast cells), in exponentially growing and in plateau-phase cells. Survival curves of SW-1573 cells are presented in Figure 3. Values of α and β derived by linear-quadratic analyses of survival curves of exponentially growing cells and plateau-phase cells are presented in Table 3. The plating conditions of the V79 cells, i.e. exponentially growing cells plating before or after irradiation (ppi or pai resp.),
Figure 3: Radiation dose-survival curves of SW-1573 human lung cancer cells exponentially growing (left) without IdUrd (open triangles) and after incubation with 4 µM IdUrd (closed triangles) and in plateau-phase (right) plated immediately after irradiation (dashed lines) and plated 24 h after irradiation (solid lines) without IdUrd (open symbols) and after incubation with 4 µM IdUrd (closed symbols). Each point represents the mean value of 3 different experiments ± sem.

Table 3: Values of the linear-quadratic parameters α and β and enhancement factors of several cell lines treated with ionizing radiation only and after sensitization with iododeoxyuridine (incubation with 4 µM IdUrd for 72 h).

| Cell line          | α (Gy⁻¹) control | β (Gy⁻²) control | α (Gy⁻¹) IdUrd-sens | β (Gy⁻²) IdUrd-sens | α-enhanc factor | β-enhanc factor |
|--------------------|------------------|------------------|---------------------|---------------------|-----------------|----------------|
| SW 1573 cells      |                  |                  |                     |                     |                 |                |
| Exp growing ip     | 0.22 ± 0.01      | 0.022 ± 0.001    | 0.83 ± 0.06         | na                  | 3.8             |                |
| SW 1573 cells      |                  |                  |                     |                     |                 |                |
| Plateau phase ip   | 0.17 ± 0.03      | 0.042 ± 0.004    | 0.31 ± 0.03         | 0.047 ± 0.005       | 1.8             | 1.1            |
| SW 1573 cells      |                  |                  |                     |                     |                 |                |
| Plateau phase dp   | 0.09 ± 0.02      | 0.046 ± 0.002    | 0.37 ± 0.04         | 0.033 ± 0.006       | 4.1             | 0.7            |
| RUCII cells        |                  |                  |                     |                     |                 |                |
| Exp growing ppi    | 0.008 ± 0.007    | 0.025 ± 0.001    | 0.06 ± 0.02         | 0.026 ± 0.001       | 7.5             | 1.04           |
| R1 cells           |                  |                  |                     |                     |                 |                |
| Exp growing ppi    | 0.23 ± 0.01      | 0.068 ± 0.003    | 0.44 ± 0.05         | 0.075 ± 0.016       | 1.9             | 1.1            |
| V79 cells          |                  |                  |                     |                     |                 |                |
| Exp growing ip     | 0.18 ± 0.02      | 0.017 ± 0.003    | 0.38 ± 0.04         | 0.023 ± 0.007       | 2.1             | 1.4            |
| V79 cells          |                  |                  |                     |                     |                 |                |
| Exp growing ppi    | 0.15 ± 0.02      | 0.013 ± 0.003    | 0.29 ± 0.03         | 0.016 ± 0.004       | 1.9             | 1.2            |
| V 79 cells         |                  |                  |                     |                     |                 |                |
| Plateau phase ip   | 0.09 ± 0.03      | 0.026 ± 0.004    | 0.17 ± 0.02         | 0.062 ± 0.005       | 1.9             | 2.4            |
| V 79 cells         |                  |                  |                     |                     |                 |                |
| Plateau phase dp   | 0.07 ± 0.02      | 0.020 ± 0.002    | 0.30 ± 0.03         | 0.024 ± 0.004       | 4.3             | 1.2            |

Means with SEM of at least three separate experiments are shown. ip=immediately plated after irradiation; dp=delayed plated after irradiation; ppi=plated prior to irradiation; na=not applicable.

and plateau phase cells plated immediately or 6-24 h delayed after irradiation (ip or dp resp.) had no influence on the factor of increase of the α-value. It is shown that the value of the linear parameter, α can be enhanced by a factor of 1.9 to 7.5 and that in general low values of α are enhanced more than higher values of α. The value of β is less enhanced and the enhancement factor ranges from 0.7 to 2.4.

The direct comparison between immediate and delayed plating of plateau-phase cells and between plateau phase and exponentially growing cells shows significant quantitative differences. The data on the linear and quadratic parameters described here provide various new insights in the interpretation of radiosensitisation of delay plated plateau-phase cells. It is demonstrated that in delay plated HP-sensitized plateau phase cells PLD is not abolished.

**Hyperthermia**

Hyperthermia refers to heat treatments if cells or malignancies in which the temperature is elevated in the range of 39°C to 45°C. It is used in combination with chemo- and/or radiotherapy since it has been shown to enhance the anti-cancer effects of both therapies [53-57]. Many in vitro studies on the combination of hyperthermia and radiation have shown a synergistic interaction between the two modalities, especially at higher temperatures (above 42ºC) [58-60]. This interaction is believed to result from inhibition of repair of radiation-induced DNA damage by hyperthermia [61,62]. The sequence of combined radiation and hyperthermia treatment is important. Optimal sensitization is obtained when radiation and hyperthermia are applied simultaneously or with a short interval [63]. In the clinic this is not always possible. In our experiments hyperthermia was applied immediately after radiation treatment.

Despite the clinical goal to reach (cytotoxic) temperatures as high as 43ºC, tumour temperature distributions are in practice heterogeneous. In large areas of the tumour temperatures are often lower than 43ºC.
Nonetheless, good results have been obtained in locally advanced cervical cancers with tumour temperatures below 43°C [54]. Mild temperatures have more subtle effects than high temperatures, such as tumour-reoxygenation [64-67]. Recently it has been shown that hyperthermia (42°C for 1h) transiently breaks down the BRCA2 protein [68]. In this paragraph the effects of hyperthermia treatment for 1h at 41 or 43°C on the linear quadratic parameters are summarized. Several different cell types have been studied.

**Effect of hyperthermia treatment on radiosensitivity of RKO cells:** The RKO cells, derived from human colon cancer, are relatively sensitive to hyperthermia treatment. Hyperthermia treatment for 1h at 43°C decreases the relative survival to less than 0.01 and combination with radiation doses in excess of 5 Gy always resulted in a situation in which no colony formation was observed. Treatment of cells with 41°C hyperthermia (1h) alone had little effect and resulted in a surviving fraction of 0.8 ± 0.1 in immediately plated (ip) cells and of 0.9 ± 0.1 in delayed plated (dp) cells. When cells were treated at 41°C for 1h immediately prior to irradiation, a significant (p < 0.001) enhancement of cellular radiosensitivity was observed both in ip (Figure 4) and dp (Figure 4) cells.

The effects of hyperthermia on the LQ parameters are summarized in Table 4. The value of the linear parameter α increased by a factor 1.7-1.8 while the value of the β parameter even increased with a factor as high as 2.5-7.0. One must bear in mind that the quadratic component in this cell line is very small and small changes can have large on the numerical values of β.

**Effect of hyperthermia treatment on radiosensitivity of SW-1573 cells:** SW-1573 cells are derived from a human lung tumour and are much less sensitive to hyperthermia treatment than RKO cells. Studies were performed to evaluate whether pretreatment with hyperthermia at 41°C or at 43°C in SW-1573 cells was able to enhance the radiosensitivity of these cells. Hyperthermia treatment at 41°C for 1h without radiation did not result in a decrease of the surviving fraction for ip and dp cells as compared to radiation alone. One hour hyperthermia treatment at 43°C decreased survival to 0.5 ± 0.1 for ip and to 0.4 ± 0.2 for dp cells. Pre-treatment of cells at 41°C for 1h did not alter cellular radiosensitivity of both ip and dp cells (Figure 5). However, 1h treatment at 43°C resulted in a significant (p < 0.001) radiation enhancement both in ip and dp cells (Figure 5). In Table 4 the values of the linear-quadratic parameters for radiation alone and for combined treatments are given. Hyperthermia treatment for 1 h at 41°C did result in an increase of the value of β by a factor 1.3-1.8 while the value of α even decreased. Hyperthermia treatment for 1 h at 43°C result in an increase of the value of α by a factor 2.3-4.4 while the value β increased with a factor 1.8-2.0.

**Discussion and Conclusion**

In most cases an increase of the α-component was observed which

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Table 4: Values of the linear-quadratic parameters α and β and enhancement factors from cells treated with ionizing radiation only and after combined radiation and hyperthermia treatment.

| Cells   | Treatment | α (Gy⁻¹) control | β (Gy⁻²) control | α-enhanc factor | β-enhanc factor |
|---------|-----------|------------------|------------------|----------------|----------------|
| RKO ip  | sham      | 0.55 ± 0.09      | 0.02 ± 0.01      |                |                |
|         | HT 41 1h  | 0.93 ± 0.09      | 0.05 ± 0.02      | 1.7            | 2.5            |
| RKO dp  | sham      | 0.47 ± 0.09      | 0.01 ± 0.01      |                |                |
|         | HT 41 1h  | 0.83 ± 0.08      | 0.07 ± 0.02      | 1.8            | 7.0            |
| SW1573 ip | sham   | 0.21 ± 0.02      | 0.06 ± 0.02      |                |                |
|         | HT 41 1h  | 0.06 ± 0.02      | 0.11 ± 0.03      | 0.3            | 1.8            |
|         | HT 43 1h  | 0.49 ± 0.04      | 0.12 ± 0.03      | 2.3            | 2.0            |
| SW1573 dp | sham  | 0.09 ± 0.02      | 0.06 ± 0.02      |                |                |
|         | HT 41 1h  | 0.05 ± 0.02      | 0.08 ± 0.02      | 0.6            | 1.3            |
|         | HT 43 1h  | 0.40 ± 0.04      | 0.11 ± 0.03      | 4.4            | 1.8            |

Sham=control is radiation only; ip=immediately plated; dp=delayed plated.
corresponds to an enhanced (potentially) direct lethal damage (PLD) at low doses. The β-component, which is assumed to depend on the interaction of sublethal lesions (SLD), was rarely affected by the studied radiosensitization agents. Moreover, it appeared that more radiosensitive cell lines were more sensitised than the radiosensitive lines. Furthermore it can be concluded that radiosensitization is also dependent on cell cycle stage like plateau or exponentially growing phase or post treatment plating conditions [69].

It is shown that cisplatin causes radiosensitization as measured by clonogenic survival, but only after allowing a potentially lethal damage repair (PLDR) time of 24 hours. These results are in agreement with those of Wilkins et al. [70] who investigated the effect of cisplatin and radiation on PLDR in confluent cultures of two different brain tumor cell lines. Wilkins et al. [71] also observed no radiosensitization by cisplatin in immediately plated cells whereas a cisplatin-induced radiosensitization was seen in cells plated eight hours after irradiation. Their results indicate that the radiosensitizing effect of cisplatin occurs through the inhibition of post-irradiation recovery. The strongest inhibition of PLDR was achieved when cisplatin was administered shortly before or after irradiation [70]. In our experiments, cells were irradiated while cisplatin was present in the medium.

Results from studies using exponentially growing cell cultures vary from a cisplatin-induced radiosensitization [11-13,71] to only an additive effect [9,11,72-74]. The effect of cisplatin treatment on radiosensitivity may depend on the cell type used. Loprevite et al. [11] observed synergism in a squamous lung carcinoma cell line when exposed to cisplatin, whereas an adenocarcinoma of the lung was not sensitized by cisplatin. Even cell lines derived from a single biopsy can differ in the response to cisplatin and radiation combination therapy [74].

Although dependence on cell cycle phase [75,76], cisplatin incubation time and the sequence of treatment modalities have been implicated [9,75,76], there is currently no consensus to account for the varying response of cells to cisplatin and radiation.

The mechanism of cisplatin induced radiosensitization might be due to the inhibition of the DNA repair, NHEJ and HR, pathways [19,24]. The Ku protein complex, which plays an important role in NHEJ, was demonstrated to show a reduced ability to translocate on DNA containing cisplatin-DNA adducts compared with undamaged DNA. This resulted in a decreased interaction between Ku and DNA-PKcs [77]. However, the biochemical processes that cisplatin undergoes in the cell are complex and the intracellular fate of cisplatin may be linked to copper transport [78]. Therefore, other processes such as the formation of peroxy complexes inside the cell might be involved in cisplatin-induced radiosensitization [14,79]. Bergs et al. [14] demonstrated an increase in the induction of apoptosis after combined treatment as compared to radiation or cisplatin alone at 24 h after treatment. This was confirmed in several other studies [80,81]. These apoptotic effects observed by Bergs et al. [14] correlated with clonogenic survival. Fujita et al. [82] also observed an inhibitory effect of the combination of cisplatin and radiation on the survival of lung tumor cells and ascribed this effect on the induction of tumor cell apoptosis.

In conclusion, a radiosensitizing effect of cisplatin on cell survival is observed in confluent cultures when cells were replated after a 24 hour incubation period during which PLD repair could take place. In contrast, cisplatin did not induce a significant radiosensitization after immediate plating.

Several studies have shown that gemcitabine is a potent sensitizer of ionizing radiation [29,34,83]. Among other proposed mechanisms of action, the effect of gemcitabine on cell cycle distributions may be the most important [35,36]. In our studies, both gemcitabine-sensitive cell lines SWp and A2780 could be sensitized to irradiation when cytotoxic gemcitabine-treatments were given. The radiosensitization was accompanied by a clear arrest of cells in early S phase which has been argued to be vital for gemcitabine-induced radiosensitization [33]. Both cell lines showed an increase in α-value indicating the efficacy of gemcitabine-induced radiosensitization in the clinically relevant dose range. Although the gemcitabine resistant cells still could be sensitized only much higher gemcitabine doses were necessary to reach an effect. In the resistant ovarian carcinoma cell line AG6000 this was demonstrated by an increase in the value of β. In contrast with this change, in the gemcitabine resistant lung tumour cell line an increase in the β-value was obtained, the α-value was not affected. In both gemcitabine-resistant cell lines the sensitivity to ionizing radiation alone was not altered. It is reported that gemcitabine resistant tumours are cross-resistant to related drugs like Ara-c [84,85]. In both gemcitabine-resistant cell line, AG6000 and SWg, this was indeed the case [40]. Moreover, the AG6000 cells were also more resistant to cisplatin and taxoids (41). However, no altered sensitivity was found in 41°C irradiated cells compared to 43°C irradiation. The same holds true for SW-g cells which were also more resistant when exposed to 41°C compared to 43°C treatment. This effect on the induction of tumor cell apoptosis.

**Figure 5:** Radiation survival curves of confluent cultures of SW-1573 cells (human lung tumour cells) plated immediately after irradiation (ip) or 24h after irradiation (dp) with or without hyperthermia pre-treatment at 41°C (left) or at 43°C (right) for 1h. Means with standard errors of at least three experiments are shown.
The HP-induced-radiosensitisation is mainly due to an increase in the linear parameter \( \alpha \). The quadratic parameter, \( \beta \), is only rarely influenced. Different mechanisms involved in the radiosensitisation induced by halogenated pyrimidines have been described [43]. Wang et al. [86] suggested that in exponentially growing cells increased DNA damage production was the major component of radiosensitisation while in plateau-phase cells radiosensitisation occurred through inhibited repair and/or enhanced fixation of potentially lethal damage. The increase of the \( \alpha \) values for exponentially growing cells as found in our study, indicates an increase in the number of directly lethal events due to the HPs. This is in agreement with observations of Webb et al. [52] and Jones et al. [87] which suggest that an important mechanism of radiosensitisation involves an increase of effective DNA double strand breaks. Miller et al. [50,51] have suggested that radiation-induced damage in cells which have HPs incorporated into the DNA after low-LET irradiation resembles the damage produced by high-LET radiation. In plateau-phase cells plated immediately after irradiation the increase of \( \alpha \) might be due to the same mechanism as involved in exponentially growing cells. In these cells also an increase of \( \beta \) was observed indicating that accumulation of sublethal lesions contributed significantly [2]. Due to the immediate plating after irradiation this sublethal damage might be fixed. Greatest increases in \( \alpha \) were found in delayed plated plateau-phase cells. This radiosensitisation can be interpreted as an enhanced fixation of potentially lethal damage due to immediate DNA damage and/or to damaged DNA repair function in these cells expressed during the interval before delayed plating. The value of \( \beta \) in these cells returned to values as found in cells not containing HPs. This demonstrates that sublethal damage has been repaired in HP-containing plateau-phase cells.

Hyperthermia is an excellent radiosensitizer which can already be effective at mild temperatures. One hour hyperthermia treatment at 41°C without radiation had only a small cytotoxic effect in both the heat sensitive and the heat resistant cell line. This is in agreement with the general idea of cell kill induction at temperatures 242°C for 1h or more Dewhirst, 2005 [64]). Hyperthermia treatment at 43°C for 1h did not have a large cytotoxic effect in heat resistant SW-1573 cells. Radiosensitisation by 41°C temperature hyperthermia was observed in RKO, but not in SW-1573 cells. The ability of mild temperatures (in the range of 40-42°C) hyperthermia to increase radiosensitivity of human tumor cells has been shown to be cell line dependent [66,67,88-93]. In a study by Xu et al. [94] 41.1°C pre-treatment of cells for 1h did not induce radiosensitization whereas treatment for 2h or more resulted in radiosensitization, in the hyperthermia resistant, but not in the hyperthermia sensitive cell line [94]. However, simultaneous treatment of the sensitive cell line with 1h 41.1°C hyperthermia and radiation did increase cellular radiosensitivity [95]. An important mechanism of mild hyperthermia induced radiosensitization in vivo is the reoxygenation of tumors by an increase in blood flow [96-98]. Recently it was demonstrated that the BRCA-2 protein is transiently inhibited by mild hyperthermia [68,99]. Also translocation of the Mre11 DSB repair protein from the nucleus to the cytoplasm has been implicated [95,100]. However, disappearance of Mre11 protein foci at the sites of irradiation induced DNA double strand breaks by 41°C pre-incubation of cells was not observed [66-68]. A role for mitotic catastrophe occurring as a result of G2/M checkpoint abrogation has also been suggested [101]. It has been shown that radiosensitisation by 41-43°C hyperthermia correlates with an increased number of chromosomal fragments, but not of color junctions, at 24h after treatment compared to radiation alone [67].

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