Differential level of DSB repair fidelity effected by nuclear protein extracts derived from radiosensitive and radioresistant human tumour cells

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Summary A cell-free plasmid reactivation assay was used to determine the fidelity of DNA double-strand break (DSB) repair in a panel of eight DSB repair-proficient human tumour cell lines. Nuclear protein extracts derived from radiosensitive tumour cells were less capable of correctly rejoining EcoRI-induced DSBs than were similar extracts from radioresistant tumour cells. Linear regression analysis suggests that there was a significant \( r^2 = 0.84, P = 0.001 \), d.f. = 6) correlation between the fidelity of DSB rejoining and the SF\(_2\) values of the cell lines studied. This cell-free assay is clearly sensitive to differences in the nuclear protein composition that reflect the clinically relevant radiosensitivity of these cell lines. The fact that our cell-free assay yielded similar results to previous studies that used intracellular plasmid reactivation assays suggests that those differences in DSB mis-rejoining frequencies in radiosensitive and radioresistant cell lines may be due to inherent differences in nuclear protein composition and are not directly attributable to differences in proliferation rates between cell lines. The underlying cause for this association between DSB mis-rejoining frequencies and radiosensitivity is presently unknown, however restriction endonuclease mapping and polymerase chain reaction (PCR) amplification analysis revealed that approximately 40% of the mis-rejoined DSBs arose as a result of the deletion of between 40 and 440 base pairs. These data raise the possibility that the radiosensitivity of DSB repair-proficient human tumour cell lines may be partly determined by the predisposition of these cell lines to activate non-conservative DSB rejoining pathways.

Keywords: \( \gamma \)-radiation; DNA repair; repair fidelity; human tumour cells

The concept that clinical radioresponsiveness is a function of the intrinsic radiosensitivity of tumour cells (Fertil and Malaise, 1980; Deacon et al, 1984) has led to considerable interest in identifying the basis for the variation in radioresistance among human tumour cells. In this regard, DNA repair-deficient cell lines have provided a valuable insight into some of the mechanisms that are important in determining the lethality of radiation-induced DNA damage. The functional inactivation of the proteins involved in the rejoining of DNA double-strand breaks (DSBs), e.g. those encoded by the XRCC4, XRCC5 (Ku86) and XRCC7 (DNA-PK\(_c\)), leads to the greatest increase in cellular radiosensitivity (Giaccia et al, 1985; Taccioli et al, 1994; Kirchgesner et al, 1995; Lees-Miller et al, 1995). Given the importance of DSB rejoining to the lethality of radiation-induced DNA damage, differences in DSB-rejoining capacity might be expected to account for the variations in cellular radioresistance in human tumour cell lines. Although there have been reports that DSB-rejoining rates correlate with radiosensitivity (e.g. Giaccia et al, 1992), there have been as many, if not more, studies that suggest that there is no correlation between these parameters (reviewed in Olive et al, 1994). Furthermore, there is no obvious relationship between the level (XRCC5, XRCC6 and XRCC7) or activity (XRCC7) of DSB repair proteins in human tumour cell lines exhibiting a range of radioresistance (Allalunis-Turner et al, 1995) that impacts on patient radioresponsiveness (West et al, 1993).

It is becoming apparent that the "DNA repair process" is considerably more complex than simply the rejoining of the DNA strands. Several cellular processes must be successfully integrated to facilitate the "biologically effective" repair of radiation-induced DNA damage. For example, the extreme radiosensitivity of ataxia telangiectasia (AT) cells is due to a mutation in a P13 kinase-like gene (Savitsky et al, 1995), which encodes a protein involved in intracellular signalling. DSB rejoining can thus no longer be considered in isolation as a determinant of radiosensitivity, but rather as a key component of an integrated cellular response pathway. Given that DSB repair capacity is a key, but not the sole, determinant of radiosensitivity, it is probably not too surprising that there is a poor correlation between DSB rejoining rates and radiosensitivity in panels of cell lines of differing radiosensitivity (reviewed in Olive et al, 1994). In randomly selected human tumour cell lines, there may be substantial differences in the cellular abilities to remove the various DNA lesions or to invoke the various DNA damage response pathways, e.g. cell cycle arrest, early-response genes or the stress-activated protein kinase pathway. Although our knowledge of these response pathways is limited, it seems likely that an optimal co-ordination of these processes would be required to produce a radioresistant phenotype. However, any defects within the various response pathways or a defective temporal co-ordination of these processes - cell cycle arrest, DNA repair, suppression of DNA synthesis, suppression of transcription and activation of early-response genes - could lead to biologically inadequate repair of the DNA damage and hence to a more radiosensitive phenotype.
The high to intermediate ('clinically important') levels of radio-resistance may thus reflect the degree to which the components of the various DNA-damage response pathways are integrated with each other and with other cellular processes.

In support of this hypothesis, the fidelity with which 'model' (i.e. restriction endonuclease induced) DSBs are repaired does indeed vary with respect to SF, values in human tumour cell lines (Powell and McMillan, 1991, 1994; Powell et al, 1992). In these studies, the most radiosensitive cell lines exhibit a higher fidelity of DSB repair than their radiosensitive counterparts (Powell and McMillan, 1991, 1994; Powell et al, 1992). However, as all of these studies determined the fidelity of DNA repair that was effected within the intact cellular environment, it is hard to establish whether the observed variations in DNA repair fidelity are a reflection of inter-cell line differences in the 'processing' of the DNA damage per se or are related to inter-cell line differences in other factors that have been shown to influence the repair of DNA damage, e.g. higher order chromatin structures (Johnston and Bryant, 1994). In an attempt to more definitively establish the basis for the higher levels of DSB mis-rejoining in radiosensitive tumour cells, we have used a cell-free plasmid reactivation assay (North et al, 1990) to establish the fidelity of DSB repair that is effected by nuclear protein extracts derived from radiosensitive and radioresistant tumour cells.

**MATERIALS AND METHODS**

**Cell lines**

The origins and establishment of the human glioblastoma M011, M059K and M071 cell lines have previously been published (Allanunis-Turner et al, 1992). All of these cell lines have detectable levels of Ku70, Ku86 and DNA-PKcs proteins (Allanunis-Turner et al, 1995).

The origins, establishment and clonal selection techniques used to isolate the human cervical tumour cell lines have been previously published (Britten et al, 1996a and b). Briefly, three primary human cervical tumour cell lines were selected at random from approximately 200 in vitro cell cultures that were frozen down after only three passages from date of establishment (Allanunis-Turner et al, 1991). After being recovered from cryopreservation and maintained for 2 weeks in exponential growth, single-cell colonies were isolated by limiting dilution. From each cell line, randomly selected clones were expanded and the resulting cell sublines maintained as monolayer cultures in Dulbecco’s Modified Eagle Medium (DMEM)/F12 media, supplemented with 15% fetal calf serum (Gibco, Grand Island, NY, USA) and antibiotics. After four further passages (subcultured every 4–5 days to ensure exponential growth), the radiosensitivity of these cloned cell lines was determined.

To facilitate inter-laboratory comparisons of the generated data, we used Chinese hamster ovary (CHO)-AA8 cells as a reference cell line. CHO-AA8 cells were maintained in monolayer culture in McCoy’s 5A medium (Hsu’s modification; Gibco) supplemented with 15% FCS and 1% penicillin–streptomycin. They were passaged every 2–3 days to ensure exponential growth.

**Radiation clonogenic cell survival assays**

The radiosensitivity of the cell lines used in this study has been previously established (Allanunis-Turner et al, 1992, 1995; Britten et al, 1996a and b). However, to ensure that the radiosensitivity of the cells recovered from frozen storage had not altered and to facilitate a better statistical comparison between radiosensitivity and DNA repair fidelity, complete radiation survival curves (over the range of 0–10 Gy) were constructed on the same batch of cell lines used to prepare the nuclear protein extract. To ensure that the radiation sensitivity determined on this occasion was not erroneous, radiation sensitivity was established a further two times in the subsequent 7 days. The radiosensitivity parameters presented in Table 1 represent the pooled data from all three survival curve experiments. In no instance did we find that the radiation sensitivity determined on the day that the protein extract was prepared varied from either historical data or from data observed in the subsequent assays. The procedure followed for all of these experiments is outlined below. The day before use, cells from exponentially growing stock cultures were detached using 0.25% trypsin/1 mM EDTA at 37°C and were placed into two 75-cm² flasks. On the day of the experiment, the media was removed, the cell monolayers washed with sterile pre-warmed (37°C) phosphate-buffered saline (PBS) and detached using 0.25% trypsin/1 mM EDTA at 37°C. The cell pellets were then washed twice with warm PBS. One pellet was used to prepare the nuclear protein extract (see below), while the other was resuspended in the DMEM/F12 media, supplemented with 15% fetal calf serum (Gibco) and antibiotics. Cells were seeded at densities of between 10⁴ and 5 X 10⁴ per 60 mm-tissue culture dish. Three replicate plates were set up for each of the five radiation dose levels. The Petri dishes were incubated for 4 h at 37°C (5% carbon dioxide/95% air) to allow for cell attachment, then placed in aluminium irradiation chambers and irradiated using a 60Co irradiator at a dose rate of 6 Gy min⁻¹. After 15 days at 37°C (5% carbon dioxide/95% air), the colonies were fixed in 70% ethanol and stained with 10% methylene blue, and those colonies containing greater than 50 cells were counted.

**Data handling and presentation**

The experimental data were fitted to the linear–quadratic equation:

\[ -\ln S = aD + \beta D^2 \]

where \( S \) is equivalent to the surviving fraction at a given dose, \( D \), and \( a \) and \( \beta \) are constants. The data were fitted to a linear–quadratic function using the non-linear regression program of the PRIZM software package (Graphpad Software, San Diego, CA, USA). \( SF_2 \) values were calculated by substitution of the derived constants into the linear quadratic equation.

**Nuclear protein extracts**

Nuclear protein was extracted (from the same batch of cells used to determine radiosensitivity) in accordance with the method of Olnes and Kurl (1994). Protein extracts were stored at −70°C (for a maximum of 6 months) until required. The level of nuclear protein obtained was determined using the Bio-Rad Protein Assay (Bio-Rad Laboratories, Richmond, CA, USA) in accordance with the manufacturer’s instructions and using bovine serum albumin (BSA) as a protein standard. Before use in the plasmid reactivation assay, the quality of the proteins was determined by separating the protein by SDS/PAGE. The protein bands were visualized by Coomassie blue staining.

**DNA repair fidelity assay**

A modified version of a cell-free plasmid reactivation system (North et al, 1990) was used to assess DNA repair fidelity. This
technique uses double-stranded pUC18 DNA containing the lacZα complementation gene and the ampicillin-resistance gene (ampR).

The DNA was isolated from HB101 bacteria containing the pUC18 plasmid using a plasmid Maxi Kit (Qiagen, Chatsworth, CA, USA). A single DSB was then introduced within the lacZα gene by the EcoRI restriction endonuclease (Pharmacia, Piscataway, NJ, USA). After heat inactivation of the EcoRI (20 min at 65°C), the plasmid was ethanol precipitated and the cutting efficiency checked by Southern analysis (see below). In the initial stages of this study, the amount of nuclear protein extract that yielded the greatest number of colonies was ascertainment. The optimum protein level was found to be 1 μg (Table 1); at higher protein levels, there was a noticeable drop in the number of bacterial transformants and a noticeable degradation of the plasmid after electrophoretic separation. This level of nuclear protein was approximately 30-fold less than that of the cytoplasmic extracts used in the earlier studies using this assay (North et al, 1990), which presumably reflects the greater concentration of DNA-processing enzymes in the nuclear vs cytoplasmic protein extracts.

The experimental conditions used to derive the data presented in this paper were as described below. Two micrograms of linearized plasmid DNA and 1 μg of nuclear protein extract were mixed in 50 μl containing 50 mM Tris-HCl (pH 7.6), 10 mM magnesium chloride, 1 mM ATP, 1 mM DTT, 50 μM dNTPs and 5% (w/v) polyethylene glycol-8000 and were incubated at 18°C for 24 h. The plasmid DNA was then isolated by spinning down the reaction mixture through Ultrafree-Probind Filter column (Millipore Corporation, Bedford, MA, USA). For quality control purposes, an aliquot was removed and treated with T4-ligase (Gibco) before being further processed.

The recovered plasmid was diluted fivefold with TE buffer (pH 7.6) and then transfected into the E. coli DH5α strain using the Gibco-BRL protocol for DH5α bacteria. The transformed cells were then mixed with 3 ml of LM medium/0.7% agar containing 67 μg ml⁻¹ ampicillin, 266 μg ml⁻¹ X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) and 266 μg ml⁻¹ of IPTG (an inducer of lacZα gene function) and were then poured onto premade LM plates containing 100 μg ml⁻¹ ampicillin. An aliquot of cells was also plated out on LM plates that contained no ampicillin to determine bacterial cell viability/plating efficiency. After overnight incubation at 37°C, the colonies were visualized, and those greater than 1 mm in diameter were counted. Correct rejoining of the DSBs produced DNA that yielded dark-blue colonies. Mis-rejoining of the DSBs resulted in a mutated lacZα sequence that produced colourless colonies. As the functional

### Table 1

Ampicillin-resistant bacterial clones generated after transfection of DH5α bacteria with EcoRI-digested pUC18 molecules after exposure to various amounts of HT137/1 nuclear protein extract

| Nuclear protein (μg) | Total no. of colonies | No. of white colonies | LacZα-deficient (%) |
|---------------------|-----------------------|-----------------------|---------------------|
| 0.1                 | 102                   | 2                     | 1.9                 |
| 0.5                 | 110                   | 2                     | 1.8                 |
| 1.0                 | 132                   | 3                     | 2.3                 |
| 2.5                 | 70                    | 1                     | 1.4                 |
| 5.0                 | 0                     | 0                     | 0                   |
| 10                  | 0                     | 0                     | 0                   |
| 25                  | 0                     | 0                     | 0                   |

*Number of colonies above 'background' level of plasmid rejoining, i.e. in the absence of nuclear protein.

### Table 2

Radiation survival curve parameters of the cell lines studied (standard error in parentheses)

| Cell line | α (Gy⁻¹) (μ) | β (Gy⁻²) (μ) | SF₂ (E1/0) (μ) | Dₐ (Gy) |
|-----------|--------------|--------------|----------------|---------|
| M011      | 0.907 (0.186) | 0.111 (0.058) | 0.10           | 2.04    |
| M071      | 0.932 (0.167) | 0.029 (0.043) | 0.14           | 2.31    |
| HT212/7   | 0.433 (0.031) | 0.005 (0.004) | 0.41           | 5.03    |
| HT212/10  | 0.323 (0.014) | <0.0001      | 0.52           | 7.11    |
| HT137/1   | 0.240 (0.033) | 0.021 (0.004) | 0.56           | 6.21    |
| HT212/4   | 0.229 (0.042) | 0.025 (0.005) | 0.57           | 6.05    |
| M059K     | 0.129 (0.093) | 0.059 (0.011) | 0.61           | 5.25    |
| CHO-AA8   | 0.154 (0.022) | 0.032 (0.002) | 0.64           | 6.41    |
| HT180/7   | 0.050 (0.032) | 0.051 (0.007) | 0.74           | 6.25    |

![Figure 1](image-url) The relationship between plasmid rejoining levels and SF₂ values (A) or Dₐ values (B) in early-passage human cervical (○, HT137/1; ■, HT212/4; □, HT212/7; □, HT212/10; ■, HT180/7) and glioma (M011, ▲; M059K, ◆; and M071, ○) tumour cells and in CHO-AA8 (△) cells. Rejoining capacity was assayed by at least three separate experiments.
Table 3  Ampicillin-resistant bacterial clones generated after transfection of DH5α bacteria with EcoRI-digested pUC18 molecules after exposure to T4 ligase or to various nuclear protein extracts

| Treatment | Total colonies produced | White colonies produced | Misrepair frequency | Total colonies produced |
|-----------|-------------------------|-------------------------|--------------------|-------------------------|
| EcoRI only | 43.44 ± 13.22 | 0.055 ± 0.055 | 0 | 46.82 ± 21.07 |
| EcoRI → T4 ligase | 4950 ± 45.92 | 0.33 ± 0.31 | 0 | 5125 ± 67.23 |
| EcoRI → nuclear protein → T4 ligase | 5744 ± 67.12 | 0.19 ± 0.08 | 0 | 5463 ± 79.19 |
| M011 | 181.5 ± 10.81 | 14.83 ± 0.75 | 8.29 ± 0.54 | 206.60 ± 11.03 |
| M071 | 241.33 ± 45.21 | 14.66 ± 0.56 | 5.16 ± 0.17 | 315.60 ± 19.84 |
| HT137/1 | 74.5 ± 5.60 | 2.33 ± 0.21 | 3.19 ± 0.29 | 99.65 ± 9.56 |
| HT212/7 | 239.33 ± 35.75 | 10.5 ± 1.61 | 4.38 ± 0.17 | 246.20 ± 38.25 |
| HT212/10 | 200.67 ± 14.09 | 8.0 ± 0.89 | 4.05 ± 0.49 | 218.50 ± 12.14 |
| HT212/4 | 132.17 ± 19.24 | 3.83 ± 0.74 | 2.83 ± 0.24 | 149.40 ± 21.81 |
| M059K | 413.25 ± 50.13 | 10.67 ± 1.57 | 2.53 ± 0.25 | 391.80 ± 42.27 |
| CHO-AA8 | 126.66 ± 21.60 | 4.33 ± 1.85 | 3.18 ± 0.85 | 164.20 ± 27.77 |
| HT180/7 | 83.33 ± 6.58 | 1.44 ± 0.24 | 1.77 ± 0.27 | 109.50 ± 8.53 |

*Number of ampicillin-resistant bacterial colonies produced per 10⁶ bacterial cells plated. *Number of ampicillin-resistant bacterial colonies produced normalized to bacterial cell viability. Values represent the mean values ± s.e.m. of data that have been corrected for background plasmid rejoining by subtraction of internal control data.

(α-complementation of β-galactosidase) inactivation of the lacZα gene had no direct effect on cellular viability, the full spectrum of DSB mis-rejoining events was probably conserved. The 'mutant' colonies were restreaked onto fresh selection media to verify the mutation in the lacZα gene.

Comparative studies of the DSB rejoining capacity/fidelity in the CHO-AA8 and human tumour cell lines were performed simultaneously using the same batch of competent cells, reagents and EcoRI-digested plasmid DNA. A minimum of three experiments were performed with each nuclear protein extract, the results presented in this paper represent the pooled data from these experiments. To eliminate the confounding effects of the presence of uncleaved plasmid and of rejoining events within the bacteria, the level of DSB rejoining and/or mis-rejoining that occurred irrespective of exposure to nuclear proteins was determined for each experiment and subtracted from the experimental data before further analysis.

Southern analysis
A 0.5-μg aliquot of plasmid was subjected to gel electrophoresis in 1% agarose gel, transferred to Hybond-N+ nylon membranes (Amer sham, Arlington Heights, IL, USA) by capillary blotting and baked at 80°C for 2 h. After prehybridization of the membranes for 4 h at 45°C in Hybrisol I (Oncor, Gaithersburg, MD, USA), the membrane was hybridized with 32P-labelled pUC18 probe for 18 h at 45°C. Unbound probe was removed by washing membranes three times in 2 × SSC/1% sodium dodecyl sulphate (SDS) at room temperature and twice in 0.1 × SSC/0.1% SDS at 60°C. The membranes were then enclosed in plastic wrap and placed in a phosphor-imager cassette for 1 h, after which they were scanned with a GS-250 Molecular Imager (BioRad, Mississauga, Ontario, Canada).

Mapping of mis-rejoined pUC18 molecules
Plasmid DNA was isolated from nine randomly selected 'white' and ten 'blue' clones generated after exposure to the nuclear protein extract of the HT137/1 cell line. The nature of the mutations in the LacZα gene after incubation with nuclear protein extracts was characterized by restriction endonuclease digest and by PCR amplification. The plasmid DNA was digested with the following restriction endonucleases (either singularly or in combination): EcoRI, NdeI, PvuI, PvuII, AafI or Psrl (Pharmacia, Piscataway, NJ, USA).

The multiple-cloning region was amplified using the M13/pUC sequencing primers (Gibco) listed below: forward, 5'-CCCGAGTCAGGTTGTAAGCG-3' (primer F); reverse, 5'-GGGGATTACCCAGGACG-3' (primer R). These primers give a 136-bp product (corresponding to nucleotides 364–500) when used on untreated pUC18 DNA.

In certain instances, the region of the pUC18 plasmid corresponding to nucleotides 150–777 was also amplified using another set of primers: forward, 5'-CGCGAAGTCAGGTTGTAAGCG-3' (primer 2F); reverse, 5'-GGGGATTACCCAGGACG-3' (primer 2R). This second set of primers gave a 628-bp product when used on untreated pUC18 DNA.

PCR was performed using a GeneAmp PCR system 9600 (Perkin Elmer) and the following conditions: 95°C for 2 min; 35 cycles of 95°C for 15 s, 60°C for 30 s, 72°C for 15 s; 72°C for 7 min. Limiting dilution analysis revealed that a PCR product was detectable when the target sequence was present in a concentration as low as 10 fg.

RESULTS
The human tumour cell lines studied exhibited a wide range of radiosensitivities. Comparative analysis of SF2 values revealed a 7.4-fold variation in clinically relevant radiosensitivity (0.10 > SF2 < 0.74). When assessed at the 10% survival level (D10), there was a 3.5-fold variation in the radiosensitivity of these cell lines, with a range of D10 values from 2.0 to 7.1 Gy. The radiation survival curve parameters of the CHO-AA8 and the eight human tumour cell lines are listed in Table 2.

After verification of the quality (i.e. no degradation) of the nuclear protein extracts by SDS/PAGE analysis, the ability of 1 μg of the nuclear protein extracts to rejoin the EcoRI-digested pUC18 (EDP) plasmid was assessed. EDP plasmid that was exposed to nuclear protein extracts yielded transformation frequencies within the range of 74–413 colonies 10⁴ competent cells (Table 2). There
was no obvious correlation between the radiosensitivity of the tumour cell line and the capacity of the nuclear protein extract to rejoin the EDP plasmid (as determined by transformation frequency), neither when expressed relative to bacterial cells plated \( r^2 = 0.05, P = 0.56 \) (SF,) and \( r^2 = 0.09, P = 0.43 \) (D_{s1}), d.f. = 6 \( (*\) (Figure 1) nor when corrected for bacterial cell viability \( r^2 = 0.11, P = 0.42 \) (SF,) and \( r^2 = 0.16, P = 0.34 \) (D_{s1}), d.f. = 6. The range of transformation frequencies after exposure of the EDP plasmid to the nuclear protein extract was between 4% and 18% of that accrued when the EDP plasmid was treated by T4 ligase (approximately 5000 colonies \( 10^4 \) competent cells) but was significantly higher than that observed when EDP that was not exposed to either nuclear protein or T4 ligase was used to transform the bacteria (32–45 colonies \( 10^4 \) competent cells) (Table 3).

T4 ligase treatment of EDP plasmid that had been previously exposed to nuclear protein extracts yielded transformation frequencies similar to that obtained after T4 ligase treatment alone (Table 3). It would thus appear that the nuclear protein extracts were capable of rejoining about 10% of the re-ligatable plasmid molecules and that there was not a significant degradation of the EDP plasmid by the nuclear protein extract. Moreover, as the frequency of DSB mis-rejoining after this treatment was not significantly different to that observed when T4 ligase was used alone (and it did not resemble that observed after nuclear protein treatment alone), it would appear that the nuclear protein-mediated increase in DSB mis-rejoining frequency was not attributable to a random degradation of the plasmid, but rather arose as a consequence of the nuclear protein-mediated rejoining of the DSBs.

There was a 4.7-fold range in the fidelity of DSB rejoining (as determined by the proportion of colonies with a non-functional lacZα gene, i.e. white colonies) achieved by nuclear protein extracts (Table 3). The CHO-AA8 cell line exhibited a low frequency of DSB mis-rejoining. The human cervical (HT212/4) tumour cell line that exhibited similar levels of radioresistance as the CHO-AA8 cells exhibited a similar DSB mis-rejoining frequency. The more radiosensitive human tumour cells exhibited higher frequencies of DSB mis-rejoining (Figure 2), while the more radioresistant HT1807 cell line exhibited a lower DSB mis-rejoining frequency than CHO-AA8 cells. Linear regression analysis indicates that there was a significant \( r^2 = 0.84, P = 0.001, \) d.f. = 6 correlation between SF₂ values and the fidelity of DSB rejoining (Figure 2) and, to a lesser extent \( r^2 = 0.62, P = 0.02, \) d.f. = 6, between D_{s1} values and DSB rejoining fidelity (Figure 2).

To assess whether the inactivation of the lacZα gene was attributable to deletional events, as previously suggested by Thacker et al. (1992), we performed restriction endonuclease mapping, as well as PCR amplification of the multiple-cloning region of the pUC18 plasmid recovered from a number of blue and white colonies generated after exposure to the nuclear protein extract of the HT137/1 cell line. The region of the pUC18 plasmid containing the relevant restriction sites and PCR primer recognition sequences is depicted in Figure 3.

The PstI and EcoRI sites were absent from the DNA obtained from four white colonies. No PCR product was generated when the first set of PCR primers was used to amplify the DNA from those plasmids that did not contain the PstI and EcoRI recognition sites. These data suggest that at least one of the primer recognition sequences has either been deleted or altered. In one of these clones (W8), we have confirmed that the recognition site for primer F was indeed deleted as this clone did not contain any of the recognition sites between nucleotide 183 (Ndel) and nucleotide 450 (EcoRI) (Table 4, Figure 3). To get a more accurate determination of the size of the deletions, a second set of PCR primers (2F and 2R) was selected whose recognition sequences corresponded to regions 5' to the Ndel site and 3' to the PstI site (nt. 628) recognition sequences. As expected, a single PCR product of 628 bp was generated in the blue clones tested. At this stage, plasmid DNA was only available from three of the nine white colonies: W1, W5 and W8. A single PCR product was generated in all three clones, however this product was approximately 628 bp, 330 bp and 180 bp in size from the W1, W5 and W8 clones respectively. In the case of the W1
clone, the inactivation of the lacZα gene would appear to be attributable to the loss of a few nucleotides from the original EcoRI restriction site. In the case of the W5 clone, it would appear that a deletion of approximately 300 bp in size accounted for the inactivation of the lacZα gene, while in the W8 clone a -448-bp deletion led to the inactivation of the lacZα gene. In the two instances for which we have sufficient data from the restriction digest and PCR analysis to accurately determine the size and location of the deletional events, it appears that, in the W8 clone, the deletions occurred preferentially 5' to the original EcoRI cut, whereas, in the W5 clone, approximately the same amount of material was lost 3' as was lost 5' to the EcoRI site (Figure 3B).

### DISCUSSION

In this study, we found that the fidelity of DSB rejoining (as detected by a cell-free plasmid reactivation assay) was significantly \( P < 0.001, r^2 = 0.84, \) d.f. = 6) correlated with the clinically relevant radiosensitivity, i.e. SF\(_2\), of the eight human tumour cell lines studied (Figure 2). It would thus appear that this assay is sensitive to differences in the nuclear protein composition of radiosensitive and radioresistant cells that reflect the clinically relevant radio-sensitivity of these cell lines. These data confirm and extend the findings of previous investigators who showed that radioresistant tumour cells rejoin restriction endonuclease-induced DSBs with a
lower fidelity than their radioresistant counterparts (Bryant and Liu, 1994; Powell and McMillan, 1994). The fact that our cell-free assay yielded similar results to these previous studies that used intracellular plasmid reactivation assays suggests that those differences in DSB mis-rejoining frequencies in radiosensitive and radioresistant cell lines (Powell and McMillan, 1994) may be due to inherent differences in nuclear protein composition and are not directly attributable to differences in proliferation rates between cell lines.

The biochemical basis for the differential level of DSB mis-rejoining in the nuclear protein extracts from radiosensitive and radioresistant tumour cells is presently unknown. It appears that cell-free plasmid reactivation assays are sensitive to DSB rejoining processes that are far more complex than simple ligation. Previous characterization of the cell-free plasmid reactivation assay used in this study suggests that multiple proteins are required in addition to the known ligases to rejoin restriction endonuclease-induced DSBs (Fairman et al., 1992). For example, fractionation of the protein extracts revealed that the level of plasmid rejoining is influenced by proteins within a specific fraction that was designated ‘rejoining enhancing proteins’ (Fairman et al., 1992). Recent data suggest that in vitro DSB rejoining assays that use restriction endonuclease-induced DSBs may detect only a subset of the pathways that are used in vivo to rejoin radiation-induced DSBs (Perrault et al., 1997). It has been suggested that restriction endonuclease-generated DSBs act as substrates for proteins that are involved in the (abnormal) conversion of DSBs into chromosomal aberrations (Bryant and Liu, 1994). This conclusion is supported by several studies on AT and hamster ird cells that exhibit no apparent defect in DSB rejoining; yet they are extremely sensitive to radiation (Taylor et al., 1975) and also exhibit high DSB mis-rejoining activities (Cox et al., 1984; Thacker and Ganesh, 1990; Ganesh et al., 1993; Powell et al., 1993; Luo et al., 1996) and enhanced levels of chromatid-type aberrations (Liu and Bryant, 1993, Bryant and Liu, 1994) in cell-free and intracellular plasmid reactivation assays. The strong correlation between the level of radiation-induced chromosomal aberrations and the radioresistivity of tumour (Lambin et al., 1994; Sasai et al., 1994) and human fibroblast (Russell et al., 1995) cells would certainly explain the correlation between the fidelity of DSB rejoining and SF values if the plasmid reactivation assay does detect the activity of enzymes involved in the processing of DSBs into chromosomal aberrations.

The identity of the enzymes that might be responsible for the differential level of DSB mis-rejoining in radiosensitive and radioresistant tumour cells remains largely speculative, however our characterization of the nuclear protein extract-induced mutations within the lacZα gene may have given some insight into their identity. Restriction endonuclease mapping and PCR amplification analysis of the mutant clones generated in our experiments suggest that the inactivation of the lacZα gene was attributable (45% of the time) to deletions of at least 40 bases and as much as 440 base pairs. These data are consistent with previous studies in which the majority of the mutations that lead to a defective lacZα gene (when cellular protein extracts from AT cells were used) was primarily deletions and insertions (Thacker et al., 1992). In those studies, these mutations were shown to arise because of illegitimate recombination, i.e. the DNA recombines at short sequence repeats that are frequently far apart (Thacker et al., 1992).

In conclusion, we have demonstrated that nuclear protein extracts from overtly DSB repair-proficient radiosensitive tumour cells are less capable of correctly rejoining EcoRI-induced DSBs than similar extracts from radioresistant tumour cells. Our data suggest that the higher level of DSB mis-rejoining in radiosensitive tumour cells is not a consequence of non-specific endonuclease activity, but rather due to an increase in non-conservative DSB rejoining activity, resulting in losses of between 40 and 440 base pairs of DNA. These data raise the possibility that the radiosensitivity of DSB repair-proficient human tumour cell lines may be partly determined by the predisposition of these cell lines to activate non-conservative DSB rejoining pathways.

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