The aim of this study was to identify potential biomarkers for radiosensitivity using the relationship between cell killing and the yield of excess chromatin fragments detected with the premature chromosome condensation (PCC) technique. This method was applied to primary cultured cells obtained from biopsies from patients. Six primary culture biopsies were obtained from 6 patients with carcinoma of the cervix before starting radiotherapy. The cultures were irradiated with two different LET carbon-ion beams (LET = 13 keV/μm, 77.1±2.8 keV/μm) and 200 kV X-rays. The carbon-ion beams were produced by Heavy Ion Medical Accelerator in Chiba (HIMAC). PCC was performed using the polyethylene glycol-mediated cell fusion technique. The yield of excess chromatin fragments were measured by counting the number of unrejoined chromatin fragments detected with the PCC technique after a 24-h post-irradiation incubation period. Obtained results indicated that cultures which were more sensitive to killing were also more susceptible to the induction of excess chromatin fragments. Furthermore there was a good correlation between cell killing and excess chromatin fragments among the 6 cell cultures examined. There is also evidence that the induction of excess chromatin fragments increased with increasing LET as well as cell-killing effect in the same cell culture. The data reported here support the idea that the yield of excess chromatin fragments detected with the PCC technique might be useful for predicting the radiosensitivity of cells contained in tumor tissue, and to predict responses to different radiation types.

**Introduction**

Carbon-ion beam radiotherapy has been used to treat cancer patients since 1994 at the National Institute of Radiological Sciences in Japan. It was realized early that it is very important to know which types of cancer were suitable for heavy-ion radiotherapy before beginning treatment. If an individual patient's radiosensitivity could be determined before treatment, heavy-ion radiotherapy could be used more effectively, and treatments could be designed for individuals. The goal of this study was to predict the radiosensitivity of individual tumors from different patients using a simple, rapid and highly sensitive biological end-point rather than cell biology methods, such as a colony-forming assay. A colony-forming assay requires time for colonies to form in culture, and the assay considers only cells which are capable of reproducing. However, in general most of the cells in a tissue may not be capable of reproduction. Chromosome aberrations in metaphase spreads are another useful method to predict cellular radiosensitivity, but these aberrations may not be always the same or comparable when cells are exposed to different types of radiation with different LET values (1,2).

Another method capable of detecting radiation-induced damage at the chromosomal level is the premature chromosome condensation (PCC) technique. Several methods are available to induce PCC, such as cell fusion-mediated and drug-induced methods. The former is carried out to transfer the maturation promoting factors of M-phase cells to interphase target cells mediating cell fusion technique, such as using Sendai virus, polyethylene glycol and electrofusion. The latter is carried out to use specific inhibitor of types 1 and 2A protein phosphatase, such as okadaic acid and calyculin A. The PCC technique is a very powerful method for detecting chromatin damage in G₁- or G₂-phase cells. It is also a very useful method for comparing responses to different radiation types, such as heavy ions and X-rays, because it overcomes a difficulty encountered when using conventional chromosome analysis in mitosis. Results found with conventional analysis are affected by different radiobiological responses caused by different types of radiation, such as cell cycle delay or interphase cell death (1,2). Induction of chromatin damage detected by the PCC technique in cells irradiated with either low- or high-LET
The expanded stock culture was frozen in liquid nitrogen until use (passage 2). Freshly thawed cells were sub-cultured once and then cells at passage 4 were used for experiments. The modal chromosome number was stably maintained in culture. The number of chromosomes (average ± standard deviation) in each culture at passage 4 was 44.0±1.7 for Cell-1, 47.2±2.2 for Cell-2, 47.3±2.4 for Cell-3, 46.8±2.3 for Cell-4, 45.9±1.1 for Cell-5 and 46.1±1.2 for Cell-6, respectively. These cell cultures displayed a fibroblast-like morphology when examined under the microscope. Mitotic XP2OS (SV40 transformed) cells, which were established from \textit{Xeroderma pigmentosum} complementation group A cells, and which were obtained from the Japanese Cancer Research Resources Bank (Cell line no. JCRB0301), were used as PCC inducers (19). The cells were cultured in Eagle's minimum essential medium (MEM) supplemented with 10% fetal bovine serum and kanamycin (60 mg/l) in a 5% CO₂ incubator at 37°C.

\textbf{Irradiation}. Carbon-ion beam irradiation was performed as previously reported (20-23). Briefly, each culture was inoculated into 25-cm² plastic flasks (BD Falcon™, 353014) for 3 days before irradiation, by which time the cells were confluent. Cells were irradiated with carbon-ion beams generated by the HIMAC accelerator at the National Institute of Radiological sciences (NIRS) in Japan. The initial energy of the carbon-ion beams was 290 MeV/n. The energy at the irradiation site was obtained by comparing the calculated and measured depth-dose distribution. The experimental dose distributions were measured with a small parallel-plate ionization chamber placed at the irradiation site (21). Two different beams with different LET values were used with Lucite absorbers of different thicknesses to change the energy of the beams. At the sample position, it was estimated that the LET∞ values (dose averaged LET) of all experiments was 13.3 keV/μm for lower LET value and 77.1±2.8 keV/μm for higher LET value (the mean value ± the standard error). The absorber of 28.5±0.1 mm in thickness was used for making a higher LET value. The dose rate of both LET beams was approximately 1.2 Gy/min. For comparison, 200 kV X-rays were used at a dose rate of 0.9 Gy/min. All of the irradiations were carried out at room temperature.

\textbf{Clonogenic cell-survival assay}. After irradiation, cells were immediately plated onto 100-mm plastic dishes (BD Falcon™, 353003) at a density which permitted the formation of 60-70 viable colonies per dish. Colonies were fixed with 100% ethanol and stained with a 5% Giemsa solution (Wako Pure Chemical Industries Ltd.) after a 14-day incubation period. Colonies consisting of more than 50 cells were scored as a

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**Table I. Information on the donors.**

<table>
<thead>
<tr>
<th>Cell</th>
<th>Age of donor</th>
<th>Primary site</th>
<th>Stage</th>
<th>Histology</th>
<th>Treatment</th>
<th>Control of primary site</th>
<th>Prognosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell-1</td>
<td>48</td>
<td>Uterine cervix</td>
<td>IVb</td>
<td>Sq</td>
<td>Radiotherapy alone</td>
<td>Controlled</td>
<td>13 mo. DOD</td>
</tr>
<tr>
<td>Cell-2</td>
<td>48</td>
<td>Uterine cervix</td>
<td>IIb</td>
<td>Sq</td>
<td>Radiotherapy alone</td>
<td>Controlled</td>
<td>98 mo. NED</td>
</tr>
<tr>
<td>Cell-3</td>
<td>43</td>
<td>Uterine cervix</td>
<td>Ib2</td>
<td>Ad</td>
<td>Radiotherapy alone</td>
<td>Controlled</td>
<td>98 mo. NED</td>
</tr>
<tr>
<td>Cell-4</td>
<td>64</td>
<td>Uterine cervix</td>
<td>Ib</td>
<td>Ad</td>
<td>Radiotherapy alone</td>
<td>Controlled</td>
<td>66 mo. DOD</td>
</tr>
<tr>
<td>Cell-5</td>
<td>71</td>
<td>Uterine cervix</td>
<td>IVa</td>
<td>Sq</td>
<td>Radiotherapy alone</td>
<td>Recurred</td>
<td>28 mo. DOD</td>
</tr>
<tr>
<td>Cell-6</td>
<td>56</td>
<td>Uterine cervix</td>
<td>IIb</td>
<td>Sq</td>
<td>Radiotherapy alone</td>
<td>Controlled</td>
<td>17 mo. DOD</td>
</tr>
</tbody>
</table>

*Sq, squamous cell carcinoma; Ad, adenocarcinoma; NED, no evidence of disease; DOD, dead of disease; mo., months.*
The plating efficiency of the 6 cell types ranged from 15 to 40% under the experimental conditions used.

Detection of chromatin breaks in G1/G0 using PCC. The experimental method used for the detection of excess chromatin fragments in G1/G0 cells with PCC was used as previously reported (19). Briefly, irradiated cells were incubated for 24 h in a 5% CO2 incubator to allow rejoining of induced chromatin breaks. Unrejoined chromatin fragments were scored as fragments remaining in G1/G0 cells with PCC. Mitotic XP2OS cells, accumulated after a 6-h incubation period in the presence of 0.1 μg/ml demecolchicine (Wako Pure Chemical Industries Ltd.), were mixed with an equal number (1x10^6) of irradiated cells in a polypropylene tube (BD Falcon™, 352059) and incubated with 0.15 ml of 50% (w/v) polyethylene glycol (PEG; M.W.=1540, cat. no. 783641, Roche Diagnostics GmbH, Germany) in 75 mM Hepes for 1 min for cell fusion. Fused cells were resuspended in 5 ml of complete medium containing 0.1 μg/ml demecolchicine, and incubated in a CO2 incubator for 1 h. These cells were subsequently treated with a 75-mM KCl solution for 20 min at room temperature, and fixed in 3:1 methanol:acetic acid. The cell suspension was dropped onto slides, air dried and stained with a 5% Giemsa solution. PCC preparations of 50 G1/G0 phase cells were scored under a light microscope. The yield of induced excess chromatin fragments per cell was estimated to be the number of remaining chromatin fragments after a 24-h post-irradiation incubation period in excess of the number of chromatin fragments found in the non-irradiated cells.

Results
Dose-response curves using a clonogenic cell-survival assay are shown in Fig. 1. These results indicate that cultures, which were sensitive to X-rays, were similarly sensitive to both low- and high-LET carbon-ion beams. Based on the survival curves, parameters for cellular radiosensitivity, such as α, β and D10 values, were calculated using the linear-quadratic model (Table II). The RBE values listed in Table III relative to 200 kV X-rays ranged from 0.96 to 1.29 for the low-LET beam, and 1.36-2.49 for the high-LET beam. Notably the RBE of both Cell-3 and Cell-5 cells for the low-LET beam was below 1 at the 10% survival level.

Fig. 2 shows the dose-response curves for the induction of excess chromatin fragments after a 24-h post-irradiation incubation period. Linear dose-responses were observed for each cell culture and each radiation type in the dose range used. The most sensitive cell line, Cell-3, with the cell survival assay showed the highest frequency of the induction of excess chromatin fragments.
chromatin fragments among the 6 cell cultures examined after exposure to either X-rays or carbon ions. The number of excess chromatin fragments observed per Gy for X-rays, the low-LET beam, and the high-LET beam were 1.50, 2.95 and 6.18 for Cell-1; 1.78, 3.21 and 7.36 for Cell-2; 1.99, 3.68 and 7.86 for Cell-3; 1.29, 1.78 and 2.29 for Cell-4; 0.99, 2.06 and 2.28 for Cell-5; and 1.10, 1.98 and 2.31 for Cell-6, respectively. RBE values calculated using the number of excess PCC breaks per Gy per cell relative to 200 kV X-rays are listed in Table III.

To clarify the relationship between radiosensitivity and excess chromatin fragments, the D 10 values for cell survival were plotted against the number of excess chromatin fragments per cell per Gy in Fig. 3. Data from a previous study using human cell lines (19) was plotted in Fig. 3, and the D 10 values of the 6 cultures established from 6 cervical carcinoma patients biopsies plotted on this curve also correlated well with the number of excess chromatin breaks per Gy per cell relative to 200 kV X-rays are listed in Table III.

Table III. Summary of RBE values for cell-killing effect and induction of excess PCC fragments per Gy per cell relative to 200 kV X-rays.

<table>
<thead>
<tr>
<th>Cell</th>
<th>Excess PCC fragmenta</th>
<th>Excess PCC fragmentb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell-1</td>
<td>1.24</td>
<td>1.97</td>
</tr>
<tr>
<td>Cell-2</td>
<td>1.17</td>
<td>1.80</td>
</tr>
<tr>
<td>Cell-3</td>
<td>0.96</td>
<td>1.85</td>
</tr>
<tr>
<td>Cell-4</td>
<td>1.29</td>
<td>1.78</td>
</tr>
<tr>
<td>Cell-5</td>
<td>0.99</td>
<td>2.06</td>
</tr>
<tr>
<td>Cell-6</td>
<td>1.10</td>
<td>1.98</td>
</tr>
</tbody>
</table>

aRBE was calculated using D 10 values (Gy). bRBE was calculated using the number of excess PCC fragments/Gy/cell.

Discussion

The PCC technique is a very powerful method for detecting chromatin damage in G1- or G2-phase cells. Notably, it offers a very useful comparison for responses to different radiation types, such as heavy ions and X-rays, because it can overcome the difficulty involved with conventional chromosome analysis in metaphase spreads. The conventional method is affected by the differences in LET-dependent and/or dose-dependent radiobiological effects produced by the different qualities of radiation, and these effects are seen as either cell cycle delays or interphase cell death (24-31). Consequently, it may be possible to underestimate the extent of chromosome damage produced by different radiation types by scoring with a single fixed sampling time using conventional analysis of metaphase spreads and not the PCC technique (1,2). Coco-Martin et al (32) demonstrated a good correlation between cell survival and the total number of chromosome aberrations detected with conventional metaphase analysis and also they suggested that combining doses markedly improved the correlation between cell survival and chromosome aberration yield. On the contrary several reports have shown that cell-killing effects did not always correlate well with the chromosome aberrations detected with conventional cytogenetic methods, especially when comparing responses after exposure to different types of radiation (1,2,33). However, investigations using the PCC technique to measure chromosomal breaks produced by low- and high-LET radiation also indicate that there is a linear dose response in the same dose range used for cell-survival experiments with different in vitro cell systems (3-10,18,19). Pantelias and Maillie (14) examined the in vitro dose-response curve of human blood mononuclear cells exposed to X-rays using the PCC technique as a biological dosimeter and found that it could be an effective system.

Several reports have shown that the number of excess chromatin fragments detected using the PCC technique in cells exposed to high-LET radiation was higher than that seen in cells exposed to low-LET radiation. Thus there was an LET-dependent pattern (5,7,9,10,18). In previous studies using normal human cells, approximately 95% of 137Cs-γ-ray induced PCC breaks were rejoined, while only 50% of 110 keV/μm- or 124 keV/μm-carbon-ion induced PCC breaks were rejoined.
after a 24-h post-irradiation incubation period (10). The percent of unrejoined breaks varied ranging from 23 to 50%, depending on the LET value of the carbon-ion beam, which ranged from 22 to 230 keV/μm (10). Some studies have also indicated that the radiosensitivity of human tumor-cell lines or normal human cells after exposure to low-LET photons and high-LET radiation correlated well with the number of excess chromatin fragments (5,10,15-18,34-36) as well as with the induction of non-repaired DNA double-strand breaks (35,37-43). However, as pointed out in the report by Dikomey et al (42), the radiation doses required to detect DNA strand breaks (~90 Gy) were much higher than those necessary to observe cell-killing (up to 9 Gy), while cytogenetic studies using the PCC technique could be completed within a dose range similar to that used for cell-killing studies. This defines another advantage for the PCC technique: it is possible to compare these different biological end-points (cell killing and PCC breaks) in a similar radiation dose range.

In a previous study, a close relationship was demonstrated between cell-killing effects and the number of excess chromatin fragments measured with G1/G0 PCC in 6 different human cell lines irradiated with different types of radiation, such as X-rays and carbon ions (19). The D10 values of 6 cultures established from 6 cervical carcinoma patient biopsies also correlated well with the number of excess chromatin fragments in a similar manner (Fig. 3). Because it will often be impossible to obtain radiosensitivity data for each individual patient using clonogenic cell-survival assays from a piece of tissue obtained from a tumor biopsy, it would be useful to be able to predict individual radiosensitivities using some other biomarker. In the present study, an attempt was made to estimate cellular radiosensitivity from cultures of 6 cervical carcinoma patient biopsies, and to compare this to the number of excess PCC fragments per Gy per cell found in each culture. To make this correlation, D10 values were plotted on the vertical axis and the number of excess PCC fragments per Gy per cell was plotted on the horizontal axis in Fig. 3. In this plot it was possible to determine the relationship between these two parameters experimentally. In Fig. 4, the same parameters were plotted and the line from Fig. 3 generated from the cell line data was added. Using this graph, the number of excess PCC fragments per Gy per cell

Figure 3. Relationship between D10 values and the number of excess chromatin fragments per Gy per cell for each radiation type in 6 different human cell lines (closed square) and 6 cultures obtained by biopsy from 6 patients (open square). The data for human cell lines was taken from Suzuki et al (19). The curve was generated from the data for 6 cell lines reported in Suzuki et al (19).

Figure 4. Estimation of D10 values for cell survival in cultures was obtained from biopsies from 6 patients using the data for excess chromatin fragments. The data for the number of excess fragments was used to read the D10 value from the Y axis by using the standard curve generated in Fig. 3 from cell line data (Suzuki et al) and plotted here.
on the X axis was used to obtain a D_{10} value for the biopsy samples by using the line generated by cell line data from Table IV. The number of excess PCC fragments per Gy per cell was used to read a D_{10} value directly on the Y axis.

Fortunately, it was possible to obtain D_{10} values experimentally from the 6-biopsy cultures used and compare them directly with the values obtained with the PCC data and the cell line standard curve shown in Fig. 4; these are summarized in Table IV. The difference in the D_{10} values between the two different protocols varied from 0 to 15.7%. However, we can predict the radiosensitivity of the cultures established from the 6 cervical carcinoma patient biopsies with a variation of less than 16% with this protocol.

In conclusion, several methods are available to induce PCC, such as Sendai virus-mediated cell fusion, polyethylene glycol-mediated cell fusion and drug-induced method. We tried to use Calyculin A for our cell system. Unfortunately, the fibroblast-mediated cell fusion and drug-induced method. We tried to use such as Sendai virus-mediated cell fusion, polyethylene glycol-less than 16% with this protocol.

The possibility of utilizing the PCC technique as a biological assay to measure the intrinsic radiosensitivity of cells contained in tumor tissue, and as a biological fundamental study for predicting the radiosensitivity of cells for clinical radiotherapy, notably radiotherapy using high-LET heavy ions.

Acknowledgements

The authors wish to thank the staff of the HIMAC for their help with the carbon-ion beam irradiations and Dr Leon Kapp at the University of California for his critical reading of the manuscript. This study was supported in part by grants from the Japanese Ministry of Education, Science and Culture, and in part by the program for ‘Ground-based Research for Space Utilization’ promoted by the Japan Space Forum.

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