G2-checkpoint abrogation in irradiated lymphocytes: A new cytogenetic approach to assess individual radiosensitivity and predisposition to cancer

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Abstract. Increased yield of chromatid breaks, following in vitro G2-phase lymphocyte irradiation, can be a marker of individual radiosensitivity and cancer predisposing genes whose role is to respond to DNA damage. Mutations or polymorphisms of genes encoding DNA repair pathways may underlie the increased chromosomal radiosensitivity. However, genes that facilitate DNA damage recognition, using signal transduction pathways to activate cell cycle arrest and preserve genomic integrity, are perhaps the most important determinant. Based on the latter hypothesis, an individual radiosensitivity parameter (IRP) is introduced, which expresses, at individual level, the G2-checkpoint potential to facilitate DNA damage recognition and repair of radiation-induced chromosomal damage during G2 to M-phase transition. Based on this parameter a new methodology for assessment of individual radiosensitivity is proposed, which involves G2-checkpoint abrogation by caffeine to obtain the IRP values. To evaluate the proposed methodology, blood samples from 52 healthy donors were taken for inter-individual radiosensitivity analysis using both the conventional G2 chromosomal radiosensitivity assay as well as the new approach using caffeine-induced G2-checkpoint abrogation. The two assays were compared in experiments using samples from 5 hypersensitive patients, 3 AT-homozygotes, 3 AT-heterozygotes, and the GM15786, GM03188A, GM09899, HCC1937 and MCF-7 cell lines. Using the G2 chromosomal radiosensitivity assay, donors are predicted as G2 radiosensitive or normal, while according to the new approach, individuals can be classified as highly radiosensitive, radiosensitive, normal, radioresistant and highly radioresistant. Overall, the new approach provides better individual radiosensitivity discrimination and intra-experimental reproducibility. Therefore, the proposed methodology using IRP values may provide a clinically applicable predictive assay for individual radiosensitivity and predisposition to cancer.

Introduction

Identification of individuals in the normal population with increased inherent radiosensitivity is of relevance for their protection from the adverse effects of radiation and other genotoxic agents, with applicability in occupational health and safety. Also, in radiation oncology, since it is known that the maximum therapeutic radiation dose is limited by the occurrence of normal tissue adverse effects (1), the individualization of radiation therapy protocols can be uniquely facilitated by the prediction of individual radiosensitivity (2-7). Furthermore, as it is well established that cellular radiosensitivity is linked to cancer predisposition (8,9), population screening for hypersensitivity to radiation is of great importance (reviewed in ref. 10).

In recent years, evidence has been obtained to support the hypothesis that possession of variants in genes, which play a role in radiation response, is predictive for individual radiosensitivity and the development of adverse effects after radiotherapy (7,11,12). Yet, in vitro irradiation of peripheral blood lymphocytes using techniques measuring DNA or chromosome damage, and in particular G2 chromosomal radiosensitivity (G2 assay), is also extensively applied for individual radiosensitivity assessment (13-17). Specifically, the cell cycle based G2 assay involves the in vitro irradiation of peripheral blood lymphocytes in G2 phase to induce DNA damage, which is processed during G2 to M-phase transition so that the residual lesions can be visualized and quantified at metaphase as chromatid breaks. Variation in the yield of chromatid breaks between individuals has been correlated to variation in radiosensitivity and predisposition to cancer (13,18-20). Current protocols for the G2 assay require stringent technical conditions (16) to generate reproducible and meaningful results, and variability in yields of induced chromatid breaks in different samples from the same individual is a recognized problem. The original method developed by Sanford et al (13) has been refined by Scott et al (20) in order to minimize problems associated with

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reproducibility and obtain better discrimination. At present, however, comparisons of the results between different laboratories are difficult since modifications in the protocols used result in variations in the yield of chromatid breaks and reproducibility (16,21). Furthermore, an important issue associated with the sensitivity of G2 assay is that, although the mean radiation-induced yield of chromatid breaks in certain groups, such as cancer patients, is significantly higher than in normal individuals, considerable overlap is found in the yield of G2 chromatid breaks (19,22-27). Therefore, a G2 radiosensitive cut-off value has to be calculated from the control population in order to assess individual radiosensitivity. In some of the studies, the 90th percentile of the distribution of results from a normal population is taken as cut-off point. This cut-off point, as introduced first by Scott et al (19), is obviously very helpful, even though it is arbitrary and very dependent on the extent of the healthy control group (17).

In the present study, a new approach is proposed for individual radiosensitivity assessment aiming to provide better discrimination of individual radiosensitivity and intra-experimental reproducibility, as well as to minimize the required stringent technical conditions of the current protocol used for the G2 assay. The methodology is based on our previous work related to the mechanisms underlying conversion of DNA damage into chromatid breaks, which is linked to cell-cycle and feedback control mechanisms and, particularly, to the cdk1/cyclin-B (MPF) complex activity that regulates G2 to M-phase transition (22,28-30). It exploits also our observation that checkpoint abrogation in G2 phase by means of caffeine compromises repair of radiation-induced chromosomal damage, which can be increased to the levels obtained in the case of highly radiosensitive ataxia telangiectasia (AT) patients (31,32). According to the new approach, two yields of chromatid breaks are obtained for each individual in response to radiation-induced DNA damage during G2 to M-phase transition of cultured peripheral blood lymphocytes. The first represents residual chromosomal damage under the effect of the G2 checkpoint, and the second under G2-checkpoint abrogation by means of caffeine. We propose that the difference between these two G2 values can be employed as an individual radiosensitivity parameter (IRP) to assess intrinsic radiosensitivity and predisposition to cancer since it reflects differences among individuals in specific mutations or polymorphisms in genes that control: a) DNA repair capacity, b) cdk1/cyclin B activity and c) G2-checkpoint activation after irradiation during the G2 to M-phase transition. Essentially, the obtained IRP value expresses, at individual level, the G2-checkpoint potential to facilitate DNA damage recognition and repair of radiation-induced chromosomal damage during G2 to M-phase transition. Consequently, the lower the IRP value, the higher the radiosensitivity of the individual and, as the IRP value approaches to zero, the predictive individual radiosensitivity will be close to that of AT patients.

Materials and methods

Blood samples and cell lines. After obtaining consent, heparinized blood samples (5 ml) were taken from 52 healthy control subjects for inter-individual variation analysis of radiosensitivity using both the conventional G2 assay, as described in previous work (22), and the new approach that, in addition, takes into consideration for each individual the yield of chromatid breaks obtained when G2 checkpoint is inactivated after irradiation, by means of caffeine. Blood samples were also taken from five cancer patients who experienced adverse radiation effects from their radiation therapy treatment, three clinically characterized AT homozygotes and three obligate AT heterozygotes. Peripheral blood lymphocytes were cultured adding 0.5 ml of whole blood to 5 ml of McCoy’s 5A medium supplemented with 10% FBS, 1% phytohaemagglutinin (PHA), 1% glutamine and antibiotics (penicillin: 100 U/ml; streptomycin: 100 μg/ml). Cultures were incubated for 72 h before their use for radiosensitivity estimation experiments. In addition, three EBV-transformed lymphoblastoid cell lines (LCL) were used: GM 15786, derived from an AT patient, GM03188A from an obligatory ATM heterozygote and GM 09899, derived from a normal individual. Cells were maintained in RPMI (Biochrom AG, Germany) supplemented with HEPES and sodium bicarbonate, 15% fetal bovine serum (FBS), 2 mM L-glutamine and 100 μg/ml streptomycin and 100 U/ml penicillin. Experiments were also carried out using breast cancer cell lines, the BRCA1-defective HCC1937 cell line, homozygous for the BRCA1 5382C mutation, and the MCF-7 human breast adenocarcinoma cell line. Both HCC1937 and MCF-7 cell lines were purchased from the American Type Culture Collection (Rockville, MD, USA). HCC1937 cells were grown in RPMI-1640 medium (Biochrom AG, Germany), while MCF-7 cells were grown in Dulbecco’s modified Eagle’s medium (DMEM) (Biochrom AG, Germany). Culture media were supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine and antibiotics. All incubations took place at 37°C in a humidified incubator in an atmosphere of 5% CO₂ and 95% air.

Irradiation conditions and the new individual radiosensitivity assay. Irradiation was carried out in a GammaCell 220 irradiator (Atomic Energy of Canada Ltd., Ottawa, Canada) at room temperature and at a dose rate of 1 Gy/min. PHA was dissolved in water at a concentration of 0.24 mg/ml. Caffeine was prepared as a 100 mM stock solution in PBS. Proliferating cells were irradiated in vitro in G2-phase with 1 Gy and the culture was divided immediately after so that, one half was treated with caffeine (4 mM) and incubated for 30 min at 37°C to allow division of cells irradiated at mitosis, while the other half was cultured for the same time period without the presence of caffeine. Colcemid was subsequently added to the cell cultures for 60 min. At 90 min post irradiation, cells were collected by centrifugation, treated in 75 mM KCl for 10 min, fixed in methanol: glacial acetic acid (3:1 v/v) and processed for cytogenetics analysis. Standard procedures were used for chromosome preparation and staining and chromosomal damage was visualized and quantified as chromatid breaks in cells at metaphase. For each experimental point, approximately 50 cells were scored for chromatid damage based on standard criteria. Chromatid breaks and gaps were scored, the latter only when longer than a chromatid width. Light microscopy was coupled with an image analysis
system (MetaSystems, Germany) to facilitate scoring. The spontaneous aberration yield was subtracted to obtain the radiation-induced G2 yield of chromatid breaks. Standard deviations of the mean values from three independent experiments were calculated. Following this protocol, chromosomes were analyzed at the subsequent metaphase and two distinct yields of chromatid breaks were obtained for each individual or cell line after G2-phase irradiation. One yield corresponded to that obtained when the conventional G2 assay was applied, and the other represented, for each individual, the maximum yield of chromatid breaks obtained when G2 checkpoint was inactivated by means of caffeine.

Results

Inter-individual variation in radiosensitivity obtained using the conventional G2 assay. For the conventional G2 assay, peripheral blood lymphocyte cultures were stimulated and 72 h thereafter cells were irradiated with 1 Gy. Chromosome preparation and yield analysis of chromatid breaks were carried out as described in the ‘Materials and methods’ section. Fig. 1 shows the inter-individual variation in chromosomal radiosensitivity as measured for 52 healthy blood donors. Experiments were carried out on a single blood sample per blood donor and standard deviations of the mean values from three independent experiments were calculated. As mentioned above, the identification of radio-sensitive individuals in the conventional G2 assay involves the calculation of a radiosensitive cut-off value above which an individual can be considered to be radiosensitive. This radiosensitive cut-off value can be statistically set at the 90th percentile, i.e., the value below which 90% of the controls lie (19). Using this approach for the 52 healthy blood donors, we obtained a cut-off value of 3.7 chromatid breaks per metaphase and 9.6% of the control population was defined as G2 radiosensitive.

Inter-individual variation in radiosensitivity as predicted using the new approach. Peripheral blood lymphocytes were irradiated in vitro in G2 phase and subsequently cultured with and without caffeine, so that two distinct yields of chromatid breaks were obtained when chromosomes were analyzed at the subsequent metaphase. While Fig. 2A shows the chromatid breaks as visualized at metaphase following the conventional G2 assay, Fig. 2B shows the chromatid breaks obtained following G2-checkpoint abrogation by means of 4 mM of caffeine. Under these experimental conditions, an increased yield of chromatid breaks similar to that obtained in AT patients can be observed. For each individual, these two yields of chromatid breaks represent residual chromosomal damage with and without G2-checkpoint activation in response to radiation-induced DNA damage. The yield obtained when G2 checkpoint is inactivated by means of caffeine is used as an internal control value and represents a maximum G2 radiosensitivity value for each individual. The difference between the two yields obtained is characteristic for each individual, as it reflects differences in specific mutations or polymorphisms in genes that control recognition and the processing of radiation-induced DNA damage during G2 to M-phase transition. We define the difference between these two yields of chromatid breaks as individual radiosensitivity parameter (IRP). Fig. 3 shows the inter-individual variation in IRP values obtained from the 52 healthy control blood donors. The inter-individual variation in radiosensitivity, as expressed by variation in the IRP values, is fitted by a normal distribution with a mean value (MV) of 4.7 chromatid breaks per metaphase, a standard deviation (SD) of 0.8, and 17% coefficient of variation (CV). According to the proposed method, individuals may be defined as normal when IRP=MV±SD, sensitive when MV-2SD≤IRP<MV-SD, highly sensitive when IRP<MV-2SD, resistant when MV+SD<IRP<MV+2SD, and highly resistant when IRP>MV+2SD. According to these criteria, 77% of healthy donors
were found to have a normal radiosensitivity response, 9% were radiosensitive, 12% were radioresistant, while 2% were highly radioresistant.

Evaluation of the proposed G2 chromosomal radiosensitivity assay. Three sets of experiments were carried out to investigate whether better discrimination in individual radiosensitivity
and intra-experimental reproducibility can be obtained by means of the proposed new approach as compared to the conventional G2 assay. In the first set of experiments, heparinized blood samples were obtained from five cancer patients who experienced adverse radiation effects from their radiation therapy treatment (radio-sensitive positive controls). Individual radiosensitivity was estimated using both the conventional G2 assay and the new approach. The G2 yields with and without G2-checkpoint activation in response to radiation-induced DNA damage during G2 to M-phase transition, as well as the IRP values obtained are shown in Table I. According to the conventional G2 assay and the calculated cut-off value of 3.7 chromatid breaks per metaphase cell, only three out of the five positive controls (radio-sensitive patients) who had G2 yields of 4.9, 5.2 and 5.8 chromatid breaks per cell (Table I) should belong to the radiosensitivity range (Fig. 1). However, using the new approach for the assessment of individual radiosensitivity and the IRP values shown in Table I, all five positive controls are predicted to be radiosensitive (Fig. 3).

In the second set of experiments, heparinized blood samples from three clinically characterized AT homozygotes and three obligate AT heterozygotes were used to study individual radiosensitivity by means of the conventional G2 assay as well as the proposed methodology using IRP values. The results are shown in Table II. Even though the AT homozygotes are predicted to be radiosensitive by means of the G2 assay, according to the proposed methodology using the IRP values they are characterized as highly radiosensitive (Fig. 3). Furthermore, when the obligate AT heterozygotes are tested for intrinsic radiosensitivity, they are predicted to be in the upper normal radiosensitivity range by means of the conventional G2 assay, whereas using the IRP values and the new methodology the AT heterozygotes (Table II) are predicted to be in the radiosensitive range (in the range of 3.1-3.9 chromatid breaks/cell).

The purpose of the third set of experiments was to test the reproducibility of the two methodologies with respect to inter- and intra-individual variation. MCF-7 cells, HCC1937 cells and three lymphoblastoid cell lines, GM15786, GM03188 and GM09899, derived from an AT patient, an obligatory AT heterozygote, and a normal individual, respectively, were used in this set of experiments. The G2 assay and the new approach using IRP values were applied to those five cell lines to measure inter- and intra-individual variation in radiosensitivity. For this purpose, two samples were taken from each cell line at three different time points. Since the MCF-7 and HCC1937 cells have a modal number of 81 and 100 chromosomes, respectively, the chromatid breaks per cell were normalized to 46 chromosomes. The results are presented in Table III, where coefficients of variation are compared between the conventional G2 assay and the new proposed methodology with IRP values, to evaluate intra-experimental reproducibility.

**Discussion**

Experimental results from patients with chromosomal instability syndromes and genetic susceptibility to carcinogenesis show similar cellular features as well as increased radiation sensitivity and have provided a link between chromosomal radiosensitivity and cancer predisposition. In fact, when the G2 chromosomal radiosensitivity assay (G2 assay) was applied to cancer-prone syndromes such as AT patients, Nijmegen breakage syndrome (NBS), and AT-like disorder (ATLD), and to many cancer patients as well, as reviewed in ref. 10, an increased chromosomal radiosensitivity was obtained (13,19,22,32,33). Early studies and data obtained using the G2 assay on lymphocytes and skin fibroblasts were carried out and reported by Sanford, Parshad and colleagues at the National Cancer Institute (NCI), USA (34-36). However, their results could not be reproduced in another laboratory (33) and a modified protocol that generated promising results was proposed (9,19,37,38). Recently, the radiosensitivity of human lymphocytes measured using a G0 or G2 assay has been linked with an individual's risk of developing normal tissue complications following radiation therapy, and a theoretical classification into three categories.
(resistant, normal and sensitive) was introduced (6,15). There was, however, no inter-assay correlation between the G0 and G2 sensitivity demonstrating that these two sensitivities depend on different genetic factors.

Although repair mechanisms such as homologous recombination and non-homologous end joining are important responses to double-strand DNA and chromosomal damage (39-43), cell-cycle regulation is perhaps the most important determinant of radiation sensitivity in G2 phase cells (32) and therapeutic radiosensitivity could be improved through modulation of the cell cycle (44). This hypothesis is corroborated by the results shown in Fig. 2B. Indeed, the presence of caffeine increases the yield of chromatid breaks in healthy donors to the levels obtained when lymphocytes from the highly radiosensitive AT patients are irradiated. Even though caffeine is known to influence radiation induced chromosomal damage and has been used to study inter-individual variation in chromosomal aberrations (47) and detection of AT heterozygotes (48), the precise mechanism of caffeine’s action is not well understood. This may be linked to the cell-cycle modulation and G2-checkpoint abrogation by means of caffeine and particularly, to the activity of the cdk1/cyclin-B complex and the subsequent changes in chromatin conformation that facilitate conversion of DNA damage into chromatid breaks. As we have already reported in previous work, following exposure to ionizing radiation, chromatin conformational changes could be observed in the DNA with the help of a specific probe. These changes are associated with a decrease in the yield of chromatid breaks in the AT patients, whereas there is no such decrease in healthy donors.

**Table II. G2 chromosomal radiosensitivity obtained for three AT heterozygotes and three AT homozygotes patients.**

<table>
<thead>
<tr>
<th>Donors</th>
<th>Active G2 checkpoint</th>
<th>G2 checkpoint abrogated by caffeine</th>
<th>Individual radiosensitivity parameter (IRP)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Chromatid breaks/cell</td>
<td>Chromatid breaks/cell</td>
<td>Chromatid breaks/cell</td>
</tr>
<tr>
<td>ATM+/-</td>
<td>A 3.5±0.6</td>
<td>7.1±1.0</td>
<td>3.6±0.3</td>
</tr>
<tr>
<td>ATM+/-</td>
<td>B 3.9±0.7</td>
<td>7.3±0.9</td>
<td>3.4±0.2</td>
</tr>
<tr>
<td>ATM+/-</td>
<td>C 2.9±0.5</td>
<td>6.8±0.9</td>
<td>3.9±0.3</td>
</tr>
<tr>
<td>ATM-/-</td>
<td>D 6.5±0.9</td>
<td>7.9±1.1</td>
<td>1.4±0.09</td>
</tr>
<tr>
<td>ATM-/-</td>
<td>E 6.9±1.0</td>
<td>7.8±0.9</td>
<td>0.9±0.08</td>
</tr>
<tr>
<td>ATM-/-</td>
<td>F 7.9±1.0</td>
<td>8.6±1.0</td>
<td>0.7±0.08</td>
</tr>
</tbody>
</table>

Conventional G2 assay was applied (active G2-checkpoint values) as well as the new approach with G2-checkpoint abrogation by caffeine, in order to obtain the IRP values to be checked against the ranges showed in Fig. 3. Using the conventional G2 assay, only one of the ATM+/- patients is predicted to be radiosensitive, whereas all three ATM+/- patients are predicted as radiosensitive when IRP values are used. All three ATM-/- patients are predicted to be radiosensitive by means of the G2 assay, whereas they are classified as highly radiosensitive according to the IRP values obtained. Mean values ± SD are calculated from three independent experiments.

**Table III. G2 chromosomal radiosensitivity for five cell lines as analyzed by means of the conventional G2 assay and the G2-checkpoint abrogation by caffeine to obtain the IRP values.**

<table>
<thead>
<tr>
<th>Cell lines</th>
<th>Active G2 checkpoint</th>
<th>G2 checkpoint abrogated by caffeine</th>
<th>Individual radiosensitivity parameter (IRP)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Chromatid breaks/cell</td>
<td>SD  CV (%)</td>
<td>Chromatid breaks/cell</td>
</tr>
<tr>
<td>MCF-7</td>
<td>6.3c</td>
<td>0.8  12.7</td>
<td>10.1c</td>
</tr>
<tr>
<td>HCC1937</td>
<td>3.1c</td>
<td>0.7  22.6</td>
<td>5.1c</td>
</tr>
<tr>
<td>GM03188</td>
<td>4.0</td>
<td>0.6  15.0</td>
<td>8.2</td>
</tr>
<tr>
<td>GM15786</td>
<td>8.9</td>
<td>1.1  12.4</td>
<td>9.8</td>
</tr>
<tr>
<td>GM09899</td>
<td>2.0</td>
<td>0.3  15.0</td>
<td>9.2</td>
</tr>
</tbody>
</table>

Better individual radiosensitive discrimination is predicted with the new approach and, moreover, the lower CV values obtained suggest better intra-experimental reproducibility when the new approach is applied. Mean values ± SD are calculated from three independent experiments carried out at various time intervals. Mean value normalized to 46 chromosomes.
ation changes play a vital role in the formation of chromatid breaks during premature chromosome condensation induction, as well as during G2 to M-phase transition (22,28-30). Therefore, the yield of chromatid breaks obtained in the presence of caffeine can be used for each individual as an internal control value that provides maximum chromosomal damage when the G2 checkpoint is inactivated. Since fewer chromatid breaks are obtained when G2 checkpoint is not inactivated by caffeine (Fig. 2A), the difference between these two yields is defined as the individual radiosensitivity parameter (IRP). This parameter expresses the cell potential for DNA damage recognition and repair of chromosomal damage during G2 to M-phase transition and its value depends on the effectiveness of the G2 checkpoint. Consequently, the lower the IRP value, the higher the radiosensitivity of the individual and, as the IRP value approaches zero, the predictive individual radiosensitivity will be close to that of AT patients.

The results obtained demonstrate that the new method provides better discrimination of radiosensitivity prediction at individual level and also intra-experimental reproducibility. Using the conventional G2 assay, a cut-off value based on the 90th percentile classifies the individual radiosensitivity in two groups, normal and radiosensitive (Fig. 1), while using the new approach, individuals can be classified as highly radiosensitive, radiosensitive, normal, radioresistant or highly radioresistant (Fig. 3). As shown in Fig. 1, a percentage of 9.6% from the 52 healthy donors can be classified as radiosensitive, whereas, according to the proposed method, a more detailed classification can be obtained. Indeed, 2% are defined as highly radioresistant, 12% as radioresistant, 77% as normal, 9% as radiosensitive, and 0% as highly radiosensitive. The advantage of the proposed methodology is clearly shown when blood samples from five cancer patients, who were clinically characterized as radiosensitive, were tested for individual radiosensitivity using both methodologies. Two out of the five positive controls (with adverse reactions) had G2 values of 3.3 and 3.5 chromatid breaks per cell (Table I) and, therefore, were predicted to be in the normal radiosensitivity range. However, using the new approach for the assessment of individual radiosensitivity and the IRP values shown in Table I, all five positive controls were predicted to be radiosensitive. In addition, when three obligate AT heterozygotes were tested for intrinsic radiosensitivity, they were predicted to be in the upper normal radiosensitivity range by means of the conventional G2 assay. However, using the new methodology, the IRP values obtained (Table II) show that all three AT heterozygotes were in the radiosensitive range. In the case of AT homozygotes patients (Table II), they were predicted radiosensitive by means of the G2 assay. Nevertheless, as they are expected to be, they were characterized as highly radiosensitive when the proposed new methodology was used. Individual radiosensitive prediction on cell lines further proved the advantages of the proposed new methodology. As can be seen in Table III, the MCF-7 cells are classified as sensitive by both methodologies while the HCC1937 cells, with a G2 yield of 3.1 chromatid breaks per cell, are classified as normal sensitive by the conventional method, since a cut-off value of 3.7 chromatid breaks per cell has been calculated. However, the HCC1937 cells, which are predicted to be normal when using the conventional G2 assay, have been reported to be highly radiosensitive (45,46), and this fact is confirmed when applying the proposed new methodology. Indeed, an IRP value of 2.0 chromatid breaks per cell is obtained and, therefore, as shown in Fig. 3, the cell line is classified as highly radiosensitive. In the case of GM03188 (AT+/-) cell line, it is characterized as radiosensitive by both methods, whereas GM15786 (AT-/+) is characterized as radiosensitive by the conventional G2 assay and as highly radiosensitive by the proposed new assay, since an IRP value of 0.9 chromatid breaks per cell was obtained. When GM09899 (control) cells were tested, they were characterized as highly radioresistant by the proposed methodology. With respect to intra-experimental reproducibility, the proposed methodology using the IRP values seems to have better reproducibility when compared to the conventional G2 assay, since lower coefficients of variation were obtained, as shown in Table III. Indeed, the fact that the IRP value is calculated for each individual as the difference between two yields of chromatid breaks obtained under exactly the same culture and irradiation conditions (culture media, pH, CO2 temperature, exact radiation dose) minimizes the stringent technical requirements needed by the conventional G2 assay.

In summary, in this study it is shown that the proposed methodology for predicting intrinsic radiosensitivity and predisposition to cancer offers several advantages when compared to the conventional G2 assay, as it provides better discrimination and minimizes problems associated with intra-experimental reproducibility. The new methodology is ultimately based on differences among individuals in specific mutations or polymorphisms in genes that control: a) DNA repair capacity, b) cdk1/cyclin B activity and c) G2 checkpoint activation after irradiation during the G2 to M-phase transition. We suggest that these differences are reflected in the individual radiosensitivity parameter (IRP), which provides the biological basis of the proposed methodology, as it predicts, at individual level, the G2 checkpoint potential to facilitate repair of radiation-induced chromosomal damage during G2 to M-phase transition. The results obtained so far are promising and this new approach may be proved to be a simple, sensitive, reliable and clinically applicable predictive assay for the assessment of individual radiosensitivity in radiation therapy and, in general, for population screening for hypersensitivity to radiation and predisposition to cancer.

References


