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Low-dose hyper-radiosensitivity of progressive and regressive cells isolated from a rat colon tumour: Impact of DNA repair

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Abstract

Purpose: To ask whether highly metastatic sublines show more marked low-dose hyper-radiosensitivity (HRS) response than poorly metastatic ones.

Materials and methods: The progressive (PRO) subline showing tumourigenicity and metastatic potential and the regressive (REG) subline showing neither tumourigenicity nor metastatic potential were both isolated from a parental rat colon tumour. Clonogenic survival, micronuclei and apoptosis, cell cycle distribution, DNA single- (SSB) and double-strand breaks (DSB) induction and repair were examined.

Results: HRS phenomenon was demonstrated in PRO subline. Before irradiation, PRO cells show more spontaneous damage than REG cells. After 0.1 Gy, PRO cells displayed: (i) More DNA SSB 15 min post-irradiation, (ii) more unrepaired DNA DSB processed by the non-homologous end-joining (NHEJ) and by the RAD51-dependent recombination pathways, (iii) more micronuclei, than REG cells while neither apoptosis nor p53 phosphorylation nor cell cycle arrest was observed in both sublines.

Conclusions: HRS response of PRO subline may be induced by impairments in NHEJ repair that targets G₁ cells and RAD51-dependent repair that targets S-G₂/M cells. The cellular consequences of such impairments are a failure to arrest in cell cycle, the propagation of damage through cell cycle, mitotic death but not p53-dependent apoptosis. Tumourigenic cells with high metastatic potential may preferentially show HRS response.

Keywords: Low-dose hyper-radiosensitivity, DNA repair, cell cycle, cell death, micrometastases

Introduction

Among each type of tumour, the required tumour lethal dose varies most commonly by a two-three factor (Friedman 1975). Radiobiological features of such intra-tumour heterogeneity have been recently highlighted by in vitro studies supporting that cells within a given tumour cell line show low-dose hyper-radiosensitivity (HRS) that is replaced by induced radiosensitivity (IRR) at a cell-line-dependent threshold of 10–30 cGy. Such a conclusion was reached after analyzing more than 40 tumour cell lines (Skov 1999, Joiner et al. 2001, Marples et al. 2004). Micrometastases elicit higher genetic instability and heterogeneity than primary tumours (Tortola et al. 2001, Klein et al. 2002). However, whether micrometastases may preferentially show more HRS response than primary tumours is not well documented. In addition, only few models of micrometastases or spontaneous metastatic clones deriving from the same parental tumor cell line are available.

By using seven highly and poorly metastatic clones derived from the same parental human melanoma cell line, we have previously shown that the more the clones show marked HRS response, the more the clones are metastatic (Thomas et al. 1997). HRS was also reported in human metastatic skin nodules (Harney et al. 2004). Hence, these findings suggest that the higher the HRS response, the higher the
tumourigenicity. Besides, it is noteworthy that HRS is not a common finding in human normal cells (Slonina et al. 2007). The fact that highly metastatic cells may preferentially show more HRS response than poorly metastatic ones suggests that HRS response should be considered more carefully in the radiotherapy of micrometastases (Thomas et al. 2007).

Although a number of molecular models have been proposed, mechanisms of HRS are still unclear (Skov 1999, Joiner et al. 2001, Marples et al. 2004). It has been suggested that HRS may depend upon changes in chromatin conformation (Joiner et al. 2001), failure of the ATM-dependent G\(_2\)/M checkpoint (Marples et al. 2004), DNA repair defects (Vaganay-Juery et al. 2000, Short et al. 2005) and/or induction of p53-dependent apoptosis (Enns et al. 2004). The DNA repair field has recently progressed, especially for DNA DSB. Notably, impairments of the major DNA DSB repair pathways, the non-homologous end-joining (NHEJ) and the RAD51-dependent recombination, result in high radiosensitivity and cancer proneness, respectively (Joubert & Foray, 2006, Joubert et al. 2008). Finally, a functional and temporal hierarchy between ATM-dependent phosphorylation events occurring early after irradiation has been demonstrated: such phosphorylations may determine success of DNA repair, cell cycle arrest and cell death (Foray et al. 2003).

Here, by using such endpoints, we examined whether HRS response is higher in tumorigenic cells with high metastatic potential than in non-tumorigenic cells. The previously described PRO and REG sublines that derived from the same parental rat colon tumour were chosen as a model (Martin et al. 1983).

Materials and methods

Cells and irradiation

PRO and REG sublines were kindly provided by Dr F. Martin (Dijon, France). PRO and REG sublines were isolated from the parental tumour cell line DHD-K12, established from dimethylhydrazine-induced colon carcinoma in syngeneic BDIX rats (Martin et al. 1983). PRO and REG sublines have been isolated according to their sensitivity to trypsin-mediated detachment from plastic surface (PRO subline is more trypsin-resistant than REG subline). PRO and REG cells show different immunogenicity and tumorigenicity in syngeneic hosts. When grafted subcutaneously in BDIX rats, REG cells produced regressive tumours disappearing within 3–4 weeks while PRO cells produced progressive tumours in 60% of animals (Martin et al. 1983). Metastases to lungs, kidney or lymph nodes were observed in more than 50% of rats in which PRO tumour was either allowed to grow for four months or excised 2–4 weeks after inoculation (Martin et al. 1983). In contrast, neither syngeneic BDIX rats nor nude mice that are inoculated with REG cells produced metastases (Martin et al. 1983). PRO and REG sublines were cultured in Roswell Park Memorial Institute (RPMI) 1640 medium with 2 mM glutamine, 10% decomponented fetal bovine serum and antibiotics (1% penicillin, streptomycin) (Gibco-Invitrogen-France, Cergy-Pontoise, France). Cells were mycoplasma-free and maintained at 37°C at 5% CO\(_2\) for no more than five passages. For all the assays described below, confluent PRO and REG cultures were softly detached with 0.025% trypsin and 0.02% ethylenediaminetetraacetic acid (EDTA) (Gibco-Invitrogen-France, Cergy-Pontoise, France) to obtain single cell suspensions. Irradiations were performed either with \(^{60}\)Co \(\gamma\)-rays at a dose rate of 0.5 Gy·min\(^{-1}\) (clonogenic assay, comet assay) or with X-rays produced by two irradiators (immunofluorescence and cell cycle assays): a Philips BV212 mobile surgical system (100 kV, 3 mA) delivering 0.1 Gy at a dose rate of 0.17 Gy·min\(^{-1}\) and a clinical irradiator (200 kV, 20 mA) delivering 6 Gy at a dose rate of 1.25 Gy·min\(^{-1}\). It is noteworthy that \(^{60}\)Co \(\gamma\)-rays, 100 kV and 200 kV X-rays induce the same DNA DSB induction rate (data not shown).

Clonogenic survival assay

Clonogenic survival was assessed as previously described (Thomas et al. 1997). Briefly, 500 cells were seeded and irradiated 2 or 24 h after plating with doses ranging from 0.05–6 Gy. Colonies were fixed and stained with standard crystal violet solution (Sigma-Aldrich-France, L’Isle d’Abeau, France) incubation without change of medium. Only colonies showing more than 50 cells were considered. Plating efficiencies of unirradiated REG and PRO cells were 30 ± 1.4% (mean ± SEM (standard error of the mean), \(n = 21\)) and 23 ± 1% (mean ± SEM, \(n = 23\)), respectively. In order to examine whether, before irradiation, cellular proliferation may impact upon HRS response, multiplicity assay was applied. To this aim, cells were fixed and stained with crystal violet 24 h after plating: in such conditions, the cell multiplicity (i.e., the number of cells per colony-forming unit) was found to be closed to one.

Survival curves analysis

Surviving fractions (SF) were fitted to three models: The one population linear-quadratic (LQ) model, the induced repair (IR) model (Lambin et al. 1994).
and the two populations linear-quadratic (2-pop LQ) model (Thomas et al. 1997) defined by, respectively:

1. \( \text{SF}(D) = e^{-(\alpha D + \beta D^2)} \)
2. \( \text{SF}(D) = e^{-\alpha (1 + (\frac{D}{x} - 1))} \cdot e^{\frac{D}{x} \cdot D} \)
3. \( \text{SF}(D) = (1 - \lambda) \cdot e^{-\alpha D} + \lambda \cdot e^{-(\alpha D + \beta D^2)} \)

The IR model is a modified version of the LQ model in which the \( \alpha \) term is dependent on dose (\( D \)): at very low doses, \( \alpha \) is large, and it decreases with increasing dose in an exponential manner at a rate determined by a constant \( d_c \). The parameter \( \alpha_s \) represents the initial slope of the curve at very low doses, \( \alpha_r \) represents the value extrapolated from the conventional high-doses response, \( d_c \) represents the dose that induced the change from HRS to IRR. The occurrence of HRS/IRR is mathematically deduced from \( \alpha_s \) and \( \alpha_r \) values that do not coincide and \( d_c \) values significantly greater than zero (Table I). With regard to the 2-pop LQ model that assumed the existence of two subpopulations of cells with different radiosensitivity, the \( \lambda \) parameter represents the proportion of radioresistant cells. The \( \alpha_s \) parameter is linked to the radiosensitive subpopulation. The \( \alpha_r \), \( \beta_r \) parameters are linked to the radioresistant subpopulation. All the survival data were fitted by using the JMP Software (version 2.0.5, SAS Institute, Cary, NC, USA).

**Alkaline comet assay**

The experimental protocol for the alkaline comet assay was previously described (Alapetite et al. 1996) and was applied here with minor modifications. Briefly, cells embedded in agarose (Sigma) were irradiated 2 h after plating on slides that were maintained on ice for at least 30 min before irradiation and protected from direct natural light during all following steps. For the assessment of initial DNA SSB, slides were immersed immediately after irradiation at doses ranging from 0.05–6 Gy for 1 h in lysis solution (2.5 M NaCl, 100 mM EDTA, 10 mM tris-hydroxymethylaminomethane, 1% N-laurylsarcosine, 1% Triton X-100, 10% dimethylsulfoxide, pH 10, 4°C) (Sigma-Aldrich-France, L’Isle d’Abeau, France). For the assessment of residual DNA SSB, slides were incubated after irradiation at 0.1 and 6 Gy for 15 min, 1 and 14 h before lysis. Slides were transferred to electrophoresis box containing alkaline buffer (300 mM NaOH, 1 mM EDTA, pH 13) (Sigma-Aldrich-France, L’Isle d’Abeau, France) for 40 min to allow DNA unwinding. Electrophoresis was performed for 20 min (25 V, 300 mA). Slides were rinsed twice for 5 min with neutralization buffer.
(0.4 M Tris, pH 7.5), stained with 8 ng·μl⁻¹ ethidium bromide (Sigma-Aldrich-France, L’Isle d’Abeau, France) and kept at 4°C. Comets were observed with Leica fluorescence microscope and analyzed with the image analysis Komet software (Kinetic imaging 4.0, Andor technology, South Windsor, CT, USA). Comets were evaluated by tail moment defined as product of tail DNA percent and tail length. The tail moment of irradiated cells was divided by the tail moment of unirradiated cells to provide the normalized tail moment. Cells were considered as highly damaged cells (HDC) if the normalized tail moment was higher than 90%. In each experiment, 15 comets per slide were randomly captured at constant gel depth, by avoiding gel edge and superimposed comets. Each experiment was replicated three times and 45 comets were considered per dose.

**Immunofluorescence assay**

This assay already described (Foray et al. 2003) was applied with minor modifications. Briefly, 5 × 10⁵ cells were seeded on slides and incubated for 24 h in complete medium at 37°C. After irradiation at 0.1 Gy, dishes were incubated at 37°C for 10 min, 1 h, 24 h and 48 h while after irradiation at 6 Gy, dishes were incubated for 24 h only. Cells were fixed in paraformaldehyde solution for 15 min at room temperature and permeabilized for 90 s at 4°C in lysis solution (20 mM 4-(2-hydroxyethyl)-1-piperazine ethanesulfonic acid (HEPES) (pH 7.4), 50 mM NaCl, 3 mM MgCl₂, 300 mM sucrose, 0.5% Triton X-100) (Sigma-Aldrich-France, L’Isle d’Abeau, France). Primary antibody incubations were performed for 40 min at 37°C. Anti-pH2AXser139 antibody (#05636; Upstate Biotechnology-Euromedex, Mundolsheim, France) was used at 1:800. Anti-pATser1981 (#2888; Abcam, Cambridge, UK), -pDNA-PKcs#ab14955; Abcam, Cambridge, UK), -pH3ser10 (#ab1525; Abcam, Cambridge, UK), -MRE11 (#56211; Abcys, Paris, France) and -P53ser15 (9284; Cell Signalling, Beverly, USA), and -RAD51 (#8349; Santa-Cruz-Biotechnologies, Santa-Cruz, USA) were used at 1:100. Incubations with anti-mouse fluorescein (green) or rhodamin (red) secondary antibodies were performed at 1:100 at 37°C for 20 min. Slides were mounted in 4’,6-diamidino-2-phenylindole (DAPI)-stained Vectashield (Abcys, Paris, France) and 100 cells per slide were examined with Olympus fluorescence microscope. DAPI staining permitted to indirectly evaluate yield of G₁ cells (nuclei with homogeneous DAPI staining), S cells (nuclei showing numerous pH2AX foci), G₂ cells (nuclei with heterogeneous DAPI staining) and metaphase (visible chromosomes). DAPI staining permitted also to quantify the percentage of cells with micronuclei by examining 100 cells at least.

**Cell cycle analysis**

PRO and REG cells were seeded at 5 × 10⁵ cells per dish and showed similar doubling time of about 36 h. Twenty-four hours after plating, cells were irradiated at 0.1 and 6 Gy. Fifteen min, 1 h, 6 h and 24 h after irradiation at 0.1 Gy and 15 min and 24 h after irradiation at 6 Gy, cells were trypsinized, fixed in 70% ethanol and stored at −20°C. At least 10⁴ fixed cells treated with 10 μg·ml⁻¹ RNAse A and 20 μg·ml⁻¹ propidium iodide (Sigma-Aldrich-France, L’Isle d’Abeau, France) were analyzed using cytometer (Beckton Dickinson, Pont-de-Clai, France). Cell cycle distribution was quantified with WinMDI software (Scripps-Research-Institute, La Jolla, USA).

**Results**

**HRS response and cell cycle distribution**

Since it was suggested that G₂ phase cells may explain the HRS response (Marples et al. 2004), cell cycle distribution and cell survival were both examined in PRO/REG cells. As a first step, the HRS response was examined by irradiating PRO/REG cells 2 h after plating. HRS response varied greatly according to the different experiments. PRO and REG cells showed HRS response in 9 among 11 and 3 among 9 independent experiments, respectively (Figure 1A, 1B). These data suggest that PRO cells display more HRS response than REG cells when cells are irradiated 2 h after plating. Twenty-four hours after plating, PRO cells also showed HRS response whereas REG cells did not. Moreover, the initial part of the survival curve (i.e., the HRS response) was similar in PRO cells whether irradiated 2 h or 24 h after plating (Figure 1C). Survival data shown in Figure 1C were fitted to the LQ, 2-pop LQ and IR models. It appeared that the HRS response in PRO cells was better described by the IR model than by the 2-pop LQ model. This was notably verified in the low-dose range for PRO cells irradiated 24 h after plating in which the \( \alpha \) parameter of the 2-pop LQ model was unable to be determined by the fitting analysis (Table I). PRO cells irradiated 2 h or 24 h after plating showed \( \alpha/\alpha_r \) ratios of 5 and 7, respectively (Table I). Thus, the HRS response in PRO cells irradiated 2 h after plating was similar to that obtained in PRO cells irradiated 24 h after plating. By contrast, REG cells did not show any significant HRS response, whatever the time after plating (Table I).
We examined thereafter whether the percentage of G₂/M cells evaluated before irradiation (i.e., 2 h or 24 h after plating) impacts upon the HRS response. Unirradiated PRO subline showed significantly less cells in G₂/M 2 h after plating and significantly more cells in G₂/M 24 h after plating than REG subline. Furthermore, since: (i) Unirradiated PRO subline showed significantly more cells in G₂/M 24 h after plating than 2 h after plating (Figure 2A, left panel); (ii) PRO cells showed similar HRS response 2 h and 24 h after plating; and (iii) no significant HRS response was observed in REG cells whatever the time after plating (Figure 1C), our data suggest that the percentage of G₂/M cells before irradiation cannot explain the difference in HRS response between PRO and REG sublines.

Cell cycle distributions were also examined in cells irradiated 24 h after plating at 0.1 Gy and 6 Gy. A low-dose of 0.1 Gy was chosen to evaluate specifically the HRS and not the IRR response since the dose inducing the transition from HRS to IRR (d_c) was 0.19 ± 0.08 Gy for PRO cells irradiated 24 h after plating (Table I). A high-dose of 6 Gy that provided the largest survival difference between PRO and REG cells was chosen to compare data obtained after irradiation at 0.1 Gy. Fifteen min, 1, 6 and 24 h after irradiation at 0.1 Gy, cell cycle distributions in PRO and REG cells remained unchanged compared to those observed in unirradiated cells, suggesting that a dose of 0.1 Gy is not sufficient to trigger significant cell cycle change whatever the subline (data not shown). By contrast, 24 h after irradiation at 6 Gy, the S fraction significantly decreased in both sublines whereas the G₂/M fraction significantly increased in PRO cells only when compared to unirradiated cells (Figure 2A, right panel).

DAPI staining and immunofluorescence performed with antibodies against phosphorylated H3 histone forms characterizing mitoses were also applied during immunofluorescence experiments in parallel to cytometry analysis. Mitotic cells represented less than 5% of cells and this feature remained unchanged after irradiation at 0.1 Gy (Figure 2B), consolidating therefore the cytometry data described above (Figure 2A). Hence, altogether, these findings suggest that the relative fraction of G₂/M cells alone cannot explain the HRS response observed in PRO/REG sublines.

Cell death

Clonogenic death is the final result of different radiation-activated death pathways such as apoptosis, mitotic death or senescence. We asked whether particular cell death pathway may explain the HRS response observed in PRO cells. By examining DAPI-stained cells with immunofluorescence, no apoptotic body was observed either between 15 min and 48 h after irradiation at 0.1 Gy or 24 h after irradiation at 6 Gy in PRO and REG cells (data not shown). Apoptosis can be mediated by p53-dependent and -independent pathways. However, since
Figure 2. Cell cycle distribution in PRO/REG cells. (A) Left panel. Relative cell cycle distribution in unirradiated PRO/REG cells before irradiation, i.e., 2 h or 24 h after plating. *$p < 0.05$ and **$p < 0.05$ for comparison of data between unirradiated PRO cells and unirradiated REG cells 2 h and 24 h after plating, respectively and *$p < 0.05$ for comparison between unirradiated PRO cells 2 h after plating to unirradiated PRO cells 24 h after plating using $t$-test. Right panel. Relative cell cycle distribution in PRO/REG cells 24 h after irradiation at 6 Gy. *$p < 0.05$ and **$p < 0.05$ for comparison of data between irradiated PRO and REG cells to unirradiated PRO and REG cells, respectively using $t$-test. Each data corresponds to the mean ± SEM of four independent experiments. (B) Immunofluorescence data obtained with antibodies against phosphorylated histone H3 in PRO cells and REG cells. Upper panel shows a representative example of mitosis stained in fluorescein (green) and clearly identifiable in DAPI. This picture was obtained with the 100 × microscope objective (the white bar represents 5 μm). Lower panel shows representative pictures with green-stained mitoses 10 min and 24 h after irradiation at 0.1 Gy in PRO/REG cells as observed in 10 × microscope objective (the white bar represents 0.05 μm).
p53-dependent apoptosis has been evoked to explain the HRS phenomenon (Enns et al. 2004), the occurrence of phosphorylated forms of p53 (pP53) was investigated by using specific immunofluorescence after irradiation at 0.1 Gy and 6 Gy. Whereas pP53 appeared essentially cytoplasmic in both unirradiated PRO and REG cells, it was nuclear in rodent control cells. Four hours after irradiation at 0.1 Gy, pP53 localization did not change in PRO and REG cells (Figure 3A). Similar conclusions were reached 24 h after irradiation at 6 Gy (data not shown). These data suggest a functional impairment of p53 in both unirradiated and irradiated PRO/REG cells, consistent with absence of p53-dependent apoptosis. At this stage, it must be stressed that these results do not necessarily mean that the p53 protein is mutated in both sublines or that p53 mutations are similar. These results only provide information about the functionality of the ATMedependent phosphorylation of p53 that impacts upon cell cycle control and p53-dependent apoptosis.

Mitotic death is characterized by micronuclei formation. No cell with micronuclei was observed in unirradiated cultures. Forty-eight hours after irradiation at 0.1 Gy, the percentage of cells showing one micronucleus was 37 ± 5% and 2 ± 0.3% (mean ± SEM, n = 3) in PRO and REG cells, respectively (Figure 3B). Such numbers are much higher than those previously obtained by Slonina et al. (2007) in irradiated primary human fibroblasts with a mean of 1.7% and a range of 0.3–3.8%. However, it is noteworthy that capacity of proliferation, distribution in cell cycle, cell type and origin are deeply different (human fibroblasts versus rodent tumour cells), which may explain, at least in part, such a difference. Corresponding surviving fraction for PRO and REG cells irradiated 24 h after plating with 0.1 Gy X-rays were 0.85 ± 0.04 (mean ± SEM, n = 6) and 1.1 ± 0.07 (mean ± SEM, n = 6), respectively. Data support that mitotic death, more likely than p53-dependent apoptosis, reflects HRS response in PRO cells. Since micronuclei result from propagation of unrepaired DNA breaks throughout cell cycle, DNA breaks in PRO/REG cells have been investigated.

**DNA SSB induction and repair**

From the analysis of 45 comets, the mean and the SEM of the tail moments in unirradiated PRO and REG cells were 66 ± 10 and 86 ± 14, respectively.
These data suggest that the yield of spontaneous DNA SSB does not discriminate the PRO/REG sublines. Normalized tail moment (NTM) was chosen as an endpoint for describing the radiation-induced initial (i.e., assessed immediately after irradiation) and residual DNA SSB. The higher the dose, the higher the yield of initial DNA SSB (Figure 4A). However, such linear relationship did not display significant difference between PRO and REG cells. Indeed, although the yield of DNA SSB assessed immediately after irradiation at 0.1 and 6 Gy in PRO cells was significantly higher than that obtained in unirradiated cells, this endpoint was not significantly different in PRO and REG cells (data not shown), suggesting that initial DNA SSB cannot explain the difference observed in HRS response between PRO and REG cells.

During comet experiments, the presence of some cells highly damaged cells (HDC) may be noticed (Olive et al. 1993). In our conditions, a substantial fraction of HDC was observed in REG cells but not in PRO cells before irradiation with a median of 7% and 0% and ranges of (0–33%) and (0–0%), respectively. HDC were observed in PRO and REG cells immediately after irradiation but not between 15 min and 24 h post-irradiation (data not shown). When HDC were omitted from NTM data, the yield of initial DNA SSB as function of dose was similar in both sublines (data not shown). The biological significance of HDC is not clear. HDC may reflect transient but significant disturbances in chromatin since comet assay is prominently influenced by chromatin organization (e.g., Kumar et al. 2002). It was also suggested that HDC may be apoptotic cells occurring rapidly after irradiation (e.g., Olive et al. 1993). However, no apoptotic body was observed between 15 min and 48 h after irradiation at 0.1 and 6 Gy in both sublines. Lastly, the number of HDC was neither correlated with radiation doses nor with surviving fractions in both sublines (data not shown). Altogether, these data suggest therefore that, like initial DNA SSB, HDC cannot explain the difference in HRS response between PRO and REG cells.

With regard to residual DNA SSB, NTM assessed 15 min after irradiation at 0.1 Gy was significantly higher in PRO cells than in REG cells while no significant difference between irradiated and unirradiated PRO/REG sublines data was observed for longer times after irradiation at 0.1 Gy (Figure 4B). Furthermore, PRO cells showed three-fold more DNA SSB 15 min after irradiation at 6 Gy than REG cells (data not shown), suggesting a production of additional DNA SSB after irradiation. Altogether, these data showed that after irradiation at 0.1 and 6 Gy, there is no significant difference in initial and residual DNA SSB between both sublines, to the notable exception of a DNA SSB burst observed 15 min after irradiation in PRO cells only.

**Induction and repair of DNA DSB processed by the NHEJ pathway**

DNA DSB induction and repair were examined by using immunofluorescence with specific antibodies...
against H2AX phosphorylation (pH2AX), an early NHEJ event (Rothkamm & Löbrich 2003). The number of spontaneous pH2AX foci was not significantly different in both sublines. Ten min and 1 h after irradiation at 0.1 Gy, the number of pH2AX foci per cell was significantly higher in PRO cells than in REG cells. Furthermore, 24 h after irradiation at 0.1 Gy, the number of pH2AX foci per cell was significantly higher in irradiated PRO cells than that obtained in unirradiated PRO cells (Figure 5A). Such a difference in DSB repair rate was confirmed 24 h after irradiation at 6 Gy (data not shown), suggesting an impairment of the NHEJ process in PRO cells but not in REG cells.

Since ATM and DNA-PK kinases are responsible for the H2AX phosphorylation, the radiation-induced kinase activity was evaluated by using antibodies against autophosphorylated ATM and DNA-PK forms that re-localize as nuclear foci (pATM and pDNA-PK, respectively). The number of pDNA-PK and pATM foci in PRO cells was similar to that observed in REG cells after irradiation at 0.1 Gy but higher to that obtained in REG cells after irradiation at 0.2 Gy (Figure 5B). Hence, these findings suggest that the HRS response observed in PRO cells after irradiation at 0.1 Gy cannot be explained by impaired ATM and/or DNA-PK kinase activities required for the DNA DSB recognition during the NHEJ process.

The MRE11 protein, associated with hyper-recombination and genomic instability (Joubert & Foray 2006; Joubert et al. 2008), was found essentially cytoplasmic in both sublines, while a rodent control showed a normal formation of radiation-induced MRE11 nuclear foci, suggesting that the HRS response in PRO cells is MRE11-independent (data not shown).

Induction and repair of DNA DSB processed by RAD51-dependent recombination pathway

While NHEJ is the major DNA DSB repair pathway acting in G1 cells, DNA DSB in S-G2/M cells are more likely repaired by the RAD51-dependent recombination. Since radiation-induced RAD51 foci are generally tiny and numerous, the percentage of cells showing RAD51 foci rather than their yield per cell was chosen as endpoint. Unlike with pH2AX assay, the percentage of cells showing spontaneous RAD51 foci was five-fold higher in PRO cells than in REG cells (data not shown). A similar conclusion was reached 4 h after irradiation at 0.1 Gy. At such dose, RAD51 remained cytoplasmic in REG cells, suggesting that this protein was activated after 0.1 Gy in PRO cells only (Figure 6A). By contrast, after irradiation at 6 Gy, 90% of REG cells and 30% of PRO cells elicited RAD51 foci (Figure 6A). Since RAD51-dependent repair is active in S-G2/M cells only, the relationship between RAD51 data and the corresponding cell cycle distributions was investigated. Before irradiation, the percentages of S-G2/M cells in both sublines were similar before irradiation (57 ± 2% and 51 ± 6% (mean ± SEM, n = 4) in PRO and REG cells, respectively) (Figure 2A). After irradiation at 0.1 and 6 Gy, a significant linear relationship was detected between the percentage of cells showing RAD51 foci and the percentage of S-G2/M cells in PRO cells but not in REG cells (Figure 6B), suggesting that unrepaired DNA DSB processed by the RAD51-dependent repair pathway may propagate through the cell cycle in PRO cells but not in REG cells.

Discussion

PRO/REG sublines: A useful model to study HRS response

PRO/REG sublines have been deliberately chosen to test whether HRS response is more marked in tumorigenic cells with high metastatic potential than in non-tumorigenic cells with no metastatic potential. PRO/REG sublines were obtained from the same parental rat colon tumour, which made possible molecular and cellular investigations on HRS response in the frame of tumour heterogeneity. Here, our data indicate that PRO cells show more HRS response than REG cells, consistently with literature showing that tumorigenic cells are generally more radiosensitive than non-tumorigenic cells (e.g., Barnetson et al. 1999). Our observations provide evidence that neither cell cycle distribution alone (both sublines elicited similar S-G2/M fraction before irradiation) nor p53-dependent apoptosis (not relevant here) nor HDC (REG cells spontaneously showed more HDC than PRO cells) may explain the fact that PRO cells show higher HRS response than REG cells.

Impact of the major DNA repair pathways upon HRS response

Induction and repair of DNA SSB and DSB were investigated by using alkaline comet and pH2AX, pATM, pDNA-PK, MRE11 and RAD51 immunofluorescence assays. Alkaline comet assay allowing the assessment of initial and residual DNA SSB has been already applied to low-dose (6 mGy) (Malyapa et al. 1998) or very-low chronic radiation dose-rate (20 μSv·h−1) (Meehan et al. 2004). The H2AX assay that allows the detection of DNA DSB has already proved its efficiency at very low-dose like 1.2 mGy (Rothkamm & Löbrich 2003).
Figure 5. Initial and residual DNA DSB processed by the NHEJ repair pathway. (A) Left panel. DSB induction and repair after irradiation at 0.1 Gy. The number of pH2AX foci per cell was plotted against the post-irradiation times. Each data represents the mean ± SEM of two independent experiments. *p < 0.05 and **p < 0.05 for comparison of data between irradiated PRO and REG cells and between irradiated PRO cells and unirradiated PRO cells, respectively using t-test. Right panel. Representative pH2AX signals provided by PRO/REG cells 10 min after irradiation at 0.1 Gy. (B) The number of pDNA-PK (left panel) and pATM (right panel) foci per cell assessed 10 min after irradiation was plotted against the indicated radiation doses in PRO/REG sublines. Each data represents the mean ± SEM of two independent experiments. Corresponding representative pictures of pATM and pDNA-PK are shown below.
With regard to spontaneous DNA damage, PRO and REG sublines elicited similar yields of DNA SSB and DSB recognized by alkaline comet and pH2AX assays, respectively. Conversely, the RAD51 data revealed more spontaneous DNA damage in PRO than in REG cells, which cannot be explained by differences in the cell cycle distribution between both sublines. Hence, these findings support that such spontaneous DNA damage are processed in PRO cells specifically by the RAD51-dependent pathway but not by the NHEJ one. The RAD51-dependent recombination pathway is able to manage SSB and DSB, and unlike NHEJ, can produce DNA SSB intermediates during the DNA DSB repair process (Raderschall et al. 1999, Schlegel et al. 2006). Hence, we propose that S-G2/M PRO cells are characterized by a lack of control in the RAD51 endonuclease function through hyper-recombination.
(e.g., Schultz et al. 2003). Whether the DNA SSB burst observed 15 min after irradiation at 0.1 and 6 Gy in PRO cells is consistent with induction of DNA SSB during repair of DNA DSB needs further investigations. However, it raises the problem of the impact of the cell cycle distribution upon the evaluation of HRS response. Indeed, any impairment of RAD51-dependent repair should not be visible in quiescent cells. Interestingly, only two among forty human primary fibroblasts elicit HRS response (Slonina et al. 2007). This last observation might be explained by the propensity of fibroblasts to be essentially in G1 phase. A direct consequence of a lack of control of RAD51-dependent repair would be also that PRO cells elicit higher chromatin fragility and genomic instability. Besides, the influence of chromatin fragility was already demonstrated in the radiosensitivity of some tumour cell lines (Chavaudiра et al. 2004) and in ataxia telangiectasia or Xeroderma Pigmentosum fibroblasts (Puvion-Dutilleul 7 Sarasin 1989).

With regard to radiation-induced DNA damage, the data presented here suggest that the HRS response in PRO cells may not be explained by an impairment in DNA DSB recognition, in agreement with data from Wykes et al. (2006). Along with this normal DNA damage recognition, PRO and REG sublines elicited after irradiation similar initial DNA SSB yields but PRO cells showed significantly more initial DNA DSB than REG cells, suggesting again higher chromatin fragility in PRO cells.

With regard to unrepaired DNA DSB, PRO cells displayed more impaired damage processed by the NHEJ pathway after irradiation at 0.1 and 6 Gy than REG cells. Such a conclusion is in agreement with the relative defect of DNA DSB repair in fibroblasts that show HRS response after irradiation at 1.2 mGy (Rothkamm & Löbrich 2003). Besides, after irradiation at 0.1 Gy, PRO cells displayed more unrepaired DNA breaks processed by the RAD51 repair pathway than REG cells, consistently with a previous report showing that HRS response is associated with an increase of the number of RAD51 foci per cell after low-dose radiation (Short et al. 2005). Altogether, the analysis of the radiation-induced DNA damage after irradiation at 0.1 Gy suggests that more numerous and less repairable DNA lesions occur in PRO cells than in REG cells. Such phenomenon concern the two major DNA breaks repair pathways: NHEJ acting in G1 and the RAD51-dependent recombination acting in S-G2/M.

Impact of the cell cycle distribution upon HRS response

It was suggested that G2 cells may dominate HRS response, which may be specific to damaged cells that fails to activate a transient G2/M checkpoint and enter into mitosis with unrepaired DNA breaks (Marple et al. 2004). Although the G2/M checkpoint was not the scope of this study, our observation that the cell cycle distribution in PRO cells after irradiation at 0.1 Gy is similar to that of unirradiated cells, may be consistent with a failure to induce G2/M arrest after irradiation at low doses. Such a conclusion is in agreement with previous reports (Marple et al. 2003, Krueger et al. 2007b), although we emphasized that the HRS response may not be dominated by the fraction of G2/M cells only. Experimental conditions may explain discrepancies in literature, e.g., irradiation was performed with cells in suspension that were plated after irradiation (Marple et al. 2003, Krueger et al. 2007b) whereas, in our study, irradiation was performed with adherent cells.

Impact of the cell death pathway upon HRS response

Such cascade of impaired events would likely result in micronuclei or apoptotic bodies. With PRO/REG cells, our data suggest that unrepaired DNA breaks induce mitotic death, but not p53-dependent apoptosis. This conclusion is not in agreement with the reports supporting that HRS response is associated with p53-dependent apoptosis in other tumour cell lines (Enns et al. 2004, Krueger et al. 2007a). Besides, whether HRS response is caused by unrepaired DNA breaks inducing apoptosis and/or mitotic death still remains controversial and, again, likely depends upon: (i) The experimental conditions (Joiner et al. 2001, Chandna et al. 2002, Rothkamm & Löbrich 2003) and notably whether cells were derived from exponential-phase or plateau-phase cultures (Krueger et al. 2007a); and (ii) the status of the genes and/or the functionality of their corresponding proteins involved in HRS, e.g., whether cells are p53-mutated or wild-typed (Enns et al. 2004, Krueger et al. 2007a), normal fibroblasts (Rothkamm & Löbrich 2003) or onco-gene-transformed or DNA repair-deficient fibroblasts (Wykes et al. 2006).

Model for HRS response

We propose that the HRS response in PRO subline is induced by both:

1. An impairment of NHEJ repair in G1 cells. It is noteworthy that impaired NHEJ is responsible for intrinsic radiosensitivity whatever the dose. Since it has been recently demonstrated that the level of NHEJ impairment is quantitatively predictive of intrinsic radiosensitivity (Joubert & Foray 2006; Joubert et al. 2008), we suggest that the level of NHEJ impairment may predict
the slope of the initial part of the survival curve \( (\alpha) \) and therefore the HRS phenomenon.

(2) A lack of control of RAD51-dependent pathway in S-G\(_2\)/M cells. However, since such a pathway is active in S-G\(_2\)/M cells only, its impact upon HRS requires both sufficient fraction of S-G\(_2\)/M cells \textit{at the time of} irradiation and failure to arrest in cell cycle to allow propagation of unrepair DNA damage in cell cycle. It is probable that such requirements are reached after irradiation at 0.1 Gy whereas irradiation at 6 Gy would limit such propagation by longer arrest in G\(_2\). Lastly, such an impairment would be responsible for the production of additional SSB, favoring the chromatin fragility and the genomic instability in PRO cells and interfering again into DNA repair and DNA damage propagation.

The cellular consequences of such repair and cell cycle deficiencies are cell death that may be either mitotic death or p53-dependent apoptosis depending on the genetic status of the irradiated cells. Alternatively, if there is no significant impairment in NHEJ and RAD51-dependent repair pathways or cell cycle arrest after irradiation, the HRS response is low or absent (Figure 7). Interestingly, such requirements for HRS response were previously evoked in the case of 19 tumour cell lines for high doses: The tumour cell lines showing the most severe radiosensitivity should elicit both DNA repair and chromatin defects. Any intermediate situation provides moderate radiosensitivity (Chavaudra et al. 2004). Further investigations are now needed to identify what kind of DNA damage would cause DNA repair deficiency and HRS response and whether other highly metastatic clones deriving from the same parental tumour cell line display similar or different HRS mechanisms.

**HRS response: Potential impact upon radiotherapy of micrometastases**

The HRS response was detected in many tumour cell lines and a few normal cell lines when irradiated at low-density (Joiner et al. 2001). Since tumour cells irradiated at high-density were protected from HRS response (Chandna et al. 2002), it is unlikely that high-density primary or secondary tumours show HRS response. An overview of radiobiological reports in which HRS response was not specifically investigated supports that inside a given tumour cell line, highly metastatic clones are more radiosensitive at high doses (i.e., higher than 0.5 Gy) than poorly metastatic ones (Fu et al. 1976, Welch et al. 1983, Rao et al. 1991, Barnetson et al. 1999). Moreover, metastatic nodules to skin elicit marked HRS response \textit{in vivo}. However, in this particular case, the mechanisms involved are still unclear since no radiosensitivity data from nodules was provided (Harney et al. 2004). We have previously shown that the more the clones isolated from a same parental human melanoma cell line are metastatic, the more they show marked HRS response (Thomas et al. 1997). To better illustrate this concept, we re-examined previous data from Thomas et al. (1997) that were fitted to the IR and 2-pop LQ models (Figure 8). Clone 4 with high metastatic potential showed HRS response while clone 1 with low metastatic potential did not. Variant 1 with intermediate metastatic potential displayed both HRS and IRR responses. However, these data were obtained from cells at early passages and no HRS response was observed at late passages (Thomas et al. 1997). Similar conclusions were also reached with PRO cells (data not shown). Our data suggest therefore that HRS response is both an inherent feature of micrometastases and a transient phenomenon, likely due to high genetic instability and heterogeneity of micrometastases (Klein et al. 2002).

HRS response may find applications in radiotherapy of micrometastases. Some early
disseminated and probably unvascularised cells may escape chemotherapy after excision of M₀ primary tumour. Evidence was provided that low-dose total body irradiation (TBI) at doses lower than 20 cGy reduces metastatic potential in rodents by immune system stimulation (Hosoi & Sakamoto 1993, Harbrecht et al. 1997) were fitted to models. Each data represents the mean ± SEM of three independent experiments. C1 survival data are well fitted by the LQ model while C4 and V1 data are well fitted by the 2-pop LQ and IR models, respectively.

Figure 8. HRS response and metastatic potential. Survival curves of clone C1 with low metastatic potential, variant V1 with intermediate metastatic potential and clone C4 with high metastatic potential. Since highly metastatic cells may display HRS response, we proposed to apply eventually a low-dose TBI after excision of M₀ primary tumour. Evidence was provided that low-dose total body irradiation (TBI) at doses lower than 20 cGy reduces metastatic potential in rodents by immune system stimulation (Hosoi & Sakamoto 1993, Hashimoto et al. 1999, Liu 2003). Since highly metastatic cells may display HRS response, we proposed to apply eventually a low-dose TBI after excision of M₀ primitive tumours to prevent micrometastases development (Thomas et al. 2001). To test such hypothesis, the smallest dose producing HRS response without increasing cancer must be found. Very low doses such as 1.2 mGy may both induce HRS response (Rothkamm & Löbrich 2003) and reduce the risk of cancer transformation compared to unirradiated cells (Azzam et al. 1996, Redpath et al. 2003, Portless et al. 2007). In addition, irradiation at 30 cGy X-rays applied to mice, whether externally or in utero, did not show significant increase of cancer (Di Majo et al. 2003). Epidemiological data suggest that the lowest dose of X- or γ- radiation showing increased cancer risk in humans may be situated between 10 and 50 mGy (Brenner et al. 2003). Thus, even if literature data consolidate the relevance of our hypothesis, the application of preventive TBI treatments at very low dose (i.e., lower than 10 mGy) raises the problem of the threshold dose in cancer incidence and requires further and careful investigations. Studies using the PRO/REG syngeneic model are in progress to secure the applicability of preventive very low-doses TBI in the radiotherapy of micrometastases.

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Low-dose hyper-radiosensitivity and DNA repair


