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Radiation dose determines the method for quantification of DNA double strand breaks

TANJA BULAT¹, OTILIJA KETA¹, LELA KORIĆANAC¹, JELENA ŽAKULA¹, IVAN PETROVIĆ¹, ALEKSANDRA RISTIĆ-FIRA¹ and DANIJELA TODOROVIĆ²

¹University of Belgrade, Vinča Institute of Nuclear Sciences, P.O. Box 522, Belgrade 11001, Serbia

²University of Kragujevac, Faculty of Medical Sciences, Svetozara Markovića 69, Kragujevac 34000, Serbia

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ABSTRACT

Ionizing radiation induces DNA double strand breaks (DSBs) that trigger phosphorylation of the histone protein H2AX (γ H2AX). Immunofluorescent staining visualizes formation of γ H2AX foci, allowing their quantification. This method, as opposed to Western blot assay and Flow cytometry, provides more accurate analysis, by showing exact position and intensity of fluorescent signal in each single cell. In practice there are problems in quantification of γ H2AX. This paper is based on two issues: the determination of which technique should be applied concerning the radiation dose, and how to analyze fluorescent microscopy images obtained by different microscopes. HTB140 melanoma cells were exposed to γ -rays, in the dose range from 1 to 16 Gy. Radiation effects on the DNA level were analyzed at different time intervals after irradiation by Western blot analysis and immunofluorescence microscopy. Immunohistochemically stained cells were visualized with two types of microscopes: AxioVision (Zeiss, Germany) microscope, comprising an ApoTome software, and AxioImagerA1 microscope (Zeiss, Germany). Obtained results show that the level of γ H2AX is time and dose dependent. Immunofluorescence microscopy provided better detection of DSBs for lower irradiation doses, while Western blot analysis was more reliable for higher irradiation doses. AxioVision microscope containing ApoTome software was more suitable for the detection of γ H2AX foci.

Key words: ApoTome software, AxioImagerA1 microscope, immunofluorescence microscopy, Western blot, γ H2AX.

INTRODUCTION

The most important direct damages induced by ionizing radiation (IR) are DNA double strand breaks (DSBs). As a consequence of DSBs the histone H2AX becomes phosphorylated (Mischo et al. 2005). Phosphorylated H2AX (γ H2AX) foci appear at the damage sites and their number is considered as one to one DNA DSB (Schmid et al.

2012). The level of γ H2AX has been used in *in vitro* studies as a tool for the estimation of cellular radiosensitivity and as a biomarker for individual studies in patient trials to measure DNA damage following radiotherapy and/or chemotherapy (Qvarnström et al. 2004).

In practice, the obtained results may be different when various detection systems are used (Ivar et al. 2010). Effects of cancer therapeutics are often analyzed by immunocytochemistry (ICC), flow cytometry and immunoblotting (Western blot)

Correspondence to: Danijela Todorović
E-mail: dtodorovic@medf.kg.ac.rs

in order to profile the activity of cell signaling proteins. However, the relative sensitivity and specificity of these methods has not yet been established (Lorincz and Nusser 2008). All of the above techniques are based on the particular chemical interaction with specific antibody. Fluorescence microscopy is frequently used in cell biology because of its sensitivity and the ability to stain and visualize specific parts of the cell, while the light microscopy is not suitable for the detection of weakly stained cellular particles (Fritschy and Härtig 2001). In fluorescence microscopy, there is no background light and the fluorescently stained object is being contrasted to a black background. Fluorescent molecules absorb the light of one particular wavelength that produces excitation of electrons, thus emitting the light of another but higher wavelength (Radek et al. 2014). The fluorescence microscope takes advantage of this difference, filtering out the excitation light. Consequently, only the emitted light is seen (Petty 2007). Immunofluorescence microscopy can be used in several microscope designs for the analysis of immunofluorescence samples. The simplest is the epifluorescence microscope, while confocal microscope is widely used. Moreover, there are powerful super resolution microscopes capable of detecting of nanostructures.

One of the crucial steps in the preparation of samples for immunofluorescence microscopy is fixation. The goal of fixation is to maintain cellular structure as close as possible to the native state (Bancroft and Gamble 2008). Once fixed, it is possible to carry on with the procedure without losing the proteins of interest and the rest of the cell. There are various fixation methods suitable for immunocytochemistry. Fixation can damage or mask antigenic sites, thereby compromising the intensity of immunostaining (Carson and Hladik 2009). Therefore, the choice of fixative and fixation protocol may depend on the additional processing steps and biological endpoints.

Flow cytometry is the method which can rapidly detect presence or absence of the protein of interest in the large cell population. Moreover, multi-parameter analysis of the cytometric data makes it possible to correlate them with other attributes of the cells, such as the cell cycle distribution (MacPhail et al. 2003, Yoshida et al. 2003, Huang et al. 2004, Huang and Darzynkiewicz 2006). However, during sample preparation one of the requirements is serial centrifugation. This can cause important cell loss, thus reducing the quantity of sample. Based on that, it is advisable to compare data obtained by this method with results obtained by other assays.

Despite its overall simplicity, Western blot has been proven to be a powerful procedure for immunodetection of proteins, especially those that are of low concentration (Geilfus et al. 2010). This procedure is used in combination with other important antibody-based methods such as immunocytochemistry to provide confirmation of the results both in research and diagnostic trials. Specificity of antibodies used for ICC is of crucial importance and therefore Western blot is essential to address antibodies specificity (Kurien et al. 2011). Literature data suggest that all mentioned methods have limitations in sensitivity but they do enable consistent characterization of protein phosphorylation and activation of specific pathways.

In this study the human HTB140 melanoma cells were chosen as a model system to disclose the formation of DNA DSBs after irradiation with γ -rays through different experimental approaches. Therefore, optimization of experimental conditions for Western blot and immunofluorescence microscopy was performed. This optimization involved two criteria: the dose and post-irradiation incubation time. Flow cytometry was not taken into account because it was not considered as the dose-time-dependent method (Yoshida et al. 2003). The results obtained by ICC and Western blot were

compared. The aim was to analyze which method is more suitable for the detection of γ H2AX in samples irradiated under the above mentioned conditions. Moreover, data obtained by two types of microscopes: AxioVision microscope comprising an ApoTome software and AxioImagerA1 microscope were compared in order to analyze influence of microscope performances on the result.

MATERIALS AND METHODS

CELL CULTURE

Human HTB140 melanoma cells were obtained from the American Tissue Culture Collection (ATCC, Rockville, MD, USA) and cultured in the RPMI 1640 cell culture medium supplemented with 10% fetal bovine serum (FBS, Sigma-Aldrich Chemie GmbH, Steinheim, Germany) and penicillin/streptomycin (Sigma-Aldrich Chemie GmbH), in a humidified atmosphere of 5% CO₂ at 37 °C (Heraeus, Hanau, Germany).

IRRADIATION CONDITIONS

Exponentially growing HTB140 cells were irradiated with γ -rays using ⁶⁰Co source at the Vinča Institute of Nuclear Sciences in Belgrade, Serbia. Applied doses ranged from 1 to 16 Gy, at the average dose rate ~ 1 Gy/min. Irradiations of cells were carried out in air at ~ 0 °C.

WESTERN BLOT ANALYSIS

Total proteins for the Western blot analysis were extracted from the HTB140 cells 30 min, 2, 6 and 24 h after irradiation. Cells were collected, washed with PBS and homogenized with buffer containing 150 mM NaCl, 50 mM Tris-HCl (pH 8.0), 1% Nonidet P-40 (NP-40), 0.1% sodium dodecyl sulfate (SDS), 0.5% sodium-deoxycholate, 1 mM ethylenediaminetetraacetic acid (EDTA), 1 mM glycol ether diamine tetraacetic acid (EGTA), 1 mM dithiothreitol (DTT), 1 mM Na₃VO₄, 5 µg/mL aprotinin, 5 µg/mL antipain, 5 µg/mL leupeptin,

0.5 mM phenylmethanesulphonylfluoride (PMSF), NaF, β -diglicophosphate and sodium-pirophosphate. The lysate was centrifuged at 1700 g for 20 min at 4 °C. In the supernatant containing the whole-cell lysate the amount of proteins was quantified spectrophotometrically (Lowry et al. 1951). The samples were mixed with denaturing buffer according to Laemmli and boiled for 5 min (Laemmli 1970). For the analysis of γ H2AX, 60 µg of proteins were loaded onto a 12% SDS-PAGE. Membranes were blocked with 5% Bovine Serum Albumin (BSA) in PBS for 1 h at room temperature and then incubated at 4 °C overnight with 1:1000 dilution of the primary anti- γ H2AX antibody (BioLegend, San Diego, USA) in PBS Tween 20 (PBST)/BSA. For normalization purposes, immunoblots were incubated overnight at 4 °C with rabbit polyclonal anti- β -actin antibody (Sigma-Aldrich Chemie GmbH) diluted 1:1000 in PBST/non-fat dry milk. After three washings in PBST, lasting 10 min each, the membranes were incubated with horseradish peroxidase-conjugated anti-mouse antibody (HRP, Cell Signaling Technology, Inc. Danvers, MA) diluted 1:5000 in PBST/BSA. Following three washings with PBST, the proteins were visualized with an enhanced chemiluminescence (Sigma-Aldrich Chemie GmbH) and exposed to X-ray film. Protein molecular mass standards (PageRuler Plus Prestained Protein Ladder, Fermentas, Vilnius, Lithuania) were used for calibration. Densitometry of protein bands on X-ray film was performed using ImageJ Analysis software.

IMMUNOFLUORESCENCE MICROSCOPY

Cells attached to coverslips were washed once with cold PBS and fixed with ice cold methanol-acetone (ratio 1:1) for 15 min, then blocked with 5% bovine serum albumin (fraction V, Sigma-Aldrich Chemie GmbH) in PBS for 60 min at room temperature then incubated with 1:500 dilution of the primary Alexa Fluor® 488 anti-H2AX-Phosphorylated (Ser139) Antibody (BioLegend), at 4 °C overnight. After

five washings in PBST, lasting 3 min each, cells were counterstained with Dapi-antifade solution (Sigma-Aldrich Chemie GmbH). The visualization of cells was done using two microscopes: AxioVision (Carl Zeiss) comprising ApoTome software and AxioImager A1 (Carl Zeiss) with a 100 x objective. All images were captured using AxioVision software. Before the observation of foci, digital images were processed with ImageJ to adjust brightness and contrast.

AUTOMATED DIGITAL IMAGE ANALYSIS

CellProfiler (www.cellprofiler.org) is an open source software that is used for a custom and automated approach to image analysis workflow (Kamentsky et al. 2011). This software enables objective and quantitative fluorescence microscopy, based on user-defined thresholds for brightness, size, shape and other parameters (Carpenter et al. 2006). The image was converted from color (Aleksa Flour 488 - green, DAPI - blue) to monochrome. Foci were identified with objects area which range from 6-30 pixels, while others were discarded. Data were exported to Excel for further analysis. Automated segmentation of foci was visually compared against manually counted foci. Relative fluorescence was measured as described previously (Burgess et al. 2010).

STATISTICAL ANALYSES

Duplicate measurements were made during each experiment and all experiments were repeated at least three times. For immunofluorescence microscopy and Western blot analysis results were presented as the mean \pm SEM (Standard Error of the Mean).

RESULTS

In order to follow the formation of γ H2AX foci, the HTB140 melanoma cell monolayers were irradiated with γ -rays. Applied dose range was from 1 to 16 Gy. Cells were maintained under standard

conditions from 30 min to 6 h after irradiation. The dose- and time- depended phosphorylation of H2AX was followed for all time points and doses. The samples were visualized and then subjected to the automated digital image analysis with CellProfiler software. Software parameters were optimized according to the properties of the cell line (size and shape of nuclei), size and shape of foci, characteristics of micrographs, etc. (Carpenter et al. 2006). In order to optimize software parameters, manual counting of γ H2AX foci in sample irradiated with 2 Gy was used as the control measurement for computer counting. All time points were taken into consideration. Data obtained by both ways of counting are shown in Figure 1.

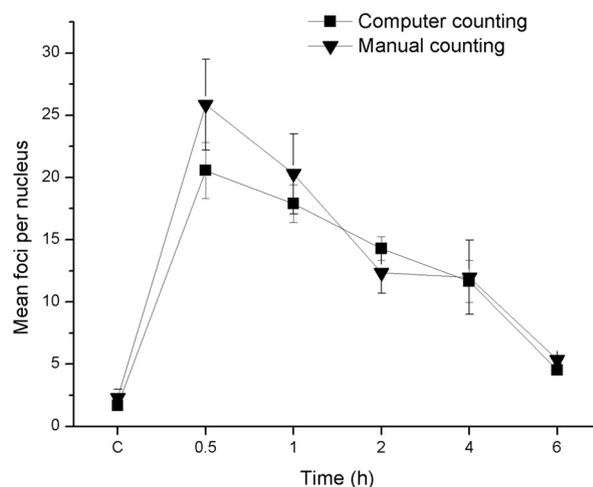


Figure 1 - Mean number of γ H2AX foci per HTB140 cell nucleus, counted by computer software (full square) or manually (full triangle), as a function of time elapsed after irradiation with 2 Gy γ -rays (C-control). Data obtained from 3 experiments are presented as mean \pm SEM.

The mean number of foci per nucleus obtained by immunofluorescence microscopy, using AxioVision microscope with ApoTome software, and AxioImager A1 microscope, are presented in Figure 2. Higher number of foci was counted with the first of the two microscopes. Both microscopes revealed smaller number of foci for the irradiation

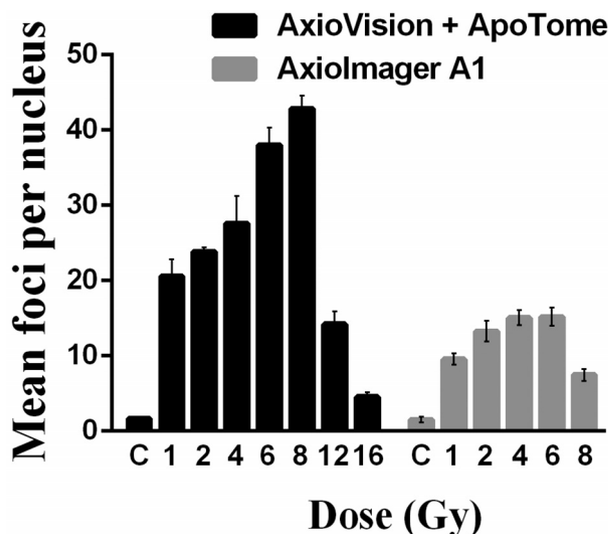


Figure 2 - Mean number of γ H2AX foci per HTB140 cell nucleus as a function of dose, 30 min after irradiation with γ -rays, measured by two types of microscopes: AxioVision comprising ApoTome software and AxioImager A1 (C-control). Data obtained from 3 experiments are presented as mean \pm SEM.

doses of 12 and 16 Gy. For these two doses values obtained by AxioImager A1 microscope were low, thus are not given in the figure.

It has been noticed that higher radiation doses have weaker effects on DNA, as observed through the number of foci per nucleus. In order to perform further analysis of the results, the relative fluorescence was measured for threshold doses. According to the results presented in Figure 2, threshold doses are 12 and 8 Gy for AxioVision containing ApoTome software and AxioImager A1, respectively. From the results presented in Figure 3, representing relative fluorescence that was evaluated for threshold doses, it can be seen that 12 and 8 Gy have present the expected higher effect on the formation of the DSBs than 1 Gy of γ -rays.

For the Western blot analysis the cells were incubated for 30 min to 24 h after irradiation. The increase of the γ H2AX started 30 min after irradiation and was detected throughout of doses applied (Fig. 4). The acquired data for other post irradiation incubation times exhibited similar dose dependence (data not presented).

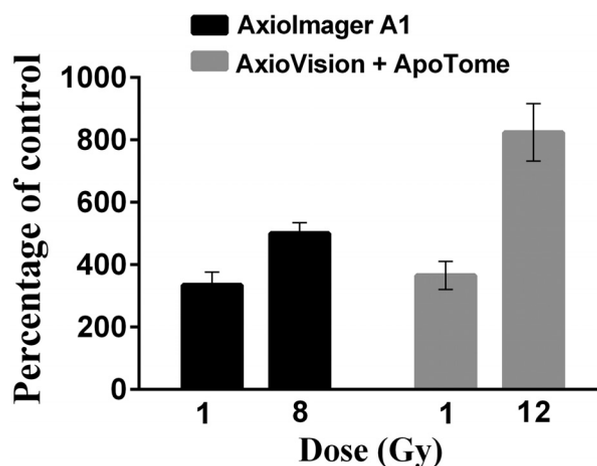


Figure 3 - Relative fluorescence, given as percentage of control, estimated with the microscopes AxioImager A1 and AxioVision comprising ApoTome software for the dose of 1 Gy each and the threshold doses of 8 Gy and 12 Gy, respectively. Data obtained from 3 experiments are presented as mean \pm SEM.

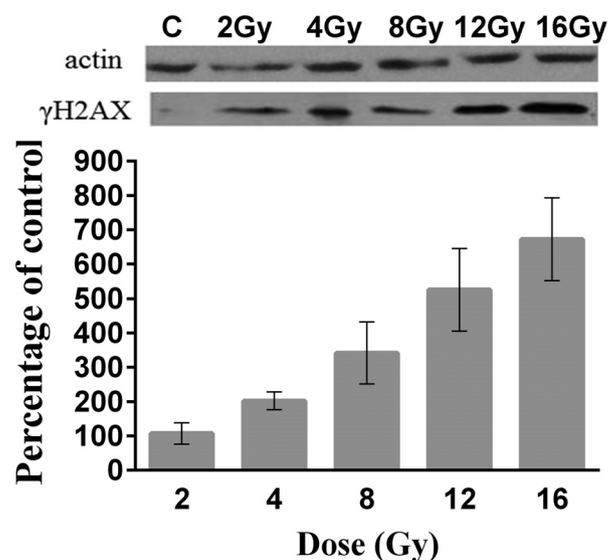


Figure 4 - Dose dependent phosphorylation of H2AX protein in HTB140 cells, given as percentage of control and analyzed by Western blot, 30 min after irradiation with γ -rays (C-control). Data obtained from 3 experiments are presented as mean \pm SEM.

Dynamic of the DNA repair in cells irradiated with 2 Gy γ -rays were analyzed by immunofluorescence microscopy using the two microscopes. The maximal number of γ H2AX

foci was attained 30 min after irradiation (Fig. 5). Again, the results obtained from AxioImager A1 resulted in lower values.

Nevertheless, for the same dose (2 Gy) Western blot analysis did not show time dependence of γ H2AX formation (appearance / disappearance). In the samples irradiated with 12 Gy the highest induction of γ H2AX was reached 2 h after irradiation, while it was significantly reduced 6 h post-irradiation (Fig. 6).

In order to investigate the unexpected weak effect for higher doses (12 and 16 Gy, Fig. 2), despite high immunofluorescence (Fig. 3), further analyses of images were performed. Among analyzed micrographs the one that was chosen is shown in Figure 7, presenting the sample irradiated with 12 Gy γ -rays. Lots of overlapping signals are most likely the reason of underestimation of the results presented in Figure 2.

CellProfiler software outlines foci and provides data that characterize each of them. The outlined γ H2AX foci in the HTB140 cell sample irradiated with 2 Gy γ -rays obtained by AxioVision

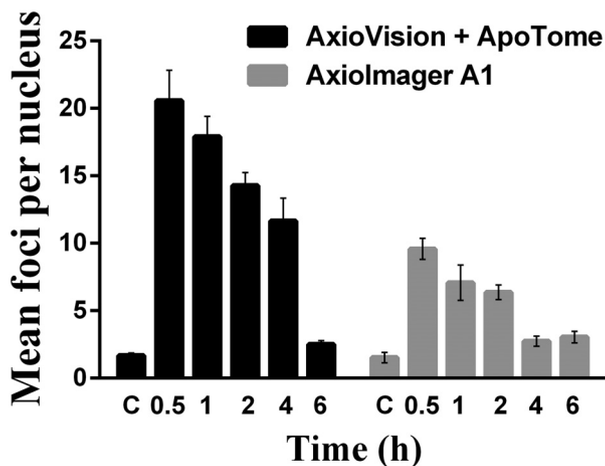


Figure 5 - Kinetics of appearance and disappearance of γ H2AX foci (time dependent mean number of foci) per HTB140 cell nucleus after irradiation with 2 Gy γ -rays (C-control). Quantification was performed with the microscopes AxioVision comprising ApoTome software and AxioImager A1. Data obtained from 3 experiments are presented as mean \pm SEM.

having ApoTome software (a) and AxioImager A1 (b) microscopes, shown in Figure 8. Due to the features of ApoTome software the real number of foci per nucleus was acquired and is presented in Figure 8a. However, on the micrograph obtained by AxioImager A1 presented in Figure 8b, only the foci that are in the focused object plane were recorded, while the others were detected as blurred structures.

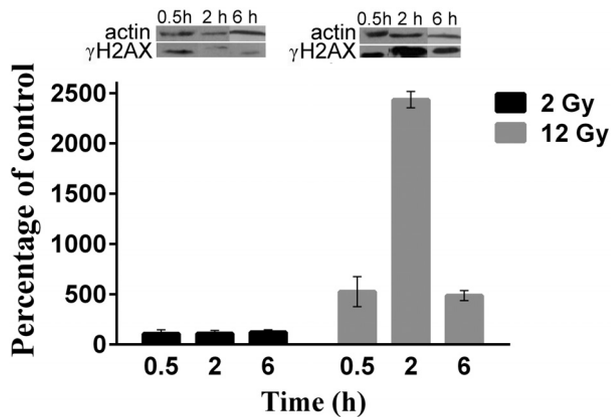


Figure 6 - Time dependent phosphorylation and dephosphorylation of H2AX protein in HTB140 cells, given as percentage of control and analyzed by Western blot, after irradiation with 2 and 12 Gy γ -rays (C-control). Data obtained from 3 experiments are presented as mean \pm SEM.

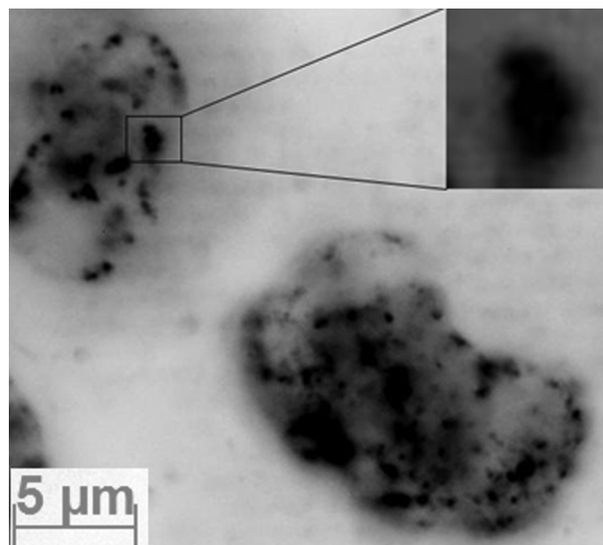


Figure 7 - Micrograph of HTB140 cell nuclei irradiated with 12 Gy γ -rays, showing overlapping foci, captured with AxioVision comprising ApoTome software.

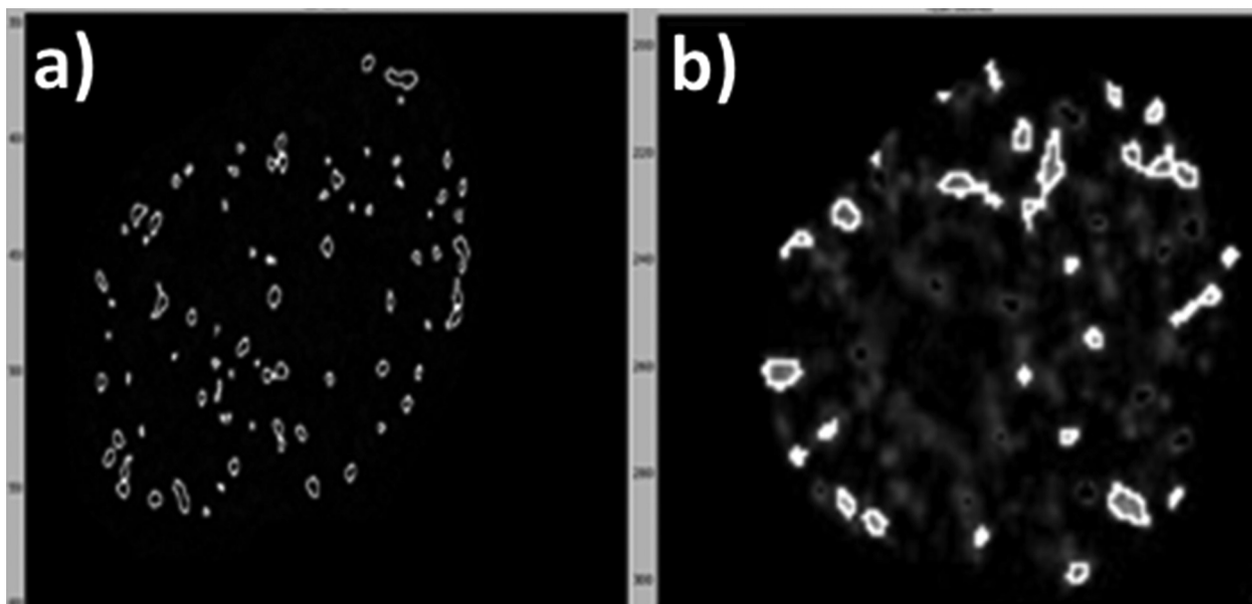


Figure 8 - Illustration of differences in counting of γ H2AX foci per HTB140 cell nucleus irradiated with 8 Gy γ -rays, obtained by the microscopes AxioVision comprising ApoTome software (a) and AxioImagerA1 (b). Objects of interest were outlined using CellProfiler image analysis software that was optimized for the analyses.

DISCUSSION

The detection of γ H2AX foci that reflect dynamics of the DNA DSB repair was analyzed after irradiation with low and high doses of γ -rays. The human HTB140 melanoma cells were used as a model system. Experimental conditions and data analyses were optimized according to the methods that were used.

Immunofluorescence microscopy enabled *in situ* visualization of each DNA DSB separately, as well as the distribution of breaks inside the nucleus of a single cell (Hu et al. 2005). Before counting, CellProfiler image analysis software was optimized. The software establishes whether the object that is analyzed is an individual one or if there are two or more clumped ones. This is a very important initial parameter which determines appropriate module by recognizing primary (nuclei) and secondary objects (foci). Most options within the module use this estimation of the size range of the objects in order to distinguish them from the noise in the image. Since pixels are used as units in CellProfiler, it is easy to zoom in on objects

and determine typical diameters (Carpenter et al. 2006). When the software parameters are correctly set the results obtained by software and by manual counting of foci are compatible (Fig. 1).

During capture of images for the analysis with AxioImager A1 microscope each image consists of signal contributions coming from the focused object plane. However it also includes contributions coming from the structures located above and below it. The blurred structures from above or below are either recognized to be out of focus or, if they are clearly outside the focal plane, they give the effect of brightening the image background (Ju et al. 2009). With the AxioVision comprising ApoTome software, a grid structure is inserted into the light field diaphragm plane of the reflected-light beam path. Advantage of this system, as compared to AxioImager A1, is its possibility to precisely make an image of the object plane. This is done by a high precision scanning mechanism that moves the grid pattern in defined steps across the sample plane. The result is a precise optical section through the specimen with no blurring and with high contrast (Conchello and Lichtman 2005).

Different performances of the microscopes explains some discrepancy in the number of foci detected with AxioVision having ApoTome software and AxioImager A1, presented in Figures 2 and 5. For higher doses, clumped objects that are shown in Figure 7 most probably cause underestimation of the number of foci. This can be misinterpreted as a weaker effect of irradiation. Image analyses through relative fluorescence (Fig. 3) indicate the need for parallel analysis on at least two different approaches for the same method, in this case for ICC. In combination with Western blot analysis, ICC confirms the assumption that results for higher doses presented in Figure 2 are underestimated, thus the doses of 8 and 12 Gy have stronger effects on the DNA level than the dose of 1 Gy of γ rays.

For analyzed cell system AxioVision with ApoTome software and AxioImager A1 are valid for up to 8 and 6 Gy, respectively. Differences in counting of foci by the two microscopes are illustrated in Figure 8 where the outlined γ H2AX foci correspond to the previously defined parameters in CellProfiler. In images captured with the AxioVision with ApoTome software there is negligible number of discarded γ H2AX foci, due to the ability of this software to make precise optical sections, thus giving higher number of foci per nucleus.

Quantification of phosphorylation of H2AX protein as a function of dose is also possible by Western blot analysis (Fig. 4), using actin for normalization purpose (Gaba et al. 2010). However, the analysis could not be fully attained with this method when using lower radiation doses at different post-irradiation incubation times (Fig. 6). In the Western blot assay, the protein to be quantified is coming from a large number of cells. Therefore, this method is not appropriate to provide the precise information about the level of γ H2AX per single cell. On the other hand, since the irradiation with doses of 1 or 2 Gy caused lower

level of phosphorylation of H2AX, the intensity of corresponding bands obtained by the Western blot do not follow the real changes in the γ H2AX kinetic (Fig. 6). Yet, the Western blot analysis was very useful for the analysis of phosphorylation of H2AX after irradiation with higher doses when the overlapping of signals disabled reliable quantification using immunofluorescence microscopy. Moreover, Western blot is a favorable method when following different cellular pathways triggered by radiation or other treatments (Wang et al. 2013).

In summary, since γ H2AX is considered to be a sensitive indicator of DNA DSBs, the development of different γ H2AX focus formation assays and the corresponding data analyses are of great experimental and even clinical importance. The described experimental techniques are becoming powerful tools for further studies of the cellular response to DNA damage. The combined use of ICC and Western blot analysis with respect to the dose level and other properties of a toxic agent, provide data about cellular processes important for the maintenance of genome stability.

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RESUMO

A radiação ionizante induz a rupturas dos filamentos duplos (DSBs) do DNA que ativam a fosforilação da proteína histona H2AX (γ H2AX). A coloração por imunofluorescência visualiza a formação de focos de γ H2AX, permitindo a sua quantificação. Esse método, em oposição ao ensaio de Western blot e citometria de fluxo, fornece uma análise mais precisa demonstrando a posição exata e a intensidade de sinal fluorescente em cada célula individualizada. Na prática, existem problemas na quantificação de γ H2AX. Esta pesquisa se baseia em duas questões: a determinação de qual

técnica deve ser aplicada, relativa à dose de radiação, e como analisar imagens de microscopia fluorescente obtidas por diferentes microscópios. Células de melanoma HTB140 foram expostas a raios- γ , com doses na faixa de 1 a 16 Gy. Efeitos das radiações a nível de DNA foram avaliadas em diferentes intervalos de tempo e após a irradiação analisada por Western blot e pela microscopia de imunofluorescência. Células coradas imuniquimicamente foram visualizadas por dois tipos de microscópios: microscópio AxioVision (Zeiss, Alemanha), que compreendem com software ApoTome, e o microscópio AxioImagerA1 (Zeiss, Alemanha). Os resultados obtidos mostram que o nível de γ H2AX é tempo e dose dependente. Microscopia de imunofluorescência forneceu uma melhor detecção de DSBs para doses de radiações mais baixas, enquanto a análise Western blot foi mais confiável para doses de radiações mais elevadas. Microscópio AxioVision contendo o software ApoTome foi mais adequado para a detecção de focos γ H2AX.

Palavras-chave: software ApoTome, microscópio Axio-ImagerA1, microscopia de imunofluorescência, Western blot, γ H2AX.

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