PhD Thesis

Signalling detection of DNA damage induced by low doses of ionizing radiation in human lymphocytes

Détection de la signalisation des cassures de l’ADN dans les lymphocytes humains par de faibles doses de rayonnements ionisants

Marco VALENTE

Directeurs de thèse : Laurence ROY, Patrick LALOI
Tutelle : Sandrine ROCH-LEFEVRE
This work is dedicated to

Cármen Valente and Fábio Valente
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SUMMARY

Individuals spontaneously present different sensitivities to ionizing radiation, measured by the severity of their post-radiotherapy side-effects. Cells from some patients with extreme clinical radiosensitivity have shown altered cellular radiosensitivity measured by different endpoints as apoptosis or DNA damage. Linking clinical and cellular sensitivity is of fundamental importance to establish a clinical test capable of predicting a person’s radiosensitivity from a sample. Easily sampled, peripheral blood lymphocytes (PBL) are an appealing cellular model to study individual radiosensitivity as they have been shown to be the most radiosensitive hematopoietic cells. DNA damages and repair can be visualized by observing the kinetics of appearance and disappearance of gamma-H2AX foci on DNA double-strand breaks through immunofluorescence microscopy. The experimental strategy chosen here was to follow lymphocyte gamma-H2AX foci kinetics in response to different levels of irradiation as delayed gamma-H2AX foci disappearance has been observed in cells of individuals with high clinical radiosensitivity.

For our initial study we irradiated *in vitro* samples of radiotherapy patients with different clinical radiosensitivities. The groups of distinct clinical sensitivities showed no corresponding differences in their cellular gamma-H2AX response. In addition, several samples were lost, mainly due to the long transportation period before being treated in our lab.

To render this method usable for clinical applications, several changes were made: after improving sample viability, speed was increased by automation of image acquisition (Metasystem) and gamma-H2AX focus scoring (freeware CellProfiler). This technique was able to detect doses as low as 0.005 Gy and gave similar results to manual focus scoring. The possibility of discriminating different lymphocyte subsets (CD4, CD8 and CD19) during analysis was added to identify among the lymphocyte subsets the one producing more gamma-H2AX foci in response to irradiation. Using the methodological tools developed during this thesis we established for the various lymphocyte subsets the relationship between radiation dose and gamma-H2AX foci frequency as well as the kinetics of appearance/disappearance of gamma-H2AX foci.
Finally, since no additional samples from patients of known radiosensitivity were available to continue the initial study with the improved protocol, we focused on radiosensitivity in another context: radio-adaptive response. This phenomenon corresponds to a lower cellular response to high dose of ionizing radiation exposure if it is preceded by an exposure to low doses. With the conditions used here we did not observe a radio-adaptive response in terms of gamma-H2AX signalling regardless of the lymphocyte subpopulation studied. However, the translocation rate of pre-irradiated CD4-positive lymphocytes was significantly different when compared to cells only irradiated acutely. This result thus indicates a differential repair of double strand breaks in lymphocytes after a radio-adaptation.
RESUME

Les individus présentent de façon spontanée une sensibilité variable aux rayonnements ionisants. Elle peut être mesurée par la gravité des effets secondaires observés post-radiothérapie. Les cellules de certains patients, présentant une hypersensibilité clinique aux rayonnements ionisants, ont montré une radiosensibilité cellulaire altérée, mesurable par différents critères tels que l’apoptose ou les dommages à l’ADN. Étudier le lien entre les radiosensibilités cliniques et cellulaires est d'une importance fondamentale pour établir un test capable de prédire la radiosensibilité d'une personne à partir d'un échantillon. Faciles à prélever, les lymphocytes du sang périphérique sont un modèle cellulaire attirant pour étudier la radiosensibilité individuelle étant qu’ils se sont avérés être les cellules hématopoïétiques les plus radiosensibles. Dans la cellule, les cassures de l’ADN ainsi que leur réparation peuvent être visualisées grâce à la cinétique d’apparition et de disparition des foyers formés par la protéine gamma-H2AX. Ils sont observés par microscopie à l’aide de technique d’immunofluorescence. La stratégie expérimentale choisie est la quantification des foyers gamma-H2AX lors de cette cinétique, dans les lymphocytes de personnes présentant une radiosensibilité clinique élevée.

Pour notre étude initiale, nous avons irradié in vitro des échantillons de patients de radiothérapie ayant des radiosensibilités cliniques différentes. L’analyse de la cinétique de réponse de la protéine gamma-H2AX n’a pas permis de mettre en évidence une corrélation directe entre le nombre de cassures et le niveau de radiosensibilité des patients. Cependant, plusieurs échantillons ont été perdus, principalement à cause du temps de transport entre le prélèvement du patient et l’arrivée de l’échantillon au laboratoire.

Ainsi, pour rendre cette méthode utilisable en application clinique, plusieurs changements ont été faits : l’amélioration de la viabilité de l’échantillon ainsi que l’augmentation de la vitesse d’analyse grâce à l’automatisation de l’acquisition des images (logiciel « Metafer4 » de MetaSystems) et du dénombrement des foyers gamma-H2AX (logiciel gratuit « CellProfiler »). Cette méthodologie a permis la détection de doses d’irradiation in vitro à partir de 0,005 Gy, tout en donnant des résultats similaires au comptage manuel.
De plus, nous avons étudié la réponse à différentes doses d’irradiation des sous-populations lymphocytaires CD4, CD8 et CD19. Les outils méthodologiques développés lors de cette thèse, nous ont permis d’établir pour les différentes sous-populations lymphocytaires, d’une part la relation entre la dose d’irradiation et la fréquence de foyers gamma-H2AX et d’autre part la cinétique d’apparition / disparition de ces foyers.


La réponse radio-adaptative est caractérisée par une réponse cellulaire, à une forte dose d’exposition, moins importante lorsqu’elle est précédée par l’exposition à une faible dose.

Dans nos conditions expérimentales, nous n’avons pas observé de réponse radio-adaptative par l’analyse de la signalisation gamma-H2AX indépendamment de la sous-population de lymphocytes étudiée. Néanmoins, le taux de translocations observé dans les lymphocytes CD4-positifs pré-irradiés est significativement différent du taux observé dans les lymphocytes irradiés directement avec la forte dose. Ce résultat nous permet de penser que la réponse radio-adaptative dans les lymphocytes impliquerait une réparation des cassures double-brin différente.
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1 INTRODUCTION

1.1 Ionizing radiation

Mankind has always been exposed to ionizing radiation (IR), mainly of natural origin (like radon gas and cosmic rays) and varying with location and altitude. Since the discoveries of Becquerel and the Curie, different types of ionizing radiation have been used in several technological fields. Therefore, since the end of the 19th century there is also an artificial component to human exposure, from certain medical treatments (radiography and cancer radiotherapy, for example), nuclear testing fallout, the Chernobyl accident, and industrial discharges.

Ionizing radiation is said to be ionizing if composed of particles that individually carry enough energy to detach electrons from atoms or molecules. Ionizing radiation is generally the result of the natural decay of unstable atomic nuclei (radioactivity) but can also be artificially generated (by particle accelerators, for instance). When particles traverse matter they transfer their energy to the medium primarily by ionizing atoms along their path. This energy is measured in electron volt (eV). A molecule of water (H\textsubscript{2}O), for example, can be ionized by approximately 14 eV. The Linear Energy Transfer (LET) is the amount of energy released by the particle over the length of its decay track. When used to describe the dosimetry of ionizing radiation in a biomedical context, LET is usually expressed in keV/\mu m. Therefore, the different types of ionizing radiation vary by their emitted particle and penetration power and can be classified into two groups according to the density of the deposited energy: High LET and Low LET radiation.

1.1.1 High LET radiation

High LET radiation typically releases 400 eV or more within target areas of approximately 5 to 10 nm (Goodhead 1989).
The spontaneous emission of **alpha particles** occurs during the radioactive decay of elements that have a mass number greater than about 150 (uranium and plutonium, for example). Alpha particles are helium atom nucleus, consisting of two protons and two neutrons. They are therefore heavy and electrically charged, rapidly depositing their energy within a short (20 to 100 µm) and almost linear track. The energy lost by these particles during their travel through matter peaks immediately before they come to rest (a phenomenon called "Bragg peak"). This happens because the total interaction cross section increases as the charged particle's kinetic energy approaches zero.

Similarly to alpha radiation, in **ion radiation** the particle is charged and also has a linear track with a Bragg peak. The particles penetration power and distance of the Bragg peak depend on the atom that generated the ion (iron, carbon, protons, etc.) and the initial energy. A variety of ion beam sources exist, from mercury vapour thrusters to particle accelerators.

**Neutrons**, as their name suggests, are not charged particles. They are not ionizing but their collisions with atom nuclei result in a chain reaction that leads to the release of charged particles (like alpha, heavy ions or protons) that do cause ionization. Therefore, the neutrons energy deposition will depend on the secondary particles produced. Neutrons are very penetrating and have a non-linear energy deposition track since free neutrons change direction with each collision.

### 1.1.2 Low LET radiation

Low LET radiation typically releases 100 eV or less within target areas of approximately 2 nm (Goodhead 1989).

#### 1.1.2.1 Beta radiation

There are two types of beta particles: electrons and positrons ("anti-electrons"). They are originated by the beta decay of atom nuclei with an excess of neutrons or protons, respectively. This radiation has a weak penetrating power, with an energy track profile that depends on the initial
energy. For higher energy values, the track starts off linear and gets more and more deviated as the energy is lost. Also to be considered is the possible ramification of this track due to secondary electrons generated by the deposition of energy. Beta decay is a common by-product in nuclear reactors and free neutrons also decay via this process.

1.1.2.2 Gamma-rays and X-rays

Gamma-rays are an electromagnetic radiation of high frequency \(10^{19}\) Hz, with energies above 100 keV and almost always less than 10 MeV) produced by sub-atomic particle interactions such as electron-positron annihilation, neutral pion decay, radioactive decay, fusion, or fission. After a decay reaction, the nucleus is often in an “excited” state. This means that the decay results in producing a nucleus which still has energy excess to get rid of. Rather than emitting another beta or alpha particle, this energy is lost by emitting a pulse of electromagnetic radiation called gamma-ray. The gamma-ray is identical in nature to light or microwaves, but of very high energy.

The upper limit of energy for such radiation is about 20 MeV, and there is effectively no lower limit (they are sometimes classed as X-rays if their frequencies are lower than \(10^{19}\) Hz). The distinction between gamma- and X-rays is not based on their intrinsic nature but rather on their origins. X-rays are emitted during atomic processes involving energetic electrons. Gamma-rays are emitted by excited nuclei or other processes involving subatomic particles. The applications of gamma-rays are much the same as those of X-rays.

Like all forms of electromagnetic radiation, gamma-rays have no mass and no charge. Gamma-rays interact with material by colliding with the electrons in the shells of atoms. They lose their energy slowly in material, being able to travel significant distances before stopping. Depending on their initial energy, gamma rays can travel from 1 to hundreds of meters in air. It is important to note that most alpha- and beta- emitters also emit gamma-rays as part of their decay process.
In the work reported here we are using 2 sources of gamma-rays: cobalt-60 and cesium-137.

1.1.3 Units used in radiobiology

The unit currently used to measure the energy absorbed is the Gray (Gy, honouring the physicist Stephen Gray). One Gy corresponds to 1 joule per kilogramme of irradiated matter. Previously the dose was expressed in RAD (for Radiation Absorbed Dose), where 100 RAD = 1 Gy. This way of measuring does not take into account the LET of the ionizing radiation but varies with both the number of ionizing particles and the atomic number of the traversed material. Dose rate is the dose absorbed per unit of time. This rate can be important as different dose rates can lead to distinct biological responses (Boreham et al. 2000). The equivalent dose is a measure of the radiation dose to tissue where an attempt has been made to allow for the different relative biological effects of different types of ionizing radiation. Equivalent dose is therefore a less fundamental quantity than radiation absorbed dose, but is more biologically significant. Equivalent dose has units of sieverts (Sv).

1.1.4 Relative biological effectiveness

Relative biological effectiveness (RBE) is the ratio of a dose of a reference radiation to the dose of the radiation of interest required to cause the same level of biological effect (chromosomal aberrations, apoptosis, etc). In human, studying radiation RBE has direct practical implications in therapy applications and in assessing risks from environmental and occupational exposures. Moreover, it provides analytic information on the underlying mechanisms of radiation biology. For many relevant effects in mammalian systems there is a general tendency to an increase in RBE with increasing LET, up to a maximum followed by a decrease at very high LET. However, the values of RBE for a given LET can vary by large amounts (even orders of magnitude) depending on other physical and biological conditions. Common general tendencies in mammalian systems are: RBE for mutations to be
greater than for cell inactivation; for particles such as protons or electrons to reach their peak at lower LET than heavy ions (such as alpha particles or carbon ions); for RBE to be larger at lower doses and dose rates; and for radiosensitive cells to show lower RBE than radioresistant cells. There are, however, many exceptions to such generalities. Therefore, we can conclude that there are a number of competing mechanisms and diverse factors that determine the effectiveness of high-LET radiations. This makes it difficult to identify the individual mechanisms and to predict RBE for individual practical application.

1.2 Biological effects of ionizing radiation

1.2.1 Radiation-induced lesions

Biological systems are structured in various levels of organization: atoms, molecules, cells, tissues, and organisms. Ionizing radiation primarily interacts with atoms and biological responses may become observable ascending to every level of organization. The probability of such observation largely depends on complex signalling within and between cells at every level of organization controlling homeostasis and adaptation of the whole system (Feinendegen et al. 2007). Proteins, cell membranes and DNA are targets of radiation exposure, and effects on these structures can be produced by direct damage as well as free radicals, including reactive oxygen and reactive nitrogen species (ROS and RNS, respectively) (Daly et al. 2007). DNA damage is formed by the ionizations/excitations of DNA or the surrounding material, mostly water within a radical-diffusion distance of 4 nm from the DNA in the cellular environment (Goodhead 1999). Various forms of DNA damage are induced by ionizing radiation. Previous publications allow the rough estimation of the yields of DNA damage in mammalian cells caused by low-LET radiation exposure: for each diploid cell, 1 Gy induces 1000 simple-strand breaks, 10000 base damage, 150 protein-DNA cross-links, 70 bulky lesions (ie, clusters of base damage), and 40 double-strand breaks (DSB) (Averbeck 2000).
Figure 1. IR-induced DSB. As a consequence of its physical properties, IR deposits a sizable proportion of its energy in the form of ionization clusters that generate clustered damage in the DNA. From such clusters of damage DSBs can form promptly, after chemical processing of sugar lesions, which in their initial form do not disrupt the sugar-phosphate backbone, or after the enzymatic processing of base damage. (Mladenov & Iliakis 2011)

In this work, we have focused primarily on DSB signalling and repair. DSB are arguably the most significant lesions since, if unrepaired it can result in cell death (Frankenberg-Schwager 1989) and if misrepaired, it can cause chromosomal translocations, an early step in the etiology of carcinogenesis (Pa Jeggo & M Lörich 2007). IR-induced DNA double-strand breaks can result from pairs of direct ionizations or pairs of nearby hydroxyl radicals or by combined direct and OH action (Goodhead 1999) (Figure 1). In addition to exposure of cells to ionizing radiation, DSB can result from other exogenous sources like topoisomerase poisons or radiomimetic drugs. Naturally occurring DSB (endogenous) are generated spontaneously during DNA synthesis when replication forks encounter damaged templates and during certain specialized cellular processes, including V(D)J recombination, class-switch recombination at the immunoglobulin heavy chain (IgH) locus and meiosis (Hartlerode & Scully 2009).
1.3 DNA Damage Response

The DNA damage response (DDR) is a signal transduction pathway that senses DNA damage and replication stress and sets in motion a choreographed response to protect the cell and ameliorate the threat to the organism (Ciccia & Elledge 2010). First, sensor proteins detect DNA damage and/or chromatin alterations that occur after damage induction. The signal is then transmitted to transducer proteins, which are usually kinases that amplify the damage signal. At the end of the chain, involved in specific pathways, are the effectors (Figure 2). Another class of DDR proteins is the mediators, which promote interaction between the other proteins. The multiple signalling cascades implicated in DDR can result in a variety of cellular responses: cell cycle arrest, induction of stress response genes, DNA repair and cell death.

The DDR is regulated by the phosphoinositide three-kinase-related protein kinases (PIKK). The PIKK primarily responsible for signalling the presence of DNA damage include ATM (Ataxia Telangiectasia Mutated), ATR (Ataxia Telangiectasia and Rad3-related protein) and DNA-PKcs (DNA Protein Kinase catalytic subunit). The signal that activates ATM and DNA-PKcs is a double-
strand break, while ATR responds to single-stranded DNA gaps. All three kinases are recruited to the DNA lesion site, which promotes kinase activation. These PIKK phosphorylate hundreds of proteins that maintain genome integrity through regulation of cell cycle progression, DNA repair, apoptosis, and cellular senescence (Lovejoy & Cortez 2009).

1.3.1 DSB signalling - Gamma-H2AX IRIF

Upon DSB induction by ionizing radiation, there is an extensive response in the chromatin flanking the break. Hundreds of molecules of multiple DNA damage response protein species accumulate at DSB sites forming large nuclear aggregates that appear as IR-induced nuclear foci (IRIF) by fluorescence microscopy (Nakamura et al. 2010) (Figure 3). Increasing evidence suggests that IRIF are required for accurate and coordinated DSB repair in the context of chromatin (Stucki et al. 2005).

![Figure 3](image-url)

*Figure 3 Recruitment of DNA sensing and repair proteins as irradiation-induced foci (IRIF) at a DSB. The DNA DSB is expanded to illustrate a histone octamer containing irradiation-induced phosphoforms of H2AX (e.g., gamma-H2AX) which can involve megabase domains surrounding the DSB. At the DSB itself, are DNA-PK (DNA-PKcs kinase and Ku autoantigen subunits form Ku70-Ku80 heterodimers which act to initiate NHEJ), RAD51-BRCA2 (if S or G2 phase to initiate HR) and the MRE11 sensing complex. Following DSB induction, MRN binds to DNA ends and facilitates recruitment and activation of ATM, the kinase responsible for the phosphorylation of H2AX. Gamma-H2AX mediates the recruitment of numerous DSB recognition and repair factors including MDC1, BRCA1, 53BP1 and ATM (Bhogal et al. 2009).*
A key regulator of IRIF formation in mammalian cells is the histone subtype H2A isoform X (H2AX), a component of the nucleosome core structure that comprises 10%-15% of total cellular H2A in higher organisms (Fernandez-Capetillo et al. 2004).

Some DDR factors have intrinsic affinity for free DNA ends. The MRN complex (Mre11/Rad50/NBS1) can detect DSB and bind to the DNA ends (M F Lavin 2007). Given its structure, the MRN complex is thought to form a microenvironment that holds the two DNA ends together (van Gent & van der Burg 2007).

ATM is then indirectly recruited to DSB and activated by the MRN complex via an interaction between the C-terminus of NBS1 and ATM (M F Lavin 2007).

One of the first substrates of ATM to be phosphorylated is histone H2AX: at the sites of DSB, H2AX becomes phosphorylated on C-terminal Ser139 (E P Rogakou et al. 1998). This modified H2AX called “gamma-H2AX” forms within seconds, and reaches its maximum level at about 30 min after irradiation (E P Rogakou et al. 1998) (Figure 4 A).

In this work, we used fluorochrome-coupled antibodies against gamma-H2AX to be able to quantify gamma-H2AX foci using fluorescence microscopy. Scoring gamma-H2AX foci has been described as an assay capable of DSB quantification induced by ionizing radiation (E P Rogakou et al. 1998). However it is important to keep in mind that a gamma-H2AX assay measures a biological response to DSB and not the actual “physical” DNA damage. There is DNA repair within the 30 min gamma-H2AX takes to peak (measured in Figure 4 B by the comet assay), given the fast kinetics of DNA repair. This implies that the DSB detected by the gamma-H2AX assay are probably an underestimation of the “real” DSB formed.

ATM seems to be the main kinase associated with gamma-H2AX formation under normal physiological conditions but all three major PIKK members, ATM, ATR and DNA-PKcs can phosphorylate H2AX (Stiff et al. 2004; Shrivastav et al. 2008; Hartlerode & Scully 2009).
This initial gamma-H2AX is required for further DDR signal amplification (Stucki & Stephen P Jackson 2004). Gamma-H2AX interacts with the mediator protein, MDC1 (mediator of DNA damage checkpoint 1) (Stucki et al. 2005), which in turn promotes the recruitment of 53BP1 (p53-binding protein 1) in the vicinity of the DSB via the ubiquitylation of histone H2A. Both MDC1 and 53BP1 interact with MRN, leading to more ATM tethering at the DSB site (Goodarzi et al. 2010). ATM phosphorylates key proteins that lead DNA repair, to cell cycle checkpoint arrest and/or apoptosis. ATM substrates include Chk1, Chk2, Rad17, NBS1, BRCA1, BLM, SMC1, 53BP1, p53 and MDC1 (Stiff et al. 2004) (Schmitt et al. 2007). It is thought that MDC1 reinforces the gamma-H2AX signal. However, the extent of spread of the signal is not dependent upon MDC1 (Hartlerode & Scully 2009).
Other DNA damage response proteins like BRCA1/BARD1 (BRCA1-associated RING domain 1) and chromatin remodeling factors accumulate on gamma-H2AX-containing chromatin (Kusch et al. 2004; A. J. Morrison et al. 2004). Chromatin architecture is of fundamental importance to proteins that need to access DNA for transcription or replication, so it is not surprising that this is also the case for proteins involved in DNA repair (Goodarzi et al. 2010) (Rossetto et al. 2010).

H2AX phosphorylation is not limited to the immediate vicinity: it spreads to a large chromatin region surrounding the DSB. It has been estimated that in mammals 0.03% of H2AX is phosphorylated per DSB. It is not clear how H2AX phosphorylation is spatially confined, but 4 Pi microscopy suggests that H2AX is not distributed randomly throughout bulk chromatin but exists in distinct clusters that define the boundaries of gamma-H2AX spreading (Kinner et al. 2008).

Other DDR proteins such as Mre11 and Rad51 exhibit a different distribution, accumulating in smaller regions at the DSB site. These data suggest that different DDR proteins may have their own territory in a DSB focus (Nakamura et al. 2010).

Although not crucial, gamma-H2AX facilitates DSB repair by either of the two main pathways (Celeste et al. 2002; Hartlerode & Scully 2009; Bassing et al. 2002).

### 1.3.2 DNA double strand break repair

The integrity of the DNA is essential for the correct information transfer and mechanical segregation of the chromosomes during cell division. Evolution has put in place several surveillance mechanisms to allow a higher fidelity of genetic material transmission from a cell to its daughter cells. The cell cycle has 4 checkpoints to verify genome integrity: the end of G1, the S phase, the end of G2 and the end of the M phase. Dividing cells after irradiation can be blocked at G1 or G2. This is due to the activation of p53 by ATM or ATR. Cells take advantage of the cycle arrest to repair the radiation-induced damage.
done to their DNA. The cell only continues through the checkpoint after having dealt with the “repairable” DNA damage.

The two major DSB repair pathways are Homologous Recombination (HR), and Non-Homologous End-Joining (NHEJ) (Figure 5). Many proteins of the initial DDR and DNA repair overlap. Furthermore, several proteins are known to be implicated in both repair pathways: Mre11, BRCA1, H2AX, PARP-1, Rad18, DNA-PKcs, and ATM. Although NHEJ factors are recruited to DSB more rapidly than HR factors, and NHEJ and HR factors are independently recruited to DSB (Kim et al. 2005), there is a significant period of time when both sets of factors are present at damage sites. This is consistent with the notion that pathway choice may be regulated by one or more proteins that act in both pathways (Shrivastav et al. 2008).

1.3.2.1 Homologous Recombination

Homologous Recombination is generally an error-free pathway of homology-directed repair. A DSB is accurately repaired by using the undamaged sister chromatid as a template for the repair of the broken sister chromatid. Therefore, HR is limited to the late S to G2 phase of the cell cycle, when sister chromatids are available.

An early step in HR is resection of DNA ends at the DSB site to yield 3’-single-stranded DNA (ssDNA) overhangs which are capable of invading duplex DNA containing a homologous sequence. This process involves the MRN complex, which possesses an endonuclease and 3’→5’ exonuclease activity. The generated single-stranded region of DNA initially becomes coated with the single strand binding protein, RPA. RPA is then subsequently replaced by Rad51, which promotes invasion of the template strand. Depending on the precise nature of the process, a Holliday junction may then be generated, followed by branch migration and finally resolution of the Holliday junction. Homologous recombination in eukaryotes is carried out by the RAD52 group genes (Rad50, Rad51, Rad52, Rad54, Rdh54/Tid1, Rad55, Rad57, Rad59, Mre11, and Xrs2), most of which were identified by their requirement for the
repair of ionizing radiation-induced DNA damage in *Saccharomyces cerevisiae* (Symington 2002).

![Figure 5 DNA DSB repair](image)

The products of the breast cancer susceptibility genes, BRCA1 and BRCA2, are also involved in the modulation of the homologous recombination. Brca2 functions in regulating Rad51 loading onto DNA and Brca1, also, most probably plays a regulatory role in the process (P. Jeggo & Markus Löbrich 2006).
1.3.2.2 Non-Homologous End Joining

NHEJ is a repair pathway that does not require a homologous DNA template, although short patches of homologous nucleotides (microhomologies of one or a few base pairs) are frequently used to align the DNA ends (van Gent & van der Burg 2007). The NHEJ pathway seems to be the main DSB repair pathway in mammals (Roth & J. H. Wilson 1985) and is not restricted to a particular phase in the cell cycle and hence DSB can be repaired via NHEJ throughout the cell cycle (Takata et al. 1998).

NHEJ repair starts with the Ku complex (the heterodimer of Ku70 and Ku80) recognizing and binding to the DSB ends of the damaged site (Mahaney et al. 2009). DNA-dependent protein kinase catalytic subunit (DNA-PKcs) is recruited to DSB ends and becomes activated. Together, the Ku complex and DNA-PKcs are referred to as DNA-PK. The activated DNA-PKcs phosphorylates itself as well as several substrates including p53, the Ku complex, DNA Ligase IV/XRCC4 (X-ray repair complementing defective repair in Chinese hamster cells 4) (Drouet et al. 2005) and replication protein A (RPA) (Hongyan Wang et al. 2001), all of which may facilitate end processing reactions (Roberts et al. 2010).

Since DSB can occur with a variety of different ends, a number of processing enzymes may be required to repair breaks. Together with the Artemis nuclease, DNA-PKcs can stimulate processing of the DNA ends (Hartlerode & Scully 2009). End processing involves the removal of damaged or mismatched nucleotides by nucleases and/or re-synthesis of ssDNA by DNA polymerases Pol-mu and Pol-lambda DNA polymerases. The end processing step can be skipped if DNA ends are compatible. It is particularly important when DSB contain unligatable ends, because all DNA ligases, catalyze the formation of a phosphodiester bond between 5’-phosphate and 3’-hydroxyl termini. DSB located within heterochromatin require additional factors for their repair prior to rejoining by NHEJ. For heterochromatic DSB, ATM phosphorylation of Kap1 allows localised chromatin relaxation facilitating repair by NHEJ. This process requires Artemis and the mediator proteins in addition to ATM (P. Jeggo & Martin F Lavin 2009).
Finally, the compatible ends are rejoined by the DNA Ligase IV/XRCC4/XLF complex (Grawunder et al. 1997; Mahaney et al. 2009).

1.3.2.3 Chromosomal aberrations - Erroneous DSB repair

The efficiency of the DNA damage repair processes determines the outcome of the cell: most commonly, the structure of DNA is repaired correctly and cellular functions return to normal. If repair is unsuccessful or incomplete cells die. It has been established a direct link between radiation-induced unrepaired DSB and cell death (N Foray et al. 1997). In the other hand, if DSB repair is imprecise, cells suffer alteration and loss of genetic information seen as mutations and chromosomal aberrations (Figure 6).

![Figure 6. Chromosomal aberrations analyzed in this work. A - The cells of interest are lymphocytes, which are in G0 (1 n) when the radiation-induced damage will take place. The yellow lines represent double strand breaks. B - Cells correctly repair most of the damages but some DSB are incorrectly resolved (orange lines). C - To observe the aberrations we stimulate the cells, which start dividing (2 n), and we block cell division at the phase where chromatin is most compacted: metaphase.](image)
The resulting translocations can activate oncogenes and lead to cancer (Agarwal et al. 2006).

1.3.2.4 Radiation-induced cell death

Unrepaired or misrepaired DSB can lead to p53-mediated apoptosis or mitosis-associated cell death (Kolesnick & Fuks 2003). The p53 is activated by ATM directly or indirectly through Chk2 phosphorylation. Apoptosis is a physiological process in which the cell initiates a highly regulated and evolutionary conserved molecular program, resulting in its own death.

An early event in apoptosis is the flipping of phosphatidylserine (PS) of the plasma membrane from the inside surface to the outside surface. In the experiments reported here, apoptotic cells were distinguished from their intact counterparts using Annexin V, which is a phospholipid-binding protein that binds selectively and with a high affinity to PS (D Vokurková et al. 2006). A late apoptotic event is DNA fragmentation, during which gamma-H2AX focus formation can occur (Emmy P Rogakou et al. 2000).

In dividing lymphocytes, exposure to ionizing radiation leads to mitotic death. This phenomenon occurs as the result of incomplete or unsuccessful segregation of chromosomes. This disturbance in chromosome segregation can induce apoptosis in the mother (or daughter) cells. It can also induce the formation of micronuclei (which are easily confused in terms of recognition, with apoptotic fragments)(Potten & J. W. Wilson 2004).

1.4 Radiosensitivity

Radiosensitivity is the relative susceptibility of cells, tissues, organs or organisms to the harmful effect of ionizing radiation. In this work, we focused on radiosensitivity at two of these levels: Clinical radiosensitivity and Cellular radiosensitivity.
1.4.1 Clinical Radiosensitivity

Radiotherapy is a very important non-surgical modality for the curative treatment of cancer. With improved cancer cure rates, the intensity of the side-effects becomes an increasingly important issue.

The clinical manifestations of radiosensitivity can be either acute (occurring during or within weeks of treatment) or late (occurring 6 months to many years later) radiation toxicity and are well documented. Acute effects are usually reversible and occur in rapidly proliferating tissues, such as skin, gastrointestinal tract and the haematopoietic system. Late side effects can be permanent and typically occur in more slowly proliferating tissues, such as kidney, heart and central nervous system. The pathogenesis includes fibrosis, atrophy and vascular damage. Other important late normal tissue side effects include hormone deficiencies, infertility and second malignancies. Generally, no clear severity correlation between acute and late radiotherapy side-effects has been found (Azria & Ozsahin 2009). Exceptions to this are some recent aggressive treatments, involving unconventional fractionation protocols or radiochemotherapy schedules (Dörr & Hendry 2001).

Exposure to therapeutic doses of ionizing radiation revealed an inter-individual variability in terms of the intensity of normal tissue responses. In this work the clinical reactions were determined in respect to acute dysphagia, dermatitis and mucositis using the Common Terminology Criteria for Adverse Events (CTCAE) scale, version 3.0, of the National Cancer Institute. Patients were subdivided according to their total average acute radiosensitivity. Patients without grade 3 dysphagia, mucositis or dermatitis were assigned to group CTC1 (mild acute radiosensitivity), patients showing at least two grade 3 toxicities were considered to be severely radiosensitive and assigned to group CTC3. Patients showing only one grade 3 reaction were considered as an intermediate category (moderate acute radiosensitivity; group CTC2).
Although a number of factors, such as patient size and variation in dose distribution, are believed to influence radiotherapy side-effects, it has been estimated that up to 70% of the inter-individual differences in clinical radiosensitivity may be due to genetic predisposition (Turesson et al. 1996). Indeed, within the 5% fraction of patients with the most severe reactions we can find individuals with known genetic diseases, frequently due to mutations in DDR or repair proteins such as ATM or NBS1 (Bentzen & Overgaard 1994)(Figure 7). In fact, clinical hyper-radiosensitivity was first reported in ataxia telangiectasia (A-T) patients treated with radiotherapy. These observations were later extended to lymphocytes from A-T patients where increased aberrations were reported post-irradiation (Khanna et al. 2001).

**Figure 7. Diseases caused by defective responses to DNA double-strand breaks.** Ataxia-telangiectasia mutated (ATM) deficiency results in a pleiotropic phenotype, the most debilitating feature of which is progressive neurodegeneration. Mre11 mutations are associated with A-T-like disease (ATLD), which mimics the neurodegenerative aspect of ataxia telangiectasia (A-T). Though NBS1 acts with Mre11 to regulate ATM activation, NBS1 hypomorphic mutations are associated with microcephaly, not neurodegeneration. The Nijmegen breakage syndrome (NBS) phenotype overlaps with ATM and rad3-related (ATR) deficiency more so than with A-T. Mutation of LIG4 is also associated with microcephaly and resembles NBS and ATR-Seckel syndrome. (Phillips & McKinnon 2007)

However there is also a subset of patients that show severe reactions to radiation therapy without presenting obvious congenital defects. It is ultimately the radiosensitivity of the few hypersensitive patients that limits
our ability to increase the treatment dose in non-sensitive patients, which in turn should increase local control and cure (Barnett et al. 2009).

Radiosensitivity syndromes illustrate that specific genes influence the radiosensitivity of tissues and cells derived from them, but even cells cultured from apparently normal populations present a wide range of cellular radiosensitivities (Little et al. 1988). Therefore, much remains to be discovered about the more moderate radiosensitivity cases and the influence of other genes, not associated with these syndromes.

1.4.2 Cellular Radiosensitivity

Different cell types have distinct radiosensitivities. In general, it has been found that cell radiosensitivity is directly proportional to the rate of cell division and inversely proportional to the degree of cell differentiation. Therefore, actively dividing cells or those not fully mature are most at risk from radiation (Bergonie & Tribondeau 1906). The same cell type can also vary in sensitivity to radiation depending on the phase of the cell cycle it was going through at the moment it was exposed. Cells are least sensitive when in the S phase, then the G1 phase, then G2 phase and the most sensitive in the M phase of the cell cycle.

Historically, cell survival has been used as a reference endpoint for determining the radiosensitivity status of the cell. The doses used in this work, however, induce low levels of apoptosis in resting lymphocytes. Therefore, we measured “radiosensitivity” by quantifying gamma-H2AX response. This is important to mention because the degree of cellular radiosensitivity can vary depending on the endpoint used to quantify the response to radiation. Some cell types respond with higher intensity with one endpoint but are considered less radiosensitive in experiments using different endpoints. Using cell survival as the endpoint, B-lymphocytes are more radiosensitive than T-lymphocytes, as their numbers decline more rapidly even after relatively low doses (Prosser 1976). However, using gamma-H2AX response as the measure of radiosensitivity, Andrievski and Wilkins found that B-cells were the lymphocytes with the lowest response to radiation.
This is logical if we consider that failure to recognise and/or repair DNA lesions can negatively influence the survival of the cell and, therefore increase its radiosensitivity status.

1.5 Radio-adaptive response

First described by Olivieri et al. (Olivieri et al. 1984), radio-adaptive response (or radioadaptation) is defined as the reduced damaging effect of a challenging radiation dose when induced by a previous low priming dose (or conditioning dose). This damaging effect has been measured using various biological endpoints in both in vitro and in vivo studies: cell lethality, gene mutations, repetitive DNA loci mutations, chromosomal aberrations and micronuclei induction, neoplastic transformation, microarrays showing gene expression changes, DNA single- and double-strand breaks, biochemical analyses of enzymatic and/or non-enzymatic antioxidant defence system (Tapio & Jacob 2007; Dimova et al. 2008) (Figure 8).

Radio-adaptive response belongs to the group of non-targeted effects that do not require direct exposure of the cell nucleus and is reported to be tightly related to cellular responses such as a bystander effect, genetic instability, and hyper-radiosensitivity (HRS) (Ojima et al. 2011; Matsumoto et al. 2009; Joiner et al. 1996).

![Figure 8. Some underlying mechanisms probably involved in radio-adaptive response. (Dimova et al. 2008)](image)

Bystander effects are radiobiological effects in unexposed cells caused by transferable factors. Genetic instability is defined as the delayed onset of de
novo aberrations and mutations or a delayed reproductive death; and
ehypersensitivity is a change in the dose-effect relationship which occurs at
doses around 0.5 Gy.
An important factor that separates radio-adaptive response from these other
responses is that there are 2 exposures to radiation in radio-adaptive
response.
Although adaptive response seems to function by an on/off principle, it has
shown to have a high degree of inter- and intra-individual variability (Leal et
al. 1995; K. J. Sorensen et al. 2002). Extensively observed in lymphocytes,
radio-adaptive response is known to have considerable diversity in these cells:
in some cases lymphocytes show additive effects or no response at all.
Furthermore, the observation of this phenomenon seems to be restricted to
somewhat specific conditions: the range of the priming doses (from 0.005 to
0.2 Gy, depending on cell type), the range of the challenging doses (0.4 to 8
Gy, depending on cell type), the dose-rate, cell cycle stage at the moment of
the conditioning, and the time between the priming and challenge (K. J.
Sorensen et al. 2002; Stoilov et al. 2007). In most of the studies done in
lymphocyte radio-adaptive response both priming and challenging irradiations
were performed on PHA-stimulated cells. Mainly because the first attempts of
inducing this type of response in G0 cells were unsuccessful (Jeffery D Shadley
et al. 1987; J D Shadley 1994). This is important since most circulating
lymphocytes are in this stage of the cell cycle. Since then, a few publications
were able to show that this response could also be observed in resting
lymphocytes (Cramers et al. 2005). Moreover, there is some reason to believe
that this protective effect of a low priming exposure can take place in vivo (J
F Barquinero et al. 1995; Mohammadi et al. 2006).
These fractioned irradiation conditions have shown to decrease chromosomal
aberrations in lymphocytes. However, there is limited information regarding
the signalling of the DNA DSB that precedes this lower frequency of
chromosomal aberrations.
Low doses, as those generally used for priming have shown produce gamma-
H2AX signaled DSB that remain unrepaid for longer periods of time than
higher doses (Rothkamm & Markus Löbrich 2003).
Histone H2AX stabilizes broken DNA strands to suppress chromosome breaks and translocations during V(D)J recombination (B. Yin et al. 2009), so it is possible that gamma-H2AX signalling has changed in a radio adapted cell, supposed to have less translocations. In this thesis, this radio-adaptive response will be studied with a particular focus on CD4-positive lymphocytes since these cells seem to have higher post-irradiation gamma-H2AX signalling and genetic modulation. In our laboratory, Gruel et al. found that 3 h after whole blood exposure to doses as low as 0.05 Gy, CD4-positive lymphocytes have several down-modulated genes - 10 times more for than for all other subsets (Gruel et al. 2008). Many of these genes were found to have a modified expression during the radio-adaptive response to a challenge dose of 2 Gy after a priming dose of 0.05 Gy in lymphoblastoid cells (Coleman et al. 2005).

1.6 Cell model used in this work: lymphocytes

Lymphocytes are a vital part of the humoral immune system. They can be divided into different classes or subsets including T-lymphocytes and B-lymphocytes. T-lymphocytes are involved in fighting intracellular infections, cancer cells, and foreign tissue. B-lymphocytes are involved in the immune response against bacterial and viral infections. These lymphocyte subsets can be discerned by the antigenic properties of cell surface (membrane) markers. **B-lymphocytes are CD19-positive** and **T-lymphocytes are CD3-positive**. T-cells can be divided further: **helper cells (CD4-positive)** and **cytotoxic cells (CD8-positive)**.

As a reference, according to website of the Clinical Laboratory of the University of Arkansas for Medical Sciences ([http://www.uams.edu/clinlab/flow.htm](http://www.uams.edu/clinlab/flow.htm)), the lymphocyte subset fractions expected in a human peripheral blood sample are as follows: 8-16% of CD19-positive cells, 64-82% of CD3-positive cells, 39-57% of CD4-positive cells and 17-31% of CD8-positive cells.
In a clinical context, peripheral blood lymphocytes (PBL) are a very attractive test material. A blood sample containing considerable amounts of these cells can be easily obtained in a standardized manner, with little medical intervention or inconvenience for the donor/patient. As such, experiments on PBL dominate the literature in comparison to any other normal tissues in assessing DNA damage.

Most predictive assay research on normal tissues to date has been based on the radiosensitivity of normal lymphocytes and skin fibroblasts. Using survival as an endpoint, radiosensitivity of lymphocytes and fibroblasts from the same individual do not correlate (Green et al. 1991). In general, the analysis of lymphocytes appears to be more promising than studies using fibroblasts (Markus Löbrich & Kiefer 2006).

Because radiosensitivity is dependent on cell cycle, for fibroblasts, this type of studies has to be performed in non-dividing (confluent) cultures (Dikomey & Brammer 2000). Lymphocytes constitute a sample of already non-proliferating normal tissue cells within the body. Because they are in the G0 phase, variations in radiosensitivity as cells move though the cell cycle should be avoided. This is an important advantage for the evaluation of gamma-H2AX focus formation since it can occur not only in cells in which DSB have been induced, but also in those undergoing DNA synthesis and mitosis (Furuta et al. 2003) or apoptotic DNA fragmentation (Emmy P Rogakou et al. 2000).

Resting lymphocytes are a viable model to study radiosensitivity since they belong to a unique population of cells that can be killed by low irradiation doses in the absence of mitosis (Woods & Lowenthal 1984).

However, different lymphocyte subsets have different radiosensitivities, which constitutes a possible source of variability. Using cell survival as the endpoint, B-lymphocytes are more radiosensitive than T-lymphocytes, as their numbers decline more rapidly even after relatively low doses (Prosser 1976). However, the T lymphocytes are a heterogeneous group made out of different subsets. CD8-positive T lymphocytes are twice as radiosensitive as CD4-positive T-lymphocytes. And B-cells have actually an intermediate radiosensitivity in terms of survival (Stewart et al. 1988). This highlights the
importance of discriminating the response to radiation of different subsets to decrease the heterogeneity of the results.

In this thesis, we discriminated subset response to determine if one of them had a particularly high response to low doses or better correlation with clinical radiosensitivity.
2 OBJECTIVES OF THIS WORK

Although much is known about the quantitative effects of exposure to ionising radiation, considerable uncertainties and divergent views remain about the health effects at low doses. The importance of low dose risk research is now recognised globally. It is accepted that there is much uncertainty on the shape of the dose-response for cancer derived from epidemiological studies below doses of 0.1 Gy and on the cellular mechanisms that determine the response, including the potential roles of non-targeted processes and of radiation sensitivity. Therefore, many questions about the effects of low-dose exposure to radiation remain unanswered and, in this context, a tool which has to meet numerous requirements is hardly needed: (i) able to detect DSB - highly misrepair-prone cellular damages, (ii) very sensitive with very low detection threshold, (iii) able to underline non-targeted effects such as bystander or radio-adaption, (iv) able to underline inter-individual radiosensitivity.

Recently, gamma-H2AX foci have been used as a biomarker for radiation-induced effects. When the gamma-H2AX foci which mark the DSB are stained, individual foci are detectible, making the assay suitable for studies requiring great sensitivity. Notably, this biomarker has been used in cell lines to detect doses of ionizing radiation as low as 0.001 Gy. The gamma-H2AX assay is also able to take into account the radiation sensitivity not available with physical modelling. Actually, it has been shown that lymphocytes from patients with clinical radiosensitivity show unusual residual gamma-H2AX foci after low-dose medical exposure to IR. This latter effect could allow determining the radiosensitivity of individuals but is however rather low and is the result of a mix of the different lymphocyte subtype responses. By isolating the most sensitive lymphocyte subtype, this would expand a specific lymphocyte response; the gamma-H2AX assay could therefore be a very sensitive marker of radiation sensitivity.

Originally, the aim of this work was to develop an in vitro assay able to evaluate individual radiation sensitivity in human peripheral lymphocytes by using gamma-H2AX quantification. This assay needed to be both sensitive and fast enough to be used on a large scale.
In the interest of sensitivity gamma-H2AX response was measured by scoring individual foci in microscope images instead of quantifying the general cellular response by a flow cytometry approach. Furthermore, in order to isolate a potentially specific low-dose effect, the gamma-H2AX response was analysed not only by considering the general lymphocyte population but also by discriminating specific lymphocyte subsets. To allow future clinical applications of this microscopy approach, the image acquisition and focus scoring steps were automated. This not only increases the speed of the results but also insures that there is no operator bias during focus scoring.

To reach our objective, this gamma-H2AX quantification assay had to be tested in lymphocytes of a large cohort of individuals with characterized clinical radiosensitivity. In mid 2008 we worked on samples from Belgian radiotherapy patients of known clinical acute radiosensitivity. We felt that a whole blood context was important to study doses known to induce non-targeted responses. However, when working with whole blood instead of isolated lymphocytes we were faced with complications due to sample aging before processing. To establish a sensitive and fast gamma-H2AX protocol to study low-dose responses in whole blood, changes were made to guarantee the viability of the blood sample. The improved protocol was validated by comparison with manual and semi-automated focus scoring. Furthermore, the gamma-H2AX response of different lymphocyte subsets was characterized after low doses of gamma-irradiation and was compared to the response observed in the general lymphocyte population.

In view of the difficulties found in obtaining additional blood samples from radiotherapy donors in France for a more complete study, the initial work plan was reoriented. Consequently we focused in lymphocyte gamma-H2AX signalling in another context of different radiosensitivity: radio-adaptive response. This adaptive response can be seen as a decrease in radiosensitivity due to a pre-exposure to a low dose. Therefore, instead of discriminating clinical radiosensitivity, we would determine if changes in cellular radiosensitivity seen using other endpoints such as chromosome aberrations and apoptosis were detectable by our gamma-H2AX assay. Given that we observed the highest radiation-induced
gamma-H2AX signalling in CD4-positive lymphocytes, we focused on this subset to study the radio-adaptive response. Our aim was to determine the potential of CD4-positive lymphocytes in distinguishing radiosensitivity through the gamma-H2AX assay.
3 MATERIALS AND METHODS

3.1 Sample Collection

For the individual radiosensitivity study, the blood samples were collected from 27 individuals treated for histologically confirmed head and neck squamous cell carcinoma (HNSCC) at the Ghent University Hospital. The blood samples were collected in heparin during the periodical medical examination in the follow-up of the treatment and were sent to Fontenay-aux-Roses (France) to be processed 24 to 48 h later. This study was approved by the Ethics Committee of Ghent University Hospital.

For the other series of experiments, the blood was collected in citrate from healthy volunteers by the "Etablissement Français du Sang" (EFS) which obtained written informed consent for all donors, according to the agreement between EFS and IRSN (reference n° 10/EFS/056). EFS provided to IRSN the anonymous blood samples used for this study in accordance with French law (L. 2004-800) on bioethics. Transportation from the medical centre to the lab was done at ambient temperature and did not exceed one hour.

3.2 Irradiation Conditions

For the individual radiosensitivity study (Ghent), the whole blood samples were irradiated without dilution upon reception.

For all other experiments, the blood was diluted (1:1) with Roswell Park Institute medium (RPMI) with L-glutamine, unless stated otherwise. The blood used to characterize and validate our protocol was sampled close to our lab. The blood was routinely diluted 1:1 with RPMI and L-glutamine shortly after sampling.
The samples were irradiated in Falcon tubes for samples with a volume of 50 mL or below. Samples with a volume above 50 mL were irradiated in culture flasks. All irradiations were performed at room temperature, at the IRSN site of Fontenay-aux-Roses. Two gamma sources were used in this work: cesium-137 and cobalt-60.

3.2.1 Irradiations with a cesium-137 gamma source

The irradiator (IBL637, CisBio, France) can be easily handled by any operator formed in radioprotection. However it has the inconvenient of being more limited in the dose-rates range. For the Ghent experiments and the samples used to validate automated scoring, the cesium-137 source at a dose rate of 0.5 Gy per min was used. Doses ranged from 0.05 Gy up to 0.5 Gy.

3.2.2 Irradiations with a cobalt-60 gamma source

The following experiments were performed with a Cobalt-60 irradiator (ICO4000) with a dose-rate of 0.05 Gy per min. This alternative offered a more precise estimation of the delivered dose, particularly important for procedures involving doses as low as 0.005 Gy. For the adaptive response studies the Cobalt-60 source was also used, but with a dose rate of 0.5 Gy per min to avoid 40-min exposures (for 2 Gy and 2.05 Gy).

3.3 Gamma-H2AX assay

3.3.1 Sample processing

After irradiation the blood was incubated at 37°C for a period of time of 30 min, 3 h, 5 h or 24 h. The blood samples were immerged in ice immediately after incubation to reduce gamma-H2AX signalling variations and DNA repair. The cold blood was carefully layered onto cold Ficoll-Histopaque and
centrifuged at 1000 g for 20 min at 0°C. The lymphocytes from the interphase were transferred and washed three times with cold phosphate buffered saline (PBS) at 400 g for 8 min at 0°C. The lymphocytes were then resuspended in PBS and spotted in DakoPen® (Dako, Glostrup, Denmark) circles of a fixed diameter onto Superfrost® (Menzel-Glaser, Braunschweig, Germany) slides for adhesion at 0°C. After an adhesion period of 30 min the cells were immediately fixed in 1% PFA for 10 min at room temperature and washed thoroughly in PBS.

3.3.2 Slide preservation

3.3.2.1 Samples from Ghent University
For the Ghent experiments, many samples arrived at a time when we were experiencing batch quality issues with the anti-gamma-H2AX antibody. Therefore, the adherent lymphocytes had to be dried and stored at -20°C for long-term preservation (up to 3 months) before staining. After gamma-H2AX staining, only 9 samples produced images acceptable for analysis, two of which were CTC3 and the 7 others were CTC1 and 2.

3.3.2.2 Samples used in all remaining experiments
The adherent lymphocytes were preserved in PBS at 4°C for 1 to 7 days before staining.

3.3.3 Immunofluorescence staining

Adherent cells were permeabilized in PBS + 0.1% Triton X-100 for 10 min. After two short PBS washes, the cells were blocked in PBS with 20% goat serum for 30 min at room temperature. Slides were incubated with monoclonal anti-H2AX Phosphorylated (Ser139) antibody (1:500 dilution, BioLegend, San Diego, CA, USA) for 60 min, washed in PBS for 5 min, and incubated with Texas Red X conjugated goat anti-mouse secondary antibody (1:1500) for 60 min, all at room temperature.
For CD3-specific (cluster of differentiation 3) staining the slides were incubated with AlexaFluor® 647-coupled anti-human CD3 (1:100 dilution) (BD Pharmingen, Franklin Lakes, NJ, USA) for 30 min. For other CD-specific staining (CD4, CD8 and CD19), FITC-coupled and Cy5-coupled antibodies were used, from the same manufacturer and at the same dilution.

Finally, the slides were washed in PBS for 10 min and mounted with cover slips; DAPI Prolong® Gold solution was used according to the manufacturer's instructions.

All the experiments that have been quantitatively compared were done using reagents from the same batches to minimise inter-experiment variations.

### 3.3.4 Image Acquisition and Processing

Slides were viewed with an epifluorescence microscope (Imager.Z1, Carl Zeiss, Oberkochen, Germany) equipped with a non-cooled CCD (charge-coupled device) camera and an external fluorescence light source (EL6000, Leica, Solms, Germany). We chose to use an alignment-free light source to minimise illumination calibration issues. The automated image acquisition used Metafer 4 software (version 3.6.0, from MetaSystemsTM, Altlussheim, Germany), which controlled the motorized microscope slide plate, filters, and digital camera. The different cell spot coordinates were noted and, for each spot, a grid of non-overlapping fields was selected for acquisition with the basic function "compute coordinates". Depending on the average cell density, 100 to 250 fields of each spot were selected and acquired by the Metafer Autocapt module, using an immersion plan Apochromat oil 63x objective (Carl Zeiss). To compile all of the 3-dimensionally distributed gamma-H2AX foci throughout the lymphocyte nuclei in one image, the Autocapt classifier was programmed to acquire 26 2D images for each field, with a 0.3 µm z-axis step between two slices.

To avoid loss of information, the images were not overexposed. Acquisition programs like Autocapt prevent overexposure when capturing images with an automatic exposure time. However, to perform rigorous comparisons, the
integration times chosen by the program were fixed and kept constant throughout the acquisition of all doses, including 0.5 Gy. The resulting fields of view (FOV) were transformed into training images (TRN) with the “Create TRN from FOV” command to allow each colour channel to be exported as an individual greyscale tiff file. All the steps described above from sample collection to image acquisition were shared by all the subsequent methods of analysis used.

### 3.3.5 Image Analysis

Both manual and automatic focus scoring were performed in uncompressed high-quality images. The manual scoring was done by 3 operators that were unaware of the doses corresponding to the images they were analysing. Each person counted a minimum of 150 foci or 250 cells per condition. The operators were told to consider every focus that they could discriminate from the background regardless of their intensity. For the automatic detection, two different programs were used: the commercial software HistoLab™ (version 7.5.2, Microvision Instruments, Evry, France) and the free cell image analysis software, CellProfiler (version 2.0, Broad Institute, MA, USA). The specific settings used for both HistoLab™ and CellProfiler are supplied in the annexed material. All image analysis parameters were kept constant throughout the duration of this study. This protocol was used in the published article of Valente and colleagues (M. Valente et al. 2011)

### 3.4 FISH-3 assay

#### 3.4.1 Sample processing

CD4-positive lymphocytes were separated from the whole blood during recovery by density gradient using the RosetteSep Human CD4-positive cell Enrichment Cocktail (StemCell Technologies, Grenoble, France).
Dynabeads® Human T-Activator CD3/CD28 for cell expansion and activation (Invitrogen) were used to stimulate the isolated CD4-positive lymphocytes during a 72-hour incubation at 37°C. Colcemid (KaryoMAX, Invitrogen, France) was added to a final concentration of 0.1 μg/mL and the cultures were continued for 2 h. After harvesting, the cells were treated by hypotonic shock (0.075 M KCl, Sigma, USA) for 8 min at 37°C and were fixed three times in 10 mL methanol:acetic acid mixture (3:1, v/v). Cells were dropped onto clean slides, air-dried and kept at -20°C until preparation of slides for analysis by FISH methods.

Simultaneously, 500 μL of whole blood from the same donor was stimulated with phytohemagglutinin (PHA, Invitrogen, France) and incubated for 46 h.

3.4.2 FISH-3 painting

FISH-3 was performed on freshly spread slides containing metaphases and incubated overnight at 37°C. Slides were pretreated with 0.05% pepsin (Roche, Switzerland) for 10 min at 37°C and subsequently washed in PBS (Invitrogen, France) for 5 min. They were incubated in a post-fixation solution, 1% formaldehyde (VWR, France) and 50 mM MgCl2 (Sigma, USA) in 1X PBS for 10 min at room temperature, and then washed in PBS for 5 min. After dehydration (70, 90, 100% ethanol for 2 min each), the slides were denatured for 2 min in 70% formamide (VWR, France) at 65°C, dehydrated (70, 90, 100% ethanol for 2 min each), and air-dried. Commercial whole chromosome probes for chromosome 2 (mix FITC:Cy3, Cambio, UK), chromosome 4 (Cy3, Cambio, UK), and chromosome 12 (FITC, Cambio, UK) were denatured for 10 min at 65°C and then incubated for 30 min at 37°C. Slides were hybridized overnight at 37°C before washing in 50% formamide at 45°C for 5 min and three times in 4X SSC plus 0.05% Tween at 45°C for 4 min. Chromosomes were counterstained using DAPI/Antifade solution (Qbiogene, USA).
Fluorescent images were captured with a Zeiss Axioplan epifluorescent microscope connected to the ISIS/M-FISH imaging system (MetaSystems, Germany).

3.4.3 Chromosome aberration scoring

Normal human metaphases have 46 chromosomes. To speed up the scoring process, the exact number of chromosomes was not checked. Therefore, any metaphase containing approximately 46 chromosomes was scored. Cells with dicentrics, rings or fragments were included in the scoring but only aberrations containing painted chromosomes were considered. We considered total translocations as the sum of apparently simple two-way and one-way translocations. Metaphases suspected to be a clone were eliminated, to avoid clonal cell bias. For CD4-positive lymphocytes from one individual we were able to score: 396 metaphases for the 0 Gy, 1290 for 0.05+2Gy and 1512 for the 2.05Gy condition. Protocol based on the work of Pouzoulet and colleagues (Pouzoulet et al. 2007).

3.5 Apoptosis assay

The apoptosis assay started with a red blood cell lysis: for each condition, 4 mL of RPMI-diluted blood was diluted in 40mL of lysis buffer (\(\text{NH}_4\text{Cl 8.32g/L, NaHCO}_3 1g/L, \text{Na}_4\text{EDTA 0.037g/L}\)) and incubated in ice for 30 min. All the manipulations that followed were performed at 0°C unless stated otherwise. The samples were centrifuged for 8 min at 350g and the resulting pellet was resuspended in 10 mL of a PBS/1%BSA (v/m) solution and transferred to a FACS tube. The cells were then washed three times by repeating the previous step. After the final centrifugation, the pellet of each condition was resuspended in 350µL of the PBS/1%BSA (v/m) solution. lGg1 FITC was used as a staining negative control in a separate tube at a concentration of 3:20. In the test tubes, FITC-coupled anti-human CD4 (1:20 dilution) (BD Pharmingen) was added to the cells for a 10-min incubation at
room temperature and in the dark. The cells were washed with 2ml de PBS per tube and concentrated with a 8-min centrifugation at 350g. The supernatant was discarded and the pallet was diluted in 100µL of annexin buffer per tube (10mM HEPES, 140mM NaCl, 2.5mM CaCl₂, pH 7.4). 5 µL of annexine V was added to each tube to incubate for 15 min at room temperature and in the dark. After adding 400µL of annexine V buffer + PI (1µg/ml) the tubes were kept on ice, awaiting flow cytometry analysis. The cells were analyzed using a FACSort (Becton Dickinson).
4 RESULTS

Because our work focused on slight radiation response differences (low-dose, cell- or individual-specific) the sensitivity of the test was paramount. Therefore, we have chosen to quantify gamma-H2AX response using microscopy based focus scoring rather than a flow cytometry approach (Figure 9).

![Image of DAPI and gamma-H2AX staining](image)

Figure 9. Typical images of DAPI and gamma-H2AX staining for a 30 min post-exposure time. All the images (both for sham-irradiated and exposed samples) were taken with the same integration time.

4.1 Using cellular DNA damage signalling as a measure of clinical radiosensitivity

Extreme clinical radiosensitivity has been observed in radiotherapy patients with ineffective cellular response to DNA damage due to hereditary mutations in DDR protein genes. To examine whether and how individual clinical
Radiosensitivity is related to cellular DNA damage response. We irradiated ex vivo samples from individuals of known clinical radiosensitivity. The intensity and kinetics of the resulting cellular DNA damage signaling was determined using gamma-H2AX focus scoring to verify if this response varied with clinical radiosensitivity.

In mid 2008 we worked on samples from Belgian radiotherapy patients of known clinical acute radiosensitivity. The clinical reactions were determined in respect to acute dysphagia, dermatitis and mucositis using the CTCAE scale. In order of increasing clinical radiosensitivity: 8 patients were CTC1, 10 were CTC2 and 9 were CTC3. Blood from the 27 individuals was sampled at Ghent and treated (irradiated then incubated at 37°C) 24 to 48 h later at Fontenay-aux-Roses, France. Gamma-H2AX foci were scored at different times post-irradiation to determine if this DSB signalling peaked (30 min) or evolved during repair (180 and 300 min) differently between the CTC groups. Samples from the same donors had been used by a Gent University lab (Werbrouck et al. 2011). After irradiating isolated T-lymphocytes, Werbrouck et al. found no difference in intensity or kinetics of gamma-H2AX signaling. Based in the previous observations of Löbrich et al. (Markus Löbrich et al. 2005) it was hypothesized that irradiating whole blood could reveal responses to radiation that are not observed when irradiating isolated T-lymphocytes. Therefore, we repeated their experiments but irradiated whole blood, to come closer to the in vivo cellular context.

After gamma-H2AX staining, only 9 samples produced images acceptable for analysis, two of which were CTC3 and the 7 others were CTC1 and 2.

4.1.1 Gamma-H2AX signalling kinetics of individuals with different clinical responses to radiation

The basal levels of gamma-H2AX in sham-irradiated samples are similar whatever the clinical radiosensitivity or time of analysis: a mean of 0.45 foci per cell (between 0.37 and 0.51) (Figure 10). Furthermore, there is no clear difference between the CTC1/2 group (low and mild clinical acute radiosensitivity) and the CT3 group (higher clinical radiosensitivity) in terms
of radiation-induced gamma-H2AX: for both groups 0.5 Gy induced an average of 3.3 foci per cell (30 min). This value decreased to 2.6 at 180 min and 1.9 at 300 min for both radiosensitivity groups, indicating similar repair kinetics. Unfortunately, with such a reduced number of individuals in each group, it was not possible to perform a statistical analysis.

Figure 10. Gamma-H2AX signalling kinetics of individuals with different clinical radiosensitivities. Blood from radiotherapy patients was sampled at Ghent (Belgium) and irradiated one day later at Fontenay-aux-Roses (France). After exposure, the samples incubated at 37°C for 30, 180, 300 min, and 24 h before being processed. Only 9 samples produced a sufficient number of well preserved cells for analysis of at least one condition. Error bars represent the standard deviation. Each condition is represented by 1 to 3 different individuals. (foci scored automatically, using CellProfiler)

4.2 Assessing the impact of the changes made to the protocol

After a one day trip between Ghent and our laboratory most of the blood samples already presented some degree of hemolysis visible during the density gradient separation within and below the lymphocyte ring (an example of this can be seen in Figure 11). After being subjected to more 24 h incubation at 37°C, this degradation was even more frequent, with a negative impact on the yield and quality of lymphocyte recovery by density gradient. In fact, we never successfully isolated lymphocytes after 24 h of incubation and for 5 tubes no lymphocytes could be retrieved for any incubation time. For
the remaining 22 samples, cells adhered badly to the microscopy slide and/or were contaminated by other white blood cells (mainly granulocytes and monocytes). The loss of quality experienced with the samples from Ghent pointed out several critical issues in our protocol that needed improving, like blood sample processing and slide storage. We took several steps to overcome these limitations and we evaluated the impact of these changes.

4.2.1 Improving lymphocyte recovery of 24-hours old blood with RPMI medium dilution

To prevent hemolysis processes we diluted a blood sample with RPMI with L-glutamine and processed it 24 h later, to compare with an undiluted sample from the same donor. Both diluted and undiluted samples waited processing

![Image](image-url)
at room temperature in order to mimic the conditions of the Ghent experiments.

After a 30 min incubation at 37°C, the undiluted samples presented more erythrocyte lysis than the diluted one: during gradient separation the lymphocyte ring of the undiluted blood was already polluted with red agglomerates of other cell types. When the samples were subjected to additional 24 h incubation at 37°C, hemolysis was apparent in both conditions but only the diluted sample had a recoverable lymphocyte ring (Figure 11). Therefore, we were able to reproduce the sample degradation encountered with the Ghent samples. RPMI dilution seems to delay hemolysis enough to allow us to recover lymphocytes at later time points for kinetics studies.

4.2.1.1 Impact of RPMI medium dilution on H2AX response

We had chosen to irradiate whole blood ex vivo to keep the physiological cellular environment and, theoretically, the response to radiation. In order to maintain this whole blood environment for as long as possible, the 24 h old blood was diluted 1:1 in RPMI medium.

To assess the impact of sample age and dilution on lymphocyte gamma-H2AX response, freshly sampled blood was irradiated and compared to the same blood irradiated 24 h later diluted or not. Foci were scored 180 min after irradiation at 0.5 Gy (Figure 12).

One-way ANOVA (analysis of variance) on ranks was used to compare foci yield results from samples after irradiation and showed statistically significant differences (p < 0.001) between fresh blood samples and samples that were 24 h old (diluted or not). On the other hand, for these older samples, the same test showed no difference (p > 0.05) between the diluted and undiluted conditions (Figure 12).
Figure 12. Impact of RPMI medium dilution on H2AX response. Blood from the same individual was treated differently after sampling. The “undiluted fresh” sample was irradiated and processed the same day as the sampling. The other samples were irradiated and processed 24 h after sampling, one of which had been diluted right after sampling. The two conditions that were irradiated and processed 24 h after sampling were indistinguishable. However, both had a statistically significant higher number of foci than the blood exposed shortly after sampling (over 440 cells scored for each condition). Foci scored with CellProfiler. Error bars represent the standard deviation (SD) considering the distribution of focus in the cells. Results from one experiment.

4.2.2 Slide preservation

It was observed from the Ghent experiments that slide freezing/thawing between the fixation and staining steps lead to damaged cells. Preservation at -20°C of fixed slides was then abandoned in favour of an immersion in PBS at 4°C for up to one week. To test this conservation change in the protocol, we proceeded to determine if staining 1 day and 7 days after processing produced comparable results. Gamma-H2AX staining performed one week apart of a sample processed 30 min after irradiation at 0.5 Gy produced statistically indistinguishable scores (t-test, n>1600 cells, p>0.5) (Figure 13).
Figure 13. Impact of storing slides in PBS at 4°C on the gamma-H2AX focus score. A blood sample from one individual was irradiated at 0.5 Gy and processed as described in the materials and methods section. Slides from the experiment were stored at 4°C in PBS after fixation and stained one week apart (7 days). The foci scores obtained were very similar between stains for both the control (0 Gy) and the irradiated (0.5 Gy) conditions. Over 1600 cells were scored for each of these 4 conditions. Experiment performed once. Foci scored using CellProfiler. Error bars represent SD.

4.2.3 Protocol speed

With the changes in sample dilution and slide preservation described above we obtained an average lymphocyte density of 25 cells per field (up to 80) for an acquisition at a magnification of x63. Currently, the time required for the process between blood sample and mounted slide is 5 h at most.

4.3 Development of automated acquisition and analysis

To increase speed and facilitate the potential clinical application of our protocol we used automated acquisition and automated analysis for cell type selection and focus scoring. A free-access program was used for automated scoring. It was compared to manual scoring and a program already in use at the lab to determine if it could be adopted as a sensible and fast alternative. Using the Metafer acquisition program the average speed of acquisition and export was 2.5 images per min. The good quality of the images allowed us to use automated focus scoring to avoid bias. However, before applying an
unsupervised image analysis to all experiments (via the freeware program CellProfiler), we validated this approach by comparison with manual and a semi-automatic scoring system already used in our lab (HistoLab™). Both programs presented similar nuclei and focus selection after the thresholds were adapted to resemble manual selection (Figure 14).

![Figure 14. Comparison of manual and automatic foci detection for the condition 30 min after irradiation at 0.5 Gy. The same images were used for the different scoring approaches. In manual scoring the operator selected every focus distinguishable from the background; with HistoLab™ we applied a TopHat threshold; and for CellProfiler the foci were detected by an Otsu algorithm (Otsu 1979; Hou et al. 2009).](image)

4.3.1 Cell type selection

If different cell types were present in the images acquired it was very important to insure a good selection of the cells of interest to perform a reliable quantitative analysis. To determine the characteristics of the cells of interest is therefore crucial. In this work, two approaches were used to select cells of interest: nucleus morphology and CD-specific membrane staining. The first one was used to distinguish the general lymphocyte population from the other cell types or the damaged cells without additional hybridization. The second one used CD-specific membrane staining for experiments that required a deeper discrimination of lymphocyte subsets (CD4, CD8 and CD19) and was also used to validate the morphological selection (CD3).
4.3.1.1 The use of nucleus morphology to discriminate lymphocytes

Because lymphocytes in general tend to have a round shape and brighter DAPI staining, their morphology allows operators to score them manually. In the current version of HistoLab™, the user has also the possibility of removing cells that have aberrant staining or morphology. Because the CellProfiler approach has no operator intervention, the cell type selection was done graphically during data analysis by the use of the nuclei “form factor” and “DAPI integrated intensity” measures. An example can be found in Figure 15 A: the two cells at the middle would not have been considered lymphocytes by the operator during manual scoring because they are not as round and have weaker DAPI staining. By quantifying these two parameters, it is possible to establish thresholds that insure that automatic scoring will also exclude them. “Form factor” is a parameter that quantifies “roundness” in the CellProfiler "Measure Object Area" module. Cells for which this parameter is closer to “1” are rounder and thus more likely to be lymphocytes. The comet-shaped dot cloud (Figure 15 B) obtained reflects the general quality of the slide: the presence of burst cells or granulocytes increases the comet tail and forms a cloud of less intense nuclei underneath the “comet”. The example presented in Figure 15 B is the extreme case of a slide considerably polluted by cells different from lymphocytes.
Figure 15. Indicators used for human lymphocyte enrichment before gamma-H2AX analysis. A - Cell DAPI intensity and "roundness" (form factor) measurements performed by CellProfiler. B - Validation of the morphological selection of lymphocytes with CD3 staining. Within the comet (i.e. excluding cells with DAPI intensity below 1000), different groups of cells were selected by choosing different form factor thresholds: 0.7, 0.8, and 0.86.
4.3.1.2 The use of membrane staining to validate morphologic selection and discriminate lymphocyte subsets

Most circulating lymphocytes have CD3-positive membrane. Therefore, we used CD3 staining to validate the morphology selection. Within the comet-shaped cloud, several form factor thresholds were tested, each stricter than the previous one to include ever rounder cells (Figure 15). With form factor threshold values closer to "1", the percentage of CD3-positive cells increased. To maximize the lymphocyte concentration of the cell population being studied, we analyzed only cells within the comet with a form factor above 0.86. This limit was selected after observing the general form factor results of several nuclei images. The selection based on form and intensity measurements presented here successfully enriched in lymphocytes the cell population studied. For the example of Figure 15, were this selection is more critical, this threshold increased the fraction of CD3-positive cells from 59% to 83%. However, with our current protocol most of the slides are very rich in analyzable lymphocytes with a shorter tail of the comet cloud. As observed in Figure 16, more than 50% of the cells have a form factor higher than 0.86 and there is no second cloud of cells of a less intense DAPI staining forming below. In slides rich in well-preserved lymphocytes applying the 0.86 threshold increases the lymphocytes considered to be CD3-positive of approximately 10%.

It is also important to mention that irradiating samples does not seem to affect this selection (Figure 16).
For several experiments the discrimination of a lymphocyte subset of interest (CD4, CD8 or CD19) was also obtained using a CD-specific staining in addition to the anti-gamma-H2AX (Figure 17). On the general lymphocyte population recovered by density gradient we found an average of 33% CD4-positive cells, 13% CD8-positive cells and 6% CD19-positive cells.

Figure 16. Effect of irradiation on morphology-based enrichment of analyzed cell population in CD3-positive lymphocytes. Above, visually, the dot cloud does not seem to change shape from the control to the irradiated condition. The histograms beneath show the percentage of CD3-positive lymphocytes before and after applying the form factor threshold. The enrichment in CD3-positive lymphocytes after applying the form factor threshold is of the same order of magnitude for both the irradiated and non-irradiated conditions. Data represented here corresponds to a 30 min post-exposure incubation time.
Figure 17. Staining gamma-H2AX foci within different lymphocyte subsets. Several Fluorochrome coupled antibodies for membrane and gamma-H2AX can be used simultaneously for more specific studies. In this example, gamma-H2AX foci are colored in magenta the membranes of CD8 lymphocytes are colored in red and CD19 lymphocytes are colored in green. In the right, a dot-plot was made in Microsoft Excel™ with the intensity measurements exported from a CellProfiler analysis to aid the operator in the selection of the thresholds.

4.3.2 Gamma-H2AX focus scoring

To validate the actual focus scoring inside the cells of interest, we analysed the same images by three different approaches: manually, with HistoLab™ and using CellProfiler. Previous experiments in the lab have shown that the kinetics of the gamma-H2AX foci score peaked around 30 min after a 0.5 Gy irradiation (Sandrine Roch-Lefèvre et al. 2010). Therefore, the dose response for three healthy individuals (the three "experiments" mentioned below) was established at this time with samples exposed at 0 to 0.5 Gy.

4.3.2.1 Scoring criteria: inclusion of foci of all intensities

The three operators who scored manually were told to consider any focus distinguishable from the background and both HistoLab™ and CellProfiler were programmed to do the same. Whatever the scoring method used (manual or automatic) it was observed that to score only the brightest gamma-H2AX foci leads to a decrease in the linearity of the dose-effect relation (Figure 18). The condition "All foci" corresponds to the threshold adopted in this work.
(threshold correction factor 3) and gives results closest to manual scoring. The condition “Only bright foci” was obtained by increasing the “threshold correction factor” to 5. This increased restriction makes CellProfiler discard dimmer foci. The resulting focus underestimation decreases the linearity of the relation between dose and number of foci. This phenomenon may be due to the fact that the gamma-H2AX staining intensity per focus tends to decrease with radiation dose. The maximum gamma-H2AX intensity per focus was significantly different between the doses 0.1 and 0.5 Gy (Mann-Whitney Test, p < 0.001).

Figure 18. Loss of linearity if only the brightest foci are scored. A - The same images from a blood sample of one human donor 30 min after exposure to 5 doses (0 Gy, 0.05 Gy, 0.1 Gy, 0.2 Gy, or 0.5 Gy) were automatically scored by CellProfiler using different threshold values. B - Distribution of gamma-H2AX foci intensity for different irradiation doses. C - The same images manually scored by a human operator twice: once considering only the brightest foci and a second time taking into account all foci. In manual scoring considering the brightest foci alone decreases the linearity of the relation between dose and number of foci. The box plot shows the minimum and maximum values, 25th and 75th percentile and median.
4.3.2.2 Correlation with manual scoring

All three scoring approaches produced very similar dose-effect correlations (Figure 19), with a linear increase with dose. After manual scoring, each operator found a linear increase of gamma-H2AX foci with radiation dose (Pearson’s test: $r^2 > 0.991$, $p < 0.0004$). The scores obtained with HistoLab™ for the three experiments revealed as well a significant linear relation between dose and foci score (Pearson’s test $r^2 > 0.981$, $p < 0.002$). Using CellProfiler, we also obtained a significant linear increase of gamma-H2AX foci number with radiation dose (Pearson’s test $r^2 > 0.971$, $p < 0.003$). Consequently, there is a significant correlation between the dose-effect curve obtained by a human operator and those resulting from both programs (Person’s test, $p< 0.001$ for either program). Nevertheless, at 0.5 Gy both programs tend to consider a lower number of foci per cell than human operators (Figure 19).

**Figure 19.** Comparison of three approaches to focus scoring: manual, semi-automatic and automatic. The same images from blood samples of 3 human donors 30 min after exposure to 5 doses (0 Gy, 0.05 Gy, 0.1 Gy, 0.2 Gy, or 0.5 Gy) were manually scored by an operator, and automatically scored by two programs (HistoLab™ and CellProfiler). A minimum of 100 foci and/or 250 lymphocytes were counted for each condition regardless of the scoring method. The number of foci increased linearly with the dose. The equations for the linear fits are: $y = 12.515x + 0.649$ for Human operator 1, $y = 11.61x + 0.569$ for Histolab and $y = 11.236x + 0.476$ for CellProfiler. Error bars represent the SD for the 3 blood donors.
4.3.2.3 Variability

The inter-operator scoring variations were estimated using images from the same experiment that were analysed manually by three different operators.

![Figure 20. Comparison of three approaches to focus scoring: manual, semi-automatic and automatic. The same images from a blood sample of one human donor 30 min after exposure to 5 doses (0 Gy, 0.05 Gy, 0.1 Gy, 0.2 Gy, or 0.5 Gy) were manually scored by three different operators, and automatically scored by two programs (Histolab and CellProfiler). A minimum of 100 foci and/or 250 lymphocytes were counted for each condition regardless of the scoring method. The number of foci increased linearly with the dose. At 0.5 Gy both programs tend to consider a lower number of foci per cell than human operators.](image)

The inter-operator results were very similar with a coefficient of variation (CV) above 10% only in the sham-irradiated condition. The inter-method scoring variations were estimated using images from the same experiment that were analyzed manually by one of the operator and also automatically by Histolab™ and CellProfiler. The scores from the three methods were very similar as well with a CV of around 10% whatever the dose condition. To estimate inter-experiment variations, images of all three experiments were scored by one of the operators manually but also automatically by Histolab™ and CellProfiler. Whatever the scoring method used, the inter-experiment variations were high for the sham irradiated condition, with a CV of at least 30%. Whereas the 0.5 Gy condition had a much lower CV: below 10% (Figure 21). In general, this variation between experiments is slightly lower for the manual scoring.
4.3.2.4 Speed and operator intervention

With the aim of developing a protocol for clinical applications, to maximize the scoring speed and to reduce the operator intervention time was a constant concern. However, reducing the time of the analysis at the cost of increasing in variability cannot be acceptable. For instance, one parameter that may negatively influence gamma-H2AX scoring variations between experiments is an insufficient number of scored lymphocytes. We therefore determined the minimal number of cells to be scored without inducing sampling biases. To address this issue we tested random partial samples of different sizes (25, 50, 100 and 250 cells) from two large samples of more than 1500 cells each, that we assumed to be representative of their respective cell populations (Figure 22). Scoring 100 cells was sufficient to provide a CV lower than 10% when these cells contained at least 150 foci. However, in a non irradiated sample where foci are rarer, at least 250 cells were required for a CV under 10% (with about 70 foci in all).

Figure 21. Inter-experiment variation according to the scoring method used. The variation in the average focus number between experiments is very elevated for low doses and tends to decrease with the dose. For all scoring methods represented here at least 250 cells were scored per experiment.
After acquiring some experience, the operators took at least 2 seconds to manually score one cell on the acquired images. Therefore, it takes 500 seconds to score 250 cells manually. Using HistoLab™, the main limitation on speed is also the operator because this program requires the presence of an operator to start the analysis of each image individually. However when we used this semi-automated approach the scoring speed increased to 2.5 cells per second. Considering an average of 25 cells per image, scoring 250 cells with HistoLab™ requires 100 seconds of “operator time”. For CellProfiler, the computers’ performance is the only limitation. With the computers we used this approach had a speed of 2.8 cells per second. However, the scoring is done in the absence of the operator, who only needs about 30 seconds to load the parameters file. For a sample of 250 cells, this approach is 16 times faster than manual scoring.

4.4 Dose response

Once the automated scoring with CellProfiler was validated, we characterized the gamma-H2AX low-dose response for different dose and time conditions.
and in lymphocyte subtypes. The gamma-H2AX signalling differences between doses were studied at later points in time, as DNA repair occurs. The response of specific lymphocyte subsets was also observed to access their different sensitivity to these low doses. The aim was to determine if this approach was sensitive enough to distinguishing responses to doses as low as 0.005 Gy from the basal levels of gamma-H2AX focus frequency of non-irradiated samples. Lymphocyte subsets can have different intensities of gamma-H2AX responses. By discriminating their responses to irradiation we increase our chances of finding a cell type more sensitive and specific response to radiation.

4.4.1 General lymphocyte population

Within the dose range tested, the signalling response increased linearly with dose (Pearson’s, $r^2=0.9824$, $p<0.000002$) (Figure 23).

![Graph showing dose response of the general lymphocyte population](image)

**Figure 23.** Dose response of the general lymphocyte population. Blood samples from different individuals were exposed to 8 doses: 0 Gy, 0.005 Gy, 0.01 Gy, 0.05 Gy, 0.1 Gy, 0.2 Gy, 0.5 Gy, and 0.55 Gy. The error bars represent the standard error. Each dose condition was repeated in at least 3 (and up to 7) independent experiments. Conditions only performed once (0.01 Gy and 0.55 Gy) were included when other doses from the same experiment were also represented here. Blood from 8 different individuals was used and each experiment used a sample from a different person.
The trend line obtained from this data predicts approximately 0.82 foci per cell for 0 Gy, which is more than the double of the actual yield (0.32) that we found for this dose. Consequently, with a yield of only 0.84, the lowest dose used (0.005 Gy) was found to be already statistically different from the sham-irradiation (t-test, p< 0.017).

Between 30 min to 1 h after exposure, gamma-H2AX signalling tends to decrease, as DNA damage repair takes place. However, it was observed from the gamma-H2AX signalling kinetics after exposure to different doses that the foci disappearance and/or repair kinetics varied with the dose (Figure 24).

![Figure 24](image)

**Figure 24.** Gamma-H2AX signalling kinetics in human lymphocytes after exposure to different doses. Blood samples from different individuals were exposed to 5 doses: 0 Gy, 0.005 Gy, 0.05 Gy, 0.1 Gy, and 0.5 Gy. The post-irradiation foci were scored after different incubation times: 30 min, 3 h and one day. The error bars represent the standard error and their presence implies that the condition was repeated in at least 2 independent experiments. Conditions only performed once (0.005 Gy, 24 h and 0.1Gy, 3 h) were included when other doses from the same experiment were also represented here. Blood from 8 different individuals was used and each experiment used a sample from a different person.

For the highest dose (0.5 Gy) almost 2 foci per cell signalled at 30 min were no longer present 3 h after exposure. This is the highest decrease in radiation induced foci (taking into account the foci detected at 0 Gy) of the doses we used: nearly 40%. Despite this initially fast decrease, the number of foci per
cell induced by a 0.5 Gy irradiation is still significantly different from the 0 Gy baseline 24 h after exposure.

For the lower doses, the foci yield varied little 3 h after exposure (0.2 foci per cell on average) since the supposed signalling peek at 30 min. Because of this difference in the signalling/repair rate among doses, their correlation with foci yield is less linear at 3 h and 24 h.

4.4.2 Lymphocyte Subsets

In many of these experiments, the response of specific lymphocyte subsets (CD4, CD8 and CD19) was also quantified. At 30 min, the post-irradiation gamma-H2AX signalling response of all the studied subsets was very similar to the one observed for the whole lymphocyte population. The CD4 positive cells, the largest fraction of the lymphocytes, showed both the slope and zero values the closest to the general lymphocyte population. On the other hand, CD19 positive lymphocytes had consistently less radiation-induced gamma-H2AX foci than CD4 positive cells at 30 min. In fact, for the average of 3 independent experiments, CD19 cells had a significantly lower number of foci than CD4 positive lymphocytes 30 min after exposure to 0.5 Gy (t-test, p<0.004) (Figure 25).

The kinetics of the 3 lymphocyte subsets studied was similar to the response observed with the general population: a decrease in number of foci from 30 min to 3 h after exposure to 0.5 Gy but not for the lower doses (Figure 26).
Figure 25. Dose responses of different lymphocyte subsets. A, B and C are the dose response curves of CD4, CD8 and CD19-positive lymphocytes respectively. D is the mean number of foci specifically for the dose 0.5 Gy. Error bars represent standard deviation and are only shown for points resulting from at least 3 independent experiments. Blood from 4 different individuals was used and each experiment used a sample from a different person. All the measurements here are from samples processed 30 min after irradiation.
Figure 26. Gamma-H2AX signalling kinetics in 3 human lymphocyte subsets after exposure to different doses. Blood samples from different individuals were exposed to 5 doses: 0 Gy, 0.005 Gy, 0.05 Gy, 0.1 Gy, and 0.5 Gy. The post-irradiation foci were scored after different incubation times: 30 min, 3 h and one day. The error bars represent the standard deviation. SD is represented when the condition was repeated in 2 or more independent experiments.
4.5 Radio-adaptive response

After characterizing the low-dose gamma-H2AX signalling response for the general lymphocyte population as well as for several subsets, we tested this signalling endpoint in the context of a radio-adaptive response. In the radio-adaptive response, a priming low-dose irradiation can alter (generally decreasing) the cellular response to an additional irradiation of a higher challenging dose performed a few h later. This apparent change in cellular radiosensitivity due to a low dose allows studying a differential cellular response to radiation without having access to samples of patients of various radiosensitivities.

4.5.1 DNA damage signalling of the general lymphocyte population

According to the gamma-H2AX dose response that we observed, 30 min after an irradiation at 2.05 Gy we should expect an average of about 20 foci per cell (Figure 23). Considering the small size of a lymphocyte, scoring such an amount of foci would be very difficult, whether automatically or not. Therefore, the gamma-H2AX response was observed 3 and 5 h after the challenging dose to consider the impact of DNA repair and the potential foci yield decrease (Figure 27).

When scoring automatically the results of at least 3 independent experiments 3 and 5 h after the challenging irradiation, both irradiation conditions (0.05 Gy + 2 Gy and 2.05 Gy) produced results that were not statistically different (t-test, p=0.99 at 3 h and p=0.47 at 5 h). Furthermore, no condition showed a consistently higher intensity or focus score: the condition “0.05 Gy + 2 Gy” produced 12% more foci per cell than “2.05 Gy” for one individual and generated 27% less foci in another individual.
Figure 27. Low-dose conditioning effect on the gamma-H2AX signalling response to a higher dose gamma-irradiation. Above, in a lighter shade of gray, the number of foci in each condition, 3 (upper left) and 5 h (upper right) after challenging dose. Score made automatically, using CellProfiler. Below, in a darker shade of gray, the mean cellular integrated intensity of gamma-H2AX staining in each condition, 3 (lower left) and 5 h (lower right) after challenging dose. Error bars represent the standard deviation of 3 independent experiments. Blood from a different donor was used for each experiment.

The doses used in these experiments are higher than those used for the validation of the automated scoring. This could lead to a biased quantification of the gamma-H2AX response. Therefore, we focused on post-irradiation times that presented a foci yield within the range of the validation performed before. Furthermore, the cellular integrated intensity of the gamma-H2AX staining was also measured. To do this, CellProfiler adds the intensity of each pixel of gamma-H2AX staining inside the nuclei, both inside and outside the foci. These are the measurements that theoretically should resemble more the flow cytometry approach, which is currently used in other works using this higher dose range (Figure 27).

Using foci yield, both irradiated conditions had responses at least 12 times higher than the non-irradiated control. In the integrated intensity approach, the irradiated cells had gamma-H2AX intensity staining values that were nearly 2 times higher than 0 Gy.
To verify whether the lack of gamma-H2AX signalling difference between the two test conditions was not due to insufficient scoring sensitivity of the automated scoring, the images of 3 of the experiments were also scored manually (Figure 28).

![Graph showing manual vs. CellProfiler scoring](image)

**Figure 28.** Low-dose conditioning effect on the gamma-H2AX signalling response to a higher dose gamma-irradiation. The same images of samples processed 3 h after challenging irradiations where scored manually and using CellProfiler. Error bars represent the standard deviation of 3 independent experiments.

The foci yield values obtained with manual scoring were approximately 3 foci per cell higher than those obtained with CellProfiler. Additionally, manual scoring had less than half of the standard deviation observed with automated scoring. Nevertheless, a Student T-test of the manual scoring results did not point to a statistical difference between the conditioned and unconditioned samples (p=0.3). This is not surprising since the conditioned sample did not have a consistently higher or lower foci frequency among individuals. This statement is relevant for both scoring methods.

Additionally, this manual approach was used to discriminate the response of specific lymphocyte subsets in two individuals (Figure 29). The 3 subsets studied presented similar focus scores: around 7 foci per cell 3 h after irradiation, with little variation amongst subsets. Furthermore, these variations amongst subsets were not consistent between two individuals: in
Figure 29, CD4-positive lymphocytes from sample 1 have a slightly higher number of foci if conditioned (0.05 + 2 Gy) but, in sample 2, the response of this subset is inverted.

There was no clear (reproducible between the two individuals) difference in the response to the two irradiation procedures amongst lymphocyte subsets.

After observing gamma-H2AX, we focused on events that are subsequent to DSB signalling: chromosomal aberrations and apoptosis. We decided to focus on the radio adaptation study in CD4-positive lymphocytes because this subset showed the most intense response to radiation in gamma-H2AX signalling (Figure 25) and in gene expression (Gruel et al. 2008).

4.5.2 Chromosomal aberrations in CD4-positive lymphocytes - the result of DNA repair

Radio-adaptive response has been described for endpoints subsequent to DNA damage repair. To verify if our irradiation protocol induced a radio-adaptive response, we examined chromosome translocations using FISH-3 painting (Figure 30).
To distinguish the radiation response of different lymphocyte subsets we isolated the cell types of interest (CD4, CD8 and CD19) after whole blood irradiation.

Simultaneously, whole blood from the same donor was stimulated with phytohemagglutinin and incubated for 46 h. The initial objective was to compare the radio-adaptive response of one subset (CD4) with the general lymphocyte population. However, the PHA culture produced fewer than 1000 metaphases per condition and was not scored.

The condition with a low dose priming irradiation (0.05 Gy + 2 Gy) produced the same frequency of cells with chromosomal aberrations than the acute

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**Figure 30.** FISH-3 painted metaphase of a CD4-positive lymphocyte. Example of a two-way translocation involving the chromosome 4 (painted in red). The chromosomes 2 and 12 are painted in yellow and green respectively.
equivalent irradiation (2.05 Gy): 22%. For the same individual, the sham-irradiation only had 2% of cells with chromosomal aberrations (Figure 31).

![Percentage of cells with chromosomal aberrations](image)

**Figure 31.** Percentage of lymphocytes with chromosomal aberrations. Results from 1 individual. 396 cells were scored for the 0 Gy, 1290 for 0.05+2Gy and 1512 for the 2.05Gy condition. Only aberrations involving painted chromosomes are considered.

However, these irradiation conditions of equivalent dose are statistically distinguishable when we look at the frequency of the specific aberrations that they induced (Figure 32).

The fractionated exposure (0.05+2 Gy) produced significantly more instable aberrations than the acute irradiation of 2.05 Gy: approximately 1.5x more dicentrics (p<0.007, chi square test) and one-way translocations (p<0.03, chi square test). However, this tendency was inverted for two-way translocations, considered to be more stable: the acute treatment produced 1.8x more than the 0.05+2 Gy condition (p<0.00006, chi square test).
The two irradiation conditions induced roughly the same frequency of incomplete translocations in CD4-positive lymphocytes.

### 4.5.3 Apoptosis

Because differences in cell death between the two conditions could have implications in the chromosomal aberration frequency, our irradiation conditions were also tested using apoptosis as a measure of radiation-induced cell death in the same samples used for the FISH study. The aim was to verify whether differences in chromosome aberration rates were due or not to differences in apoptosis rates. Three hours after the challenging doses were administered, approximately 1.5% of the general lymphocyte population and 4% of the CD4-positive cells were going through apoptosis (i.e. were annexin V positive). There was no detectable difference amongst irradiation conditions (control, conditioned and unconditioned).
Twenty-four hours after challenging irradiation, there was a slight decrease in the fraction of apoptotic CD4-positive lymphocytes to values between 1.5% and 3.4% (Figure 33).

Figure 33. Low-dose conditioning impact on High-dose induced Apoptosis. Above is the apoptotic response (annexin V) of the general lymphocyte population and CD4-positive lymphocytes 3 h after the 3 different irradiation conditions. The apoptotic response of the CD4-positive cells was observed again 24 h after the challenging irradiation. All the results here represented are from the same sampling of one individual.
5 DISCUSSION

5.1 Establishing and validating a fast and sensible gamma-H2AX protocol

Gamma-H2AX signalling has been described as an assay capable of quantifying DSB induced by ionizing radiation (E P Rogakou et al. 1998). In a clinical context, most attempts of developing a fast gamma-H2AX assay, used a flow cytometry approach (Olive & Banáth 2004; Porcedda et al. 2008; Bourton et al. 2011; Ismail et al. 2007; Muslimovic et al. 2008). This technique has the advantage of analyzing a high number of cells but measures the relative intensity of the gamma-H2AX staining instead of scoring the actual number of foci. Actually, manual scoring of gamma-H2AX foci with a microscope gives the true number of foci per cell allowing the detection of ionizing radiation doses almost ten times lower than with flow cytometry (Ismail et al. 2007). Scoring gamma-H2AX foci has proven to be an approach sensitive to the point of detecting doses as low as 0.001 Gy (Rothkamm & Markus Löbrich 2003). Since our work focused on subtle low-dose responses, we needed a tool that could detect small changes in focus scores and that was fast enough to be used on a large scale. To be able to analyze microscope images as quickly as possible image acquisition of the slides as well as focus scoring were therefore done automatically.

5.1.1 Speed of processing and analysis

Manual scoring of gamma-H2AX foci through microscope eyepieces is time-consuming and may become tiresome if many samples need to be analyzed. The alternative used in this work was to acquire z-stacked images automatically and score on images instead of slides avoiding issues of slide prolonged conservation and fluorescence fading. The automation of image acquisition, although not essential for the focus scoring that follows, will render it less time consuming for the operator. Automatic acquisition is
becoming a standard function in many systems of computer-controlled microscopes. The acquisition software that we used is relatively common in cytogenetic labs and several alternatives exist, both commercial and open source, like micro-manager (http://micro-manager.org/). Also in the interest of shortening acquisition and analysis time, 250 cells were established as the minimum number of cells to score in order to reach CV values below 10%. With our current configuration, acquisition took under 8 min per condition (500 cells acquired to insure the analysis of 250).

The number of scored cells varies greatly across publications and very few describe their scoring limits. Similarly to this work, Beels et al. scored manually on images gamma-H2AX foci of 200 to 250 T-lymphocytes (Beels et al. 2010). However, other authors establish their limits by the number of foci. Kuefner et al. scored manually until reaching 40 foci (Kuefner et al. 2010), which, with our conditions, would not have been enough since at least 70 foci needed to be counted to insure a CV below 10%.

In terms of scoring speed, we were able, after some training, to score manually 0.5 cells per second on images. The automated alternatives (HistoLab™ and CellProfiler) used in this study were at least 5 times faster than manual scoring on the same images. However, these two automatic scoring systems are not equivalent since they vary in terms of “operator time”. The use of HistoLab™ allows the user to associate the unbiased automatic cell and focus detection with a visual supervision during the manual loading of each image. On the other hand, the analysis step is still dependent on the operators’ availability, increasing the “operator time” needed for the analysis. Our goal was to develop a complete automated image cytometry with an acceptable compromise between operator time and scoring quality (i.e. a protocol that requires operator intervention time as low as possible, while keeping a good scoring sensitivity). With CellProfiler, the user only takes about 30 seconds to launch the analysis, which will continue in his absence. Moreover, if several conditions need to be analysed several computers can work simultaneously, multiplying the speed analysis by the number of machines available.
Given our current protocol, which uses CellProfiler, it is possible to obtain the analysis results of an irradiated blood sample in less than 6 h.

5.1.2 Characterization and validation of the automated analysis

As long as the blood sample is in good condition to allow lymphocyte isolation and processing, the slides obtained using our method are very rich in analyzable lymphocytes. However in degraded experimental conditions lymphocytes are frequently polluted by granulocytes and other cell debris; it is therefore needed to insure that the analysis of the resulting images is limited to the cells of interest. Because lymphocytes tend to be rounder and to have brighter DAPI staining, their morphology allows operators scoring manually to count them alone. In Histolab™, after all the nuclei in the image are detected, the user has also the possibility to manually remove cells which have aberrant staining or morphology. CellProfiler is a freeware image analysis program that offers a great number of measurement and detection options. The general nuclei detection in the images used an Ostu algorithm, as seen in other works on fibroblasts (Hou et al. 2009). However, in our case, a more specific selection was required to discard non-lymphocytes and damaged cells from the analysis. Therefore we used these measurements to choose only the brighter and rounder cells therefore limiting the analysis to lymphocytes. This morphology-based lymphocyte selection was validated by comparison with a CD3 staining and can be performed in the absence of human intervention.

For gamma-H2AX foci analysis, we chose to score all detectable foci whatever the scoring method used. The coherence of such selection was validated by the linear relationship of gamma-H2AX foci rate with radiation dose. Actually, restricting the selection to the brightest foci reduced the linearity of this correlation. The results obtained in our sham-irradiated condition have shown that the background number of foci per cell was of 0.31. This is acceptable as there are great variations concerning this background level across the different labs working with lymphocytes. The mean yield of gamma-H2AX in
non-irradiated lymphocytes has been described to be as low as 0.09 (Sandrine Roch-Lefèvre et al. 2010) or as high as 0.67 (Beels et al. 2010). Given the dose response that we observed, our protocol estimates that an irradiation of 1 Gy should induce an average of 10 foci per cell. This is well within the range of the published works on radiation induced gamma-H2AX signalling response in lymphocytes. As examples: Beels et al. obtained a relation of 8.6 foci per cell per Gy (Beels et al. 2010), Vilasová et al. obtained 15 foci per cell per Gy (Vilasová et al. 2008) and Löbrich et al. obtained 20 foci per cell per Gy (Markus Löbrich et al. 2005).

5.1.3 Comparison with other approaches in the literature

The number of image cytometry applications that are capable of automated scoring is increasing. In the publications that applied them to gamma-H2AX focus scoring in lymphocytes, all seem to present good correlations to manual scoring once the good parameters are inputted (Jost et al. 2009; Garty et al. 2010; Sandrine Roch-Lefèvre et al. 2010; Ivashkevich et al. 2011). We believe that the choice of the ideal program probably depends more on the type of experiment rather than the sensitivity of the program itself. The scope of the comparison was reduced to the free alternatives to the programs used in our work.

A free alternative that resembles HistoLab™ is FociCounter (Jucha et al. 2010): it was developed specifically to score foci, with a simple interface and few parameters to change (faster to set up). Both programs are ideal for simple scoring of a reduced number of cells/conditions, where the quality of the images/slides frequently requires operator intervention (to eliminate aberrant objects, for example). Since these programs require user intervention to select nuclei, the time of the analysis is their main disadvantage.

CellProfiler is a freeware image analysis program that offers a great number of measurement and detection options. The main drawback of CellProfiler is a less user-friendly graphical user interface that may render the vast number of
detection algorithms and measurements overwhelming to a beginner. However, once the pipeline is set up, it requires very little operator intervention for analysis. We do not compare it to imageJ (Jost et al. 2009) since a CellProfiler module has been recently created to allow the user to run ImageJ macros and plugins as part of a CellProfiler image processing pipeline (http://cellprofiler.org/CPmanual/RunImageJ.html).

In a recent publication, Ivashkevich et al. (2011) present a new image cytometry program that will become freely available. The details on the measurements that will be possible to obtain with this software are not yet disclosed, so we cannot fairly compare it with CellProfiler. However, the computational approach that they present is a good reference to help researchers in selecting the parameters of other programs of this type (Ivashkevich et al. 2011).

5.1.4 Potential and limitations of the gamma-H2AX protocol

The loss of viability of the samples from Ghent showed an important limitation to consider: the fast sample degradation of whole blood. After changing and characterizing our protocol, we believe it to be well adapted for experiments using whole blood irradiations preformed 24 h or less after sampling. Furthermore it possess the sensitivity required for the low-dose context.

Besides the experiments here reported, this method could be used to obtain potentially relevant information only retrievable by a microscope approach. A good example can be found in the work of Böcker and Iliakis, where they use a microscope-based automated analysis to quantify gamma-H2AX and Rad51 foci colocalisation (Böcker & Iliakis 2006). To measure data of gamma-H2AX foci can also be useful to study focus spatial distribution and bystander effect after microbeam irradiations (Ivashkevich et al. 2011). Another possible application is to compare conditions in terms of focus intensity, as done in this work to reveal the bias of scoring only the brightest foci.

However, this technique has also limitations that need to be mentioned. It is better adapted to experiments involving a reduced number of foci per cell.
Foci overlap occurs quite easily in small cells like lymphocytes. Here it is possibly the factor behind the small foci underestimation at 0.5 Gy for automated scoring. Others have also reported underestimations when using automated detection. Jost et al. found twice more foci when scoring manually lymphocytes irradiated at 1 Gy than their ImageJ module (Jost et al. 2009). When using this protocol to study colocalization, it is important to be attentive to possible bias due to antibody competition or fluorochrome bleed-through (Rénier et al. 2007).

In conclusion, the protocol presented here can combine automated acquisition with high-throughput image cytometry to produce a focus score similar to that of manual scoring, with typical kinetics and a linear correlation with dose. This method has proven to be fast and sufficiently sensitive to detect doses as low as 0.005 Gy. Given our current protocol conditions it is possible to obtain the analysis results of an irradiated blood sample in less than 6 h. Our automatic scoring technique produced results fast enough to be seriously considered as an alternative to flow cytometry, for studies that need more detailed information about foci number, distribution, size or intensity. Moreover, any kind of microscopically visible nuclear domains can be quantified in the same manner. Although our research focused more on gamma-H2AX foci, its flexibility allows this method to be adapted to other human lymphocyte-related projects or even other cell types.

5.2 Low-dose response

The gamma-H2AX scoring results of both HistoLab™ and CellProfiler were similar to manual scoring and sensitive enough to statistically distinguish 30 min after exposure the sham-irradiated condition from the 0.05 Gy dose. In fact, using CellProfiler to score foci, it was possible to detect doses as low as 0.005 Gy.

In terms of repair kinetics, we found that low doses (0.005 to 0.05 Gy) induced foci that disappeared at a slower rate than those induced by higher doses (0.1 to 0.5 Gy). This is also in accordance with the findings of Löbrich et al. in lymphocytes and scoring manually: his group found that 5 h after X-ray
irradiation, only 10% of the foci induced by 0.5 Gy were present whereas still half of the foci induced by 0.005 Gy were present at the same time (Markus Löbrich et al. 2005).

5.2.1 Variation of lymphocyte subset response

By using CD-specific membrane staining we were able to discriminate the response of the different lymphocyte subsets. CD4-positive lymphocytes were the subset that had the highest radiation-induced gamma-H2AX signalling. CD19-positive lymphocytes were the cell fraction that presented the lowest gamma-H2AX focus score. These results are consistent with the work of Andrievski and Wilkins. Using flow cytometry, they observed that CD4-positive lymphocytes had a response 1.10 times higher than CD8-positive and 1.56 times higher than CD19-positive cells after being irradiated at 2 Gy (Andrievski & Wilkins 2009). However, at 2.05 Gy, we did not find a consistent difference of response among the subsets.

Given that different lymphocyte subsets could have distinct gamma-H2AX signalling responses, we discriminated the response of lymphocyte subsets to determine if that reduced variation in some of our radio adaptation response experiments. Nevertheless, at the conditions we used to test lymphocyte radio adaptation, we did not find any subset of consistent response between two individuals.

Given the generally higher gamma-H2AX response of CD4-positive lymphocytes at low-dose, it would be interesting to observe more closely this subset using doses below 0.005 Gy to improve even further the sensitivity of the test.

5.3 Individual Clinical radiosensitivity

At Ghent University, Joke Werbrouck and colleges used gamma-H2AX focus scoring to evaluate DNA damage signalling kinetics of HNC patients of
characterized clinical acute radiosensitivity (using the CTCAE classification) (Werbrouck et al. 2011). They irradiated isolated T-lymphocytes and processed them at different times post-irradiation. After analyzing 31 samples, no statistically different gamma-H2AX response was found among the 3 CTCAE radiosensitivity groups. It was hypothesized that the absence of difference in gamma-H2AX signalling kinetics could be due to the irradiation conditions: isolated lymphocytes. Indeed, most of the few published works that tried to link post-irradiation gamma-H2AX response in lymphocytes to clinical radiosensitivity (both acute and late) irradiated isolated lymphocytes and found no correlation (Olive et al. 2008; Vasireddy et al. 2010; Werbrouck et al. 2010). Furthermore, another research group had reported that irradiating whole blood from a highly radiosensitive individual produced similar gamma-H2AX results to in vivo irradiation (CT scan) though in vitro irradiation of isolated lymphocytes from the same person resulted in a normal gamma-H2AX signalling response (Markus Löbrich et al. 2005).

To verify this hypothesis, we received samples from the same patients to repeat the Ghent experiments irradiating whole blood instead of isolated lymphocytes. However, after the travel between the two labs, many blood samples became unusable and no statistical comparison of the results could be done neither with the work of Joke Werbrouck nor between radiosensitivity groups. Given the small amount of analyzable samples, we had to join the results of the CTC1 and CTC2 patients to have an equivalent quantity of data to compare to the patients of highest clinical acute sensitivity (CTC3). We failed to see a difference in DNA damage signalling between these two radiosensitivity groups (CTC1+2 and CTC3).

Presently, it is not possible to conclude whether there was a difference in gamma-H2AX signalling in this study group to be found. The irradiation conditions were not necessarily impairing the test’s sensitivity: Rübe et al. (2010) irradiated both whole blood and isolated lymphocytes and demonstrated that, using the gamma-H2AX foci approach, patients at high risk of developing high-grade life-threatening or lethal toxicities after radio-/chemotherapy could be identified (C. E. Rübe et al. 2010). Moreover, Bourton et al. (2011), only irradiated isolated lymphocytes to follow DSB repair with
gamma-H2AX and was able to discriminate 3 groups: cancer patients with extreme secondary overreactions, cancer patients with none to mild secondary effects to radiotherapy and a non-cancer control group (Bourton et al. 2011). They attributed their successful results to the selection of extremely radiosensitive cases for their study groups. This is coherent with other works that have been able to point out particularly overreacting patients inside their more radiosensitive groups (but not all) or individuals with known DSB repair deficiencies (like ATM-/- or Ligase IV -/-) using gamma-H2AX (Vasireddy et al. 2010; Markus Löbrich et al. 2005; Werbrouck et al. 2011).

These results show that gamma-H2AX has promise in detecting quickly and sensitively individuals with an atypical response to DSB and, therefore, potentially more susceptible to cancer or life-threatening reactions to radiotherapy. Assays like this may therefore help doctors tailor radiotherapy treatments in the future. However, many radiosensitive patients cannot be detected this way, particularly those with moderate radiosensitivity. Furthermore, studies only focussing on the extreme radiosensitivity cases are not representative of the general population. A better understanding of the link between DSB repair and clinical sensitivity is necessary in order to improve the predictive power of these tests. A possible starting point would be to include the co-localization of gamma-H2AX with other DNA repair proteins like ATM or MRE11 (Joubert et al. 2008).

5.4 Radio-adaptive response

In the final part of this work, we turned to a low-dose response currently known as radio-adaptive response in which an exposure to a low-dose (0.005 to 0.2Gy) reduces the damages that result from a higher dose (0.4 to 8 Gy) delivered some time after (Sheldon Wolff 1998; Tapio & Jacob 2007). Here, we report the results of experiments where whole blood from the same individual was irradiated in two distinct ways to achieve a total dose of 2.05 Gy: either an acute (single) exposure or a fractionated irradiation of 0.05 Gy followed by 2Gy 3 h later.
It was not possible to measure the maximum induction of DSB (that would be detected around 30 min after the challenging dose) since too many overlapping foci would result of these doses. So we cannot draw any conclusions regarding the initial radiation-induced DNA damage between the two tested conditions. Nevertheless, the similar foci yield obtained at 3 and 5 h post-challenging indicate that the two conditions we were comparing had very similar DSB repair kinetics. In terms of gamma-H2AX signalling, we found no statistical difference between the two conditions. To insure that these results were not due to a limitation of the automated scoring to separate overlapped foci, we confirmed the results by scoring manually some individuals. Even though manual scoring detected more foci per cell than automated scoring, the final result did not change. There was no detectable difference in gamma-H2AX response between the conditioned and unconditioned samples. This is consistent with previous findings of gamma-H2AX response in fibroblasts. With a 0.1 Gy conditioning dose given 4 h before challenging, they observed no difference in gamma-H2AX fluorescence between conditioned and unconditioned cells, for challenging doses up to 3 Gy. No differences in the kinetics of disappearance of foci were found either (Cramers et al. 2005).

Because different lymphocyte subsets can have distinct gamma-H2AX signalling responses at low doses, we discriminated the radio-adaptive response for the 3 lymphocyte subsets. We found no specific changes in gamma-H2AX response due to priming irradiation for any lymphocyte subset.

We then focused in the response to radiation of CD4-positive lymphocytes. Given their generally higher gamma-H2AX and gene modulation responses (Gruel et al. 2008) to low-doses (in the dose range of the priming dose), this subtype seemed to be the more adequate study the differences in DNA damage that arise due to a pre-irradiation.

Although we did not find any difference between these two conditions in gamma-H2AX signalling at 3 and 5 h, when we used FISH painting to see the specific result of this DNA repair, we found indications that DNA repair had in
fact been changed by the priming dose. The lymphocytes conditioned by the low dose exposure had almost half the two-way translocations of the unconditioned counterparts. This cannot be called a radio-adaptive response since the conditioning also increased significantly the number of dicentrics and one-way translocations.

This increase in dicentrics is not consistent with the previous works of Barquinero (1995) and Sasaki (2002). Barquinero worked with peripheral blood lymphocytes isolated from a group of twelve hospital workers occupationally exposed to X- and gamma-rays. Their cells were considered to be primed in vivo and showed lower frequencies of dicentrics after in vitro irradiation at 2 Gy than the non-exposed control group (J F Barquinero et al. 1995). Sasaki induced adaptive response with a 0.02 Gy pre-irradiation of embryonic CB17 mouse cells. After the 3 Gy challenging dose, the conditioned cells exhibited less dicentrics than the unconditioned controls (Masao S Sasaki et al. 2002).

There are a few possible explanations for the results reported here. For the acute 2.05 Gy irradiation, it would be probabilistically expected to generate the same number of apparently simple translocations (one way, two-way, and incomplete) and apparently simple dicentrics (Duran et al. 2009), however this condition has quite less dicentrics. This can be explained by the fact that we used 72 h of culture and dicentrics are more negatively selected by division than translocations (considered more stable). Cell division could also be the reason why the conditioned sample had more dicentrics than the unconditioned sample: if the fractioned irradiation induced more (or longer) cell cycle arrest than the acute 2.05 Gy irradiation, the higher division rate of the latter would account for its lower dicentrics frequency.

Another factor of importance to explain the differences found between 0.05+2 Gy and 2.05 Gy is apoptosis. It is important to mention that the apoptosis test used here was performed in lymphocytes that were not activated by magnetic beads. Given the low apoptotic rate induced by both conditions and the fact that cell proliferation is known to rescue human peripheral blood lymphocytes from radiation induced apoptosis (Carloni et al.
2001) it is relatively unlikely that apoptosis would be the main pathway behind these different chromosomal aberration frequencies.

Given the known variability in radio-adaptive response and the unexpected ratio of dicentrics versus two-way aberrations in the unconditioned sample these experiments need to be repeated before concrete conclusions can be drawn. Moreover, gamma-H2AX signalling response should be observed at later times to have a wider study of DNA repair kinetics. There has been recently observed that the lower priming irradiation increases cell survival by stimulation of NHEJ repair, with consequences to H2AX signalling kinetics. Apoptosis should be quantified in cultured cells as well as non-cultured cells, to better understand the potential effect in radio-adaptive response.

To better understand these results, this experiment should be repeated with some changes. Firstly, gamma-H2AX response should be observed at later post-irradiation times to better follow the progression of DNA repair. At times intervals were the number of foci has lowered enough automatic scoring should be able to produce more accurate results.

To assess if pre-irradiating induces a longer cell cycle arrest, we could determine a simple mitotic index at different post irradiation times, before and after the 72-hour point presented here. An alternative would be to do a giemsa colouring at 72 h and compare the secondary metaphase frequencies of the two test conditions.

Additionally, apoptosis and micronuclei assays should be performed for the same times and using cultured cells, to better assess the impact of cell death in the resulting chromosomal aberration frequencies.
General conclusions and perspectives

The first objective of this work was to develop a sensitive and fast in vitro assay to evaluate individual radiation sensitivity in human peripheral lymphocytes by using gamma-H2AX quantification.

To reach this aim, we used a microscopy approach to score foci and applied automation to the image acquisition and focus scoring steps to reduce the operator time, making the procedure more viable to be used in a large scale. We were able to distinguish from the sham-irradiated condition doses as low as 0.005 Gy, 30 minutes after exposure. In terms of speed, our approach allows us to obtain the focus yield of a sample in less than 24 h, with only 30 seconds of operator time for the focus scoring step.

Several steps were taken to characterize and validate this protocol. However more are needed before it can be used routinely in an analysis lab:

- more individuals need to be added to the dose-effect curves established
- doses below 0.005 Gy should be included to better establish the detection threshold
- the intra-individual variability should be assessed

Additionally, the specific response of different lymphocyte subsets was discriminated. Different subsets respond to radiation with different intensities. Therefore, by focusing on the response of one subset, it should be possible to develop clinical tests that are potentially more specific and/or more sensitive. We found a higher gamma-H2AX response in CD4-positive lymphocytes, which makes it the most promising in this regard.

More doses and individuals should also be included in the dose-effect curve of CD4-positive lymphocytes, to determine if the detection threshold is improved by focusing on this subset.

To determine the potential of this approach to predict clinical radiosensitivity, an initial version of the protocol was applied to determine de foci yield of samples from Belgian radiotherapy patients of different known
radiosensitivities. The samples were irradiated *in vitro* and the foci yield of the general lymphocyte population was determined for different post-exposure times. No noticeable difference was detected in the foci yield of patients with different clinical radiosensitivities.

Since this experiment, our approach has been improved and validated. So much so that we believe that another test should be performed with samples from individuals of different clinical radiosensitivities. For this new test, however, we would focus on CD4-positive lymphocytes and include a few extreme cases of clinical radiosensitivity.

In view of the difficulties found in obtaining additional blood samples from radiotherapy donors in France for a more complete study, the initial work plan was reoriented. Consequently we focused in lymphocyte gamma-H2AX signalling in another context of different radiosensitivity: radio-adaptive response. This phenomenon has been primarily seen using other endpoints like cell survival and chromosomal aberrations. Therefore, we performed apoptosis and FISH assays to determine if our experimental conditions induced this response before quantifying the gamma-H2AX response. Our preliminary results show a radio-adaptive response in terms of the two-way translocations induced but not for the apoptosis or the gamma-H2AX assays. Furthermore, the dicentrics frequency increased in the fractionated irradiation instead of presenting a radio-adaptive response.

These experiments need to be repeated in at least two other individuals before concrete conclusions can be made. Furthermore, the apoptosis and gamma-H2AX responses need to be quantified at later post-exposure times (between 24 and 72 hours), to better understand the observations made with the FISH assay.
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1.1 Detailed parameters used for automated focus scoring

1.1.1 Histolab

The current version of Histolab is “semi-automatic” since it still requires that the images of gamma-H2AX foci be loaded separately by the user during analysis. Histolab detects both nuclei and, foci by applying a “top hat” threshold to the greyscale images of the DAPI and gamma-H2AX staining respectively.

1st Step: Loading the nuclei images

The Operator loads all DAPI images simultaneously with the button “Read image folder”.

2nd Step: Identify Nuclei

The Operator chooses “nuclei detection” and clicks on the button “identify objects”.
The “nuclei detection” has been programmed with the following parameters:
  - Fixed threshold
    - Intensity from 70 to 255
  - Filtering
    - Macro
      - Fill holes in identified objects
      - Discard objects touching the border of the image
  - Criteria (size restrictions to retain an object as a nucleus)
    - Minimum Object diameter equal or inferior to 3 µm
    - Maximum Object diameter equal or superior to 7 µm
At this point, aberrant nuclei can be manually discarded by the operator if necessary.

3rd Step: Loading the foci images

The Operator loads one by one Texas Red-X image with the menu “download image”.

4th Step: Identify Foci

The Operator chooses “foci detection” and clicks on the button “identify objects”.
The “foci detection” has been programmed with the following parameters:
  - Adaptive threshold (TopHat filter)
    - Bright tones
Maximum diameter: 0.989 µm
Minimum contrast: 45

Filtering
None

Separation
Fine

Criteria (size restrictions to retain an object as a focus)
- Minimum Object diameter equal or inferior to 1.5 µm
- Maximum Object diameter equal or superior to 0.24 µm

The 3rd and 4th steps are repeated until the end of the analysis

Supplemental Table I: Measurements made by Histolab with our current analysis parameters.

<table>
<thead>
<tr>
<th>Nuclei</th>
<th>Foci</th>
</tr>
</thead>
<tbody>
<tr>
<td>Location Center X</td>
<td>Location Center X</td>
</tr>
<tr>
<td>Location Center Y</td>
<td>Location Center Y</td>
</tr>
<tr>
<td>Major Axis Length</td>
<td>Major Axis Length</td>
</tr>
<tr>
<td>Minor Axis Length</td>
<td>Minor Axis Length</td>
</tr>
<tr>
<td>Median size</td>
<td>Median size</td>
</tr>
<tr>
<td>Thickness</td>
<td>Thickness</td>
</tr>
<tr>
<td>Diameter</td>
<td>Diameter</td>
</tr>
<tr>
<td>Perimeter</td>
<td>Perimeter</td>
</tr>
<tr>
<td>Area</td>
<td>Area</td>
</tr>
<tr>
<td>Mean Intensity</td>
<td>Mean Intensity</td>
</tr>
<tr>
<td>Integrated Intensity</td>
<td>Integrated Intensity</td>
</tr>
<tr>
<td>Gamma-H2AX foci Count</td>
<td></td>
</tr>
<tr>
<td>Average Area of the foci inside</td>
<td></td>
</tr>
<tr>
<td>Total Area of the foci inside</td>
<td></td>
</tr>
<tr>
<td>Average Intensity of the foci inside</td>
<td></td>
</tr>
<tr>
<td>Integrated Intensity of the foci inside</td>
<td></td>
</tr>
</tbody>
</table>
CellProfiler

For the analysis with CellProfiler an IdentifyPrimAutomatic module used the Otsu adaptive algorithm on the DAPI pictures to detect nuclei with diameters from 42 to 87 pixels (each pixel = 0.1024 µm). Inside these nuclei, an IdentifyPrimAutomatic module used the Otsu Adaptive per Object algorithm on the H2AX pictures pre-subjected to a Top Hat treatment (using the "Enhance or suppress features" module) to detect foci with diameters from up to 20 pixels. The last modules measured the area and intensity of both nuclei and foci. When CD-specific membrane staining took place, CellProfiler also measured the mean intensity of the associated colour channel inside each cell.

1\textsuperscript{st} Module: LoadImages

File type to be loaded: individual images
Number of channels per group: 2 (DAPI images and H2AX staining images)

2\textsuperscript{nd} Module: IdentifyPrimaryObjects (identify Nuclei)

Select the input image: DAPI staining
Name the primary objects to be identified: Nuclei
Typical diameter of objects, in pixel units (Min, Max): 42, 90
Discard objects outside the diameter range?: Yes
Try to merge too small objects with nearby larger objects?: No
Discard objects touching the border of the image?: Yes
Select the thresholding method: Otsu Global
Threshold correction factor: 2
Lower and upper bounds on threshold: 0, 1
Approximate fraction of image covered by objects?: 10%
Method to distinguish clumped objects: Shape
Method to draw dividing lines between clumped objects: Intensity
Size of smoothing filter: 10
Suppress local maxima that are closer than this minimum allowed distance: 5
Speed up by using lower-resolution image to find local maxima?: Yes
Name the outline image: None
Fill holes in identified objects?: Yes
Automatically calculate size of smoothing filter?: Yes
Automatically calculate minimum allowed distance between local maxima?: Yes
Manual threshold: 0.0
Select binary image: Otsu Adaptive
Retain outlines of the identified objects?: No
Automatically calculate the threshold using the Otsu method?: Yes
Enter Laplacian of Gaussian threshold: .5
Two-class or three-class thresholding?: Three classes
Minimize the weighted variance or the entropy?: Weighted variance
Assign pixels in the middle intensity class to the foreground or the background?: Foreground
Automatically calculate the size of objects for the Laplacian of Gaussian filter?: Yes
Enter LoG filter diameter: 5
Handling of objects if excessive number of objects identified: Continue
Maximum number of objects: 500
Select the measurement to threshold with: None

3rd Module: EnhanceOrSuppressFeatures (TopHat filter)
Select the input image: H2AX staining
Name the output image: Fociplus
Select the operation: Enhance
Feature size: 19
Feature type: Speckles
Range of hole sizes: 1, 10

4th Module: Crop
Select the input image: Fociplus
Name the output image: CropFociPlus
Select the cropping shape: Objects
Select the cropping method: Coordinates
Apply which cycle's cropping pattern?: Every
Left and right rectangle positions: 0, end
Top and bottom rectangle positions: 0, end
Coordinates of ellipse center: 500, 500
Ellipse radius, X direction: 400
Ellipse radius, Y direction: 200
Use Plate Fix?: No
Remove empty rows and columns?: No
Select the masking image: None
Select the image with a cropping mask: None
Select the objects: Nuclei

5th Module: IdentifyPrimaryObjects (identify Foci)
Select the input image: CropFociPlus
Name the primary objects to be identified: h2ax
Typical diameter of objects, in pixel units (Min, Max): 1, 50
Discard objects outside the diameter range?: Yes
Try to merge too small objects with nearby larger objects?: No
Discard objects touching the border of the image?: No
Select the thresholding method: Otsu PerObject
Threshold correction factor: 3
Lower and upper bounds on threshold: 0.04, 0.2
Approximate fraction of image covered by objects?: 0.9
Method to distinguish clumped objects: Intensity
Method to draw dividing lines between clumped objects: Intensity
Size of smoothing filter: 2
Suppress local maxima that are closer than this minimum allowed distance: 4
Speed up by using lower-resolution image to find local maxima?: No
Name the outline image: None
Fill holes in identified objects?: Yes
Automatically calculate size of smoothing filter?: Yes
Automatically calculate minimum allowed distance between local maxima?: No
Manual threshold: 0.057
Select binary image: Otsu PerObject
Retain outlines of the identified objects?: No
Automatically calculate the threshold using the Otsu method?: Yes
Enter Laplacian of Gaussian threshold: .5
Two-class or three-class thresholding?: Three classes
Minimize the weighted variance or the entropy?: Weighted variance
Assign pixels in the middle intensity class to the foreground or the background?: Background
Automatically calculate the size of objects for the Laplacian of Gaussian filter?: Yes
Enter LoG filter diameter: 5
Handling of objects if excessive number of objects identified: Continue
Maximum number of objects: 500
Select the measurement to threshold with: None

6th Module: MeasureObjectSizeShape

Select objects to measure: Nuclei
Select objects to measure: h2ax
Calculate the Zernike features?: No

7th Module: MeasureObjectIntensity

Select an image to measure: H2AX staining
Select objects to measure: h2ax
Select an image to measure: DAPI staining
Select objects to measure: Nuclei

8th Module: RelateObjects

Select the input child objects: h2ax
Select the input parent objects: Nuclei
Calculate distances?: None
Calculate per-parent means for all child measurements?: Yes
Supplemental Table II: Measurements made by CellProfiler with our current analysis parameters.

<table>
<thead>
<tr>
<th>Image</th>
<th>Nuclei</th>
<th>Foci</th>
</tr>
</thead>
<tbody>
<tr>
<td>Count Nuclei</td>
<td>Area</td>
<td>Area</td>
</tr>
<tr>
<td>Count Gamma-H2AX foci</td>
<td>Eccentricity</td>
<td>Eccentricity</td>
</tr>
<tr>
<td>Area Retained After Cropping Foci</td>
<td>Euler Number</td>
<td>Euler Number</td>
</tr>
<tr>
<td>Original Image Area Crop Foci</td>
<td>Extent</td>
<td>Extent</td>
</tr>
<tr>
<td>Execution Time of all Modules</td>
<td>Form Factor</td>
<td>Form Factor</td>
</tr>
<tr>
<td>Final Threshold Nuclei</td>
<td>Major Axis Length</td>
<td>Major Axis Length</td>
</tr>
<tr>
<td>Final Threshold foci</td>
<td>Minor Axis Length</td>
<td>Minor Axis Length</td>
</tr>
<tr>
<td>Orig Threshold Nuclei</td>
<td>Orientation</td>
<td>Orientation</td>
</tr>
<tr>
<td>Orig Threshold foci</td>
<td>Perimeter</td>
<td>Perimeter</td>
</tr>
<tr>
<td>Sum Of Entropies Nuclei</td>
<td>Solidity</td>
<td>Solidity</td>
</tr>
<tr>
<td>Sum Of Entropies foci</td>
<td>Gamma-H2AX foci Count</td>
<td>Integrated Intensity</td>
</tr>
<tr>
<td>Weighted Variance Nuclei</td>
<td>Integrated Intensity</td>
<td>Integrated Intensity of the Edges</td>
</tr>
<tr>
<td>Weighted Variance foci</td>
<td>Integrated Intensity of the Edges</td>
<td>Lower Quartile Intensity</td>
</tr>
<tr>
<td></td>
<td>Lower Quadrile Intensity</td>
<td>Mass Displacement</td>
</tr>
<tr>
<td></td>
<td>Mass Displacement</td>
<td>Max Intensity Edge</td>
</tr>
<tr>
<td></td>
<td>Max Intensity Edge</td>
<td>Mean Intensity</td>
</tr>
<tr>
<td></td>
<td>Mean Intensity</td>
<td>Max Intensity Edge</td>
</tr>
<tr>
<td></td>
<td>Mean Intensity Edge</td>
<td>Mean Intensity</td>
</tr>
<tr>
<td></td>
<td>Median Intensity</td>
<td>Min Intensity Edge</td>
</tr>
<tr>
<td></td>
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<td>Min Intensity Edge</td>
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<td></td>
<td>Min Intensity</td>
<td>Std Intensity Edge</td>
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<td>Std Intensity</td>
<td>Upper Quadrile Intensity</td>
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<td></td>
<td>Std Intensity Edge</td>
<td>Location Center X</td>
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<tr>
<td></td>
<td>Std Intensity</td>
<td>Location Center Y</td>
</tr>
<tr>
<td></td>
<td>Std Intensity</td>
<td>Location Center Y</td>
</tr>
<tr>
<td></td>
<td>(Average of all Measurements made to foci inside each nucleus)</td>
<td></td>
</tr>
</tbody>
</table>
Publications
Quantification of $\gamma$-H2AX Foci in Human Lymphocytes: A Method for Biological Dosimetry after Ionizing Radiation Exposure

Sandrine Roch-Lefèvre, a, Tania Mandina, b Pascale Voisin, a Gruel Gaëtan, a Jorge Ernesto González Mesa, b Marco Valente, a Pierre Bonnesoeur, a Omar García, b Philippe Voisin a and Laurence Roy a

a Institut de Radioprotection et de Sûreté Nucléaire, 92262 Fontenay-aux-Roses, France; and b Centro de Protección e Higiene de las Radiaciones, AP 6195 C. Habana, Cuba

INTRODUCTION

The yield of chromosome aberrations such as dicentrics or translocations has long been used in biological dosimetry in cases of known or suspected exposure to ionizing radiation (1). The dicentric assay is considered to be the reference method for biodosimetry: it is specific to ionizing radiation and is stable enough to be used for dose estimation several months after exposure. However, it has several limitations. It requires skilled personnel and is time consuming. Moreover, one major application of biodosimetry is the identification, in the event of a large-scale radiation emergency, of the most severely exposed individuals. In this situation, a reliable bioassay is needed for population triage during the first few hours. The dicentric assay requires a 48-h culture period to obtain metaphases before chromosome scoring. For faster dose estimation, current protocols for triage assessment call for the analysis of only 50 metaphases. This reduces the delay somewhat but substantially increases the confidence interval and consequently decreases sensitivity to 1 Gy (2, 3). The development of efficient new biodosimeters could overcome the limitations of dose sensitivity and avoid the lymphocyte culture step inherent to the dicentric assay.

Potential candidates include several proteins that are involved in the early steps of cellular response to ionizing radiation and specifically to DNA damage (4). H2AX is one of these. Histone H2AX is phosphorylated at serine 139 soon after double-strand breaks (DSBs) in DNA and generates $\gamma$-H2AX foci. The presence of $\gamma$-H2AX foci was evaluated in human peripheral blood lymphocytes exposed ex vivo to $\gamma$ rays in a dose range of 0.02 to 2 Gy. We analyzed the response of $\gamma$-H2AX to ionizing radiation in relation to dose, time after exposure, and individual variability. We constructed dose–effect calibration curves at 0.5, 8 and 16 h after exposure and evaluated the threshold of detection of the technique. The results show the promise of automatic $\gamma$-H2AX scoring for a reliable assessment of radiation doses in a dose range of 0.6 Gy to 2 Gy up to 16 h after exposure. This $\gamma$-H2AX-based assay may be useful for biodosimetry, especially for triage to distinguish promptly among individuals the ones who have received negligible doses from those with significantly exposures who are in need of immediate medical attention. However, additional in vivo experiments are needed for validation.

1 Address for correspondence: Institut de Radioprotection et de Sûreté Nucléaire/DRPH/SRBE, BP17, 92262 Fontenay-aux-Roses, France; e-mail: srochlefevre@gmail.com.
In Vitro Irradiation and Blood Sample Processing

Twenty-two healthy French donors and six healthy Cuban donors provided blood samples in heparinized tubes. Informed consent was obtained for each donor. Blood samples were collected in accordance with French law (L. 2004-800) on bioethics. The blood was split into several aliquots of 1 ml in dry tubes and irradiated at room temperature with a cobalt-60 source at a dose rate of 0.5 Gy/min for a total of 16 h between exposure and blood sampling. The dose–response curve associated with H2AX responses in relation to dose, time since exposure, and individual variability.

In this study, we assessed the usefulness of automatic quantification of γ-H2AX in peripheral human blood lymphocytes as a new and reliable method of biological dosimetry. Using blood samples irradiated ex vivo, we quantitatively analyzed the formation and loss of γ-H2AX foci at various times after γ-ray exposure, especially the inter- and intraindividual reproducibility. We constructed dose–effect calibration curves and calculated the uncertainty associated with them to determine the technique’s sensitivity up to 16 h after exposure to γ rays. Finally, to check the applicability of this technique in conditions simulating accidents, we tested different ways of inhibiting γ-H2AX signal loss.

MATERIALS AND METHODS

In Vitro Irradiation and Blood Sample Processing

Slides were viewed with an epifluorescence microscope (Provis AX70, Olympus) with a motorized z-stage capacity to map the 3D distribution of γ-H2AX foci through the lymphocyte nucleus. We used a 63× PlanApo objective and an uncooled CCD camera to acquire images. Fields were selected on the basis of DAPI-counterstained nuclei. After acquisition of the DAPI image and application of monochrome thresholding, lymphocyte nuclear areas were defined automatically with Histolab™ software (Microvision Instruments). The resulting image was used as a mask to define focus structures in the nuclei. For focus capture, eight FITC 2D images were acquired with a 1-μm step size between two slices to map the entire nucleus. An FITC 3D image was generated with deconvolution software (Cartograph™, Microvision Instruments). To allow comparisons, FITC image amplification, z-steps and magnification parameters were kept constant during the study.

Focus Analysis

Histolab™ was used for the detection of foci and scoring in FITC images. After thresholding with both object size and object contrast as detection parameters (“Top Hat” filtering), the focus signal was extracted from the noise. To maintain the same sensitivity for focus detection, all images were processed with the same detection parameters of object size and contrast.

Data and Statistical Analyses

To test whether the basal yield of γ-H2AX foci for each individual was significantly higher than the mean value, we applied a one-way test for Poisson variable, which allows the comparison of a scored value to a reference value. For a determined number of scored cells, this test allows calculating the number of γ-H2AX foci as significantly higher with a 95% confidence interval (13). To test for variations between experiments, at least 400 cells from three different experiments were scored for each data point. The mean number of foci per cell (yield, Y), the standard deviation (SD) and the coefficient of variation (CV) of at least three independent measures were calculated. The relationships between two variables were calculated by their correlation coefficient (r), with Pearson’s test, and their two-tailed probability values (P) obtained with a table of critical value for the Pearson’s test. Friedman’s nonparametric ANOVA was used to compare two groups of samples treated under different conditions.

Dose–Effect Calibration Curves and Associated Uncertainties

Dose–effect curves were constructed with delays of 30 min, 8 h and 16 h between exposure and blood sampling. The dose–response curve for γ-H2AX is generally assumed to be linear and therefore was fitted with the equation $Y = \alpha + \beta D$, where $Y$ is the yield of γ-H2AX foci and $D$ is the dose. For the dose–effect curve at 30 min, seven doses in the range 0.02–1 Gy were used. For the dose–effect curves at 8 h and 16 h, three doses in the range 0.5–2 Gy were used. The curves were fitted with linear regression so that the value of the coefficient $\beta$ fit the data points best and the curve passed through the coefficient $\alpha$, which corresponds to the level of γ-H2AX in unirradiated lymphocytes. As already described, we considered that the distribution of γ-H2AX foci among the analyzed lymphocytes follows Poisson’s law (11). To test this hypothesis, we performed a U test, which uses the variance and the mean of each distribution (14). For each dose, the estimated error
(EE) was calculated by combining three uncertainties: the uncertainty of the dose delivered to the blood, the uncertainty from experimental points derived from Poisson’s law, and the uncertainty of the curve fit. The upper and lower curves were then fitted by linear regression such that the value of the coefficient $b$ best fit the curve points $\pm$ EE.

**Calculation of Uncertainty on Dose Estimation**

We expressed the uncertainty as 95% confidence intervals of the dose (95% CI) as described in IAEA Trs no. 405 (1).

**Calculation of Detection Threshold**

The threshold of detection of the method was defined as the lowest dose ($D_{LD}$) significantly different from 0 with a 95% CI. The $D_{LD}$ was calculated according to our coefficient curves as the minimum dose for which the lower confidence limit was greater than zero.

**RESULTS**

**Basal Levels of $\gamma$-H2AX in Human Lymphocytes**

To study the variation in baseline $\gamma$-H2AX levels among individuals, we analyzed the mean number of foci per cell in nonirradiated lymphocytes from 21 healthy donors from the French population (Fig. 1, nos. 1–22). The mean yield of $\gamma$-H2AX $\pm$ SD was 0.09 $\pm$ 0.05 with a range of 0.01 to 0.17 focus per lymphocyte. Data were obtained from six Cuban control subjects (Fig. 1, nos. 23–28). The observed values were similar to those of the French group, i.e., a mean yield of $\gamma$-H2AX $\pm$ SD of 0.07 $\pm$ 0.05 and a range of 0.01 to 0.15 focus per lymphocyte. We also determined for each donor whether the yield of spontaneous $\gamma$-H2AX foci was significantly higher than the mean value calculated with the data from the 27 individuals. We observed that five donors had a basal yield of $\gamma$-H2AX foci that was significantly higher than the mean value yield. We looked at the age and gender of the five donors, but we could not determine any effect of these factors on the basal levels of $\gamma$-H2AX. Unfortunately, we were unable to obtain information about smoking habits for all individuals.

**Kinetics of $\gamma$-H2AX Foci in Lymphocytes after Exposure to Ionizing Radiation**

We investigated the dynamics of focus induction and loss over 5 h after a 0.5 Gy $\gamma$ irradiation (Fig. 2). The number of foci increased rapidly and reached 5.1 ($\pm$ 0.2) foci/cell at 30 min. This number then decreased so that 18.5% of the $\gamma$-H2AX foci present at 30 min were still present at 5 h after irradiation (Fig. 2A). An incubation period of 30 min at 37°C was optimum for determining the maximum yield of $\gamma$-H2AX for the irradiated whole blood. The SD peak at 60 min after $\gamma$ irradiation may be explained by the distribution of the foci at each time. Despite the very similar average yields of $\gamma$-H2AX at 30 and 60 min after exposure (Student’s $t$ test, $P > 0.82$), the distribution of foci in the cells differed substantially (Fig. 3). The distribution of $\gamma$-H2AX foci showed great overdispersion at 60 min, which may reflect higher interindividual variability of kinetics of DNA repair at this time (Fig. 2A).

**Individual Variability**

To have some indications about intraindividual variability, the $\gamma$-H2AX yield was measured in lymphocytes from a single individual in three separate blood
FIG. 2. Panel A: Induction of γ-H2AX foci automatically counted in lymphocytes and assessed from 5 min to 5 h after irradiation with 0.5 Gy. The error bars represent the SD (n = 10). Panels B–D: γ-H2AX foci in lymphocytes. There is good correlation between the intensity of automatically segmented foci (right) and visually apparent foci (left). Panel B: γ-H2AX foci in lymphocytes 5 min after exposure to 0.5 Gy of γ rays. Panel C: γ-H2AX foci in lymphocytes 30 min after exposure to 0.5 Gy of γ rays. Panel D: γ-H2AX foci in lymphocytes 60 min after exposure to 0.5 Gy of γ rays.

FIG. 3. Distribution of γ-H2AX foci counted automatically in lymphocytes assessed before and from 5 min to 5 h after irradiation with 0.5 Gy. Data from four independent experiments were combined. NE: not exposed.
To quantify interindividual variability, we measured the γ-H2AX focus yield in the lymphocytes again before exposure for 27 individuals and at each of the times after exposure for 11 individuals (Table 1B). In nonexposed lymphocytes, the interindividual variation of the γ-H2AX yield was two to three times higher than the intraindividual variation, with a CV = 51% when individuals nos. 1–28 were taken into account and a CV of 31% when only individuals nos. 14–22 were taken into account. On the other hand, the interindividual variations of the γ-H2AX yield after irradiation were similar to the intraindividual variation. The CV was the lowest (<15%) at more than 30% at 30 min and peaked at 5 min and 60 min. This result confirmed the lower interindividual variability observed at 30 min (Fig. 2A).

### Dose–Effect Calibration Curves

Three dose–effect calibration curves were constructed: at 30 min, when the lowest interindividual variation was observed and at 8 and 16 h after exposure, times at which blood samples might be taken after unexpected radiation exposure. The yield of γ-H2AX foci measured 30 min after *ex vivo* exposure of whole blood increased linearly with the radiation dose up to 1 Gy (Fig. 4A). Beyond this dose, the curve was distorted, probably due to overlapping foci (data not shown). When we limited the highest dose to 1 Gy and assumed linear dose dependence, we obtained the following dose–effect relationship:

\[
Y = 0.09(\pm 0.15) + 10.7(\pm 0.5)D,
\]

where \(Y\) is the yield of γ-H2AX foci and \(D\) is the dose \((r = 0.999, P < 0.001)\).

For the later times after exposure, the yield of γ-H2AX foci increased linearly with the radiation dose up to 2 Gy (Fig. 4B and C). Curve distortion was observed beyond 2 Gy. When we limited the highest dose to 2 Gy and assumed linear dose dependence, we obtained the following dose–effect relationships: 0.13 (±0.14) + 0.88 (±0.05) \(D\) at 8 h and 0.15 (±0.16) + 0.50 (±0.03) \(D\) at 16 h after exposure \((r = 0.999, P < 0.001\) and \(r = 0.996, P < 0.005\), respectively).

### Thresholds of Detection

The threshold of detection is the dose at which the yield of γ-H2AX foci is significantly different from the yield of γ-H2AX foci in nonirradiated lymphocytes. Since the level of γ-H2AX varies with time after irradiation, the sensitivity of its detection is necessarily dependent on time. We calculated the threshold of detection at each time for which we constructed a dose–effect calibration curve. Given our curve coefficients \(a\) and \(b\), the calculated \(D_{15}\) value was 0.05 Gy (95% CI

---

**Note:** Part A reports intraindividual variability; individual no. 16 provided 3 blood samples taken on three different days (numbered 16A–C). Part B reports interindividual variability. NE: not exposed. ND: not determined.

**Notes.**

- "At least 50 foci scored in nonexposed cells.
- "CV less than 15% (Table 1A). The CVs after irradiation were less than 15%.

### TABLE 1

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>5</th>
<th>30</th>
<th>60</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>A</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>NE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16A'</td>
<td>0.16</td>
<td>3.96</td>
<td>3.26</td>
</tr>
<tr>
<td>16B'</td>
<td>0.17</td>
<td>4.03</td>
<td>3.65</td>
</tr>
<tr>
<td>16C'</td>
<td>0.13</td>
<td>2.17</td>
<td>4.22</td>
</tr>
<tr>
<td>Mean</td>
<td>0.15</td>
<td>3.39</td>
<td>3.71</td>
</tr>
<tr>
<td>SD</td>
<td>0.02</td>
<td>1.05</td>
<td>0.48</td>
</tr>
<tr>
<td>CV</td>
<td>0.14</td>
<td>0.13</td>
<td>1.94</td>
</tr>
<tr>
<td>No. of nuclei scored</td>
<td>1011</td>
<td>421</td>
<td>402</td>
</tr>
</tbody>
</table>

**Note:** Part A reports intraindividual variability; individual no. 16 provided 3 blood samples taken on three different days (numbered 16A–C). Part B reports interindividual variability. NE: not exposed. ND: not determined.

**Notes.**

- "At least 50 foci scored in nonexposed cells.

**Mean **

- "CV less than 15% (Table 1A). The CVs after irradiation were less than 15%.

### TABLE 1

Yields of γ-H2AX Foci Obtained in Nonexposed Lymphocytes or at Various Times after a 0.5-Gy γ-Ray Dose and Subsequent Incubation at 37°C

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time (min)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>A</strong></td>
<td>NE</td>
<td></td>
</tr>
<tr>
<td><strong>B</strong></td>
<td>NE</td>
<td></td>
</tr>
<tr>
<td>16A'</td>
<td>0.16</td>
<td></td>
</tr>
<tr>
<td>16B'</td>
<td>0.17</td>
<td></td>
</tr>
<tr>
<td>16C'</td>
<td>0.13</td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>0.15</td>
<td></td>
</tr>
<tr>
<td>SD</td>
<td>0.02</td>
<td></td>
</tr>
<tr>
<td>CV</td>
<td>0.14</td>
<td></td>
</tr>
<tr>
<td>No. of nuclei scored</td>
<td>1011</td>
<td>421</td>
</tr>
</tbody>
</table>

**Note:** Part A reports intraindividual variability; individual no. 16 provided 3 blood samples taken on three different days (numbered 16A–C). Part B reports interindividual variability. NE: not exposed. ND: not determined.

**Notes.**

- "At least 50 foci scored in nonexposed cells.

**Mean **

- "CV less than 15% (Table 1A). The CVs after irradiation were less than 15%.
Focus Signal Stabilization

To use $\gamma$-H2AX scoring for dose determination after unplanned exposure, the signal loss must be inhibited during transport to the expert laboratory. We therefore tested blocking the repair process by incubating the whole blood on ice at $0^\circ$C. After the 0.5-Gy $\gamma$-ray dose and an optimal period of focus induction of 30 min at $37^\circ$C, we added 1.5 ml of cold RPMI-1640 medium (at $0^\circ$C) to 1 ml of whole blood. We compared the yields of $\gamma$-H2AX foci obtained 30 min after irradiation and incubation at $37^\circ$C to the yields obtained with the same treatment and additional incubation of 30 min, 2 h, 4 h, 30 min, and 23 h 30 min at $0^\circ$C on ice, or 30 min, 2 h, 4 h 30 min, and 23 h 30 min at $37^\circ$C (Fig. 5A). We observed that the yield of $\gamma$-H2AX foci decreased significantly during incubation at $37^\circ$C (ANOVA, $P < 0.04$) as DNA repair continued, whereas the number of $\gamma$-H2AX foci remained constant for up to 24 h after irradiation when the blood was incubated at $0^\circ$C in ice (ANOVA, $P < 0.45$). For incubation periods longer than 16 h at a temperature of $37^\circ$C, we were unable to quantify $\gamma$-H2AX foci correctly because of the poor quality of lymphocyte isolation.

To avoid the use of ice, two other possible methods to maintain $\gamma$-H2AX signals were examined. We tested whether calyculin A, a well-known protein phosphatase 1 and 2A inhibitor, could maintain H2AX phosphorylation in vitro, as reported previously (15). After we added calyculin A to the whole blood we observed the same kinetics of $\gamma$-H2AX focus induction and loss, with in addition a very high rate of apoptotic lymphocytes (data not shown). Finally, we blocked the focus signal loss by adding paraformaldehyde (PFA) directly to the whole blood before isolating the lymphocytes and compared the yields of $\gamma$-H2AX foci obtained 30 min after irradiation and incubation at $37^\circ$C with the yields obtained with the same treatment and additional incubation for 90 min in 1.3% PFA at room temperature or 90 min at $37^\circ$C without PFA (Fig. 5B). We observed that the yield of $\gamma$-H2AX foci was decreasing at 120 min after incubation at $37^\circ$C, whereas in the presence of PFA, the number of $\gamma$-H2AX foci remained constant for 120 min. However, we were unable to isolate lymphocytes sufficiently well for the PFA incubation times exceeding 150 min.

DISCUSSION

This study demonstrates that automatic scoring of $\gamma$-H2AX foci in human lymphocytes provides reliable information about ionizing radiation dose for blood samples analyzed up to 16 h after exposure. Using whole blood irradiated ex vivo, we verified that the number of foci induced is linearly associated with the radiation dose and that the yield of $\gamma$-H2AX foci peaks at 30 to 60 min after exposure. Furthermore, we showed that the technique’s sensitivity is 0.05 Gy at 30 min after irradiation and still 0.6 Gy at 16 h after exposure. Finally, incubation of the whole blood at $0^\circ$C allows the stabilization of focus signals for up to 24 h after sample collection before analysis.
After ionizing radiation induced γ-H2AX foci, we observed induction dynamics consistent with both in vivo and in vitro data from the literature (9, 11, 12). Our findings confirmed that 30 min after irradiation is the optimum time for determining the maximum yield of γ-H2AX foci for human lymphocytes, as reported for the HF19 human cell line (9, 10). Measuring foci induced by γ radiation at 30 min after irradiation, we found a mean yield of 10.7 foci/cell Gy⁻¹. We compared this value to those of three studies where γ-H2AX foci were scored manually. Although our value is very close to the 10.4 foci/cell Gy⁻¹ reported in a study after in vitro irradiation of human lymphocytes (12); it is lower than the yield (≈ 10–15 foci/cell Gy⁻¹) reported in two other studies (9, 11). We cannot exclude the possibility that the observed discrepancies are due at least partly to differences in scoring protocols between laboratories.

**Technical Limitations of γ-H2AX Focus Scoring for Dose Estimation**

We identified some technical limitations of our methodology for high and low doses of radiation and also for in vitro incubation times longer than 16 h after exposure. When we used Histolab™ software for automatic γ-H2AX focus detection, the linearity of the dose-effect relationship seemed to disappear at doses above 1 Gy at 30 min after exposure and above 2 Gy at 8 h after exposure. This was probably due to an overlap of foci that occurs as the number of foci per nucleus increases with the dose of ionizing radiation. Some measurement methods use algorithms that attempt to correct for focus overlap. In any case, focus overlap should not exceed 20% for accurate focus scoring (16). Conventional cytogenetic methods such as the dicentric assay can provide accurate dose assessment for doses exceeding 1 Gy when 500 metaphases are scored (17).

The lower limit of dose detection by γ-H2AX focus scoring will depend on its reproducibility. With the automatic scoring method we used here, reproducibility was good in irradiated lymphocyte samples from the same individual, and 0.05 Gy was the lowest detectable dose 30 min after exposure. This threshold of detection may be disappointing compared with previous studies that used scoring of γ-H2AX foci for very low dose estimation in human lymphocytes (9, 11). However, the reduced sensitivity in our study comes from the fact that pre-exposure levels of γ-H2AX are considered to be unknown as would be the case for accidental overexposure to ionizing radiation. We then constructed dose–effect calibration curves using values from at least three different individuals to calculate individual doses. Then the curve coefficients α (representing the mean level of γ-H2AX foci in nonirradiated lymphocytes from all individuals) and β (representing the slope of the dose–effect curve) were used to calculate the dose significantly different from 0 with a 95% confidence interval. The dose sensitivity of the γ-H2AX assay is therefore directly linked to the slope of the calibration curve as well as the basal level of γ-H2AX foci. Finally, we observed that incubating whole blood in vitro (even diluted with an appropriate medium) for more than 16 h at 37°C could lead to a lymphocyte preparation of lower quality and consequently to a poorer γ-H2AX focus staining quality. As a result, we could study the dose–effect relationship only until 16 h after exposure. This problem was not observed when lymphocytes were isolated before irradiation.

**Relevance of the γ-H2AX Assay as a New Method of Biodosimetry**

To apply the γ-H2AX assay, for biodosimetry, it is essential to check whether the yield of γ-H2AX foci
observed in the patient's lymphocytes differs significantly from the background level in control subjects. After assessing the basal levels of γ-H2AX foci in 21 French and 6 Cuban control individuals, we obtained mean yields of 0.09 and 0.07 focus/cell, respectively, which were similar to those observed in nonexposed human lymphocytes after manual scoring (0.06 focus/cell) (9, 11). Regardless of country, however, high variation in base levels between individuals was observed. When only 8 to 20 foci were scored (in 200 cells), the variation between individuals was about 50%, whereas after scoring of 50 foci (in 500 cells), the variation was lower but was still 30%. This means that a part of the interindividual variability could be due to an insufficient number of scored cells. The Poisson distribution of γ-H2AX foci in lymphocytes implies that the more cells scored for each sample, the smaller the uncertainties. Nevertheless, even if enough cells are scored, the interindividual variation in base levels is high and can be explained by the fact that the formation of γ-H2AX foci may be induced by a variety of physical, chemical and biological factors (18), including heat (19). Therefore, to take individual variability of physiological parameters into account, γ-H2AX foci must be scored in large control population cohorts. Moreover, considering the differences in lifestyles from one country to another, it is important to compare basal yields of γ-H2AX foci in various countries. To avoid artifactual scoring bias between laboratories, an automatic system for γ-H2AX focus detection allowing the scoring of foci based on objective criteria of focus size and intensity must be employed.

In view of the overall uncertainty of the γ-H2AX assay, a sensitivity of about 0.3 to 0.6 Gy was assumed in a period that allows blood sampling in case of unplanned overexposure to ionizing radiation. This is similar to the sensitivity of the dicentric assay when only 50 metaphases per individual are observed (20).
dicentric assay is the reference method for biological dosimetry in cases of recent overexposure to ionizing radiation. Using γ-H2AX as a biodosimeter may reduce the delay of dose estimate, since this method does not require the 48-h culture period. This may be very useful especially in the case of a large-scale nuclear event, because it can discriminate individuals who were and were not exposed to radiation (4). Furthermore, γ-H2AX focus scoring can be automated to facilitate dose estimates for larger cohorts. Several studies of software-based γ-H2AX focus scoring reported data validated in human cell lines (16, 21–23). Currently, there are several software/freeware programs that can allow automatic focus scoring with a similar kind of approach (e.g., Image-Pro Plus, Cell Profiler, Metacyte, Scan R Analysis). The use of different software in biodosimetry laboratories would still allow laboratory intercomparisons since “training” images can be swapped between laboratories to adjust the parameters used for focus detection to obtain the same scoring results.

In many accident situations, radiation casualties would have only partial-body exposures to ionizing radiation. For the dicentric assay, the IAEA manual recommends either the contaminated Poisson method or the Qdr method (1). Both methods rely on the distribution of cells containing dicentrics in the body compared to the expected Poisson distribution. Since the distribution of foci containing cells follows Poisson law, these methods could be applied to the γ-H2AX assay in the same way. This would allow both the calculation of the dose received by the irradiated part of the body and the estimated volume of exposed lymphocytes. However, this will require in vivo experiments for validation.

**Strategy in Case of Accidental Exposure to Ionizing Radiation**

The most critical issue for the application of the γ-H2AX assay to biological dosimetry appears to be focus signal loss. The number of γ-H2AX foci can decrease from 1 h after γ irradiation; the slope of the dose–effect calibration curve should therefore decrease with γ-H2AX signal loss. The time between exposure to ionizing radiation and blood sampling must therefore be known to be able to relate the yield of γ-H2AX foci to the dose–effect curve established after the same delay after irradiation (T1, Fig. 6). Here dose–effect curves for γ-H2AX were constructed at 8 h and 16 h after irradiation. These times may allow the collection of blood samples from victims in the event of accidental exposure. We determined the slope at each time and the associated lower doses for which the level of γ-H2AX foci was significantly different from 0 with a 95% confidence interval. The repair process must be inhibited during blood sample transport. For incubation of the whole blood on ice, inhibition of γ-H2AX signal loss was possible for up to 24 h. This implies that transport and analysis of the blood should be done within 24 h (T2, Fig. 6).

A strategy that takes into account the constraints inherent to γ-H2AX can be designed to provide reliable dose estimation after unplanned overexposure as efficiently as possible. During the first 16 h after the potential overexposure, the dose estimation should use the γ-H2AX assay for the rapid determination of doses in a dose range of 0.3 to 2 Gy. This could quickly distinguish individuals exposed to doses below 2 Gy from those exposed to higher doses of ionizing radiation, who require emergency medical treatment. In case of overexposure that occurred more than 1 day before blood sampling, only the dicentric assay is currently possible.

In summary, this article presents a method of biodosimetry based on automatic scoring of γ-H2AX foci in human lymphocytes and validates it in a period from 30 min to 16 h after exposure to γ rays and in a dose range of 0.05 to 2 Gy. It generates results comparable to those from the literature obtained by manual γ-H2AX analysis but avoids constraints associated with manual scoring. One potential application of the method described here could be rapid population triage in case of a large-scale nuclear accident. Our study shows that automatic counting of γ-H2AX foci in peripheral human blood lymphocytes may be useful for biodosimetry. However promising, this method will require additional in vivo experiments for further validation, especially in the case of partial-body exposure, which represents the majority of accidental overexposures.

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Authors: M. Valente, P. Voisin, P. Laloi, L. Roy, S. Roch-Lefèvre

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Automated gamma-H2AX focus scoring method for human lymphocytes after ionizing radiation exposure

M. Valente\textsuperscript{a}, P. Voisin\textsuperscript{a}, P. Laloi\textsuperscript{a}, L. Roy\textsuperscript{a} and S. Roch-Lefèvre \textsuperscript{a,*}

\textsuperscript{1} Institut de Radioprotection et de Sureté Nucléaire (IRSN), DRPH, SRBE, LDB, BP 17 92262 Fontenay-aux-Roses, France

* Corresponding author: Sandrine.roch-lefevre@irsn.fr
Abstract

The purpose of this study was to develop a microscopy-based foci quantification protocol in human lymphocyte capable of supplying useful data for radiation sensitivity assays. Human peripheral blood was exposed to gamma-rays and isolated lymphocytes were stained with fluorochrome-coupled anti-gamma-H2AX (Histone 2AX phosphorylation of serine 139) antibodies. Microscopy slides were automatically acquired and the resulting images were subjected to 3 focus scoring methods: manual, semi-automated and fully automated. All scoring methods were sufficiently sensitive to detect an irradiation of at least 0.05 Gy with low variation between experiments. For higher doses, both automated approaches tend to detect fewer foci than manual scoring but still obtaining a linear correlation (lowest $r^2 > 0.971$). Compared with manual scoring on the images, the automated approaches are at least 5 times faster with minimum operator intervention needed. We can conclude that our method is able to obtain the foci score of a blood sample in less than 6 hours. In addition to the foci score the programs used perform several cell and foci measurements of potential biological importance.

Keywords: lymphocytes; gamma-H2AX foci; automated scoring
1. Introduction

Humans can be exposed to several sources of ionizing radiation (IR), from cosmic rays to medical scans and different types of radiotherapy. Since the discovery of radioactivity, much has been learned about the effects of high-dose exposure and of radiosensitizing diseases such as ataxia telangiectasia. However, many questions about the effects of low-dose exposure and of hypersensitivity to radiation remain unanswered, perhaps because no tool is both sensitive and fast enough to be used on a large scale, in epidemiological studies for example.

One of the primary cellular effects of ionizing radiation is the induction of DSB (double-strand breaks). Following DSB induction, hundreds of histone H2AX molecules are phosphorylated (gamma-H2AX) in the chromatin flanking the DSB site (Rogakou et al., 1998). Specific antibodies against gamma-H2AX allow the visualization of a "focus" at each DSB site. The scoring of nuclear gamma-H2AX foci is widely accepted as a very sensitive method for quantifying radiation-induced DSB.

Several protocols of automated image acquisition and/or analysis are used to study gamma-H2AX foci in human cellular models (Bocker and Iliakis, 2006; Costes et al., 2006; Hou et al., 2009; Leatherbarrow et al., 2006; Mistrik et al., 2009). Recently, this approach has also been used to score gamma-H2AX foci in lymphocytes (Roch-Lefevre et al., 2010) but without taking advantage of any other type of measurements. Here we describe the optimization of this approach for human lymphocytes isolated from blood exposed ex vivo to different doses of radiation. We used resources common in cytogenetic labs for automatic acquisition and both a commercial and a freeware program for image analysis. In addition to focus counting, the latter was used to measure DAPI (4’,6-diamidino-2-phenylindole) intensity and morphology allowing the operator to enrich the lymphocyte fraction of the analyzed cell population. Based on our extensive experience with microscopy, we optimized and validated this morphological cell selection, an alternative less expensive than using a dedicated cellular membrane staining. Then, to further accelerate the protocol, we determined the minimum number of cells to analyze. The scoring results obtained automatically were very similar to manual scoring. Furthermore, we could verify that gamma-H2AX focus frequency increased linearly with ionizing radiation dose from 0 to 0.5 Gy with a detection threshold of 0.05 Gy at 30 minutes post-exposure.
2. Materials and methods

2.1 Ethics and Sample Collection
The blood was collected from healthy volunteers by the "Etablissement Francais du Sang" (EFS) which obtained written informed consent for all donors, according to the agreement between EFS and IRSN (reference n° 10/EFS/056). The blood samples were collected in citrate and were processed shortly after sampling.

2.2 Irradiation Conditions and Lymphocyte Separation
The dose-response curve was established with blood samples from 3 healthy donors 30 minutes after exposure to gamma-rays. Five doses were used: 0, 0.05, 0.1, 0.2 and 0.5 Gy. Total blood was irradiated at room temperature (RT) with a cesium-137 source (IBL637, CisBio, France) at a dose rate of 0.5 Gy per minute, and then incubated at 37°C to induce gamma-H2AX formation. All subsequent procedures until cell fixation were performed on ice to reduce variations in gamma-H2AX by preventing ongoing DNA double-strand break repair.

The lymphocytes were isolated using Ficoll-Histopaque (Invitrogen, Carlsbad, CA, USA) and washed three times with cold phosphate buffered saline (PBS) (Invitrogen, Carlsbad, CA, USA). The lymphocytes were then spotted by adhesion in DakoPen® (Dako, Glostrup, Denmark) circles onto Superfrost® slides (Menzel-Glaser, Braunschweig, Germany) at 0°C. Then, the cells were immediately fixed in 1% PFA for 10 minutes at RT and washed in PBS. Finally, the adherent lymphocytes were preserved in PBS at 4°C for one to three days before staining.

2.3 Immunofluorescence Staining
Adherent cells were permeabilized in PBS + 0.1% TritonX-100 for 10 minutes. The cells were blocked in PBS with 20% goat serum for 30 minutes at RT. Slides were incubated with monoclonal anti-H2AX Phosphorylated (Ser139) antibody (1:500 dilution, BioLegend, San Diego, CA, USA) for 60 minutes, washed in PBS for 5 minutes, and incubated with Texas Red X conjugated goat anti-mouse secondary antibody (1:1500) for 60 minutes, all at RT.

For CD3-specific (cluster of differentiation 3) staining the slides were incubated with AlexaFluor® 647-coupled anti-human CD3 (1:100 dilution) (BD Pharmingen, Franklin Lakes, NJ, USA) for 30 minutes.
Finally, the slides were mounted with cover slips; DAPI Prolong® Gold solution (Invitrogen, Carlsbad, CA, USA) was used according to the manufacturer's instructions.

2.4 Image Acquisition and Processing

Slides were viewed with an epifluorescence microscope (Imager.Z1, Carl Zeiss, Oberkochen, Germany) equipped with a non-cooled CCD (charge-coupled device) camera and an alignment-free external fluorescence light source (EL6000, Leica, Solms, Germany). The automated image acquisition used Metafer 4 software (version 3.6.0, from MetaSystems™, Altusheim, Germany). 100 to 250 fields of each spot were selected and acquired by the Metafer Autocapt module, using an immersion plan Apochromat oil 63x objective (Carl Zeiss). To compile all of the 3-dimensionally distributed gamma-H2AX foci throughout the lymphocyte nuclei in one image, 26 2D-images for each field were acquired with a 0.3 μm z-axis step between two slices. The resulting fields of view (FOV) were transformed into training images (TRN) with the "Create TRN from FOV" to allow each colour channel to be exported as an individual greyscale tiff file.

All the steps described above from sample collection to image acquisition are shared by all the subsequent methods of analysis used.

2.5 Image Analysis

Both manual and automatic focus scoring were performed in uncompressed high-quality images. Even the manual scoring mentioned in this work was performed in the images and not through the eyepieces. The manual scoring was done by 3 operators that were unaware of the doses corresponding to the images they were analysing. The lymphocytes were identified under the microscope by morphological criteria thanks to their roundness and their bright DAPI staining. Each operator counted a minimum of 150 foci or 250 cells per condition. The operators were told to consider every focus that they could discriminate from the background regardless of their intensity. For the automatic detection, two different programs were used: the commercial software HistoLab™ (version 7.5.2, Microvision Instruments, Evry, France) and the free cell image analysis software, CellProfiler (version 2.0, Broad Institute, MA, USA) (Carpenter et al., 2006). For the CellProfiler analysis the PC specifications were: windows XP Professional, Processor Intel Core 2 Duo 2 GHz, 512 Mo RAM.

The specific settings used for both HistoLab™ and CellProfiler are supplied in Table 1. All image analysis parameters were kept constant throughout the duration of this study.
2.6 Statistical analysis

Statistical analysis was performed using SigmaPlot (Version 11, from Systat Software Inc, San Jose, California) and Microsoft Excel (Version 2003, Redmond, Washington, USA). To test the sensitivity of the scoring approaches the foci frequency of 0 Gy (3 individuals) was compared with the foci frequency of 0.05 Gy (3 individuals) with a Students’ T-test.
Results
Here, we describe a microscopy-based method for gamma-H2AX foci scoring in human lymphocytes after gamma-ray exposure. The images automatically acquired using Metafer™ were analysed by three different methods: manual, semi-automatic (HistoLab™) and automatic (Cellprofiler). We compare for each method: the selection of the lymphocytes to be analyzed, the linearity of gamma-H2AX dose response, the sensitivity, the measurement variations, and the time of operator intervention needed for analysis.

3.1 Sample Processing and Image Acquisition
Using the protocol described in the “Material and methods” section, the time required for the process, from the blood sample to the mounted slide, is 5 hours at most. With our experimental conditions, we obtained an average lymphocyte density of 25 cells per field (up to 80) for an acquisition at x63 magnification. The average speed of image acquisition and export with our conditions was 2.5 images per minute. The images acquired this way were the source of the comparison between that follows.

3.2 Image Analysis: Cell Type Selection
Operators scoring manually were able to exclude monocytes and granulocytes from the analysis. With the current version of HistoLab™ the user has the possibility of removing manually the cells with aberrant staining or morphology. Because the CellProfiler approach has no operator intervention, the cell type selection was done graphically using two measurements: the nuclei "form factor" that quantifies "roundness" and the "DAPI integrated intensity". To maximize the lymphocyte concentration of the cell population we studied, we analyzed only the cells with a form factor above 0.86 and a DAPI-integrated intensity above 1000 (Figure 1A). To verify that this morphologic selection by CellProfiler was able to increase efficiently the proportion of lymphocytes in the cell group, we quantified the proportion of CD3-positive lymphocytes. The selection based on form and intensity measurements presented here successfully enriched in CD3-positive lymphocytes the cell population from 59% to 83% (Figure 1B). It is also important to mention it was verified that irradiating samples does not influence this selection (data not shown).

3.3 Image Analysis: Gamma-H2AX Foci scoring
To evaluate the potential of CellProfiler to score gamma-H2AX foci, this automatic method was compared to the manual one and a semi-automatic scoring system already being used in
our lab (HistoLab™, (Roch-Lefevre, et al.)). The same automatically acquired images using Metafer™ were analysed by all three approaches.

A dose response was established using the three different scoring methods (Figure 2). After manual scoring, each operator found a linear increase of gamma-H2AX foci with radiation dose (Pearson's test: $r^2 > 0.991$, $p < 0.0004$). The results obtained with HistoLab™ and CellProfiler revealed as well a very linear relation between dose and foci score (Pearson's test $r^2 > 0.981$, $p < 0.002$ and $r^2 > 0.971$, $p < 0.003$, respectively). For all scoring methods (manual or automated) the average number of foci per cell scored for the three blood samples was significantly different between sham irradiation and the dose of 0.05 Gy (Students’ T-test, $p < 0.02$).

The inter-operator scoring variations were estimated using images from the same experiment which were analysed manually by three different operators. The inter-operator results were very similar with a coefficient of variation (CV) above 10% only in the sham irradiated condition. The inter-method scoring variations were estimated using images from the same experiment that were analyzed manually by one of the operator and also automatically by HistoLab™ and CellProfiler. The scores from the three methods were very similar as well with a CV of around 10% whatever the dose condition. To estimate inter-experiment variations, images of all three experiments were scored by one of the operators manually but also automatically by HistoLab™ and CellProfiler. Whatever the scoring method used, the inter-experiment variations were high with a CV of at least 30% for the sham irradiated condition and a CV below 10% only for the 0.5 Gy condition.

We determined the minimum number of cells to be scored without inducing sampling bias that may increase gamma-H2AX scoring variations between experiments. To address this issue we tested random partial samples of different sizes (25, 50, 100 and 250 cells) from two large samples of more than 1500 cells each, which we assumed to be representative of their respective cell populations. We observed that scoring 100 cells was sufficient to provide a CV inferior to 10% when these cells contained at least 150 foci. However, in a non irradiated sample where foci are rarer, at least 250 cells were required for a CV under 10% (with about 70 foci in all) (Figure 3).

After some training, it takes 500 seconds to score 250 cells manually (0.5 cell per second). With HistoLab™, the operator has to load foci images one by one. Nevertheless, this semi-automated approach increased scoring speed to 2.5 cells per second. With CellProfiler, the scoring is done in the absence of the operator and the computers' performance is the only limitation. This approach needs about 30 seconds of operator time to load the parameters file.
This is more than 16 times less than scoring manually if we consider only one sample of 250 cells (8.33 cells per second)
3. Discussion

Gamma-H2AX focus scoring is a highly sensitive technique of DSB quantification and thus of great interest for the development of radiosensitivity and repair capacity assays. In a clinical context, where the user is particularly interested in quickly quantifying the gamma-H2AX signalling response, most attempts of developing a fast H2AX assay used flow cytometry (Ismail et al., 2007). This technique measures a relative intensity of the gamma-H2AX staining instead of scoring the actual number of foci and is known to have a large level of inter-individual variation (Ismail et al., 2007; Hamasaki et al., 2007; Andrievsky and Wilkins, 2009). In manual scoring of gamma-H2AX, the exact number of foci can be obtained increasing the detection sensitivity to the point of detecting doses as low as 0.001 Gy (Rothkamm and Lobrich, 2003). The speed of manual scoring through the eyepieces can be relatively fast for a well-trained scorer, but this approach quickly becomes tiresome if many samples need to be analysed.

The alternative used in this work was to acquire z-stacked images automatically and score on the images. Automatic acquisition is becoming a standard function in many systems of computer-controlled microscopes. The acquisition software we used is relatively common in cytogenetic labs and several alternatives exist, both commercial and open source, like micro-manager (http://micro-manager.org/). Also in the interest of shortening acquisition and analysis time, 250 cells were established as the minimum number of cells to score in order to reach CV values below 10%. With our current configuration, acquisition took under 8 minutes per condition (500 cells acquired to insure the analysis of 250).

The gamma-H2AX scoring results of both HistoLab™ and CellProfiler were similar to manual scoring and sensitive enough to statistically distinguish the sham-irradiated condition from the lowest dose tested (0.05 Gy). Whatever the focus scoring approach used (manual, semi or automatic) similar foci frequencies have been reported: the formation of approximately 12-14 gamma-H2AX foci per cell per Gy found here is consistent with other recent studies on lymphocytes after gamma-rays exposure (Sak et al., 2007; Beels et al., 2010; Roch-Lefèvre et al., 2010).

In terms of scoring speed we were able, after some training, to score manually 0.5 cells per second on images. The automated alternatives (HistoLab™ and CellProfiler) used in this study were at least 5 times faster than manual scoring on the same images. However, these two automatic scoring systems are not equivalent since they vary in terms of “operator time”. The use of HistoLab™ allows the user to associate the unbiased automatic cell and focus
detection with a visual supervision during the manual loading of each image. On the other hand, the analysis step is still dependent on the operators’ availability, increasing the “operator time” needed for the analysis. Our goal was to develop a complete automated image cytometry that requires operator intervention time as low as possible with a good scoring quality. With CellProfiler, the user only takes about 30 seconds to launch the analysis, which will continue in his absence. Moreover, if several conditions need to be analysed several computers can work simultaneously, multiplying the speed analysis by the number of machines available. Furthermore, CellProfiler is a freeware image analysis program that offers a great number of measurement and detection options. This software makes possible to quantify specifically object edge intensity, granularity and roundness. The last has proven to be particularly useful to limit analysis to lymphocytes. This morphology-based lymphocyte selection was validated here by comparison with a CD3 staining. The advantages of choosing lymphocytes in general by their morphology, rather than using specific antibodies, are both economical and practical: economical because general lymphocyte response experiments need no additional antibodies, and practical since this leaves the researcher with a wider set of choices of antibodies that can now be used in conjunction with anti-gamma-H2AX.

The number of free (and paid) automated scoring applications is increasing. All seem to present good correlations to manual scoring once the right parameters are inputted. The free alternative that resembles most HistoLab™ is FociCounter (Jucha et al., 2010): it was developed specifically to score foci, with a simple interface and few parameters to change (faster to set up). These programs are ideal for simple scoring of a reduced number of cells/conditions, where the quality of the images/slides frequently requires operator intervention (to eliminate aberrant objects, for example). Since these programs require user intervention to select nuclei, the time of the analysis is their main disadvantage. CellProfiler is a freeware image analysis program that offers a great number of measurement and detection options. The main drawback of CellProfiler is a less user-friendly graphical user interface that may render the vast number of detection algorithms and measurements overwhelming to a beginner. However, once the pipeline is set up, it requires very little operator intervention for analysis. A free equivalent is difficult to find. We do not compare it to imageJ as a CellProfiler module has been recently created to allow the user to run ImageJ macros and plugins as part of a CellProfiler image processing pipeline (http://cellprofiler.org/CPmanual/RunImageJ.html). In a recent publication, a new image cytometry program that will become freely available was presented (Ivashkevich et al., 2011). The details on the measurements that will be possible to obtain with this software are not yet
disclosed, so we cannot fairly compare it with CellProfiler. However the computational approach they present is a good reference to help researchers select the parameters of other programs of this type (Ivashkevich et al., 2011).

4. Conclusion
In conclusion, the protocol presented here can combine automated acquisition with high-throughput image cytometry to produce a focus score similar to that of manual scoring, with a typical linear correlation with dose. This method has proven to be fast and sufficiently sensitive to detect doses as low as 0.05 Gy. Given our current protocol conditions it is possible to obtain the analysis results of an irradiated blood sample in less than 6 hours. Our automatic scoring technique produced results fast enough to be seriously considered as an alternative to flow cytometry, for studies that need more detailed information about foci number, size or intensity. Moreover, any kind of microscopically visible nuclear domains can be quantified in the same manner. Although our research was more gamma-H2AX foci oriented, its flexibility makes it easy for this method to be adapted to other human lymphocyte-related projects.
References


Figure 1: Indicators used for human lymphocyte enrichment for the gamma-H2AX analysis. A) Cell DAPI intensity and "roundness" (form factor) measurement. The two cells at the middle would not have been considered lymphocytes by the operator during manual scoring because they are less round and have weaker DAPI staining. By quantifying these two parameters, it is possible to establish thresholds that insure that automatic scoring will also exclude them. B) Validation of the morphological selection of lymphocytes with CD3 staining. Within the comet (i.e. excluding cells with DAPI intensity below 1000), different groups of cells were selected by choosing different form factor thresholds: 0.7, 0.8, and 0.86. Choosing fractions of the dot cloud with higher form factor values increases the proportion of CD3-positive cells.

Figure 2: Comparison of the correlation between dose and focus score found by three different scoring methods. The same images from blood samples of three different human donors (three experiments) exposed to 5 IR doses were scored manually by one human operator, semi-automatically by HistoLab™ and automatically by CellProfiler. A minimum of 100 foci and/or 250 lymphocytes were scored for each condition of each donor regardless of the scoring method. The error bars correspond to the standard deviation of the three different donors.

Figure 3: Impact of sample size on the variation of focus score. Focus score results presented here were obtained using CellProfiler. Different sized samples (25, 50, 100, 250 and 500 cells) were run 100 times each for two conditions: sham-irradiated and 0.2 Gy. A box plot that displays the variation of the focus score according to sample size. When 250 cells or more are scored the CV is below 10% for both irradiated and control conditions.

Table 1: Steps for the automated and semi-automated approaches used in this work. Each step includes the parameters for both programs, when applicable. Highlighted in grey are the steps that require operator intervention.
Table 1 - Steps for the automated and semi-automated approaches used in this work.

<table>
<thead>
<tr>
<th>Analysis steps</th>
<th>Cell Profiler</th>
<th>Histolab</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Loading images</td>
<td>The operator loads the pipeline with all images (H2AX and DAPI) parameters and measures.</td>
<td>The operator loads all DAPI images simultaneously</td>
</tr>
<tr>
<td>2. Identifying Nuclei</td>
<td>In the DAPI staining images, identify nuclei (as primary objects) with a diameter between 42 and 90, using an Otsu adaptive threshold.</td>
<td>Fixed threshold: Intensity from 70 to 255 with diameter criteria: 3 to 7 µm.</td>
</tr>
<tr>
<td>3. Loading the foci images</td>
<td>Not applicable (Already loaded in step 1).</td>
<td>The operator loads foci images one by one.</td>
</tr>
<tr>
<td>4. Identifying Foci</td>
<td>Enhance speckles (Top Hat) in the H2AX staining images with a feature size of 19. Crop nuclei (objects) in the enhanced H2AX images. Use the Otsu PerObject threshold method in the cropped H2AX images to identify foci (also as primary objects) between 1 and 50 pixels. Correct the threshold by a factor of 3, between 0.04 and 0.2. Use “Intensity” to distinguish clumped objects and also to draw the dividing lines.</td>
<td>For each field the operator chooses “foci detection” and clicks on the button “identify objects”. Foci detection parameters: Adaptive threshold (Top Hat) Bright tones. Maximum diameter: 0.989 µm. Minimum contrast: 45. Filtering: None. Separation: Fine. Criteria: Minimum Object diameter equal or inferior to 1.5 µm. Maximum Object diameter equal or superior to 0.24 µm.</td>
</tr>
<tr>
<td>6. Filtering Cells of interest</td>
<td>If necessary, the operator may filter only cells with a Form Factor above 0.86 to enrich in lymphocytes.</td>
<td>Not applicable (Selection already made in step 2)</td>
</tr>
<tr>
<td>7. Export data to spreadsheet</td>
<td>The module “export to spreadsheet” exports all data at the end of the analysis.</td>
<td>Data exported manually as a text file.</td>
</tr>
</tbody>
</table>
Number of foci per cell

0.2 Gy

CV: 5.8%  CV: 3.2%  CV: 3.4%

0 Gy

CV: 15.7%  CV: 7.8%  CV: 3.7%

Number of cells scored

25 Cells  50 Cells  100 Cells  250 Cells  500 Cells
# List of abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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</thead>
<tbody>
<tr>
<td>ANOVA</td>
<td>ANalysis Of VAriance</td>
</tr>
<tr>
<td>A-T</td>
<td>Ataxia Telangiectasia</td>
</tr>
<tr>
<td>ATM</td>
<td>Ataxia Telangiectasia Mutated</td>
</tr>
<tr>
<td>ATR</td>
<td>ATM- and Rad-3 related</td>
</tr>
<tr>
<td>BARD1</td>
<td>BRCA1-Associated RING Domain 1</td>
</tr>
<tr>
<td>BLM</td>
<td>BLooM’s syndrome protein</td>
</tr>
<tr>
<td>53BP1</td>
<td>p53 Binding Protein 1</td>
</tr>
<tr>
<td>BRCA1/2</td>
<td>BReast CAncer 1/2</td>
</tr>
<tr>
<td>CCD</td>
<td>Charge-Coupled Device</td>
</tr>
<tr>
<td>Chk1/2</td>
<td>Checkpoint Kinase 1/2</td>
</tr>
<tr>
<td>CTC(AE)</td>
<td>Common Toxicity Criteria (for Adverse Events)</td>
</tr>
<tr>
<td>2D</td>
<td>2 Dimensions</td>
</tr>
<tr>
<td>DAPI</td>
<td>4‘,6-DiAmidino-2-Phenylindole</td>
</tr>
<tr>
<td>DDR</td>
<td>DNA Damage Response</td>
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<tr>
<td>DNA</td>
<td>DeoxyriboNucleic Acid</td>
</tr>
<tr>
<td>DNA-PKcs</td>
<td>DNA-dependent Protein Kinase catalytic subunit</td>
</tr>
<tr>
<td>DSB</td>
<td>Double- Strand Break(s)</td>
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<tr>
<td>EFS</td>
<td>Etablissement Français du Sang</td>
</tr>
<tr>
<td>eV</td>
<td>electron Volt</td>
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<tr>
<td>FISH</td>
<td>Fluorescence In Situ Hybridization</td>
</tr>
<tr>
<td>FITC</td>
<td>Fluorescein IsoThioCyanate</td>
</tr>
<tr>
<td>G0/1/2</td>
<td>Gap 0/1/2 phase of the Cell Cycle</td>
</tr>
<tr>
<td>Gamma-H2AX</td>
<td>Histone subtype H2A isoform X phosphorylated at ser-139</td>
</tr>
<tr>
<td>Gy</td>
<td>Gray</td>
</tr>
<tr>
<td>HNSCC</td>
<td>Head and Neck Squamous Cell Carcinoma</td>
</tr>
</tbody>
</table>
HR - Homologous Recombination
HRS - Hyper-RadioSensitivity
Hz - Hertz
IgH - Immunoglobulin Heavy
IR - Ionizing Radiation(s)
IRIF - Ionizing Radiation Induced Focus
IRSN - Institut de Radioprotection et de Sûreté Nucléaire
keV - kilo electron Volt
LET - Linear Energy Transfer
Lig4 - Ligase 4
LNT - Linear-Non-Threshold
M - Mitosis phase of the Cell Cycle
MDC1 - Mediator of DNA damage Checkpoint 1
Mdm2 - Murine double Minute 2
MeV - Mega electron Volt
µg - micro (µ) gram
µm - micro (µ) meter
MRN - Mre11/Rad50/NBS1 complex
mSv - mili Sieverts
NBS1 - Nijmegen Breakage Syndrome protein 1
NHEJ - Non-Homologous End Joining
nm - nano meter
PARP-1 - Poly(ADP-Ribose) Polymerase 1
PBL - Peripheral Blood Lymphocytes
PBS - Phosphate Buffered Saline
PFA - ParaFormAldehyde
PHA - PhytoHaemAgglutinin
PS - PhosphatidylSerine
PIKK - Phosphoinositide three-Kinase-related protein Kinases
RAD - Radiation Absorbed Dose
RBE - Relative Biological Effectiveness
RNS - Reactive Nitrogen Species
ROS - Reactive Oxygen Species
RPA - Replication Protein A
RPMI - Roswell Park Institute Medium
RT - Room Temperature
S - Synthesis phase of the Cell Cycle
SD - Standard Deviation
SEM - Standard Error of the Mean
SMC1 - Structural Maintenance of Chromatine 1
SSB - Single-Strand Break(s)
ssDNA - single stranded DNA
Sv - Sieverts
V(D)J - Variable (Diverse) Joining
v/m - volume/mass
XLF - XRCC4-Like Factor
XRCC4 - X-ray Repair Complementing defective repair in Chinese hamster cells