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Gamma-H2AX biodosimetry for use in large scale radiation incidents: comparison of a rapid lyse/fix protocol with a routine method

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Following a radiation incident, preliminary dose estimates made by γ-H2AX foci analysis can supplement the early triage of casualties based on clinical symptoms. Sample processing time is important when many individuals need to be rapidly assessed. A protocol was therefore developed for high sample throughput that requires less than 0.1 ml blood, thus enabling finger prick sampling. The technique combines red blood cell lysis and leukocyte fixation in one step on a 96 well plate, in contrast to the routine protocol, where lymphocytes are separated by Ficoll density gradient centrifugation with subsequent washing and fixation steps. The rapid lyse/fix method reduced the estimated sample processing time for 96 samples to about 4 h compared to 15 h using the routine protocol. However, scoring 20 cells in 96 samples prepared by the rapid protocol took longer than for the routine method (3.1 versus 1.5 h at zero dose; 7.0 versus 6.1 h for irradiated samples). Similar foci yields were scored for both protocols and reliable dose estimates were obtained for coded samples, with mean absolute differences from the actual doses of 0.26 and 0.27 Gy for the routine and lyse/fix method, respectively. The lyse/fix protocol can therefore facilitate high throughput processing for γ-H2AX biodosimetry for use in large scale radiation incidents, at the cost of somewhat longer foci scoring times.
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INTRODUCTION

The γ-H2AX assay has been shown to be a reliable and sensitive indicator of radiation-induced DNA double-strand breaks (Rothkamm & Löbrich, 2003; Ivashkevich et al, 2012). In addition, several ex vivo studies (Horn, Barnard & Rothkamm, 2011; Horn & Rothkamm, 2011; Roch-Lefevre et al, 2010; Mandina et al, 2011), have shown the potential of the γ-H2AX assay as a useful biodosimetry tool from hours to ~3 days post exposure. The γ-H2AX assay can produce dose estimates within a few hours of receiving a blood sample (Rothkamm et al, 2013a; Ainsbury et al, 2013) making it a good candidate for high through-put biodosimetry in the case of a mass casualty event. One option has been to develop fully automated systems such as the RABIT system (Garty et al, 2010; Turner et al, 2011) capable of processing several thousand samples a day. A drawback of this approach is that it requires highly sophisticated robotic equipment and tailored process control software, which may be unavailable to some laboratories. An alternative option to increase through-put is to have an assistance network of laboratories, such as those involved in the European funded MULTIBIODOSE and RENEB projects, and to reduce the processing time required for the γ-H2AX assay using routinely available equipment.

Here a method is presented for the assessment of γ-H2AX foci for biological dosimetry, that uses a lyse/fix buffer to lyse red blood cells and fix leukocytes in one step from small ‘finger prick’ blood samples. Potentially, the lyse/fix method would be a faster and more scalable technique for high sample through-put compared to the routine protocol used to process samples for γ-H2AX foci scoring, which involves the isolation of lymphocytes from at least 2 mL of blood using Ficoll density gradient centrifugation with subsequent washing and fixation steps.

MATERIALS AND METHODS

Blood sampling and irradiation

Heparinised venous blood was taken with written informed consent and the ethical approval of the Berkshire research ethics committee (Ref 09/H0505/87) from a panel of healthy donors. Whole blood was exposed to 0.5 or 1.0 Gy of 250 kVp x-rays (with Cu/Al filtration) at a dose rate of 0.5 Gy / minute (min). Zero radiation dose control samples were also included. Following irradiation, the blood was held at 37 °C for 0.5 or 1 hour to simulate in vivo repair. Samples were then processed for the assessment of γ-H2AX foci by the routine histopaque or rapid lyse/fix method. A record was made of the time taken for each step in the process. To test the suitability of the lyse/fix method for biological dosimetry, samples were exposed to x-ray doses up to ~4 Gy and incubated for 2 hours at 37 °C. Coded samples were processed using both methods, scored and used to produce estimates of dose. Positive and negative reference samples (0 and 1 Gy) were also included.

Rapid lyse/fix method

Aliquots of 90 µL whole blood were placed into a 96 deep well plate. To each well, 1x lyse/fix buffer (BD Biosciences, UK) pre-warmed to 37 °C was added to give a dilution of 1:20, thoroughly mixed and incubated at 37 °C for 14 min. After centrifugation at 250 g for 5 min the supernatant was removed using an 8 channel aspirator. Following two washes in cold phosphate buffered saline (PBS) the cell pellet was re-suspended in ~50 µL of PBS. An aliquot of 25 µL per sample was spotted onto a teflon/silane coated 14-well slide (Tekdon Incorporated, USA) and the cells were allowed to adhere for ~15 min. When the slides were almost dry they were placed in a moist chamber and incubated in 0.3% Triton-X (Sigma, UK) in PBS for 5 min. The slides were
then drained and incubated for 10 min in blocking solution (BS), containing 1% bovine serum albumin (Sigma, UK) in PBS. Once any excess liquid had been removed, 100 µL of mouse γ-H2AX antibody (BioLegend, UK), diluted 1:500 in BS, was applied to each slide. After incubating the slides in the dark for 40 min at room temperature they were washed 3 times in BS. Slides were then incubated in the dark with 100 µL of goat anti-mouse AlexaFlour 488 secondary antibody (Abcam, UK), diluted 1:500 in BS, together with 0.2 µg/mL 4',6-diamidino-2-phenylindole (DAPI) for 30 min at room temperature. Finally the slides were washed three times in PBS and mounted in Vectashield anti-fade solution (Vector Laboratories, UK). A detailed description of the lyse/fix protocol, sample requirements and materials can be found in Supplementary material: A Rapid Protocol for γ-H2AX Processing.

Routine histopaque method
The histopaque method has been described in detail by Rothkamm et al (2013b). In brief, 2 mL of whole blood was mixed 50:50 with PBS and layered onto 4 mL Histopaque 1077 (Sigma, UK). After centrifugation at 1200 g for 5 min, the buffy coat layer was washed twice in PBS and the cell pellet re-suspended in ~0.1 mL of PBS. 10 µL of the cell suspension was spotted onto a multi-well slide. Cells were allowed to adhere to the slide for ~15 min and then fixed in 2% formaldehyde in PBS for 5 min, followed by 5 min in 0.25% Triton-X100 + 0.1% glycine in PBS. At this stage the blocking step and the immuno-staining was carried out in the same manner as the lyse/fix method described above.

Manual scoring of γ-H2AX foci
All the slides were examined at x600 magnification using a Nikon Optiphot 2 fluorescence microscope, equipped with separate filters for DAPI and fluorescein isothiocyanate (FITC). Manual scoring was timed for three unirradiated and three 1 Gy/1 h incubated samples on slides produced by both processing methods. A total of 50 lymphocytes were scored per sample and the time taken to do this was recorded every ten cells. To ensure the lyse/fix processing itself did not affect foci levels, samples irradiated at 0.5 Gy/repair time 30 minutes and prepared using both protocols were used to assess the number of foci per cell seen in a panel of donors. 50 lymphocytes were scored in each of the reference samples and the foci numbers were used to adjust the calibration coefficients and the associated standard errors (Rothkamm et al, 2013b) of the laboratory’s calibration curve (Horn, Barnard & Rothkamm, 2011). Whole body dose estimates for the unknown samples were produced by scoring up to 50 lymphocytes or 200 foci per sample.

Automated scoring of γ-H2AX foci
The performance of automated scoring was also assessed on samples processed by the two methods. For each processing method and the two irradiation conditions, 10 fields of view were scored automatically. Maximum projection images of seven z planes at 1 µm step size were captured for γ-H2AX and DAPI staining (only one central plane imaged) using a x40 objective. Foci analysis was performed with the TRI2 program, which contains batch processing functionality and automatic sorting of images. The method used for foci counting and analysis was the Compact Hough and Radial Map (CHARM) algorithm aimed at faint and ill-defined shapes (Barber et al, 2007). The foci analysis software has been fully described in Rothkamm et al. (2012).

RESULTS AND DISCUSSION

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The time taken to process 16 samples using the lyse/fix protocol and 8 samples by the histopaque method is very similar, 179.5 min and 171 min respectively (Figure 1). The time for each step in both protocols has been recorded and an estimate made of the timings for handling of 96 samples. Scaling up the timings for preparing 96 samples by the lyse/fix or histopaque method gives an increase in the estimated processing time of a factor of ~1.4 (256 min total) and ~ 2.5 (439 min total) respectively. Most of the difference between the two protocols is the time taken to process the samples through the first 3 steps. Provided that multi-channel pipettes and aspirators for liquid handling are used, one person could complete the first three steps of the lyse/fix protocol for 96 samples in 72 min. If one person were to process 96 samples by the histopaque method up to and including step 3 this would take more than three times as long (226 min). In reality, given the long processing time required for one person to get 96 samples to step 4 of the histopaque protocol, which may adversely affect the blood separation or lower foci levels, either several persons would need to work together or the samples be divided into batches. Assuming four batches of 24 samples each, the histopaque processing time for 96 samples would be 880 minutes (14.7 h) for one operator (Figure 1).

Usually in the routine histopaque protocol cell suspensions are spread onto individual slides, but for a large number of samples this would be time consuming. For the purposes of this comparison, cells prepared by both techniques were spotted onto 14-well slides. The time taken to complete steps 5 to 11 would be similar for both protocols, 169 (lyse/fix) and 198 min (histopaque). Multi-well slides also enable one person to carry out the protocol after lysis and fixation, as the time between the addition of a solution to the first well and the last is not so great as to adversely affect any step of the staining protocol. However, this is not the case for the histopaque protocol, which requires cells to be fixed after they have been placed on the slide, and this is a time critical step. Again, several persons would be needed to process all the slides together; alternatively they could be stained in batches by one person.

Histopaque separation of whole blood produces a cell suspension containing lymphocytes. In contrast, the lyse/fix protocol produces a cell suspension made up of lymphocytes and other white cell types, which can make foci analysis more challenging. Figure 2 shows images of one field of view of cells processed by the lyse/fix and histopaque methods from irradiated and unirradiated samples. By comparing the images it can be seen that the lyse/fix method gives fewer lymphocytes per field of view (i.e. round cells) and more debris. In addition the foci are not as clearly visible as in samples processed by the histopaque method. Despite differences in the appearance of samples produced by the two methods, manual scoring produced similar foci yields in lymphocytes from a panel of donors exposed to 0.5 Gy x-rays, as shown in Figure 3. Samples processed by the histopaque and lyse/fix method gave similar average foci counts of 6.83 ± 0.80 and 7.12 ± 0.51 respectively. These values are consistent with previous studies (Horn, Barnard & Rothkamm, 2011; Chua et al, 2011; Rothkamm et al, 2007).

In order to estimate the time required to analyse 96 samples, three slides for each processing method and dose point were scored. For each slide the time taken to score a batch of 10 cells was recorded and then repeated a further 4 times. The average time taken to score each batch of 10 cells from the three slides was then used to estimate the time it would take one person to score 20 cells from 96 samples. Previous work has demonstrated that scoring 20 cells can produce dose estimates that will reliably place samples into the correct exposure categories required for triage (Rothkamm et al, 2013a). As expected the mean time taken to score a 0 Gy sample is less than for a 1 Gy sample at 1 h post exposure for both preparation methods. However, for the 0 Gy sample the estimated time taken to score 20 cells in 96 samples prepared by the lyse/fix protocol was
about twice as long as for those produced by the histopaque method; 3.1 h compared to 1.5 h. At
the higher dose of 1 Gy the difference in scoring time between the two processing methods was
not as great, (7.0 h compared to 6.1 h).

Simply comparing the lyse/fix protocol with the histopaque method for processing and scoring all
96 samples suggests the time taken would be similar for both irradiation conditions, 7.4 h versus
8.8 h (unirradiated) and 11.3 h versus 14.4 h (irradiated) respectively. However, the histopaque
protocol requires at least two people to achieve this compared to one for the lyse/fix method. A
more realistic comparison involves the 96 samples being split into 4 batches for the histopaque
method to be carried out by one person. When this is done the estimated time taken to process
and score 96 samples by the histopaque method is 16.2 h (unirradiated) and 20.8 h (irradiated).
This is approximately twice the time required for sample processing and scoring using the
lyse/fix protocol. Furthermore, reagent costs for the histopaque method (~£116/96 samples) are
twice as high as those for the lyse/fix protocol (~£57/96 samples).

The more challenging analysis procedure required for samples produced by the lyse/fix method
may be aided by an automated scoring approach. Image analysis software using a form factor
parameter would allow the positive discrimination of round lymphocyte nuclei, thereby enriching
their number in the analysed cell population (Valente et al, 2011). Here, TRIL2 software was used
to identify round cell nuclei and score foci automatically (Rothkamm et al, 2012). Figure 4 shows
the automated scoring results of 10 fields of view for each processing method and the two
irradiation conditions. Automated foci counts are comparable between methods, although ten
times more scorable cells per field of view are seen with the histopaque protocol. The lyse/fix
protocol produced 3 false negative fields out of 10 compared to 1 for the histopaque method,
likely caused by high background fluorescence due to excessive debris in lyse/fix samples. These
results suggest that if automated scoring were to be used to analyse samples, at least 2
(histopaque) and 3 (lyse/fix) fields of view need to be scored and results checked to ensure
consistency. Further optimisation of the software may reduce the number of false negative fields.

To determine whether the rapid lyse/fix protocol can still provide dose estimates with similar
accuracy as the routine histopaque protocol, an intercomparison exercise was performed using
uniformly X-irradiated, coded samples. Figure 5 shows that the dose estimates obtained for the
two different protocols correlated very well with each other. Mean absolute differences from the
actual doses were 0.26 and 0.27 Gy for the histopaque and lyse/fix protocols, respectively.

CONCLUSION

The lyse/fix protocol enables a high sample throughput, with an estimated processing time for 96
donors of just over 4 hours, using finger prick-sized quantities of blood. This can be achieved by
one person, without the need for highly sophisticated equipment. The manual microscope based
analysis of samples produced by the lyse/fix protocol is more challenging, which increases the
scoring time, but has no adverse effect on dose estimation. Overall the rapid lyse/fix method
allows one person to process and score 96 samples in about half the time taken using the routine
protocol.

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Figure 1

Sample processing times for the lyse/fix versus the routine histopaque method for different numbers of samples.

Steps 1 – 11: 1 = lyse/fix or isolation; 2 = washing; 3 = transfer; 4 = adhere; 5 = permeabilise (and fixation for histopaque method); 6 = blocking; 7 = stain1; 8 = wash; 9 = stain2; 10 = wash; 11 = mount.
**Figure 2**

γ-H2AX foci in human blood leukocytes prepared with the lyse/fix or histopaque method.

Fluorescence microscopic maximum projection images (x40 objective) show γ-H2AX foci (green) and DNA counterstain (blue) following exposure to 0 or 1 Gy X-rays and incubation for 1 hour.
Figure 3

Comparison of manual γ-H2AX foci counts.

Different volunteer blood samples were processed by the lyse/fix or histopaque method following 0.5 Gy X-irradiation and 30 minutes incubation at 37 °C.
Figure 4

Comparison of automated $\gamma$-H2AX foci counts.

Ten fields of view were analysed using Tri2 software (Barber et al, 2007) for each processing method and two irradiation conditions (0 or 1 Gy plus 1 hour incubation at 37 °C). Open triangles: 0 Gy + lyse/fix; solid triangles: 1 Gy + lyse/fix; open squares: 0 Gy + histopaque; solid squares: 1 Gy + histopaque.
Figure 5

Comparison of whole body $\gamma$-H2AX dose estimates.

Foci yields were determined for X-irradiated and 2 hours incubated blood samples processed with the lyse/fix versus histopaque protocol. At least 50 cells or 200 foci were manually scored per coded sample. Foci yields were converted to dose estimates using calibration data from Horn et al, (2011). Actual doses were 0, 0.2, 0.6, 1.1, 1.2, 2.1 and 4.3 Gy. The line indicates the ideal 1:1 relationship.