Methodology article

A modified and automated version of the 'Fluorimetric Detection of Alkaline DNA Unwinding' method to quantify formation and repair of DNA strand breaks

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Abstract

Background: Formation and repair of DNA single-strand breaks are important parameters in the assessment of DNA damage and repair occurring in live cells. The 'Fluorimetric Detection of Alkaline DNA Unwinding (FADU)' method [Birnboim HC, Jevcak JJ. Cancer Res (1981) 41:1889–1892] is a sensitive procedure to quantify DNA strand breaks, yet it is very tedious to perform.

Results: In order (i) to render the FADU assay more convenient and robust, (ii) to increase throughput, and (iii) to reduce the number of cells needed, we have established a modified assay version that is largely automated and is based on the use of a liquid handling device. The assay is operated in a 96-well format, thus greatly increasing throughput. The number of cells required has been reduced to less than 10,000 per data point. The threshold for detection of X-ray-induced DNA strand breaks is 0.13 Gy. The total assay time required for a typical experiment to assess DNA strand break repair is 4–5 hours.

Conclusion: We have established a robust and convenient method measuring of formation and repair of DNA single-strand breaks in live cells. While the sensitivity of our method is comparable to current assays, throughput is massively increased while operator time is decreased.

Background

Formation and repair of DNA single-strand breaks in live cells is an important functional parameter in the assessment of genotoxicity, and therefore the reliable and convenient assessment of DNA strand breakage is of crucial importance for a wide range of basic and translational biological research fields including genetic or environmental toxicology (mechanistic research; screening for genotoxins); DNA repair research (basic biochemical studies; molecular epidemiology); cancer research; pharmacology and drug development; ageing research and many more. The measurement of DNA strand breaks by
FADU as described by Birnboim & Jevcak [1] is based on the partial denaturation ("unwinding") of double-stranded DNA under controlled alkaline conditions. Briefly, after infliction of DNA damage, cell lysis was performed. DNA strand breaks are sites where controlled unwinding of DNA can start under controlled conditions of pH and temperature. To terminate unwinding, a neutralising solution was added. To quantify the amount of DNA remaining double-stranded after the alkali incubation, ethidium bromide was added as a fluorescent probe. Low fluorescence intensities indicated a large number of DNA strand breaks present at the time of lysis. In practical terms, the following samples are processed in parallel: T samples are representative of the total amount of DNA present and are obtained by adding neutralisation solution prior to the alkaline unwinding solution. As a consequence, the critical alkaline pH needed for DNA denaturation will never be reached and no unwinding will occur. On the other hand, P₀ samples do undergo alkaline unwinding at the endogenous DNA strand breaks and at replication forks, thus reflecting physiological conditions. In contrast, in B samples DNA will denature completely due to the massive induction of DNA breaks, e.g. by sonication or shearing, and therefore only background fluorescence is observed. P₁ samples (P₀, P₂, P₃ etc) are samples in which to measure DNA damage-induced DNA strand breaks, and R₁ samples (R₀, R₂, R₃ etc) are samples where cells are post-incubated to allow repair.

Since the original publication by Birnboim & Jevcak [1], a number of assay modifications have been described that led to reduction of the numbers of cells needed and faster completion of the assay [2-7]. The modified and automated version of the FADU assay described in the present paper enables measurement of DNA strand breaks and DNA repair in a very reliable and convenient manner by exploiting the power of a liquid handling device (LHD) in terms of its extremely high level of control of various parameters and the reproducibility of dispensing small volumes. In particular the transfer at sub-millimetre precision within the three dimensions of the LHD’s workspace as well as precise control of timing and rate of liquid delivery are extremely useful features, as the creation of a separate layer of alkaline solution on top of the lysis buffer without any mixing is critical [cf. [1]]. Protecting the samples from light by encasing the LHD also greatly adds to assay sensitivity, as the genomic DNA liberated from histones by the urea present in the lysis buffer is prone to DNA breaks induced artificially by visible light.

**Methods**

**Plasmid**

For a set of pilot experiments we used the plasmid pEGFP-N1 (size: 4.7 kb; BD Biosciences, Heidelberg, Germany). The plasmid was restricted or not with EcoRI (NEB, Frankfurt am Main, Germany). To obtain linear DNA, 8 μg of plasmid DNA was digested in 60 μl distilled H₂O, 8 μl 10× buffer and 4 μl EcoRI at 37°C for 1 h. To stop digestion the plasmid solution was incubated at 65°C for 20 min. Completeness of the digestion was verified by agarose gel electrophoresis.

**Primary cells**

Ethical clearance for obtaining peripheral venous blood from healthy adult volunteers was obtained from the University of Konstanz Ethics Committee. Peripheral blood mononuclear cells (PBMC) were purified from freshly drawn blood. All donors were healthy adult volunteers. For the separation of PBMC we used Percoll (Pharmacia Biotec AB, Uppsala, Sweden) gradient centrifugation. Briefly, 7 ml of whole blood was drawn with heparin as anti-coagulant. The blood was diluted 1:2 with PBS, layered on the top of 15 ml Percoll solution and centrifuged at 800 g for 10 min at room temperature in a swing-out rotor without break. Mononuclear cells harvested from the gradient were washed by resuspension in PBS and centrifugation at 200 g for 10 min at 0°C.

**Cell lines**

Jurkat T cells were cultured in suspension in tissue culture flasks (Corning, Schiphol-Rijk, The Netherlands) in RPMI medium (Sigma-Aldrich, Deisenhofen, Germany) supplemented with 10% FCS and antibiotics (100 U/ml penicillin, 100 μg/ml streptomycin).

HeLa cells were cultured as monolayer in non-pyrogenic tissue culture flasks (Corning, Schiphol-Rijk, The Netherlands) in DMEM medium (Sigma-Aldrich, Deisenhofen, Germany) supplemented with 10% FCS and antibiotics. Prior to any analyses by FADU assay, monolayers were trypsinized, and detached cells were resuspended in complete medium (DMEM plus FCS plus antibiotics), followed by centrifugation at 200 g for 10 min. Pellets were resuspended in DMEM without FCS (see below, 3 – DNA Repair).

**96-well plate**

A 96-well plate master block (2 ml) with V-shaped bottom (Greiner-Bio One GmbH, Frickenhausen, Germany) was used, which was cut horizontally to hold 1 ml in order to enable measurement of the fluorescence intensity in a commercial fluorimeter for standard plates.

After use, the 96-well plates were washed with 1 M NaOH for 1 h, rinsed with water, incubated in DMSO (Merck, Darmstadt, Germany) for 5 min, once again rinsed with water and finally washed in 70% ethanol for 5 min.
Liquid handling device and accessories
For the automated steps of the new FADU protocol (see below) a TECAN Genesis RSP 100 liquid handling device (LHD) from TECAN AG (Hombrechtikon, Switzerland), controlled by GEMINI V4.0 software from TECAN software GmbH (Berlin, Germany) was used.

For the precise positioning of the cells in the X-irradiation device, a custom-made ice box, made of plastic and comprising an aluminium mould, was used. It can accommodate a 96-well-plate or Eppendorf tubes and maintains the samples at 0°C. This box can also be inserted at a defined position in the working area of the LHD to allow transfer of the samples to a custom-made cooling device mounted in the LHD’s working space. The cooling device accommodates a 96-well plate and is connected to a cooling water bath for temperature control.

The eight nozzles of the LHD find their target with regard to X, Y, Z coordinates with a precision of 0.2 mm. Solutions to be transferred by the LHD are stored in trays, and those for ice-cold solutions are placed in a temperature-controlled holder. All buffers are accessible to the nozzles from the top. For protection against light the LHD is accommodated in a closed box with a roller shutter at its front.

Steps of the modified and automated FADU protocol
This new version of the FADU assay comprises a total of 11 steps described below. It should be noted that steps 2 and 3 are alternatives, depending on whether or not DNA repair is to be analysed. Steps 1, 2, 3, 10 and 11 were performed manually whereas the critical steps 4–9 have been automated (Fig. 1A). It should be noted that in some experiments performed for protocol optimisation (Fig. 2A), different conditions were used, as indicated.

*T samples* represent the total amount of DNA present.

*B samples* represent the background fluorescence observed when DNA is denatured completely.

*P*₀ samples reflect DNA strand interruptions present under physiological conditions. *P*₁ samples (P₃, P₂, etc) are from cells to which DNA damage had been inflicted in order to induce DNA strand breaks, and *R*₂ samples (R₁, R₂, R₃ etc) are from cells allowed to repair DNA damage.

Steps 1 and 2 – Preparation of Cells and DNA Damage by Irradiation
For the analysis of DNA strand break formation only (without repair) cells were resuspended in suspension buffer (0.25 M meso-inositol; 10 mM sodium phosphate, pH 7.4; 1 mM magnesium chloride) or in medium without FCS. For the irradiation we used an X-ray generator from CHF Müller (Hamburg, Germany). The dose to be delivered was applied by variation of irradiation time at a fixed dose rate. The irradiation parameters were the following: 70 keV energy, 1.25-mm aluminium filter, and 9.4 mA current. The dose was measured using a PTW Universal Dosimeter UNIDOS E [D545.151.00/02] from PTW-Freiburg (Freiburg, Germany). Seventy µl of cell suspension (at 100 cells/µl) in a 96-well plate was irradiated on ice in a custom-made ice box described above.

Steps 3 and 4 – DNA Repair and Cell Transfer
For the assessment of DNA repair, cells were resuspended in DMEM without FCS or other supplements at a concentration of 6 × 10⁵ cells per ml. Aliquots of 100 µl were cooled down in Eppendorf tubes and irradiated on ice. To allow DNA strand break repair, the damaged cells were incubated at 37°C for various time periods in an incubator. To stop repair 500 µl ice-cold suspension buffer was added to 100 µl cell suspension, respectively. Seventy µl/well of this cell suspension (i.e. 7,000 cells) were transferred to the 96-well plate, which had been placed in the cooling device described above.

Step 5 – Lysis
The 96-well plate holding the 70 µl cell samples was positioned in the working space of the robot and kept at 0°C in the dark. Then automated dispensing of 70 µl of lysis solution (9 M urea; 10 mM NaOH; 2.5 mM cyclohexylidiamine-tetraacetate; 0.1% sodium dodecyl sulphate; kept at RT) to each sample at a rate of 150 µl/s was initiated.

Step 6 – Alkaline solution
After 12 min of lysis (temperature of cooling device set to 0°C), 70 µl of alkaline solution (0.425 parts lysis solution in 0.2 M NaOH, pre-cooled to 0°C) was added on the top of the cell lysate in such a way as to form a second layer, thus avoiding any mixing with the lystate. To do this the nozzles of the LHD were positioned precisely 1.5 mm above the level of the lystate and the alkaline solution was dispensed at the very low rate of 10 µl/s. Then diffusion of alkaline solution into the lystate was allowed to occur for 15 min. Temperature readings showed that the content of the microwells displayed approximately 8°C during this time period. A pH value of 12.5 was reached under these conditions. Thereafter the temperature setting of the cooling device was shifted to 30°C and kept constant for 60 min in order to allow alkaline unwinding of the DNA.

Step 7 – Neutralization solution
Prior to the addition of 140 µl of neutralization solution (14 mM β-mercaptoethanol; 1 M glucose) at a rate of 200 µl/s the temperature was shifted to 22°C. For *T*-samples (see below), an internal standard representing cells with 100% double-stranded DNA, 140 µl of neutralization solution was added prior to the alkaline solution.
Steps 8 and 9 – SybrGreen® addition and mixing
To determine the amount of double-stranded DNA, SybrGreen (10,000×) from MoBiTec (Göttingen, Germany) was used. After dispensing 156 μl diluted SybrGreen (1:8,333 in H2O), respectively, the samples were mixed by pipetting a volume of 400 μl up and down once at a rate of 100 μl/s.

Steps 10 and 11 – Read-out and interpretation of data
Samples were analysed in a 96-well-plate fluorescence reader at 492 nm excitation/520 nm emission immediately after SybrGreen addition. Statistical significance was evaluated by using Student’s t-test. In some experiments B-samples were included as an internal standard representing lysates with no double-stranded DNA left after alkali exposure as a result of extensive DNA breakage induced by passing the lysate 20 times through a 0.5 mm cannula. In course of the work it was noted that, using SybrGreen as a fluorescent probe, there was a stable ratio between T and B values, and since the B values were quite low, B samples were omitted in the final FADU protocol.
Results

Optimisation of the FADU protocol

In the original FADU protocol, Birnboim & Jevcak had used ethidium bromide as a fluorescent probe to quantify the amount of DNA that remained double-stranded under the experimental conditions chosen [1]. When establishing our modified protocol, we first compared SybrGreen, a more recently developed fluorescent compound with high specificity for double-stranded DNA, with ethidium bromide. Our data indeed showed that SybrGreen produced much less background fluorescence (Fig. 2A) and thus increases assay sensitivity. Therefore for all subsequent analyses SybrGreen was used.

To characterise further the alkaline unwinding process we used purified DNA of plasmid pEGFP-N1, which was linearised or not with EcoRI, as a model substrate. While covalently closed circular plasmid DNA did not undergo unwinding under our assay conditions as revealed by the high P0 value, the linear form of the plasmid could unwind starting from its ends. Our data showed that only about 20% of DNA remained double-stranded (Fig. 2B). Agarose gels revealed that undigested plasmid preparations typically comprised a small fraction of nicked circular DNA (data not shown). This form of DNA can unwind under the conditions chosen, and this explains the slightly lower fluorescence signal of the circular DNA P0 samples compared to T values.

To assess whether the time interval between SybrGreen addition and fluorescence reading is critical, we systematically measured fluorescence as a function of time after SybrGreen addition. All the samples showed saturation immediately after addition of SybrGreen. The fluorescence signal remained constant during 20 min. We conclude that the fluorescence determination yields stable results if readings are performed within 20 min after SybrGreen addition (Fig. 3).

In order to determine the minimal number of cells required, we irradiated cells at three different cell titres. The ratio of fluorescence signals of P0/T and P1/P0 using 297, 132 and 90 cells/μl varied less than 2.4% and of P2/P1 less than 7%, indicating that the automated FADU assay yields robust results over a range of cell titres. Interestingly, standard deviations tended to be smaller with lower cell titres (Table 1).

Definitive assay protocol

The essential steps of the definitive assay protocol for the modified automated FADU assay are summarised in Fig. 1A and the detailed protocol is described in Materials and Methods.
To assess assay reproducibility we measured the dose-dependent unwinding induced by X-irradiation of cells in several independent experiments. DNA unwinding increased with increasing radiation dose in all experiments, and the results were highly reproducible, both in a high (Fig. 1B) and in a low irradiation-dose range (Fig. 1C). The lowest dose detected with statistical significance was 0.13 Gy (Fig. 1C).

We also tested the applicability of the automated FADU assay in several different cell systems. Induction of DNA strand breaks and repair were measured in PBMC and two human cell lines, i.e. the T-cell derived cell line Jurkat and the cervical carcinoma cell line HeLa. To measure DNA repair, the damaged cells were post-incubated at 37°C for 40 min. During this time period cells were able to repair the DNA breaks to the level of P₀ samples, as is apparent from the recovery of SybrGreen fluorescence (Fig. 4). Analysis of the time course of repair in 5-min intervals revealed the steady progression of repair with time (Fig. 5).

Table 1: Influence of cell titre on fluorescence signal intensity.

| Cells/μl | P₀/T | P₁/P₀ | P₂/P₁ | Standard deviation [%] |
|----------|------|-------|-------|------------------------|
| 297      | 0.82 | 0.94  | 0.77  | 5.5                    | 10.6 | 12 | 12.7 |
| 132      | 0.84 | 0.96  | 0.68  | 3.3                    | 4.5  | 4.9 | 5.4  |
| 90       | 0.86 | 0.95  | 0.71  | 2.9                    | 4.6  | 4.2 | 3.2  |

P₁ and P₂ samples were irradiated with 0.47 or 2.3 Gy, respectively. Standard deviation was evaluated for 8 replicates, respectively, and expressed as % of mean values.
Discussion

A sensitive, robust and convenient assay for the detection and quantification of DNA damage and repair is useful not only for basic scientific research on DNA damage and DNA repair, but also for medical diagnostics (for example to assess individual genotoxic exposures and biological responses) and toxicological screening in the chemical and pharmaceutical industry.

The two current methods to measure DNA damage and repair with a high sensitivity are the "comet" assay and the conventional, manually performed FADU [1]. Detection limits of 0.03 Gy for the comet assay and 0.1 Gy for the FADU assay have been reported [8]. The principles of these two kinds of assays differ. The comet assay is based on different migration velocities of DNA fragments of different sizes in agarose gels. In an electrical field, fragmented DNA migrates out of the nucleus into the agarose gel and can be visualised by staining with a fluorescent dye. Viewed microscopically, the combination of the DNA that has stayed in the nucleus and the "tail" of DNA that has migrated makes the cells look like a "comet". The length and intensity of the comets in relation to the signal of the non-migrating nuclear DNA compared to the controls can be quantified with the help of software. Being based on a microscopic readout, the comet assay can also detect heterogeneity within a cell population, which is not possible with any lysate-based assay including FADU. The comet assay is, however, very tedious and requires a large number of steps to be performed by highly trained personnel. Likewise, the FADU assay in its original, manually operated version [1] is technically very demanding, labour-intensive and requires large numbers of cells.

The automated version of the FADU assay described here relies on a properly configured LHD equipped with some custom-made accessories. It is operated in a 96-well format and is able to measure DNA strand breaks and repair with a high reproducibility. We have been able to reduce the number of cells necessary for the assay dramatically. While Birnboim & Jevcak used 1–2 × 10^6 cells per data point [1], we were able to reduce the number of cells required by more than 100-fold to 4,900–8,400 cells. Temperature control is also part of the automated opera-

Figure 4
DNA damage and repair in several different cell systems. HeLa S3 cells (A), Jurkat cells (B) and human PBMC (C) were damaged with 2.3 Gy of X-irradiation (P_1). To allow repair, cells were incubated at 37°C for 40 min (R). T and P_0 samples were controls. In all cases the fluorescence signal of the P_1 samples was lower than the control value P_0, and those of the R values higher than the damaged samples P_1. Error bars represent SDs from 8 replicates.

Figure 5
Time course of DNA strand break repair of Jurkat cells in intervals of 5 min. Cells were X-irradiated on ice with 6.8 Gy (basal line). To measure DNA strand break repair, the damaged cells were incubated at 37°C for different time periods as indicated (grey columns). The black column represents level of SybrGreen fluorescence obtained in undamaged cells (P_0 values). Error bars represent SDs from 3 replicates.
tion. Furthermore samples are completely protected from light throughout the duration of the automated steps. Using this system we have been able to measure DNA strand breaks and repair in a variety of cell systems. In our hands the lowest dose of X-ray-induced DNA strand breaks detected with statistical significance is 0.13 Gy. Collectively, the automated procedure including all the critical steps (Fig. 1A) takes 2 hours. Usually a complete FADU experiment for the assessment of DNA repair can be performed within 4–5 hours. While the FADU assay cannot detect cellular heterogeneity, determination of the average level of strand breaks will be sufficient for many purposes, and here the automated FADU is clearly superior due to its higher throughput and much lower labour utilisation.

Since the introduction of the comet assay, many modifications have been introduced to increase its performance [9] and enhance throughput [10-12]. A method using fluorescence analysis of DNA unwinding to measure DNA damage was also modified to decrease processing time [3,6]. However, the method described in the present paper is fully automated and has higher throughput than previous methods for measuring DNA damage and repair. It should be noted that we have typically used 8 replicates per T/P0/Px/Rx value, respectively. Depending on the magnitude of the effect to be studied, however, statistically significant results can be obtained with only half the number of replicates (data not shown), thus further increasing throughput.

Another clear-cut advantage of the automated FADU assays compared with the comet assay is the better definition of time of lysis. While in the FADU assay lysis occurs instantaneously at the time of addition of lysis solution, this may not be the case in the comet assay as lysis solution has to diffuse through the soft agar, in which the live cells are embedded. This ill-defined situation is likely to decrease precision of the comet assay especially when assessing the rapid phase of DNA repair.

It is known that at pH values >13, alkali-labile sites are converted to DNA single-strand breaks and thus give rise to additional initiation sites for unwinding. Our pH measurements, however, showed that upon addition of the alkaline unwinding solution, the pH is 12.5, and therefore we have conditions that will not or only marginally generate strand breaks at alkali-labile sites [13].

**Conclusion**

In conclusion, we have established a robust and convenient method for the quantitative assessment of formation and repair of DNA single-strand breaks in live cells. While the sensitivity of our method is comparable to the assays most often used currently, the throughput is massively increased while the operator time decreased.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

MMV and RP contributed to designing, performing, evaluating and interpreting the experiments shown in this paper. They also contributed to the set-up of the hardware and master protocol of the assay and to drafting the manuscript. TS, AL and MM performed in preliminary experiments and contributed to the set-up of the hardware and master protocol of the assay. TBLK contributed to directing the project and writing the manuscript. AB conceived and led the project. All authors read and approved the final manuscript.

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