Cells and Stripes: A novel quantitative photo-manipulation technique

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Laser micro-irradiation is a technology widely used in the DNA damage response, checkpoint signaling, chromatin remodeling and related research fields, to assess chromatin modifications and recruitment of diverse DNA damage sensors, mediators and repair proteins to sites of DNA lesions. While this approach has aided numerous discoveries related to cell biology, maintenance of genome integrity, aging and cancer, it has so far been limited by a tedious manual definition of laser-irradiated subcellular regions, with the ensuing restriction to only a small number of cells treated and analyzed in a single experiment. Here, we present an improved and versatile alternative to the micro-irradiation approach: Quantitative analysis of photo-manipulated samples using innovative settings of standard laser-scanning microscopes. Up to 200 cells are simultaneously exposed to a laser beam in a defined pattern of collinear rays. The induced striation pattern is then automatically evaluated by a simple algorithm, which provides a quantitative assessment of various laser-induced phenotypes in live or fixed cells. Overall, this new approach represents a more robust alternative to existing techniques, and provides a versatile tool for a wide range of applications in biomedicine.

Confocal laser scanning microscopy (LSM) enables precise spatio-temporal photo-manipulations based on LSM-embedded lasers in defined regions of interest (ROIs). These techniques include induction of DNA damage in cell nucleus1,2 and various photo-bleaching/activating procedures such as fluorescence recovery after photobleaching (FRAP)3.

Among the research areas that exploit these techniques, live-cell monitoring of factors involved in cellular responses to DNA damage has been particularly prominent. Compared to standard conventional methods commonly employed in the DNA damage field, laser-induced DNA damage provides multiple advantages, such as the ability to damage DNA in a certain time-point and in only part of the nucleus while the surrounding DNA stays intact, a feature especially attractive for dynamic analyses of protein recruitment4–8. Arguably the most deleterious lesions formed after laser micro-irradiation are DNA double strand breaks (DSB), a response to which involves a cascade of carefully orchestrated events, eventually leading to recruitment of factors participating in repair of DSBs via either homologous recombination or nonhomologous end joining pathways9. Among the main factors involved in the stepwise recruitment and assembly of the DSB response proteins is MDC1, a scaffold protein10,11 the GFP-tagged version of which was used as one of the DNA damage reporters in live-cell imaging experiments performed as part of our present study. MDC1 is one of the early-response proteins that forms a platform for recruitment of other important DDR factors (eg. RNF8, 53BP1, BRCA1, etc)12–14. The other protein we used as a GFP-tagged reporter is 53BP1, a protein that plays an important role in DSB repair pathway choice15. Finally, as a representative of factors involved in DNA repair of non-DSB lesions, we also employed FANCD2, a protein that becomes activated by monoubiquitination and plays a central role in the so-called Fanconi anemia (FA) pathway implicated in repair of DNA interstrand crosslinks, translesion DNA synthesis, nucleolytic incision and homologous recombination16.

In general, current approaches to the photo-manipulation techniques involve manual definition of the ROI, which is then exposed to a particular laser resulting in the respective photo-effect. The ROI is followed over time

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and its signal intensity is measured. The process is laborious and commonly requires constant attention for active repositioning of the readout area due to movement of cells over time. Here we propose a method to circumvent limitations of the manual approach by simultaneously exposing many cells to a co-linear pattern of laser rays. This is achieved by a particular set of LSM settings likely universally compatible with microscopes of various producers (tested on two types of LSM systems so far).

Results
Standard LSM systems can be adapted for photo-manipulation in a predefined striation pattern covering the entire acquisition area. Once the LSM system is set to a very low resolution and slow scanning speed, it generates a specific striation pattern. In principle, the entire acquisition area is scanned with small resolution e.g. only 32 pixels in each dimension (x and y). Laser scanning microscope does not switch off the lasers during the bi-directional scanning process between points nor lines. Such settings then result into a pattern of 32 horizontal lines, rather than $32 \times 32$ dots. The total dose of light energy for each of the lines can be adjusted by altering the scanning speed (pixel dwell time) and further by repetitions of the scanning cycles (iterations). This effect can be visualized for example on a microscopic slide covered with a thin layer of fluorescent paint which is bleached by the laser only within the scanning lines (Fig. 1a). If the used laser wavelength and power has DNA-damaging properties, the same spatially defined DNA damage striation pattern can be induced in live cells and visualized for example by translocation of tagged responsive proteins towards the DNA lesions (Fig. 1b). If the laser is set to a bleaching mode, a similar negative striation pattern is achieved (Fig. 1c). A large number of cells (limited only by the numbers of cells that can grow within the acquisition area) can be exposed in a very short time and the induced striation pattern of known parameters is amenable to an automatic software evaluation.

In the LSM-based laser-induced DNA damage applications, the amount of recruited and/or modified responsive proteins (DNA damage recognition) and their subsequent release, degradation or de-modification (processes reflecting DNA repair) can be measured. This is applicable to immunofluorescence (IF) as well as live-cell imaging. IF enables simultaneous detection of multiple proteins, including protein modifications (Supplementary Fig. S1), but has the disadvantages of an end-point assay. Live-cell analysis based on reporter cell lines tracks individual cells over time. The evolution of the striation pattern is followed providing a unique readout for various dynamic studies (Fig. 1d).

Principles of the signal quantification. The induced striation pattern is described by two pre-defined parameters: the stripe width and the inter-stripe distance (gauche). The stripe width is defined by the used hardware and cannot be manipulated by the user. On the other hand, the inter-stripe distance can be manipulated by changing the scanning resolution or objective magnification defining the total number of stripes per field (Supplementary Fig. S2). Simple software was developed for quantitative evaluation of the induced striation pattern. The algorithm first recognizes individual nuclei. In each nucleus, stripe(s) are automatically identified (see Fig. 1e and Methods for details and Supplementary information for the software code). A measure of striation (MS) is proposed to capture the excess signal in the stripes for each nucleus. Such an approach is useful for DDR quantification directly compared to a commonly used method measuring mean fluorescence signal per nucleus. For fixed samples involving protein modifications at sites of DNA damage such as lesion-associated phosphorylation of histone H2AX, the MS provides better discrimination between positive and negative controls (Supplementary Fig. S1). For assessment of reporter protein recruitment, such as for MDC1-GFP accumulation at sites of DNA damage, only the MS value enables meaningful analysis of the DDR (Supplementary Fig. S1).

Method optimization. In live-cell reporters involving GFP-tagged DNA-damage responsive proteins, the dependence of MS on time can be captured by plotting the median MS for each time-point. Such dependence can be visualized for example by translocation of tagged responsive proteins towards the DNA lesions (Fig. 1b). If the laser is set to a bleaching mode, a similar negative striation pattern is achieved (Fig. 1c). A large number of cells (limited only by the numbers of cells that can grow within the acquisition area) can be exposed in a very short time and the induced striation pattern of known parameters is amenable to an automatic software evaluation.

Method validation for DNA damage analysis. Once properly set, the striping method together with the software solution were tested on a small panel of known chemicals to prove its potential utility for applications involving large numbers of cells such as high-content screens for compounds and/or factors relevant for DNA integrity maintenance (potential cancer therapeutics). The compounds we selected are known to interfere with processes contributing to genomic stability, DNA damage signaling and/or repair pathways. The experiment was performed using a 96-well plate format in a semi-automatic regimen with autofocus and automatic repositioning.
and acquisition of striped regions in three different reporter cell lines. The tested compounds were added 2 hours before the experiment. For each compound, the cellular response at multiple time points after laser irradiation was compared with the mock treated sample. Four numerical parameters describing the MS curve (see Methods for details) were statistically evaluated (Fig. 2a). Notably, some of the compounds scored differentially for the MDC1-GFP, 53BP1-GFP and FANCD2-GFP reporters, respectively (Fig. 2a,b). Among the tested compounds only one (a PLK1 inhibitor) did not score in any of the measured parameters, in any of the reporter cell lines.
Method validation for FRAP analysis. The presented method including the software solution is easily adaptable for other photo-manipulation techniques. As an example of its utility, the FRAP method was tested in U-2-OS H2B-GFP reporter cells. The cells were pre-sensitized by 0.5 μM BrdU for 24 h and bleached by the co-linear stripes with either 488 nm laser or 355 nm laser. MS evaluation was performed along the same principle as for live-cell analysis after DNA damage. Despite the signal was initially bleached to the same level for both lasers, the H2B-GFP signal recovery was significantly faster after the UV-A laser bleaching (Fig. 2c,d). Apart from their technical merit, these results with the UV-A laser induced damage are novel and consistent with a published...
report that DNA repair of lesions induced by UV-C lasers accelerates turnover of histones including H2B\textsuperscript{12}, suggesting that accelerated histone turnover is a common feature shared by multiple DNA repair pathways.

**Discussion**

The method presented in this study provides several advantages over currently used techniques, and offers a broad applicability in the area of LSM-based photo-manipulations in biomedicine. For the first time, a precisely defined striation pattern was used for simultaneous micro-irradiation of dozens of cells. Known parameters of striation patterns simplify software-based quantification of cellular responses. This enables an easy and unbiased evaluation of numbers of cells sufficient for robust statistical testing. Moreover, the method we present meets all requirements for full automation and application in high-content screens for new drugs and/or factors modulating diverse aspects of the DNA damage response. Several compounds that positively scored in our test require approx. 6 times higher laser irradiation dose for the 355 nm, and a 10 times higher dose for the 405 nm laser (Supplementary Fig. S3). Thus either laser power or exposure time introduce serious limiting factors for presensitizer-free experiments if performed in a standard manner of manual laser path definition. Given all the advantages of the method we report here, we believe this new approach utilizing standard, widely used equipment could facilitate a broad range of biological and biomedical applications.

**Methods**

**Microscopic devices.** Microscopic station Zeiss Axioimager Z.1 with laser scanning LSM780 module. Lasers: UV-A 355 nm 65 mW (used for DNA damage induction, bleaching experiments and acquisition of Hoescht signal), argon-neon 458, 488, and 514 nm (used for acquisition of GFP and Alexa Flour 488 signals) and 568 nm (used for acquisition of Alexa Flour 568 signal). 40× objective (C-Apo, 1.2 DICIII, water immersion) was used for all experimental procedures unless stated otherwise in figure description. All other tested objectives are listed in Supplementary Fig. S2. Microscopic station Leica DMI 6000B with laser scanning TCS SP5 AOBS TANDEM module. Lasers: UV-A 405 nm 100 mW (used for DNA damage induction), argon-ion 458, 488, and 514 nm (used for acquisition of GFP signal). Objective: 40× objective (HC PL APO 40×1.30 OIL CS2, LP/0.17/D). Both microscopic stations were equipped with temperature-controlled incubator for live-cell support.

**LSM settings for stripping procedure.** ZEN 2011 software was used to control the Zeiss LSM780 device. For micro-irradiation 355 nm laser at maximum power (65 mW, 100% output) was used. To obtained desired striation pattern scan mode of very low resolution (size i.e. resolution 32×32, 64×64 or 128×128 pixels) and the lowest possible scanning speed (pixel dwell time 709.27\(\mu\)s for 32×32 pixels) was used. Striping at pixel resolution 32×32 was chosen for most experiments with the 40× objective as it is causing approx. 1–2 stripes per cell nucleus. The scan was bidirectional, zoom 0.6. Number of iterations (means scanning cycles) is indicated in the figure legends (experiment dependent). Autofocus before each micro-irradiation was performed with 488 nm laser (the same settings as for respective GFP signal acquisition). For image acquisition in live-cell experiments with GFP-tagged reporters 488 nm laser was used. Images were acquired in 16 bit depth, zoom 0.6, five z-stack planes (0.6\(\mu\)m apart). To minimize bleaching the autofocus for image acquisition was based on reflected light of 561 nm laser which allows visualization of a thin contact layer between cells and the glass cultivation surface (also known as backscatter image). For bleaching experiments either 488 nm laser was used (100% output), or 355 nm laser (100% output). Autofocus was performed in the same manner as for micro-irradiation experiments.

**Visualization of the striation pattern using microscopic slide covered by fluorescent paint.** Standard microscopic glass was covered by a homogeneous paint layer using red permanent marker (type: Permanent 8566, ink color 04, Centropen) and air dried. The paint layer was mounted by water and covered by a standard cover glass (0.17 mm). Signal was visualized using 561 nm laser and emission spectrum 570–710 nm. Micro-irradiation (stripping) procedure included 355 nm laser, 100%, 32×32 pixels, 1 iteration, pixel dwell time 709.27\(\mu\)s, bi-directional scan.

**Cell lines and treatments.** All reporter cell lines were stably expressing the respective protein with a GFP tag. U-2-OS-MDC1-GFP, U-2-OS-53BP1-GFP and PD20F-FAACD2-GFP were obtained from Danish Cancer Society Research Centre, Copenhagen, Denmark\textsuperscript{8,21}, MRC-5 cells were obtained from ATCC and McCoy mouse fibroblasts were obtained from R-D Biotech. All cell lines were incubated under standard cultivation conditions (5% CO\textsubscript{2}, 37\(^\circ\)C), in DMEM (Gibco) supplemented with 10% FBS (Invitrogen). Cells were pretreated by 0.5\(\mu\)M BrdU (Sigma) for 24 h before micro-irradiation unless stated otherwise (see figures description). For micro-irradiation procedures cells were seeded into 96 well plates with glass bottom (Corning) 18 hours before...
laser irradiation. 2 hours before laser irradiation the medium was changed to DMEM CO₂ - independent medium without phenol red (Gibco) supplemented with 10% FBS and the tested chemicals. Cells were seeded in amounts to ensure sub-confluent density (proper cell cycle progression enables BrdU incorporation). After seeding the cells into 96 well plates the specimens were first placed on equilibrated bench for 20 min at RT to ensure equal cells distribution and then placed into incubator.

**U-2-OS-H2B-GFP clone construction.** Commercial plasmid pBOS-H2BGFP Vector was used. Cells were electroporated with Neon Transfection System (Life Technologies) with 15 μg of plasmid per 1.10⁶ cells (pulse voltage - 1230 V, pulse width 10 ms, pulse number 2). 72 h after transfection, GFP-positive cells were sorted for bulk culture stably expressing H2B-GFP. After proper cell culture propagation, 1 cell per well sorting was done to obtain individual clones (BD FACS Aria).

**Quantitative analysis of the striation patterns.** An in-house specialized software routine for the evaluation of the striation patterns was developed and implemented in MatLab. The first step of the routine consists of the segmentation of the images to find the individual nuclei. To be selected, a nucleus must satisfy certain conditions on its size and solidity. The process begins by segmenting the first image, acquired at the beginning of the experiment (t = 0, before laser irradiation). Standard thresholding algorithms (e.g. the Otsu’s method) do not perform well here because there is a large variance in the signal intensity among the individual nuclei (a common problem in most live-cells based reporter systems). Consequently, a cascade of decreasing local thresholds is used. After the cascade has finished, usually all the nuclei of desirable characteristics are identified in the image. For the segmentation of the subsequent images, a tracking algorithm is used. The algorithm attempts to track each nucleus by searching the vicinity of its positions in the previous image. This tracking approach is necessary because the signal intensity in the inter-stripe regions of the nuclei may decrease all the way to the background level and thus become completely “invisible” for any simple thresholding procedure.

Several measures were proposed and tested to quantify the evolution of the striation patterns induced by the laser within each recognized and over the time tracked nucleus. In the end, the relative excess signal in the stripes, hereafter called the measure of striation and abbreviated “MS”, proved to be the most useful. It is constructed as follows: Let us consider an image of an individual cell nucleus at time t. We start by rotating the image so that the (possible) stripes become vertical. This is done by taking the Radon Transform of the image and finding its maximum. If there are no stripes in the image, the nucleus gets usually rotated by a random angle.

Next, the position of the stripes and their diameter is found. This is done by fitting the stripes, i.e. maximizing a function of two variables \( F(D,P) \) where \( D \) stands for the distance from the nuclei to the center of the range of the stripes and \( P \) stands for the horizontal offset of the stripes (i.e. the position of the center of the first stripe from the rightmost boundary point of the nucleus, see Fig. 1e). The function \( F \) measures the ratio of the mean intensity of the signal in the stripes to the mean intensity of the signal in the entire nucleus. The width of the stripes is a user-defined constant and it is not fitted. Although both the parameters \( D \) and \( P \) are known from the initial setting of the laser, the optimization (i.e. fitting) step is necessary due to elasticity of the nuclei and their movement in time. Without this optimization step, the measure of striation would automatically decrease in time as the nuclei move around and change shape. This would create a systematic error – a false decrease in the intensity of the stripes. The minimization is implemented by means of a trivial algorithm that fully evaluates all plausible values of \( D \) and \( P \) and selects the highest value for \( F \). The parameter \( D \) is allowed to take only integer values from a pre-specified range and the parameter \( P \) is not allowed to depart too much from the point where the vertical sum of the signal reaches its maximum. More advanced optimization procedures may naturally be used, however, this optimization step takes much less time than the cell-tracking part of the algorithm. Moreover, the full evaluation of a single image (which may contain around 100 nuclei) takes only few seconds on a standard PC.

Once the position of the stripes is found, the measure of striation is computed according to equation (1):

\[
MS = \frac{J_{\text{meas}} - J_{\text{expt}}}{J_{\text{expt}}}
\]

**Equation (1):** \( J_{\text{meas}} \) (measured) stands for the actual signal integrated density of regions inside the stripes, and \( J_{\text{expt}} \) (expected) stands for the expected value of the signal integrated density in region of the stripes, if the nucleus was perfectly homogeneous and no stripes were present. \( J_{\text{expt}} \) is computed according to equation (2):

\[
J_{\text{expt}} = \frac{\text{Area inside the stripes}}{\text{Total area of the nucleus}} \times \text{SID of the nucleus}
\]

**Equation (2):** SID stands for signal integrated density.

The measure \( MS \) thus captures the relative excess signal in the stripes. The baseline is zero which means there is no excess signal in the stripes, i.e. the stripes contain only as much signal as corresponds to their area, i.e. the signal is homogeneously distributed throughout the nucleus, i.e. there are no visible stripes. As the fluorescently labeled DNA-damage responsive proteins get recruited to the places of laser-induced damage (i.e. to the stripes), the measure of striation grows. After some time, the proteins are released, they begin to spread out again over the entire nucleus, and/or undergo degradation and the measure of striation decreases again. Thus, the measure of striation captures the dynamics of the DNA-repair processes. The construction of MS ensured that it is independent of the total SID of the nucleus. This is necessary because the total SID may vary greatly among individual nuclei.
DNA damage repair (DDR) curve parameters. By a DDR curve we mean the dependence of MS on time for a particular nucleus. To compare different DDR curves, the following three parameters were defined (Fig. 1f,g):

1. The amplitude of the response (Amp) which measures the maximum amount of the protein recruited to the stripes. It is computed as the maximum of MS over all observed time points.
2. The time to peak response (Tpeak) measures the speed of the protein recruitment. It is computed as the time (in minutes) from the micro-irradiation until the MS reaches its maximum.
3. The relaxation speed (Relax) which reflects the dynamics of the repair process after the peak response. It is calculated as the slope of the line fitted to the function $MS(t) = MS_{\text{peak}} - MS_{\text{relax}}$ (where $MS_{\text{peak}}$ is the peak of MS and $MS_{\text{relax}}$ is the relaxation speed).

These parameters are computed for every nucleus within a sample and differences among samples are tested for statistical significance by means of the Kruskal-Wallis test. The significance level of all tests was set to 0.05. For illustrative graphs the median MS at each timepoint was computed and these medians were connected by lines (Figs 1f,g and 2b).

Screen for compounds potentially affecting DNA damage response. Panel of compounds including VE-82119 (Selleck Chem.), LY260361820 (Selleck Chem.), Caffeine21 (Sigma), NU702622 (Sigma), Olaparib23 (Selleck Chem.), KU5593324 (Sigma), BI256325 (Selleck Chem.), MIRIN33 (Sigma) was tested in 3 reporter cell lines. Each compound was added to cells 2 hours before micro-irradiation in final concentration of 10 μM with the exception of BI2563 (100 μM). Respective DDR protein recruitment to sites of damage was evaluated as MS in each timepoint for every single cell. Four parameters characterizing the DDR curve (see above) were calculated and statistically tested against mock treated control (Fig. 2a). Every sample was tested in technical duplicate (i.e. in two separate wells on the same 96 well plate). Results from duplicates were pooled and tested against pooled mock treated sample. At least two independent biological replicates were performed for each tested compound. Only significant effects scored similarly in both biological replicates were marked in the table by a color change (Fig. 2a). Minimum of 100 cells were scored in each experiment.

FRAP experiment. Cells were seeded and treated in the same way as for the micro-irradiation experiments, including BrdU pre-sensitization. The FRAP curve (i.e. the evolution of MS in time) was constructed in the same way as the DDR curve. The first two DDR curve parameters (Amp, Tpeak) were defined similarly as above (taking the minimum of MS instead of a maximum). Tpeak was not used as the minimum MS (i.e. maximum striation) is always reached immediately after the bleaching. The relaxation speed was measured in a slightly different way. The parameter Relax30 denotes the slope of the line fitted to the FRAP curve between $t = 1$ min and $t = 30$ min, the parameter Relax30 denotes the slope of the line fitted to the FRAP curve between $t = 1$ min and $t = 60$ min (Fig. 1g). Experiment was performed in two biological replicates. Minimum of 150 cells were scored in each experiment.

Immunofluorescence. Cells were seeded and treated with BrdU in the same way as described for the live-cell experiments (see above). At indicated timepoints after micro-irradiation, cells were fixed with 10% buffered formalin (Sigma) and permeabilized with 0.5% Triton X for 5 min. After 20 min blocking in 1% BSA in PBS, the samples were incubated with primary antibodies at 4 °C for 18 h, followed by 1 h incubation with secondary antibodies at RT. DNA was stained by Hoechst33342 (Invitrogen) for 5 min. Following antibodies were used: phospho-H2AX (Ser139) JBW301 (Millipore, 500 μM), cyclin A (Leica, 500 μM), cyclin A (Leica, 50 μM), AlexaFluor568 (Invitrogen, 1000 μM). Slides were mounted with FluorountM with the exception of BI2563 (100 μM). Respective DDR protein recruitment to sites of damage was evaluated as MS in each timepoint for every single cell. Four parameters characterizing the DDR curve (see above) were calculated and statistically tested against mock treated control (Fig. 2a). Every sample was tested in technical duplicate (i.e. in two separate wells on the same 96 well plate). Results from duplicates were pooled and tested against pooled mock treated sample. At least two independent biological replicates were performed for each tested compound. Only significant effects scored similarly in both biological replicates were marked in the table by a color change (Fig. 2a). Minimum of 100 cells were scored in each experiment.

Cell cycle analysis. Cells were seeded and treated in the same way as for the micro-irradiation experiment. After 24 h incubation with indicated concentrations of BrdU, cells were fixed with 10% buffered formalin (Sigma) and stained with Hoechst33342 5 μg/ml in PBS (Invitrogen) for 10 min. Cell cycle was evaluated using OlympusBX71 inverted microscope and ScanR Acquisition and Analysis software (Olympus).

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Author Contributions
M.M. and E.V. wrote the manuscript and performed most cell work and stripping experiments, T.F. designed the software script and performed all statistical analysis, H.H. performed parallel stripping experiments on LeicaSP5 LSM system, I.F. prepared and characterized the U-2-OS H2B-GFP cell line, J.G. and D.M. performed stripping experiments and contributed to data evaluation, J.B. wrote the manuscript and contributed to data evaluation.

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