RESEARCH NOTE

The study of protein recruitment to UV-induced DNA lesions can be distorted by photoconversion of DNA dyes like Hoechst or DAPI [version 1; peer review: 1 approved, 2 approved with reservations]

Verena Hurst, Susan M. Gasser

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Abstract

A common approach used to assess DNA repair factor binding in mammalian cells is to induce DNA damage with a UV laser and follow the movement of GFP-tagged proteins to the site of damage. Often these measurements are performed in the presence of the blue DNA intercalating dye Hoechst or DAPI, which is used to label nuclear DNA. A UV-induced switch of Hoechst and DAPI from a blue-light to a green-light emitter will give a false positive signal at the site of damage. Thus, photoconversion signals must be subtracted from the overall green-light emission to determine true recruitment. Here we demonstrate the photoconversion effect and suggest control experiments to exclude false-positive results.

Keywords

Photoconversion, Hoechst, DAPI, UV laser, DNA repair
Corresponding author: Susan M. Gasser (susan.gasser@fmi.ch)

Author roles: Hurst V: Data Curation, Writing – Original Draft Preparation; Gasser SM: Writing – Review & Editing

Competing interests: No competing interests were disclosed.

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Abbreviations
DAPI: 4', 6-diamidino-2-phenylindole; UV: ultraviolet light; U2OS: human bone osteosarcoma epithelial cells; YOYO-1: tetracationic homodimer of Oxazole Yellow; GFP: Green fluorescent protein; 53BP1: Tumor suppressor p53-binding protein 1; XRCC1: X-ray repair cross-complementing protein 1; FEN-1: Flap endonuclease 1; PARP-1: Poly [ADP-ribose] polymerase 1; Ku70: 5’-deoxyribose-5-phosphate lyase Ku70/X-ray repair cross-complementing protein 6, MDC1: Mediator of DNA damage checkpoint 1; SMARCA5: SWI/SNF-related matrix-associated actin-dependent regulator of chromatin subfamily A member 5

Introduction
A variety of DNA binding dyes, such as DAPI, Hoechst, Vybrant DyeCycle Violet and YOYO-1, can change their optical properties upon exposure to light. Whereas the induction of YOYO-1 blinking is an intended change that is exploited for super resolution microscopy \(^1\), the photoconversion of DAPI, Hoechst or Vybrant Dye Cycle Violet during multicolor fluorescence microscopy is unexpected and can lead to false-positive signals.

Upon UV exposure or a low pH, the emission spectra of DAPI, Hoechst and Vybrant DyeCycle Violet shift from the blue to the green wavelength with detectable signals in the yellow and orange\(^2\)-\(^4\). This shift makes the signal indistinguishable from the emission of other standardly used fluorescent proteins such as GFP. An experimenter expecting the DNA dyes to emit in the blue can misinterpret the green signal as that arising from another probe in the sample. This risk has been raised previously\(^3\)-\(^5\), yet the artefact is rarely controlled for.

With respect to these findings, a microscopic setup like the one used to study the localization of repair proteins to a UV laser-induced zone of DNA damage can be particularly problematic. Very commonly, cell nuclei are stained with DAPI or Hoechst and a restricted part of the nucleus is exposed to a strong UV laser. The protein of interest is detected in the green channel thanks either to its fusion to GFP or else through an antibody labelled with a green light-emitting fluorophore. Unfortunately, photoconversion of the DNA dye is usually not checked\(^7\)-\(^12\). Here will illustrate the problem and suggest necessary controls.

Results
To study the recruitment of a potential DNA damage related protein, we made use of a previously established setup in which cell nuclei are sensitized with Hoechst, DNA damage is induced with a UV laser, and the recruitment of a protein of interest is measured over time by fluorescence microscopy. Unexpectedly, cells stained with Hoechst that did not express any GFP-tagged protein showed a similar increase in the green channel at the UV damage site, as cells expressing the GFP-tagged protein (Figure 1). The detected increase in signal was not due to protein recruitment to the damage site, since there was no GFP-tagged protein in the cell. Moreover, in cells expressing the GFP-tagged protein that were not stained with Hoechst, there was no increase in signal intensity at the UV damage site. This demonstrates conclusively that the increase in fluorescence in the green channel was a false-positive result. Raw images are available on figshare\(^13\).

Discussion
We illustrate here that one should avoid exposing DAPI or Hoechst to a strong UV laser if one is imaging green light emitting probes such as GFP or a secondary antibody coupled to fluorescein/Alexa488. This is because photoconverted Hoechst and DAPI emit strongly in the same channel. As an alternative nuclear marker, we suggest employing a fluorescently tagged protein that localizes at the nuclear periphery and does not interfere with the experimental process.

If Hoechst is employed as a sensitizing agent, we suggest using the minimum dye concentration and laser power necessary and to combine it with probes/secondary antibodies of a color that is well separable from photoconverted DAPI/Hoechst. For instance, far red emission is compatible with photoconverted DAPI/Hoechst\(^1\). Yet, quantitation of the signal of the investigated protein requires normalization to a background control that is obtained by performing the laser experiment on DAPI/Hoechst-stained but otherwise native cells and acquiring signal with the same channel and exposure conditions, as used for the experimental probe. However, there is evidence showing that DNA sensitization prior to laser exposure is not required: DNA repair proteins such as 53BP1\(^14\)-\(^16\), XRCC1\(^15\), FEN-1\(^15\), PARP-1\(^15\), Ku70\(^15\), MDC1\(^16\), and SMARCA5\(^16\) are recruited to sites of damage without previous sensitization by Hoechst.

![Figure 1. Representative U2OS cell nucleus before and after UV-induced photoconversion of Hoechst.](image-url)
We note that in addition to particular situations in which one induces damage, the photoconversion of DAPI can occur during standard dual color microscopy. To minimize artefacts one should be careful about the order in which dyes are observed, and visualize the green channel prior to exposing to short-wave light.

Methods

U2OS cells (a gift from Prof. Primo Leo Schaer, Department of Biomedicine, University of Basel) were incubated with 1.5 μg/ml Hoechst 33342 (Thermo Fisher Scientific, H1399) for at least 30 minutes prior to photoconversion. Photoconversion was induced with a VisiFRAP module (Visitron) mounted on the backport of the microscope and equipped with a 405 nm laser (Toptica, illumination power at the objective 12.8 mW). Confocal images were acquired with an Olympus IX81 microscope equipped with a PlanApo 100x/1.45 TIRFM oil objective, a CSU-X1 scan-head (Yokogawa), an Evolve 512 EMCCD camera (Photometrics), a 491 nm laser (Cobalt Calypso 100), a 488/568 dichroic (Semrock Di01-T488/568-25x15x0.5), a band-pass 525/40 emission filter (Semrock Kalypso 100), an epi-fluorescence setup, and a band-pass 675/75 emission filter (Semrock FF01-525/40-25) and controlled with the Visiview Software (Visitron). Images in Figure 1 show maximum intensity projections of stacks covering 7 μm.

Data availability

Raw images of the stacks taken during this study are available on figshare. DOI: https://doi.org/10.6084/m9.figshare.7583960.

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Media

The three available avi files, C1 green, C2 blue and composite, represent a time series of maximum intensity projections showing the UV-induced emission change of the DNA intercalating dye Hoechst from the blue to the green region of the visible spectrum. Under live conditions, a Hoechst-stained cell nucleus was irradiated with 405 nm UV laser light along a predefined pattern (#). A time series of image stacks was acquired (25 equally spaced time points over 65s, stacks covering 7-μm sample depth) in two channels (C1 “green”: 491/525 nm, C2 “blue”: 405/450 nm). DOI: https://doi.org/10.6084/m9.figshare.7583960.

Grant information

This study was funded by the Swiss National Science Foundation and Novartis Research Foundation.

References


Open Peer Review

Current Referee Status: ✅ ? ?

Version 1

Referee Report 26 February 2019

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Jerzy Dobrucki
Department of Cell Biophysics, Faculty of Biochemistry, Biophysics, and Biotechnology, Jagiellonian University, Krakow, Poland

The paper "The study of protein recruitment..." by Hurst and Gasser is a very useful and timely technical report. It touches upon an important but often overlooked methodological aspect of studies of recruitment of repair factors to DNA lesions. When unnoticed, photoconversion of UV-excited DNA fluorescent probes can constitute a pitfall leading to incorrect interpretation of imaging data.

General comments:

“UV-induced lesions”
In the title and the text of this report the authors use a term “UV-induced DNA lesions”. The use of this term is misleading, since UV-induced lesions are generally defined as pyrimidine dimers and photoproducts. However, the authors refer to DNA lesions inflicted by 405nm light (which is not UV, see below) in the presence of a DNA-bound fluorescent dye Hoechst, and this leads to induction of a host of various lesions, not only “UV-induced”. When Hoechst 33342 has been introduced into live cells and exposed to 405nm focused laser light, it is (marginally) excited and acts as a photosensitizer. This leads to induction of oxidative damage and DNA breaks, i.e. not typical UV-induced lesions.

“UV laser”
The term ‘UV laser’ is used throughout the manuscript. This term appears incorrect, since the authors used a laser emitting 405nm wavelength light. By definition, 405 nm is visible light (and indeed it is readily visible by human eye, as opposed to UVA, UVB or UVC). I suggest to refer to the Toptica laser, which the authors used, as “blue laser”, as in the paper [Kong et al. 2009] which the authors cite.

Hoechst photosensitization vs DNA labeling
Fluorescent DNA probe Hoechst is described in two disguises in this paper – as a photosensitizer and as a counterstain. These two roles and the ensuing problems are not clearly distinguished and explained in the paper. If Hoechst or DAPI are added to live cells prior to microirradiation with a focused beam of light of 405nm wavelength, the dye acts as a photosensitizer (as correctly stated in Results). However, in the Abstract the authors state that “the measurements are performed in the presence of the blue intercalating dye Hoechst or DAPI which is used to label nuclear DNA”. This statement appears incorrect for two reasons: 1. Hoechst is usually added to live cells in order to photosensitize and yield massive, readily detectable damage, not to just label DNA (if Hoechst were to be used as a label to mark DNA, it could be added after microirradiation); 2. Hoechst and DAPI are not DNA intercalators, they are rather minor groove binders (the mode of binding is complex and depends on a number of factors, including the type of
DNA and a DNA/dye ratio).

Minor comments:

It is unclear what the authors mean by referring to photoconversion of Hoechst, DAPI and VybrantDyeCycle and, in the same sentence and context, to blinking of YoYo described as “a change in optical properties”. What change of optical properties of YoYo do the authors have in mind?

The authors state (in Discussion): “As an alternative nuclear marker, we suggest employing a fluorescently tagged protein that localizes at the nuclear periphery and does not interfere with the experimental process.” Why use such a complicated approach? It is much simpler to detect a transmitted light image, preferably using phase contrast or Nomarski interference contrast, and mark the outline of the nucleus in image overlay.

The authors state: “However, there is evidence showing that DNA sensitization prior to laser exposure is not required...” This is true, but again the way the authors put it is somewhat confusing and they refer to research in which UVA as well as visible light was used. Indeed, photosensitisation by exogenous DNA-binding compounds is never required – regardless of the type of light, UV or VIS. Moreover, adding a photosensitizer to the experimental system influences the type of damage. Thus, I suggest to distinguish two cases:

1. using UV-excited dyes and UV or near-UV (405nm) light, and
2. using dyes excited by visible light and visible light excitation.

In the case (1) the exciting light (UV or 405) will induce typical UV damage (PP, PD). Adding a photosensitizer prior to microirradiation will result in photodynamic effect, and cause induction of more types of lesions, and more extensive damage. There is vast literature about the action of UV alone and photodynamic effect type I and II. Some reference to this field of knowledge should be made in this paper.

Contrary to general belief, in the case (2) the exciting visible light alone, without any exogenous photosensitisers, will also induce DNA damage – single- and double-strand breaks, and recruitment of repair factors (Solarczyk et al., 2012, DNA Repair).

In summary, indeed various types of DNA damage can be induced without adding photosensitisers to cells prior to microirradiation, not only when UV, or 405 nm light is used, but also when visible light is applied. I suggest to clarify these facts in the paper.

A few inaccurate statements should be straightened out:

- “A common approach used to assess DNA repair factor binding in mammalian cells is to induce DNA damage with a UV laser and follow the movement of GFP-tagged proteins to the site of damage”. To be precise, induction of (local) DNA damage by a focused laser beam is used to detect recruitment of repair factors, but not their binding per se.

- “Movement of GFP-tagged proteins to the site of damage”. Movement of GFP-tagged proteins is not detected (this can be done by FCS) – only local increase of a concentration of a fusion protein is detected, and this arises from recruitment.

The authors state: “exposing DAPI or Hoechst to a strong UV laser” - light intensities and doses of energy delivered to the exposed region of the nucleus (area?) should be given.
Full information (dose) is lacking but it appears that the authors used an excessive power of 405nm light (12.8 mW at objective). DNA damage, especially in the presence of a photosensitizer, can be expected when using only microJ of energy (microW in the laser beam, seconds of exposure). This means that the photoconversion the authors describe most likely would have been less prominent, had a lower intensity and dose of energy been used. Applying excessive energy during microirradiation leads to such an extensive damage that relevant physiological studies may be impossible (Note that some images in the paper [Kong et al.] show microirradiation tracks in phase contrast images; the power which was used in this study was very high). I suggest that the authors state that the intensities (and doses of energy) they used may have been too high to induce DNA damage on the level encountered under typically encountered physiological conditions.

Typo:
"Here will illustrate the problem and suggest necessary controls." Here we will.

Hope this helps.

References

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Not applicable

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Cell Biophysics, DNA repair, microscopy

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Verena Hurst, Friedrich Miescher Institute Basel, Switzerland
A: Thank you for sharing your expertise. Your comments helped to significantly increase the quality of the article.

R: “UV-induced lesions”
In the title and the text of this report the authors use a term “UV-induced DNA lesions”. The use of this term is misleading, since UV-induced lesions are generally defined as pyrimidine dimers and photoproducts. However, the authors refer to DNA lesions inflicted by 405nm light (which is not UV, see below) in the presence of a DNA-bound fluorescent dye Hoechst, and this leads to induction of a host of various lesions, not only “UV-induced”. When Hoechst 33342 has been introduced into live cells and exposed to 405nm focused laser light, it is (marginally) excited and acts as a photosensitizer. This leads to induction of oxidative damage and DNA breaks, i.e. not typical UV-induced lesions.

A: We added information on the types of damage generated at different wavelengths and photosensitizing agents.

R: “UV laser”
The term ‘UV laser’ is used throughout the manuscript. This term appears incorrect, since the authors used a laser emitting 405nm wavelength light. By definition, 405 nm is visible light (and indeed it is readily visible by human eye, as opposed to UVA, UVB or UVC). I suggest to refer to the Toptica laser, which the authors used, as ‘blue laser”, as in the paper [Kong et al. 2009] which the authors cite.

A: You are right. We corrected that.

R: Hoechst photosensitization vs DNA labeling
Fluorescent DNA probe Hoechst is described in two disguises in this paper – as a photosensitizer and as a counterstain. These two roles and the ensuing problems are not clearly distinguished and explained in the paper. If Hoechst or DAPI are added to live cells prior to microirradiation with a focused beam of light of 405nm wavelength, the dye acts as a photosensitizer (as correctly stated in Results). However, in the Abstract the authors state that “the measurements are performed in the presence of the blue intercalating dye Hoechst or DAPI which is used to label nuclear DNA”. This statement appears incorrect for two reasons: 1. Hoechst is usually added to live cells in order to photosensitise and yield massive, readily detectable damage, not to just label DNA (if Hoechst were to be used as a label to mark DNA, it could be added after microirradiation); 2. Hoechst and DAPI are not DNA intercalators, they are rather minor groove binders (the mode of binding is complex and depends on a number of factors, including the type of DNA and a DNA/dye ratio).

A: We corrected and clarified that point. Actually using Hoechst helps identify the placement of the laser beam.

R: Minor comments:
It is unclear what the authors mean by referring to photoconversion of Hoechst, DAPI and VybrantDyeCycle and, in the same sentence and context, to blinking of YoYo described as “a change in optical properties”. What change of optical properties of YoYo do the authors have in mind?
A: In order to focus on Hoechst and DAPI we removed the comments on the other two dyes.

R: The authors state (in Discussion): “As an alternative nuclear marker, we suggest employing a fluorescently tagged protein that localizes at the nuclear periphery and does not interfere with the experimental process.” Why use such a complicated approach? It is much simpler to detect a transmitted light image, preferably using phase contrast or Nomarski interference contrast, and mark the outline of the nucleus in image overlay.

A: We and other labs standardly use fluorescent labels of the nuclear envelope without complications. We are not sure transmitted light images are compatible with rapid time lapse imaging since filters have to be changed but we mentioned this possibility in the text.

R: The authors state: “However, there is evidence showing that DNA sensitization prior to laser exposure is not required:…” This is true, but again the way the authors put it is somewhat confusing and they refer to research in which UVA as well as visible light was used. Indeed, photosensitisation by exogenous DNA-binding compounds is never required – regardless of the type of light, UV or VIS. Moreover, adding a photosensitizer to the experimental system influences the type of damage. Thus, I suggest to distinguish two cases:

1. using UV-excited dyes and UV or near-UV (405nm) light, and
2. using dyes excited by visible light and visible light excitation.

In the case (1) the exciting light (UV or 405) will induce typical UV damage (PP, PD). Adding a photosensitizer prior to microirradiation will result in photodynamic effect, and cause induction of more types of lesions, and more extensive damage. There is vast literature about the action of UV alone and photodynamic effect type I and II. Some reference to this field of knowledge should be made in this paper.

A: We have added a section discussing the types of damage generated under different conditions.

R: Contrary to general belief, in the case (2) the exciting visible light alone, without any exogenous photosensitisers, will also induce DNA damage – single- and double-strand breaks, and recruitment of repair factors (Solarczyk et al., 2012, DNA Repair1).

A: We noticed this phenomenon in our own experiments, and have now commented upon this in the text and have cited the reference suggested.

R: In summary, indeed various types of DNA damage can be induced without adding photosensitisers to cells prior to microirradiation, not only when UV, or 405 nm light is used, but also when visible light is applied. I suggest to clarify these facts in the paper.

A few inaccurate statements should be straightened out:

- “A common approach used to assess DNA repair factor binding in mammalian cells is to induce DNA damage with a UV laser and follow the movement of GFP-tagged proteins to the site of damage”. To be precise, induction of (local) DNA damage by a focused laser beam is used to detect recruitment of repair factors, but not their binding per se.

A: True. Corrected.

- R: “Movement of GFP-tagged proteins to the site of damage”. Movement of GFP-tagged proteins is not detected (this can be done by FCS) – only local increase of a concentration of a fusion protein is detected, and this arises from recruitment.
The authors state:
“exposing DAPI or Hoechst to a strong UV laser” - light intensities and doses of energy delivered to the exposed region of the nucleus (area?) should be given. Full information (dose) is lacking but it appears that the authors used an excessive power of 405nm light (12.8 mW at objective). DNA damage, especially in the presence of a photosensitizer, can be expected when using only microJ of energy (microW in the laser beam, seconds of exposure). This means that the photoconversion the authors describe most likely would have been less prominent, had a lower intensity and dose of energy been used. Applying excessive energy during microirradiation leads to such an extensive damage that relevant physiological studies may be impossible (Note that some images in the paper [Kong et al.] show microirradiation tracks in phase contrast images; the power which was used in this study was very high). I suggest that the authors state that the intensities (and doses of energy) they used may have been too high to induce DNA damage on the level encountered under typically encountered physiological conditions.

A: See response to Anna Fortuny, who raised this issue as well. Now we state that we use high laser power in order to demonstrate the effect and provide further information on the laser conditions. Furthermore, we refer to a study detecting and minimizing such an effect in their setup.

R: Typo:
“Here will illustrate the problem and suggest necessary controls.” Here we will.

A: Corrected.

R: Hope this helps.

A: Yes. Thank you!

**Competing Interests:** No competing interests were disclosed.
The authors should be more specific when referring to UV laser or UV light. They should specify “UVA”, as other UV wavelengths such as UVC, also used to introduce local DNA damage (Dinant et al., 2007), may not have the same effect.

They could suggest using BrdU instead of Hoechst to pre-sensitize cells to UVA light as done in a number of studies (e.g. Lukas et al., 2003).

The authors illustrate the problem with GFP-tagged proteins but the issue would be similar with YFP-tagged proteins. They explain that far-red emission is compatible with photoconverted DAPI/Hoechst. How about red emission?

There are 55 files on figshare, which seems excessive, and they are difficult to navigate through so in the end it is not very useful.

Result section: “cells expressing the GFP-tagged protein (Figure 1)”. These cells are not shown on the figure and which GFP-tagged protein is it?

References

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Not applicable

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Partly

*Competing Interests:* No competing interests were disclosed.

*Reviewer Expertise:* epigenetics, UV damage repair, imaging

We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however we have significant reservations, as outlined above.
Verena Hurst, Friedrich Miescher Institute Basel, Switzerland

Thank you very much for your comments! Version 2 of this research note accommodates the changes that you suggested unless stated otherwise.

- The authors refer to a “strong UV laser”. It is not clear which conditions are required for DAPI/Hoechst photoconversion (laser power, exposure time) and how they compare to classical exposure times for DAPI visualization and to UV laser settings for laser damage induction. It is important to clarify this point to determine if we face a marginal or a general problem.

Indeed, we use higher laser power than used in most studies in order to demonstrate the photoconversion effect. The purpose of this research note is to increase the awareness of the phenomenon and highlight that a negative control is particularly essential in this experimental setup. Even small amounts of photoconverted dye can change image quantitation and may alter apparent recruitment dynamics. We do think that this effect is particularly relevant to the study of uncharacterized proteins. Explorative studies may tempt the researcher to vary laser power and exposure time because the degree of damage required for protein recruitment is not known. This bears the risk of a false positive result. A preventive measure is to generate the damage without using Hoechst.

We now cite a study in which photoconversion was detected and minimized, indicating the relevance of this note.

A recent report on photoconversion in standard multiple color microscopy applications specifies exposure times for photoconversion.

- The authors should be more specific when referring to UV laser or UV light. They should specify “UVA”, as other UV wavelengths such as UVC, also used to introduce local DNA damage (Dinant et al.,2007), may not have the same effect.

We added a section on the type of damage generated +-sensitizing agents and cited Dinant et al.

- They could suggest using BrdU instead of Hoechst to pre-sensitize cells to UVA light as done in a number of studies (e.g. Lukas et al.,2003).

Now we mention BrdU as a sensitizing agent and specify which type of damage is generated in its presence (double strand breaks). However, we favor omission of sensitizing agents altogether.

- The authors illustrate the problem with GFP-tagged proteins but the issue would be similar with YFP-tagged proteins. They explain that far-red emission is compatible with photoconverted DAPI/Hoechst. How about red emission?

We have used GFP in our setup and therefore mainly discuss GFP. However, according to the publications we cited the effect may apply to YFP and other proteins with emission in the orange and near red as well. This is now mentioned.

- There are 55 files on figshare, which seems excessive, and they are difficult to navigate through so in the end it is not very useful.

We were asked to deposit the raw data on figshare. As mentioned in our note we also deposited avi files of timelapse MIPs in both channels as well as a channel merge. These files are easy to handle.

- Result section: “cells expressing the GFP-tagged protein (Figure 1)”.

We changed the position of the reference to Fig. 1 in this sentence to avoid implying that we will show cells expressing a GFP-tagged protein. Rather than naming the protein we want to highlight that this type of false-positive result can apply to any GFP-tagged protein tested, which is not recruited to the damage site.
References

**Competing Interests:** No competing interests were disclosed.

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**Referee Report 29 January 2019**

https://doi.org/10.5256/f1000research.19537.r43632

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**Vincent Dion**
UK Dementia Research Institute, Cardiff University, Cardiff, UK

This short report by Hurst and Gasser exposes an important experimental detail that is often, but not always, overlooked: Hoechst-sensitized cells irradiated with UV light produce a signal in the same region of the spectrum as GFP. Consequently, without the proper controls, it may lead to false positives. That is, that it may be falsely concluded that GFP-tagged proteins are recruited to laser-induced damage.

In the discussion, they point out potential controls to that would prevent an experimenter to make this mistake. Specifically, they propose leaving out Hoechst or DAPI altogether, using the appropriate control cells that do not express the GFP-tagged protein of interest, using another nuclear marker, such as a protein localizing to the nuclear periphery, or using a different fluorescent protein. Those are all good ways of getting around the reported problem. I would also add the use of a UVC source (Dabin *et al.*, 2018).

**References**

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**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Not applicable
Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** DNA repair, chromatin, expanded trinucleotide repeat disorders

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 31 Mar 2019

**Verena Hurst,** Friedrich Miescher Institute Basel, Switzerland

Thank you for your comments! In version 2 we have added a section discussing the different types of damage generated at different UV/VIS wavelengths.

**Competing Interests:** No competing interests were disclosed.

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