The plant-specific DDR factor SOG1 increases chromatin mobility in response to DNA damage

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**Abstract**

Homologous recombination (HR) is a conservative DNA repair pathway in which intact homologous sequences are used as a template for repair. How the homology search happens in the crowded space of the cell nucleus is, however, still poorly understood. Here, we measure chromosome and double-strand break (DSB) site mobility in Arabidopsis thaliana, using lacO/LacI lines and two GFP-tagged HR reporters. We observe an increase in chromatin mobility upon the induction of DNA damage, specifically at the S/G2 phases of the cell cycle. This increase in mobility is lost in the sog1-1 mutant, a central transcription factor of the DNA damage response in plants. Also, DSB sites show particularly high mobility levels and their enhanced mobility requires the HR factor RAD54. Our data suggest that repair mechanisms promote chromatin mobility upon DNA damage, implying a role of this process in the early steps of the DNA damage response.

**Keywords** Arabidopsis; chromatin mobility; DNA damage; DSBs; SOG1

**Subject Categories** Chromatin, Transcription & Genomics; DNA Replication, Recombination & Repair

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**Introduction**

Genome integrity is constantly threatened by internal and external stressors. Therefore, in response to DNA damage, eukaryotic organisms evolved elaborate DNA-damage response (DDR) systems that comprise DNA-damage signaling processes, DNA repair, and other responses such as cell death and control of cell division (Yoshiyama et al., 2013b). Among the different types of DNA damage, double-strand breaks (DSBs) are particularly harmful to cells, leading potentially to chromosome rearrangements or loss of entire chromosome arms (van Gent et al., 2001). DSBs can be repaired by two main pathways, nonhomologous end joining (NHEJ) and homologous recombination (HR; Jackson, 2002; West et al., 2004). NHEJ is achieved by stabilization and re-ligation of broken DNA ends, often with loss or mutation of bases. HR is a more complex and more conservative mechanism in which intact homologous sequences are used as a template for repair. HR most commonly occurs in S/G2 phases of the cell cycle in eukaryotic cells when sister chromatids are present, although homologous donor templates present elsewhere in the genome can also be used (Johnson & Jasin, 2000; Li & Hoyer, 2008; Goldfarb & Lichten, 2010). Despite the vast knowledge about the molecular players involved in DNA repair via HR, the mechanisms behind the search and recognition of homologous sequences (“homology search”) are still not well understood. In yeast, large-scale movements of DSBs have been identified following DSB induction (Oza et al., 2009; Dion et al., 2012; Ryu et al., 2015; Schrank et al., 2018). Yet, the precise functions of these movements remain unclear.

Plants are potentially subject to particularly high levels of DNA damage resulting from dependence on sunlight for energy and exposure to environmental stresses (Ries et al., 2000; Lee et al., 2012; Kawarazaki et al., 2013; Küpper & Andresen, 2016; Zhao et al., 2018). Moreover, plant development is mostly postembryonic with a late germline differentiation. Therefore, it is particularly interesting to understand the mechanisms that allow these organisms to cope with the constant assaults on their genome integrity. Indeed, plants have evolved a distinct DDR master regulator—SUPPRESSOR OF GAMMA RESPONSE 1 (SOG1). This transcription factor initiates a repair response by inducing genes involved in cell cycle arrest and repair, as well as in programmed stem-cell death in response to DNA damage (Yoshiyama et al., 2009, 2013a; Bourbousse et al., 2018). While the molecular processes involved in DDR pathway have been extensively characterized also in plants, little has been done to address how chromatin mobility changes in response to DNA damage and in particular to DSBs. Here, we have used locus tagging systems and HR reporter lines to study chromatin mobility upon genotoxic stress with the DSB inducer agent zeocin. We observed that in the presence of DSBs, both damaged and potentially undamaged loci increase the volume that they explore within the nuclear space. We showed that this increase in chromatin mobility occurs specifically during the G2 phase of the cell cycle and depends on the plant-specific DDR master regulator SOG1, implying...
an important role for chromatin mobility during the early steps of the DNA damage response.

Results and Discussion

To measure chromatin mobility in plant cells, we used the lacO/LacI-GFP locus-tagging system (Matzke et al., 2010, 2019; Fig 1A). For simplicity and to be consistent with previous studies, we will refer to the term chromatin mobility, even though we are measuring the mobility of individual foci at a given time. Foci mobility measurements were carried out using a mean square displacement (MSD) analysis. This analysis robustly measures the mobility of diffusing, fluorescently tagged chromosomal loci and provides kinetic parameters describing loci motion (Horigome et al., 2015; Meschichi & Rosa, 2021). We first tested our setup by measuring “steady-state” chromatin mobility levels for cells in the division versus differentiation zones of the Arabidopsis thaliana (Arabidopsis) root (Fig 1B). Measurements of histone exchange had previously shown that cells at the division zone have a more dynamic chromatin state as compared to differentiated cells (Rosa et al., 2014; Arai et al., 2017). Consistently, we observed that chromatin mobility is also higher in cells from the division zone compared to cells from the differentiation zone (Fig 1C). The radius of constraint (Rc), which indicates the nuclear volume within which a fluorescent spot can move, was therefore significantly higher in cells from the division zone (Fig 1C). These results confirmed that our setup is suitable to unpick differences in chromatin mobility between cells. In Arabidopsis root, differences in nucleus size are often evident, not only between nuclei from the division and differentiated zones but also within the meristematic region. As such, we thought to verify if our MSD measurements would be affected by differences in nucleus size. Within the meristematic region from the root, cells have the same ploidy level (diploid), but nuclei of atrichoblast cells are considerably bigger than that of trichoblast cells (Fig 1D). Nevertheless, these two cell types show the same chromatin mobility and radius of constraint (Fig 1E), ruling out that the nuclear volume per se could affect overall chromatin mobility levels.

Because HR requires pairing of the broken DNA molecule with a homologous intact template, we tested whether Arabidopsis cells actively regulate chromatin mobility in response to DSBs. We induced DNA damage by incubating 6-day-old seedlings with the DSB inducer zeocin for 24 h (Fig 2A). This treatment led to the upregulation of the DDR-responsive genes PARPB1, RAD51, and BRCA1, indicating that the HR was effectively stimulated (Appendix Fig S1A). This provided us with a system to induce different levels of DNA damage. We further focused our analysis on cells within the division zone since previous studies showed that the principal actors of HR, RAD51 and RAD54, are mainly expressed in these cells (Da Ines et al., 2013; Hirakawa et al., 2017). MSD analysis revealed that lacO/LacI foci mobility was not changed upon low concentrations or shorter times of zeocin incubation but increased significantly with high concentrations of zeocin for 24 h (Fig 2B; Appendix Fig S2). We acknowledge that 170 μM zeocin is a very high concentration, and many damage sites are induced simultaneously (Appendix Fig S3). While we cannot rule out that effects at other cellular processes may take place at this concentration, similar doses have been applied in other systems and are required to bring the level of damage above a certain threshold that triggers the increase in chromatin mobility or to increase the probability of inducing a break near to lacO transgene (Seeber et al., 2013). Importantly, the effect seen at the higher concentration was not due to DNA damage-induced programmed cell death as tested by PI staining (Appendix Fig S4). Only stem cells and their early descendants, which are known to be highly sensitive to DNA damage (Fulcher & Sablowski, 2009), showed PI-positive staining but not the epidermal cells used in our chromatin mobility analysis. We also tested another DSB inducer chemical, mitomycin C (MMC). A similar increase in chromatin mobility was observed in response to MMC treatment (Appendix Fig S5), showing that this is a general response to DSB induction.

In order to verify if the increase in chromatin mobility observed upon zeocin treatment was specific for the particular lacO insertion site in line112 or a response at the global chromatin level, we analyzed additional lacO/LacI lines with insertions at different chromosomal locations (Fig 2C). In control conditions, line 26 shows the same chromatin mobility as line 112, whereas line 107 showed significantly lower chromatin mobility and Rc (Appendix Fig S6). The lower mobility in line 107 could be linked to the transgene insertion at the subtelomeric region, known to physically interact at the nucleolar periphery in Arabidopsis (Armstrong et al., 2001; Fransz et al., 2002; Pontvianne et al., 2016; Fig 2C). Upon treatment with high zeocin concentration, all lines showed a significant increase in chromatin mobility and Rc (Fig 2D and E), indicating that chromatin mobility increases globally in the nucleus in response to DNA damage. We also tested whether these results could be an artifact of the lacO/LacI system itself. For that, we performed the same experiments using another locus tagging system—the ANCHOR system (ParB-parS; Meschichi et al., 2021; Fig 2F). The ANCHOR line showed a similar increase in chromatin mobility (Fig 2G).

Existing evidence in several systems shows that cell cycle arrest upon DNA damage is used by cells to facilitate DNA repair before cell division (Johnson & Jasmin, 2000; Weimer et al., 2016; Chang et al., 2017; Hustedt & Durocher, 2017). Since DNA content and cohesion differ in different cell cycle phases, we sought to test if changes in cell cycle dynamics (i.e. the proportion of cells in different cell cycle phases) could explain the increased chromatin mobility observed in response to DNA damage. To test this hypothesis, we crossed the lacO/LacI (line 112) with the S/G2 reporter CDT1a::RFP (Yin et al., 2014; Fig 3A and B). To first verify that this setup was working as expected, we quantified the ratio of cells in S/G2 in root epidermal cells treated with Hydroxyurea (HU), a drug known to block cells in S phase (Cools et al., 2010; Singh & Xu, 2016). Indeed, we observed that there was a higher proportion of cells in S/G2 in HU samples (Fig 3C). Consistent with previous studies (Chen et al., 2017), treatment with 10 μM zeocin significantly increased the number of cells in S/G2 (Fig 3C). However, with the highest concentration of zeocin (170 μM), the ratio of cells in S/G2 phase decreased to half in comparison with control conditions (Fig 3C), suggesting an accumulation of cells in G1. Thus, it became important to determine if G1 cells had different chromatin mobility compared to S/G2 cells. MSD analysis revealed that cells in the S/G2 phase (CDT1a-RFP positive cells) showed lower chromatin mobility than G1 cells (Fig 3D). Similarly, HU-treated cells showed lower chromatin mobility, most likely due to cells being arrested in
the S/G2 phase (Fig 3E). These results revealed that an accumulation of cells in G1 could potentially explain the increased mobility observed in response to DSBs. If this is the case, we hypothesized that we should not see differences when comparing cells at the same stage of the cell cycle with or without zeocin. We, therefore, measured the chromatin mobility specifically at G1 and S/G2, in control conditions and upon treatment with different concentrations of zeocin. Consistent with previous studies in yeast (Dion et al., 2012; Cheblal et al., 2020), we observed a significant increase in chromatin mobility in cells at S/G2 after zeocin treatment, whereas cells in G1 did not show any significant change (Fig 3F and G). We concluded that the increased mobility observed in response to DNA damage at high zeocin concentrations (170 µM) could be both a result of an accumulation of cells in G1 and a specific increase in chromatin mobility.
mobility at S/G2 phase. This observation is consistent with the idea that HR is particularly relevant in G2 when sister chromatids have been synthesized and suggests that increased chromatin mobility may be important during this stage.

In yeast, as in plants, studies have shown that HR is executed mainly during S/G2 phases of the cell cycle (Ferreira, 2004; Weimer et al., 2016). Because the increase in chromatin mobility upon zeocin treatment was specific to S/G2, we decided to investigate the mobility of actual break sites (DSBs) during HR. Homologous recombination is divided into two main phases: the presynaptic phase, which includes 5’-end resection and homology search, and the synaptic phase, which includes the strand invasion for homologous strand pairing (Fig 4A; Wright et al., 2018). Two main actors of HR, RAD51 and RAD54, function in the initiation of the strand invasion and at the strand exchange reaction that finalizes the repair (Solinger & Heyer, 2001). We wanted to investigate how the increase in chromatin mobility is placed in relation to these two phases. By performing an 8-h time course experiment on RAD51-GFP and

Figure 2. DNA damage increases chromatin mobility.
A Scheme illustrating the experimental setup for Arabidopsis seedling treatment.
B Left: MSD analysis of lacO/LacI line 112 based on time-lapse experiments of nuclei in different zeocin (Zeo) concentrations. Control (n = 116 nuclei); 10 μM (n = 97 nuclei); 170 μM (n = 93 nuclei). Right: Radius of constraint was calculated from MSD curves. Values represent means ± SEM. Letters indicate one-way ANOVA followed by Bonferroni’s correction (P < 0.05).
C Chromosomal positions of lacO/LacI lines as reported previously (Matzke et al., 2010, 2019). Line 26, line 107, and line 112 are respectively inserted in chromosomes 2, 3, and 5. The ANCHOR construct is inserted in chromosome 5. The NORs are marked as black circles and centromeres as light gray circles.
D Left: MSD analysis of lacO/LacI line 107 based on time-lapse experiments of nuclei in control conditions and plants treated with 170 μM zeocin. Control (n = 53 nuclei), 170 μM zeocin (n = 48 nuclei). Right: Radius of constraint calculated from MSD curves. Values represent means ± SEM. Student’s t-test, *P < 0.05.
E Left: MSD analysis of lacO/LacI line 26 based on time-lapse experiments of nuclei upon zeocin treatment. Control (n = 52 nuclei), 170 μM zeocin (n = 52 nuclei). Right: Radius of constraint calculated from MSD curves. Values represent means ± SEM. Student’s t-test, *P < 0.05.
F Left: Schematic representation of the ANCHOR system. parS-ParB-GFP interactions and oligomerization along the flanking genomic region. ParB-GFP can directly bind to parS sequence as a dimer and along the flanking genomic region. Right: Representative image of epidermis nuclei in the division zone. Scale bar, 5 μm.
G Left: MSD analysis of ANCHOR line based on time-lapse experiments of nuclei upon zeocin treatment. Control (n = 54 nuclei), 170 μM zeocin (n = 22 nuclei). Right: Radius of constraint calculated from MSD curves. Values represent means ± SEM. Student’s t-test, *P < 0.05.
RAD54-YFP lines after induction of damage with 10 μM zeocin, we were able to visualize the appearance of foci with accumulations of these proteins at DSB sites in the nucleus (Fig 4B and C). The tagged version of RAD51 forms the nucleofilament at DSBs but is defective in recombination and repair (Kobayashi et al., 2014). This defect leads to very high nucleoplasmic fluorescence with high zeocin concentrations thereby preventing individual foci from being visualized. Therefore, for these experiments, a low concentration of zeocin (10 μM) was required. RAD51-GFP foci were formed approximately 1 h 30 min after DSB induction, whereas RAD54-YFP foci appeared later, at around 5 h after treatment (Fig 4B). From this experiment, we can infer that RAD51 interacts first with DSBs, while RAD54 comes in later. To investigate the mobility of foci tagged with these proteins, we treated RAD51-GFP and RAD54-YFP plants with 10 μM zeocin (Fig 4C). The MSD analysis revealed that only RAD51 showed significantly higher mobility than lacO/LacI foci (Fig 4D), showing that high mobility levels seem to happen at early HR stages. Additionally, the high levels of mobility observed with RAD51-GFP could be due to the fact that these correspond to DSB sites, whereas lacO/LacI foci mobility most likely corresponds to measurements at undamaged loci. RAD54 foci mobility however seemed to be at the level of lacO/lacI at 10 μM zeocin (Fig 4D). Previous studies have shown that RAD54 foci relocate to the nuclear periphery after c-irradiation (Hirakawa & Matsunaga, 2019). Therefore, our MSD results for RAD54 may correspond to a mixture of foci located at the nuclear periphery and non-periphery. To test if RAD54 at the different nuclear compartments behaved differently, we determined the MSD for RAD54 at these two nuclear locations (Fig 4E).
**Figure 4. The mobility of DSB sites via tagged HR factors.**

A Schematic representation of the critical steps of homologous recombination. RAD51 (purple) assembles onto the single-stranded DNA (ssDNA) formed after the resection of DNA double-strand break (DSB) ends to form a filament, which is known as the presynaptic filament. After searching for DNA homologous sequence, the presynaptic filament binds the DNA template to form the synaptic structure with RAD54. The ssDNA invades the homologous region in the duplex to form a DNA joint, known as the displacement (D)-loop promoted by Rad54 (green).

B Time-lapse experiment of the formation of RAD51-GFP and RAD54-YFP foci in Arabidopsis nuclei, imaged every 30 min. Timeline of RAD51 and RAD54 foci formation for 8 h. The middle line in the box shows the mean, and the whiskers represent the standard error. At least four roots were counted for each line.

C Representative images of root epidermal cells showing foci formation in RAD51-GFP and RAD54-YFP plants after 10 µM zeocin treatment for 48 h. PI staining (red).

D Left: MSD analysis of RAD51-GFP (n = 64 nuclei) and RAD54-YFP (n = 64 nuclei) foci and lacO/LacI (line 112; n = 109 nuclei) plants upon 10 µM zeocin. Right: Radius of constraint calculated from MSD curves. Values represent means ± SEM. Letters indicate one-way ANOVA followed by Bonferroni’s correction (P < 0.05)

E Representative image of a root epidermal nucleus with RAD54-YFP foci located on the nuclear periphery (p) and non-periphery (n). Scale bar, 5 µm.

F Left: MSD analysis of RAD54 foci in the periphery (n = 24 nuclei) and non-periphery (n = 30 nuclei) upon 10 µM zeocin. Right: Radius of constraint calculated from MSD curves. Values represent means ± SEM. Student’s t-test, ***P < 0.001.
results showed that non-peripheric RAD54 foci have much higher mobility than the foci at the periphery (Fig 4F), revealing that RAD54 foci can have mobilities similar to those of RAD51. Moreover, these results highlighted that large changes in DSB site mobility occur during the repair process—a strong increase in DSB mobility is observed in the early HR phase, with a subsequent drastic drop in mobility associated with the relocation of DSBs to the nuclear periphery. This relocation to the nucleus periphery has been associated with different possible roles—to bring homologous sequences together, thereby reducing the 3D search to a 2D scale (Seeber & Gasser, 2017); or due to the fact that the repair machinery may specifically interact with nucleopores (Nagai et al., 2008). These observations support the hypothesis that higher DNA movement is induced at early steps of HR presumably to facilitate the 3D search.
Tracking chromatin movement, using DNA labeling tools and HR reporter lines, showed an increase in mobility upon DNA damage. Next, we wanted to determine whether the increase in mobility was actively regulated by the DDR pathway. For that, we quantified lacO/LacI (line 112) mobility in sog1-1 mutant, in which DDR is abolished. MSD analysis in sog1-1 mutant revealed no increase in mobility upon treatment with high zeocin concentration, indicating that the increase of mobility seen in the WT (SOG1+/− progeny from the F1) was dependent on SOG1 and thus on DDR activation (Fig S5A and B). However, it is important to rule out that the lack of response to zeocin treatment was not due to a change in the cell cycle dynamics in this mutant. Indeed, in sog1-1 the cell cycle arrest upon DNA damage is compromised (Yi et al., 2014; Chen et al., 2017; Mahapatra & Roy, 2021) and a loss of G1-arrested cells could potentially explain the results observed. We used EdU staining to check if, under our zeocin treatment conditions, sog1-1 cells were not being arrested in G1 (Fig SC-E; Appendix Fig S7). The results showed that also in sog1-1 there is a substantial reduction in EdU staining upon zeocin treatment, indicating that cells are also being accumulated at G1 although to a lesser extent than in the WT. We, therefore, decided to further analyze chromatin mobility in G1 and S/G2 in sog1-1 mutant. Given the complexity of this line, with several T-DNA insertions, instead of crossing it with CDT1α::RFP reporter we used nuclear area as a proxy for cell cycle stage taking as a reference CDT1 labeling (Appendix Fig S8). This analysis revealed that in sog1-1 at both G1 and S/G2 stages of the cell cycle, there is no increase in mobility upon zeocin treatment (Fig SF-H). These results demonstrate that SOG1 is required for the increase in chromatin mobility induced by zeocin treatment, indicating that this phenomenon is actively regulated during the early steps of the response to DNA damage and not a physical by-product from extensive DNA “fragmentation.” Previous studies in yeast and animals have shown that increased chromatin mobility is dependent on the protein kinases ATM/ATR (Dion et al., 2012; Becker et al., 2014), which have a primary role in DDR and promote SOG1 activation (Yoshiyama et al., 2009, 2013a). It will now be relevant to define which molecular players downstream of SOG1 are involved in the increased chromatin mobility in plants. RAD51 and RAD54 are two factors that act downstream of SOG1 and are essential for HR. RAD51 forms filaments on ssDNA, while RAD54 is a Snf2-type ATPase with translocase activity. As recruitment of RAD51 to DSBs does not depend on RAD54 (Hernandez Sanchez- Rebato et al., 2021), we generated a RAD51-GFP line containing a mutation on RAD54 (rad54-2) and followed the dynamics of RAD51 foci on this mutant line. Rad54-2 mutation resulted in a significant decrease in RAD51-GFP foci mobility to levels similar to those of undamaged sites (Fig S1 and J). Thus, we concluded that this important component of the HR machinery (RAD54) is involved in DSB mobility in plants. These results are in agreement with studies in yeast (Dion et al., 2013), which show that the mobility of damaged DNA increases in a RAD54-dependent manner. Whether RAD54 has a role beyond DSB mobility and is required for global changes in chromatin mobility remains to be addressed. However, previous studies in Arabidopsis have shown the involvement of RAD54 in pairing of lacO/LacI foci (Hirakawa et al., 2015), suggesting a potential role at the global chromatin level.

Our analysis of global chromosome and double-strand break (DSB) site mobility, using lacO/LacI lines and RAD51-GFP and RAD54-YFP reporters, has revealed that an increase in chromatin mobility occurs in response to DNA damage in Arabidopsis. Similar responses have been observed in yeast and animal cells, pointing towards a general mechanism of response to DSBs across kingdoms (Dimitrova et al., 2008; Dion et al., 2012; Krawczyk et al., 2012; Miné-Hattab & Rothstein, 2012). Although the exact function of such an increase in mobility has not been fully uncovered, some studies suggest it could increase the probability of an encounter between the break and the repair template (Barzel & Kupiec, 2008; Gehlen et al., 2011). However, this hypothesis needs to be tested with experiments that directly link chromatin mobility and HR efficiency. Even though the DNA repair machinery is highly conserved among eukaryotes, some of the most important regulators in animals, such as the tumor suppressor p53, are absent in plants. Its function is instead served by the plant-specific DDR master regulator SOG1. Interestingly, we have been able to show that in plants the increase in chromatin mobility is dependent on SOG1 function. These results suggest that the increase in chromatin mobility was conserved in evolution, as a response to DNA damage potentially through the action of different molecular players. A deeper understanding of the mechanisms downstream of SOG1 directly responsible for the increased chromatin mobility upon DNA damage in plants is needed in future studies.

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**Figure 5.** SOG1 and RAD54 regulate mobility in response to DNA damage.

A Left: MSD analysis of Arabidopsis lacO/LacI line 112 crossed with sog1-1 (Control (n = 83 nuclei); 170 μM zeocin (n = 91 nuclei)). Right: Radius of constraint calculated from MSD curves.

B Left: MSD analysis of SOG1+/− lacO/LacI progeny from crossing with sog1-1 (Control (n = 59 nuclei); 170 μM zeocin (n = 29 nuclei)). Right: Radius of constraint calculated from MSD curves.

C Schematic representation of the experimental setup for EdU labeling.

D Schematic representation of cell cycle progression, with the EdU signal displayed in S/G2 phase.

E The proportion of EdU-labeled cells in root tips of Col-0 and sog2-1, in control conditions and upon treatment with 170 μM zeocin. For each condition, between 5 and 6 roots were analysed (n = more than 50 nuclei).

F MSD analysis of G1 cells from lacO/LacI line 112 crossed with sog2-1 mutant in control conditions (n = 23 nuclei) and in plants treated with 170 μM zeocin (n = 25 nuclei).

G MSD analysis of cells in S/G2 from lacO/LacI line 112 crossed with sog2-1 (Control (n = 10 nuclei); 170 μM zeocin (n = 12 nuclei)).

H Radius of constraint calculated from MSD curves depicted in F and G.

I MSD analysis for RAD51-GFP foci in wild-type plants (n = 14 nuclei), RAD51-GFP foci in rad54-2 mutant background (n = 31 nuclei) and lacO/LacI (line112, n = 109 nuclei) plants treated with 10 μM zeocin.

J Radius of constraint calculated from MSD curves depicted in I.

Data information: Values represent means ± SEM. Student’s t-test, *P < 0.05, ***P < 0.001. Letters indicate one-way ANOVA followed by Bonferroni’s correction (P < 0.05).
Material and Methods

Plant lines and growth conditions

Mutants and transgenic lines used in this study come from the following sources: sog1-1 (Yoshiyama et al., 2009), rad54-2 (Hernandez Sanchez-Rebato et al., 2021), RAD51-GFP (Da Ines et al., 2013), RAD54-eYFP (Hirakawa & Matsunaga, 2019), Cytrap line (Yin et al., 2014), lacO/LacI lines (Matzke et al., 2010), and ANCHOR line (Meschichi et al., 2021). All mutants and transgenic lines are in Columbia background.

To visualize S/G2 cells in the lacO/LacI line 112, we crossed this line with Cytrap line. The resulting F2 plants were selected on MS plates containing 50 mg/l of kanamycin (Sigma-Aldrich, catalog number K1377). Because the G2/M-marker CYCB1;1 is strongly expressed during DNA damage (Culligan et al., 2006), the selected F2 were screened only for LacI-GFP and CDT1a-RFP.

Seeds were sterilized in 5% v/v sodium hypochlorite for 5 min and rinsed three times in sterile distilled water. Seeds were stratified at 4°C for 48 h in the darkness. Seeds were then plated on Murashige and Skoog (MS) solid medium and then grown in 16/8 h light/dark cycles at 22°C in vertically oriented Petri dishes. The roots were observed after 6 to 7 day of incubation, depending on the experiment.

Genotoxic treatment

To induce DNA damage response, 5- to 6-day-old seedlings were transferred in solid MS medium without or with 100 μM mitomycin C (MMC); 2, 10, 50, 100, or 170 μM zeocin or 10 mM hydroxyurea (HU) and treated for 2, 6, or 24 h. Each chemical was obtained respectively from Fisher Scientific (catalog number 2980501), Invitrogen (catalog number R25001), and Sigma-Aldrich (catalog number H8627-1G).

Microscopy

For root staining with propidium iodide (PI), 6- to 7-day-old seedlings were mounted in water between slide and coverslip and sealed with 0.12-mm-thick SecureSeal Adhesive tape (Grace Bio-Labs) to reduce drift drying during imaging.

For EdU staining and immunostaining, samples were imaged using a Zeiss LSM800 inverted microscope, with 63× water objective (1.20 NA) and Microscopy Camera Axiocam 503 mono; fluorescence of Alexa488 was detected using a 450–470 nm. For RAD51-GFP and RAD54-YFP time course experiments, samples were imaged using a Zeiss LSM780 inverted microscope, with 63× water objective (1.20 NA). Imaging was performed every 30 min for 8 h. Z-stacks of 19.5 μm size and 0.5 μm z-step were collected. Image size was 595 × 512 pixels with a zoom factor of 2x. RAD51-GFP and RAD54-YFP signals were detected using a 488 nm excitation line and collected between 493–598 nm.

Mean square displacement

For all MSD experiments, time-lapse imaging was performed every 6 s, taking a Z-stack of 3 μm spread through 1 μm slices for 5 min, with a 512 × 512 pixels format with a 1–2× zoom factor. All images were analyzed using Fiji software (NIH, Bethesda, MD, http://rsb.info.nih.gov/ij/; Sage et al., 2005) and with the plugin SpotTracker 2D (obtained from http://bigwww.epfl.ch/sage/soft/spottracker). All our time series were corrected for XY drift and nuclear movement using the plugins Stackreg and SpotTracker 2D, respectively. All the details of our image analyses pipeline were done as described in Meschichi and Rosa (2021).

Expression analysis using real-time RT-PCR (qPCR)

Seedlings grown for 7 days were harvested, and total RNA was extracted using the TRIzol reagent (Invitrogen). A total of 1 μg of RNA was treated with TURBO DNase (Life Technologies) and used for cDNA synthesis (Superscript IV; Life Technologies). The resulting cDNA was diluted 10 times and used for quantitative PCR using a Bio-Rad iCycler Thermal Cycler IQ5 Multicolor Real-Time and HOT FIREPol® EvaGreen® qPCR Mix Plus (Solis BioDyne). For data normalization, the data were first normalized to the PP2A2 reference gene, and the values from two independent samples were normalized to the average Delta Ct value Col-0 level or control condition (2-ΔΔCt Method). The final values presented are given as the mean ± SD from three independent samples. Minus RT (no reverse transcriptase control) controls were set up to make sure the values reflect the level of RNA and not DNA contamination. The standard Student’s t-test was used to determine the statistical significance of the results. The primers used are listed in Appendix Table S1.

EdU labeling

Five-day-old seedlings were grown on solid medium, treated with 170 μM zeocin for 24 h and incubated on solid medium containing 20 μM EdU and 170 μM zeocin during the last 6 h before imaging. Roots were fixed in 4% paraformaldehyde (PFA) for 30 min and washed three times with 1 × PBS. The roots were transferred to slide and covered by a glass cover slip, then squashed, and immediately dipped in liquid nitrogen for few seconds. The cover slips were removed, and the roots were left to dry at room temperature for 30 min. The samples were washed with PBS + BSA (Bovine Serum Albumin) 3% (w/v) and incubated with a Clickit Buffer (PBS 1× pH7.4, CuSO4 100 mM, Ascorbate 1 M, Alexa fluor azide, 2 μM) solution in the dark for 15 min. Samples were washed once in 1× PBS + BSA 3%, followed by DAPI staining for 15 min in the dark. Samples were washed twice with PBS 1× pH 7.4 and mounted in vectashield (Vector Laboratories).

Immunofluorescence

Roots were fixed in 4% PFA for 30 min and washed three times with 1 × PBS. The roots were transferred to slides and covered by a glass coverslip, then squashed, and immediately dipped in liquid nitrogen for few seconds. The coverslips were removed, and the roots were left to dry at room temperature for 30 min. Samples were then rinsed three times with 1× PBS solution and incubated with the enzyme mix (5% Driselase, 2.5% Cellulase, 5% Macerozyme in 1× PBS) for 15 min in a humid chamber at 37°C. Each slide was incubated overnight at 4°C with 50 μl rabbit, anti-plant γ-H2AX antiserum diluted 1:500 in fresh blocking buffer (0.5% BSA, in 1× PBS) and washed three times in 1×...
PBS solution. Slides were incubated for 2 h in a humid chamber at 37°C in 50 μl blocking buffer consisting of Alexa 488-conjugated goat anti-rabbit (1:1,000 Agrisera, catalog number: AS09633) secondary antibodies. Finally, slides were washed three times for 5 min in blocking buffer followed by DAPI staining for 15 min in the dark. Samples were washed twice with PBS 1× and mounted in vectashield (Vector Laboratories). Three-dimensional image stacks were captured, and γ-H2AX foci were counted manually.

The primary antibody was provided by C. White and used as performed in a previous study (Charbonnel et al, 2010).

Statistical analysis

For statistical analysis, we used the GraphPad Prism 8.3 software. Data sets were tested for normality using the Shapiro–Wilk test. Statistical significance was determined by using the standard Student’s t-test (two-tailed) and one-way ANOVA (multiple comparisons with Bonferroni correction). All experiments were performed in several nuclei as mentioned in figure legends.

Data availability

Data from this study are not deposited in external repositories, but can be requested from the corresponding author.

Expanded View for this article is available online.

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Anis Meschichi: Conceptualization; formal analysis; investigation; writing – original draft; writing – review and editing. Lihua Zhao: Investigation; formal analysis. Svenja Reeck: Investigation; formal analysis. Charles White: Methodology; writing – review and editing. Olivier Da Ines: Methodology; writing – review and editing. Adrien Sicard: Conceptualization; supervision. Frederic Pontvianne: Investigation; methodology. Stefanie Rosa: Conceptualization; formal analysis; supervision; funding acquisition; investigation; writing – original draft; project administration; writing – review and editing.

Disclosure and competing interest statement

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