Rad10 exhibits lesion-dependent genetic requirements for recruitment to DNA double-strand breaks in *Saccharomyces cerevisiae*

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Received December 12, 2008; Revised August 6, 2009; Accepted August 9, 2009

ABSTRACT

In the yeast *Saccharomyces cerevisiae*, the Rad1–Rad10 protein complex participates in nucleotide excision repair (NER) and homologous recombination (HR). During HR, the Rad1–Rad10 endonuclease cleaves 3' branches of DNA and aberrant 3' DNA ends that are refractory to other 3' processing enzymes. Here we show that yeast strains expressing fluorescently labeled Rad10 protein (Rad10-YFP) form foci in response to double-strand breaks (DSBs) induced by a site-specific restriction enzyme, I-SceI or by ionizing radiation (IR). Additionally, for endonuclease-induced DSBs, Rad10-YFP localization to DSB sites depends on both *RAD51* and *RAD52*, but not *MRE11* while IR-induced breaks do not require *RAD51*. Finally, Rad10-YFP colocalizes with Rad51-CFP and with Rad52-CFP at DSB sites, indicating a temporal overlap of Rad52, Rad51 and Rad10 functions at DSBs. These observations are consistent with a putative role of Rad10 protein in excising overhanging DNA ends after homology searching and refine the potential role(s) of the Rad1–Rad10 complex in DSB repair in yeast.

INTRODUCTION

DNA double-strand break repair (DSBR) is highly conserved between the yeast *Saccharomyces cerevisiae* and *Homo sapiens* (1,2). In both organisms, double-strand breaks (DSBs) are resected, generating 3' single-stranded regions that are coated with replication protein A (RPA). Subsequent strand invasion of single-stranded end(s) into a homologous duplex requires that Rad51 recombinase displaces RPA and forms a nucleoprotein filament. In mammalian cells, displacement of RPA and activation of Rad51 is mediated by BRCA2 (3). In budding yeast, Rad52 protein facilitates formation of Rad51 filaments by displacing RPA protein and recruiting proteins of the *RAD52* epistasis group (2,4). Among these, Rad52 functions downstream of the Mre11–Rad50–Xrs2 (MRX) complex (5). Hence, Rad52 function is required in advance of the functions of many proteins participating late in homologous recombination (HR) (downstream of key nuclease processing events), including those required for homology searching, and excision of extraneous DNA following homology searching and DNA ligation.

The Rad1–Rad10 endonuclease complex in *S. cerevisiae* participates both in HR and in nucleotide excision repair (NER), recognizing junctions between single- and double-stranded DNA, and cleaving in the double-stranded region a few nucleotides 5' to the junction to remove 3' nonhomologous ends (6,7). Recent genetic studies have demonstrated that the complex is also required for microhomology-mediated end joining (MMEJ), a mode of DSB repair that functions independently of Rad52. MMEJ is also Ku-independent (Yku70–Yku80), and is thus distinct from nonhomologous end-joining (NHEJ) (8–10). In human cell lines ERCC1 protein, the human homolog of Rad10, also facilitates a mode of homology-mediated DSB repair that is Ku86-independent (11,12). On the basis of existing evidence, the most likely model is that the Rad1–Rad10 endonuclease functions downstream of Rad52, cleaving nonhomologous DNA ends.
prior to gap filling and DNA ligation of the recombination site. However, the precise timing of Rad1–Rad10 recruitment to DSB sites relative to other DSB repair proteins has not been directly demonstrated. Additionally, recent work has revealed that Saw1 recruits the Rad1–Rad10 complex to DSBs in the context of single-strand annealing (SSA), one mode of DSB repair (13). Other than for SSA, the mechanism by which Rad1–Rad10 is recruited to DSBs remains largely unclear (13).

Recent studies have demonstrated the utility of fluorescence microscopy in monitoring biochemical events in live yeast cells (5,14). Three advantages of this experimental approach are especially noteworthy. First, the gene that encodes a given fluorescent protein under study is in the same chromosomal location as the native gene, and is also under control of its endogenous promoter. Thus, physiologically irrelevant overexpression associated with the use of ectopic genes expressed from non-native promoters is avoided. Second, this technique avoids fixation of cells prior to staining with fluorescent antibodies, as is typically the case in mammalian fluorescence microscopy experiments. Thus, the dynamics of the proteins under study are observed in real time in live cells, allowing for the accurate analysis of temporal relationships (14). Third, HR is monitored at the single-cell level, which allows for the analysis of individual repair events separately. Using this technology, the present study demonstrates that yeast Rad10 tagged with yellow fluorescent protein (Rad10-YFP) is recruited directly to DSB repair foci.

When a site-specific DSB is induced by the restriction endonuclease I-sceI in a DNA sequence context lacking direct repeat sequences, recruitment of Rad10-YFP to DSB foci requires functional RAD51 and RAD52 genes, but not MRE11. In contrast, when DSBs are induced by ionizing radiation (IR) during which DNA damage can occur in random DNA sequence contexts including those with direct repeats (6), we find that recruitment of Rad10-YFP to Rad52-CFP foci is RAD51-independent. These observations provide the first direct evidence that Rad1–Rad10 functions downstream of Rad52 during DSB repair and that its recruitment to DSB sites has different genetic requirements depending on the method by which DSB induction occurs and therefore, probably, the location of the DSB within the genome and the level of DNA sequence homology flanking the DSB. Moreover, the results support a model of HR in which the primary role of Rad1–Rad10 is cleavage of DNA ends following homology searching.

MATERIALS AND METHODS

Engineering of yeast strains Rad10-YFP and rad10A

The *S. cerevisiae* RAD10 gene was genetically fused in frame at the chromosomal locus with the DNA coding region of YFP to prepare the Rad10-YFP strain in the haploid W303-1A genetic background by the adaptamer-mediated PCR method as described previously (15). The resulting strain was backcrossed to the W3646-11D strain to produce WPF006-4C, which was used in microscopy experiments. The presence and identity of the YFP tag were confirmed by PCR, fluorescence microscopy and DNA sequencing which showed no mutations.

Expression of the full-length Rad10-YFP polypeptide was confirmed by immunoblotting of yeast whole cell extracts (WCEs) from appropriate strains using an α-Rad10 antibody. The Rad10 polyclonal antibody was prepared by overexpression of Rad10 in *E. coli*, purification and rabbit immunization. SDS PAGE was conducted with NuPAGE® Novex 4–12% Bis-Tris (Invitrogen) protein gels with 2-(N-Morpholino) ethanesulfonic acid (MES) running buffer used according to manufacturer’s recommendations to separate proteins prior to immunoblotting. Bands were sized with Benchmark™ protein standards (Invitrogen).

An isogenic disruption mutant of *RAD10* (PF033-6C) was prepared by PCR amplification of the KanMX deletion cassette from the YML095C strain (#6492) from the BY4741 deletion library and transformation into the W303-1A background. Transformants were selected on YPD plates containing 400 mg/ml G418 (Gibco) and backcrossed to wild-type strain W1588-4C to give PF033-6C. The presence of the deletion cassette was confirmed by PCR. All other strains used in this study were prepared by genetic crosses. The rad51A and rad52A genotypes were confirmed by gamma-irradiation (400 Gy) sensitivity.

UV survival studies

The UV sensitivity of the Rad10-YFP strain was tested using standard methodologies in parallel with isogenic wild-type and deletion mutant controls (16). Triplicate experiments were conducted and the results averaged and reported along with the corresponding standard error.

SSA/gene conversion assays

SSA/gene conversion (GC) assays were conducted essentially as previously described (17,18). Strains were co-transformed with the pJF6 reporter plasmid and a plasmid containing the HO endonuclease under the control of the GAL promoter, pFH800. Transformants were cultured in YEP-Lactate to a cell density of 5 × 10^6 to 1 × 10^7 cells/ml, induced by addition of 1/10 volume of 20% aqueous galactose (w/w) and incubated with shaking (2–4h). Plasmid DNA was isolated, digested with *Hind*III, *Pst*I and *Smal* restriction enzymes, resolved on a 0.8% agarose gel (1 × TAE, 50 V, 21h), and transferred to Immobilon-Ny+ Transfer Membrane (Millipore). Following transfer, the membrane was hybridized (12h, 68°C) with a digoxigenin-labeled (Roche) DNA probe prepared by PCR amplification of an ~800 bp fragment using primers 5'-d(CGTCATAAGCATAAGG) and 5'-d(CGTCGGGGATAGTTTTCTTGC). The probe was detected using a standard α-digoxigenin/CSPD protocol and manufacturer’s instructions (Roche). Quantity One image analysis software (BioRad) was used for densitometric calculations.
General microscopy

Cultures for microscopy experiments were propagated in SC medium supplemented with 200 μg/ml adenine (SC + ade) at 23°C unless otherwise indicated. Stationary phase cultures were freshly diluted (to 0.1 OD_{600}) and incubated (3 h). Aliquots of the cultures were then centrifuged (3600g), resuspended in residual medium and admixed on glass microscope slides with an equal volume of identical medium at 42°C containing 1% (w/v) low-melting agarose (Fischer). Microscopy was performed on a Zeiss AxioImager M1 microscope with a Plan–Apochromat 100×, 1.4 numerical aperture (NA) objective oil immersion lens, a motorized Z-drive and automated shutters. The fluorescent light source was an X-Cite 120 IRIS Fluorescent Light Source. IR experiment images for Figure 6 were acquired on a DM5500B Leica fluorescence microscope with a 100×, 1.46 NA objective and a mercury fluorescence light source. For both microscopes, images were captured with a Hamamatsu ORCA-ER digital camera (CA4742-80-12AG). Images at different focal planes through the entire thickness of the cells were captured at 0.3 μm intervals along the Z-axis (a Z-stack). Filter sets employed were Yellow GFP, Chroma #41028, (λex = 500/20, λ dichroic = 515, λ em = 535/30), CFP, Chroma #31044V2 (λex = 436/20, λ dichroic = 455, λ em = 480/40) and NAR/EX DsRed, Chroma #41035, (λex = 546/11, λ dichroic = 560, λ em = 605/75). Integration time for image acquisition was 800 ms for Rad10-YFP, 400 ms for Rad52-CFP and 400 ms for TetR-RFP. For both setups, microscopes and cameras were controlled by the Volocity software package (Improvision v 3.7.0 for the Zeiss or v 5.0.2 for the Leica). Foci were counted by inspecting images from each focal plane of the Z-stack. As is commonly the case in fluorescence microscopy work, with each fluorophore it was necessary to contrast enhance the original images, which was optimized using the gain, offset and gamma contrast parameters of Volocity. The same contrast enhancement settings were used for all images of a given fluorophore. For publication quality images, the settings were re-optimized so that foci could be seen when printed and, for merged images, the opacity and fill settings were changed so that the various colors could be viewed in a single image.

Cells were classified with respect to phase in the cell cycle based on their appearance in both the fluorescence images and in differential interference contrast (DIC) images. Unless otherwise noted, for each trial of each strain, at least 100 cells were counted. Graphs of the data report averages of at least three independent experiments with the corresponding standard error. In the case of mre11A data, P-values were calculated using a two-tailed Student’s t-test, to determine whether mre11A data were statistically significantly different from wild-type (19). Images were prepared for publication using the Adobe Photoshop and Adobe Illustrator software packages (Adobe Systems, Mountain View, CA). For images showing triple colocalization, only single-plane images were captured and the exposure time for Rad10-YFP was adjusted to 3 s. For single-plane image experiments, at least 600 galactose-induced cells (total) were analyzed for the appearance of colocalized foci from two independently conducted experiments of at least 300 cells each.

Induction of DSBs by I-SceI endonuclease

Strains for DSB induction experiments bear chromosomally integrated copies of the Tetracycline repressor protein (TetR) fused to RFP (TetR-RFP) and 224 copies of the Tetracycline operator (tetO) repressor binding site abutted with one copy of the I-SceI cut site at the iYER186 intergenic region on chromosome V as described (20). The site is cut by the I-SceI endonuclease with 60–70% efficiency in asynchronously growing cells (20,21). The strains used in I-SceI DSB induction experiments included PF025-7A, PF030-49A, PF023-15A, WP0F019-26C and PF034-11D. Strains were transformed with plasmid pWJ1320, containing the I-SceI gene under control of the GAL1 promoter and containing the ADE2 selectable marker as described (4). Transformants were plated on SC–ade medium containing 2% raffinose (SC–ade w/raff). Cultures for DSB induction experiments were propagated in SC–ade w/ raff liquid medium at 23°C. Overnight cultures were diluted to 0.1 OD_{600} and incubated (3 h). DNA DSBs were subsequently induced by adding galactose to the medium to a final concentration of 2% and incubating the cultures (4 h) prior to processing cells for microscopy as described above.

DAPI-stained images

Cell cultures were grown as described under ‘General Microscopy’ and centrifuged. The cell pellet was resuspended in fixative solution (2.5% formaldehyde, 30 mM sodium phosphate, pH 7) and incubated (5 min). Cells were centrifuged (3600g), washed twice with water, resuspended in 1 μl residual water and admixed on glass microscope slides with DAPI mounting medium [SC + ade, 1% (w/v) low-melting agarose (Fischer), 50 ng/ml 4’,6-diamidino-2-phenylindole (DAPI)]. In addition to DIC and YFP images, cells were imaged using the YFP filter set and a DAPI/Hoechst/AMCA filter set. Chroma #31000v2, (λex = 350/50, λ dichroic = 400, λ em = 460/50). YFP exposures were 3 s and DAPI exposures were 20 ms.

Induction of DSBs by gamma irradiation

Gamma induction experiments were conducted as described above for I-SceI induction experiments except for the following experimental alterations. Strains WP0F019-26C and W8757-12A which did not contain the plasmid pWJ1320 were cultured in SC + ade medium, diluted to an OD_{600} = 0.1 and grown (3 h), irradiated with gamma radiation (40 Gy, 60Co source), incubated (2 h) and imaged using the same conditions as for I-SceI experiments.
Table 1. Strains used in this study

| Name      | Genotype                                                                 | Published in |
|-----------|--------------------------------------------------------------------------|--------------|
| SX46a     | MATa ade2-1 trpl-289 his3-532 ura3-2                                   | (34)         |
| BJRrad10Δ | MATa rad10::URA3 leu2 trpl-32 ura3-1 pep4-1 prb1-1122 prec1-407          | From EC Friedberg laboratory strain collection |
| W303-1A   | MATa ade2-1 lys2A trpl-1 can1-100 his3-11,15 leu2-3,112 ura3-1 rad5-535   | (35)         |
| W1588-4C  | MATa ade2-1 lys2A trpl-1 can1-100 his3-11,15 leu2-3,112 ura3-1            | (36)         |
| W3646-11D | MATa lys2A TRP1 can1-100 his3-11,15 bar1::LEU2 ura3-1                    | (35)         |
| WPF006-4C | MATa lys2A trpl-1 can1-100 his3-11,15 leu2-3,112 ura3-1 Rad10-YFP        | This manuscript |
| PF033-6C  | MATa rad10::His3 ade2-1 can1-100 his3-11,15:YFP-lacI-R197L::HIS3       | This manuscript |
| PF044-2A  | MATa ade2-1 rad10::Rad10-YFP rad51-CFP                                 | This manuscript |
| WPF033-6C | MATa ade2-1 rad10::KanMX his3-11,15 leu2-3,112 trpl-1 ura3-1             | This manuscript |
| ML47-1D   | MAT::HIS3 ade2-1 trpl-1 can1-100 his3-11,15:YFP-lacI-R197L::HIS3       | A derivative of strains described in supplementary section of (14) |
| WPF025-7A | MATa ade2-1 trpl-1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |
| WPF021-4A | MATa ade2-1 trpl-1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |
| WPF030-49A| MATa ade2-1 trpl-1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |
| WPF023-15A| MATa ade2-1 trpl-1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |
| WPF019-26C| MATa ade2-1 trpl-1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |
| WPF034-11D| MATa ade2-1 trpl-1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |
| W8757-12A | MATa ade2-1 TRP1 can1-100 his3-11,15 leu2-3,112 TetR-mRFP (YGL119W)  | This manuscript |

All strains in this study are haploid and derivatives of W303-1A and W303-1B (35) except SX46a and BJRrad10Δ. Additionally, all strains are wild-type for the ADE2 and RAD5 genes unless otherwise noted.

RESULTS

Rad10 protein in the yeast S. cerevisiae was tagged at its C-terminal end by integrating the gene encoding the yellow-shifted version of green fluorescent protein from Aequorea victoria in-frame at the RAD10 chromosomal locus, using adaptamer-mediated PCR as previously described (15). The resulting yeast strain expresses a Rad10-YFP fusion protein (Rad10-YFP) that migrates as a 24 kDa band (compare lane 2 and 4 with lanes 5–7 in Figure 1A). A control lane containing GFP protein alone (lane 1) exhibits no signal with α-Rad10 antibody, and extracts from a rad10 deletion mutation (rad10Δ, lane 3) exhibit only background signal of the antibody. As expected, by fluorescence microscopy, the Rad10-YFP strain exhibits yellow fluorescence in the nucleus as indicated by colocalization of the YFP signal with DAPI-stained DNA in the nucleus (Figure 1B).

Next, the functionality of the Rad10-YFP protein in vivo was examined. A plasmid-based assay was used to test Rad10-YFP function in SSA and GC (Figure 2A). Densitometric analysis of genome blot results (Figure 2B) shows that the efficiencies of SSA and GC are somewhat lower (86 and 54%, respectively) in the YFP-tagged strain compared with wild-type. However, the fluorescent label does not compromise the function of Rad10 protein during NER, as evidenced by examining the UV radiation sensitivity of the Rad10-YFP strain.

Indeed, the Rad10-YFP strain exhibits wild-type levels of survival following exposure to UV-C irradiation (Figure 2C).

To determine whether Rad10 protein is recruited to DSBs, such lesions were generated in vivo in a strain containing both Rad10-YFP and a unique DSB labeling system previously reported (Figure 3A) (4). Specifically, a yeast strain was prepared that expresses the coding sequence for the TetR fused to monomeric red fluorescent protein (TetR-RFP). The strain additionally contains a DNA cassette with 224 copies of the TetR binding sequence abutted with one copy of the I-SceI restriction endonuclease site. I-SceI enzyme was introduced into the strain via a selectable extrachromosomal plasmid under the control of the GAL1 promoter (pWJ1320) (4). Subsequent induction of I-SceI generates a single DSB in close physical proximity to many copies of TetR-RFP (DNA-RFP) within the nucleus (Figure 3B). Such induction of DSBs results in increased punctate yellow foci (Figure 3B, Rad10-YFP panel), indicating the presence of a high local concentration of Rad10-YFP. A significant subset of Rad10-YFP foci colocalized with DNA-RFP foci in DSB-induced cells, indicating that Rad10-YFP protein is indeed recruited to DSB sites (Figures 3B and 4). Rad10-YFP foci were also observed spontaneously in uninduced cells, but in fewer numbers and they rarely colocalized with the DSB site (see below). Some of these
spontaneous events likely reflect the recruitment of the tagged protein to spontaneous DSBs, as previously reported for proteins of the RAD52 epistasis group, suggesting that Rad10 is recruited to both endonuclease-induced and naturally occurring DSBs (4). Indeed, cells co-expressing Rad10-YFP and Rad52-CFP exhibited spontaneous Rad52-CFP foci that colocalized with Rad10-YFP (Figure 6). Since spontaneous Rad10 foci were consistently observed at a higher frequency (70–80%) than spontaneous Rad52 foci (5–10%), many of the Rad10 foci likely reflect the involvement of Rad10 in other DNA repair pathway(s), such as NER and MMEJ.

To determine the cell cycle dependence of Rad10 foci, cells were scored not only for Rad10-YFP and DSB-RFP foci, but also for cell cycle phase (S, G2/M or G1, see ‘Materials and Methods’ section). S phase cells manifested the greatest magnitude of DSB-colocalized Rad10 foci, exhibiting approximately a 12-fold increase in DSB-induced cells over that in uninduced controls (Figure 4, wild-type). Colocalized foci were also induced in G2/M and G1 cells, but to a much lesser extent (Figure 4), suggesting that the repair of DSBs by HR transpires primarily during S phase of the cell cycle, presumably due to ongoing functions of Rad10 such as repair of spontaneous DSBs and NER. The great majority (70–80%) of uninduced cells contained at least one Rad10-YFP focus. However, upon DSB induction the increase in the numbers of Rad10-YFP foci matched the increase in the numbers of Rad10-YFP foci colocalized to DSB sites.

We next examined the genetic requirements of recruitment of Rad10-YFP to DSBs. Isogenic strains deleted for the MRE11, RAD52 or RAD51 genes (mre11A, rad52Δ or rad51A) were transformed with the 1-3βGal expression plasmid, pWJ1320. DSBs were induced and the formation
of DSB-colocalized Rad10-YFP foci was monitored in parallel with wild-type controls. The frequency of DSB-colocalized Rad10-YFP foci observed in the \( mre11 \Delta \) strain was not appreciably altered relative to the wild-type, ( \( P \)-values > 0.19 in all phases of the cell cycle) indicating that Mre11 protein is not required to recruit Rad10-YFP to DSBs (Figure 4, compare black wild-type and \( mre11 \Delta \) bars). Hence, while visual examination of Figure 4 suggests a small reduction in the number of Rad10-YFP foci colocalized with DSB-RFP foci in the \( mre11 \Delta \) background, this difference is not statistically significant under the experimental conditions employed.

In contrast, \( rad52 \) and \( rad51 \) deletion mutants exhibited very few Rad10-YFP foci colocalized to sites of DSBs upon I-SceI induction (Figure 4, compare wild-type, \( rad52 \Delta \) and \( rad51 \Delta \) bars).

To determine whether Rad10 colocalizes with Rad52 or Rad51 at DSB sites, DSB induction experiments were conducted using triple-labeled Rad10-YFP/DSB-RFP/Rad52-CFP or Rad10-YFP/DSB-RFP/Rad51-CFP strains (Figure 5A). Triple-labeled experiments were imaged as single focal planes owing to photobleaching of the Rad10-YFP chromophore which precluded the formation of Z-stack images (see ‘Materials and Methods’ section). Accordingly, the overall percentage of cells containing Rad10-YFP is reduced in single-plane images (Figure 5B) versus Z-stack images (Figure 4), since a single focal plane reveals only a portion of the total thickness of the average cell.

In both strains, colocalization of all three labels (YFP, RFP and CFP) was observed, indicating that Rad10-YFP is present at DSB sites at the same time as Rad52-CFP (Figure 5A) or Rad51-CFP (data not shown).

Furthermore, \( \sim 75\% \) of the Rad10-YFP foci that colocalized with the DSB-RFP marker were also colocalized with Rad52-CFP, both in S and G2/M phase cells, while essentially no G1 cells exhibit similar triple colocalization (Figure 5B). Hence, most of the Rad10-YFP foci localized to the induced DSBs also had Rad52-CFP present indicating considerable temporal overlap of the functions of Rad10 with Rad52. Altogether, we conclude that Rad10 protein colocalizes with Rad51 and Rad52 proteins at DSBs during S and G2/M phases of the cell cycle.

Since the repair of DSBs induced by a restriction enzyme can differ from the repair of breaks induced by IR (20.21), a series of experiments were done using a gamma source to induce DSBs. Experiments were conducted in both \( RAD51 \) and \( rad51 \Delta \) strains containing Rad10-YFP and Rad52-CFP. Two hours post-irradiation, individual and colocalized foci were counted and compared with results from non-irradiated controls. Following IR, the numbers of cells containing at least one Rad10-YFP focus increased similarly in both the \( RAD51 \) and \( rad51 \Delta \) strains relative to non-irradiated controls (1.3- and 1.7-fold, respectively; data not shown). The numbers of cells containing at least one Rad10-YFP/Rad52-CFP colocalized focus also increased in both strains relative to non-irradiated controls (6.2-fold for \( RAD51 \) and 4.2-fold for \( rad51 \Delta \), respectively; data not shown). As with the I-SceI experiments, most of the foci were observed in the S, G2 or M phases of cell cycle. Upon IR induction, the number of S, G2 or M phase cells containing at least one colocalized...
Rad10-YFP/Rad52-CFP focus increased 4.8-fold for \textit{RAD51} cells and 2.5-fold for \textit{rad51}D cells (Figure 6). Thus, IR induction of colocalizing Rad10 and Rad52 foci is \textit{RAD51}-independent unlike that observed following I-SceI induction. Interestingly, in the absence of IR induction, the number of S, G2 and M phase cells containing at least one spontaneous colocalized Rad10-YFP/Rad52-CFP focus was 2.5-fold higher in \textit{rad51}D cells than in \textit{RAD51} cells (Figure 6).

**DISCUSSION**

Here we monitored in real time, in living yeast cells, several proteins involved in DNA repair, which are expressed from their native chromosomal locations and endogenous promoters. This approach provides a more physiologically relevant aspect to \textit{in vitro} biochemical experiments involving cell free extracts or overexpression systems. Our experiments demonstrate that Rad10-YFP foci are induced in response to the formation of DNA DSBs, and that such foci are localized specifically to such sites in \textit{S. cerevisiae}. The total number of cells that contained at least one Rad10-YFP focus was high (70–80%) even without induction of DSBs and many of the foci observed are not localized to the induced DSBs. These foci presumably reflect other roles of the Rad1–Rad10 complex in yeast, notably NER, MMEJ and spontaneous mitotic recombination, and indicate that Rad1–Rad10 is active in a majority of cells at all phases of cell cycle. The observation that the number of spontaneous Rad10-YFP foci is reduced in G1 cells when compared with S and G2/M phase cells suggests that a significant fraction of foci observed is associated with DNA replicative activity, such as spontaneous mitotic recombination events.

Although the Rad10-YFP protein is regulated by its natural promoter and from its native site, it appears slightly compromised in SSA function (86% of the wild-type level) and GC function (54% of the wild-type level), the two DSB repair processes in which Rad10 plays a role (Figure 2B and data not shown). We suspect that this attenuated functionality is the result of a kinetic lag in the recruitment of Rad10-YFP to the repair site, the YFP tag occluding the binding of other accessory factors, or the YFP tag causing faster dissociation from DSB sites, thus limiting the time available to execute Rad1–Rad10 function. However, the fact that the YFP tag does not compromise Rad10 function in NER as
measured by UV resistance (Figure 2C) suggests that the Rad1–Rad10 endonuclease activity itself is fully functional. Indeed, the functionality of the Rad10-YFP fusion protein is comparable to other YFP-tagged DNA repair and checkpoint proteins (4).

Previous genetic studies have implicated RAD1 and RAD10 with RAD51 and RAD52 in DSB repair (6). However, this study constitutes the first direct experimental evidence for the involvement of either RAD51 or RAD52 in recruiting Rad10 to DSBs. It is well established that RAD1 and RAD10 are required for several modes of HR in yeast, including SSA, MMEJ and synthesis-dependent strand annealing (SDSA) (6). Here we show that two ways of creating of DSBs (I-SceI and IR) give rise to the formation of Rad10-YFP foci and recruitment to sites of DSBs. However, the genetic requirements observed for Rad10 recruitment are dependent on how and, hence, where we create the break in the genome and likely reflect which modes of repair are being used.

Following IR, Rad10-YFP/Rad52-CFP colocalized foci are induced (Figure 6), suggesting that Rad10 is recruited to IR-induced DSBs. The extent of induction is approximately the same whether or not the RAD51 gene is present (Figure 6). These observations auger well with existing genetic and biochemical evidence for the several modes of DSB repair requiring RAD10, not all of which require RAD51 (6). For example, ribosomal DNA, which makes up almost 10% of the genome, is a multiple tandem array containing 150–200 repeats. When a DSB occurs in one of these repeats, it is largely repaired by SSA (6,17,22). Similarly, events initiated in other repetitive sequences such as Ty elements or delta sequences can also be repaired by SSA (23). SSA requires RAD52 but does not strictly require RAD51 (22,24,25). Since DSBs induced by IR occur in a broad range of DNA sequence contexts throughout the genome, many sequences flanked by DNA repeats are likely repaired by SSA (Figure 7). Therefore, we explain the 2.5-fold increase in the percentage of rad51A cells containing spontaneous Rad10-YFP/Rad52-CFP colocalized foci relative to that observed in RAD51 cells (Figure 6) by a shift in repair of spontaneous damage from RAD51-dependent GC to RAD51-independent SSA.

In contrast, the I-SceI-induced DSBs in this study (Figures 4 and 5) are not flanked by DNA repeats and therefore not repaired by SSA. We find that RAD51 is required for recruitment of Rad10 to the I-SceI sites, consistent with SSA-independent repair. It is unlikely these breaks are repaired via MMEJ, another mode of DSB repair known to require both RAD1 and RAD10 since MMEJ has an absolute requirement for MRE11 (8,10) and we show that Rad10 foci form independently of MRE11. The break-induced replication (BIR) pathway, which requires RAD52 and has both RAD51-dependent and RAD51-independent modes could, in principle, be consistent with the I-SceI-induced repair observed in this study (26,27). However, we do not think it is the preferred pathway since a strict requirement for RAD1 or RAD10 in BIR has not been established (28–30).

To explain the RAD52 and RAD51 dependence we observe for Rad10 focus formation after I-SceI-induced DSBs, we suggest that the breaks are repaired by SDSA (Figure 7). In a dividing cell where one sister chromatid has been cleaved by I-SceI and the other is still intact, SSA would be initiated by RAD1 and RAD51-dependent strand invasion. The branched structure may be recognized by Rad1–Rad10 (Figure 7). Indeed, genetic evidence for the requirements of RAD52, RAD51 and RAD10 in SDSA is consistent with this notion (6,31,32).

Results from experiments in which triply labeled strains containing Rad10-YFP/Rad51-CFP/DSB-RFP or Rad10-YFP/Rad52-CFP/DSB-RFP were induced with I-SceI show that a significant number of Rad10-YFP/DSB-RFP foci also contained either Rad51-CFP or Rad52-CFP suggesting that significant temporal overlap occurs in the functions of Rad10 with either Rad51 or Rad52 (Figure 5B). The observation that I-SceI-induced
DSBs require Rad51 to recruit Rad10 leads us to speculate that Rad51 strand-exchange activity may be required for efficient recruitment of Rad10 to I-SceI-induced DSBs. Conceivably, the strand-exchange reaction generates a DNA structure in which a region of duplex DNA is abutted next to a region lacking homology in the 3’ direction (i.e. a 3’ nonhomologous end). It is possible that Rad51 or Rad52 recruits the Rad1–Rad10 complex at the point-in-time when the junction between homologous and nonhomologous DNA is encountered. It will be interesting in the future to determine whether a rad51 mutant deficient in strand-exchange activity can still recruit Rad10 to I-SceI-induced DSBs and to determine the extent to which Rad51 and/or Rad52 precedes Rad10 temporally during a repair event. Nonetheless, our existing data support a model in which the Rad1–Rad10 complex functions after Rad51 and Rad52 helping to execute the final processing of DNA ends by trimming DNA ends prior to ligation.

In humans, Rad52 protein participates in a three-protein complex with XPF/ERCC1 endonuclease, the human counterpart of Rad1–Rad10 (33). However, it remains to be determined whether an analogous three-protein complex involving Rad52 and the Rad1–Rad10 endonuclease is generated in yeast, and, if so, whether Rad52 protein recruits Rad1–Rad10 to DSB sites via a direct physical interaction. The triple-labeled colocalization experiments presented herein (Figure 5) show that Rad10 and Rad52 are both present at DSB sites at the same time, but do not provide evidence for a direct protein–protein interaction since other proteins may ultimately ‘bridge’ direct physical interaction between Rad52 and the Rad1–Rad10 complex as has been shown for Saw1 in processing recombination intermediates (13). It will be of interest to see whether a direct Rad52–Rad1–Rad10 complex forms in vitro, and if so, whether this interaction is required for the recruitment of Rad10.

ACKNOWLEDGEMENTS

The authors thank Dr Neal Sugawara and Dr Jim Haber for kindly providing plasmids and strains for the SSA assay. They thank Jeff Wagner and the Radiation Safety Department at UCLA for generously providing access to their 137Cs gamma irradiator in order to genotype rad51Δ and rad52Δ strains.

FUNDING

California State University Northridge’s Competition for Research, Scholarship and Creative Activity Awards [to P.L.F.], the National Institutes of Health [5-R01-ES011344 to E.C.F., SC2GM081155 to P.L.F., GM50237, GM67055 to R.R.], the Danish Agency for Science, Technology and Innovation [to M.L.], the Villum Kann Rasmussen Foundation [to M.L.] and Fundación Alfonso Martín Escudero [to S.G.-B.]. Funding for open access charge: National Institutes of Health (SC2GM081155).

Conflict of interest statement. None declared.

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