Synthetic viability genomic screening defines Sae2 function in DNA repair

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Abstract

DNA double-strand break (DSB) repair by homologous recombination (HR) requires 3’ single-stranded DNA (ssDNA) generation by 5’ DNA-end resection. During meiosis, yeast Sae2 cooperates with the nuclease Mre11 to remove covalently bound Spo11 from DSB termini, allowing resection and HR to ensue. Mitotic roles of Sae2 and Mre11 nuclease have remained enigmatic, however, since cells lacking these display modest resection defects but marked DNA damage hypersensitivities. By combining classic genetic suppressor screening with high-throughput DNA sequencing, we identify Mre11 mutations that strongly suppress DNA damage sensitivities of sae2Δ cells. By assessing the impacts of these mutations at the cellular, biochemical and structural levels, we propose that, in addition to promoting resection, a crucial role for Sae2 and Mre11 nuclease activity in mitotic DSB repair is to facilitate the removal of Mre11 from ssDNA associated with DSB ends. Thus, without Sae2 or Mre11 nuclease activity, Mre11 bound to partly processed DSBs impairs strand invasion and HR.

Keywords Mre11; Sae2; suppressor screening; synthetic viability; whole-genome sequencing

Introduction

The DSB is the most cytotoxic form of DNA damage, with ineffective DSB repair leading to mutations, chromosomal rearrangements and genome instability that can yield cancer, neurodegenerative disease, immunodeficiency and/or infertility (Jackson & Bartek, 2009). DSBs arise from ionising radiation and radiomimetic drugs and are generated when replication forks encounter single-stranded DNA breaks or other DNA lesions, including DNA alkylation adducts and sites of abortive topoisomerase activity. DSBs are also physiological intermediates in meiotic recombination, being introduced during meiotic prophase I by the topoisomerase II-type enzyme Spo11 that becomes covalently linked to the 5’ end of each side of the DSB (Keeney et al, 1997). The two main DSB repair pathways are non-homologous end-joining (NHEJ) and homologous recombination (Lisby et al, 2004; Symington & Gautier, 2011). In NHEJ, DNA ends need little or no processing before being ligated (Daley et al, 2005). By contrast, HR requires DNA-end resection, a process involving degradation of the 5’ ends of the break, yielding 3’ single-stranded DNA (ssDNA) tails that mediate HR via pairing with and invading the sister chromatid, which provides the repair template.

Reflecting the above requirements, cells defective in resection components display HR defects and hypersensitivity to various DNA-damaging agents. This is well illustrated by Saccharomyces cerevisiae cells harbouring defects in the Mre11–Rad50–Xrs2 (MRX) complex, which binds and juxtaposes the two ends of a DSB (Williams et al, 2008) and, through Mre11 catalytic functions, provides nuclease activities involved in DSB processing (Furuse et al, 1998; Williams et al, 2008; Stracker & Petrini, 2011). Once a clean, partially resected 5’ end has been generated, the enzymes Exo1 and Sgs1/Dna2 are then thought to act, generating extensive ssDNA regions needed for effective HR (Mimitou & Symington, 2008; Zhu et al, 2008). Notably, while Mre11 nuclease activity is essential in meiosis to remove Spo11 and promote 5’ end resection, in mitotic cells, resection is only somewhat delayed in the absence of Mre11 and almost unaffected by mre11-nd (nuclease-dead) mutations (Ivanov et al, 1994; Moreau et al, 1999), indicating the existence of MRX-nuclease-independent routes for ssDNA generation.

Another protein linked to resection is S. cerevisiae Sae2, the functional homolog of human CtIP (Sartori et al, 2007; You et al, 2009). Despite lacking obvious catalytic domains, Sae2 and CtIP have been reported to display endonuclease activity in vitro (Lengsfeld et al, 2007; Makharashvili et al, 2014; Wang et al, 2014), and their functions are tightly regulated by cell cycle- and DNA damage-dependent phosphorylations (Baroni et al, 2004; Huertas et al, 2008; Huertas & Jackson, 2009; Barton et al, 2014). In many ways, Sae2 appears to

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function together with MRX in DSB repair. For instance, *mre11-nd* as well as *mre11S* and *rad50S* hypomorphic alleles phenocopy SAE2 deletion (sae2Δ) in meiosis, yielding unprocessed Spo11–DNA complexes (Keeney & Kleckner, 1995; Nairz & Klein, 1997; Prinz et al, 1997). Furthermore, recent findings have indicated that Sae2 stimulates Mre11 endonuclease activity to promote resection, particularly at protein-bound DSB ends (Cannavo & Cejka, 2014). Also, both sae2Δ and *mre11-nd* mutations cause hypersensitivity towards the anti-cancer drug camptothecin (Deng et al, 2005), which yields DBSs that are repaired by HR. Nevertheless, key differences between MRX and Sae2 exist, since *sae2* leads to persistence of MRX at DNA damage sites (Lisby et al, 2004) and hyperactivation of the MRX-associated Tel1 protein kinase (Usui et al, 2001), the homolog of human ATM, while MRX inactivation abrogates Tel1 function (Lisby et al, 2011). These findings, together with sae2Δ and *mre11-nd* cells displaying only mild resection defects (Clerici et al, 2005), highlight how Sae2 functions in HR cannot be readily explained by it simply cooperating with MRX to enhance resection.

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Notably, however, mre11-H37R rescued the hypersensitivity of sae2Δ cells to etoposide, which produces DSBs bearing 5'0 DNA ends bound to Top2 (Supplementary Fig S2B; deletion of ERG6 was used to increase permeability of the plasma membrane to etoposide), suggesting that significant differences must exist between the repair of meiotic and etoposide-induced DSBs.

Next, we examined the effects of mre11SUPsae2Δ alleles on Sae2-dependent DSB repair by single-strand annealing (SSA), using a system wherein a chromosomal locus contains an HO endonuclease cleavage site flanked by two direct sequence repeats. In this system, HO induction produces a DSB that is then resected until two complementary sequences become exposed and anneal, resulting in repair by a process that deletes the region between the repeats (Fishman-Lobell et al., 1992; Vaze et al., 2002; Fig 3C). Despite displaying only mild resection defects (Clerici et al., 2006), we observed that sae2Δ cells were defective in SSA-mediated DSB repair and did not resume cell cycle progression after HO induction as fast as wild-type cells, in agreement with published work (Clerici et al., 2005). Notably, mre11-H37R did not alleviate these sae2Δ phenotypes (Fig 3D and E).

Finally, we examined the effect of the mre11-H37R mutation on telomere-associated functions of the MRX complex and Sae2. It has been established that simultaneous deletion of SGS1 and SAE2 results in synthetic lethality/sickness, possibly due to excessive telomere shortening (Mimitou & Symington, 2008; Hardy et al., 2014). To test whether mre11-H37R can alleviate this phenotype, we crossed a sae2Δ mre11-H37R strain with a sgs1Δ strain. As shown in Supplementary Fig S2C, we were unable to recover neither sgs1Δsae2Δ nor sgs1Δsae2Δmre11-H37R cells, implying that mre11-H37R cannot...
supress this phenotype. In agreement with this conclusion, the mre11-H37R mutation did not negatively affect Mre11-dependent telomere maintenance as demonstrated by Southern blot analysis (Supplementary Fig S2D).

Together, the above data revealed that mre11SUPsae2 alleles suppressed sae2Δ DNA damage hypersensitivities but not sae2Δ meiotic phenotypes requiring Mre11-mediated Spo11 removal from recombination intermediates, nor mitotic SSA functions that have been attributed to Sae2-mediated DNA-end bridging (Clerici et al., 2005). Subsequent analyses revealed that suppression did not arise largely through channelling of DSBs towards NHEJ because the key NHEJ factor Yku70 was not required for mre11-H37R or mre11-H37Y to suppress the camptothecin sensitivity of a sae2Δ strain (Fig 3F). In addition, this analysis revealed that the previously reported suppression of sae2Δ-mediated DNA damage hypersensitivity by Ku loss (Mimitou & Symington, 2010; Foster et al., 2011) was considerably less effective than that caused by mre11-H37R or mre11-H37Y. Also, suppression of sae2Δnuclease activity by mre11SUPsae2Δ alleles did not require Exo1, indicating that in contrast to suppression of sae2Δ phenotypes by Ku loss (Mimitou & Symington, 2010), mre11-H37R and mre11-H37Y did not cause cells to become particularly reliant on Exo1 for DSB processing (Fig 3G). Further characterisations, focused on mre11-H37R, revealed that while not suppressing camptothecin hypersensitivity of an xrs2Δ strain (Fig 3H), it almost fully rescued the camptothecin hypersensitivity of a strain expressing the rad50S allele, which phenocopies sae2Δ by somehow preventing functional Sae2–MRX interactions that are required for Sae2 stimulation of Mre11 endonuclease activity (Keeney & Kleckner, 1995; Hopfner et al., 2000; Cannavo & Cejka, 2014; Fig 3I).

**H37R does not enhance Mre11 nuclease activity but impairs DNA binding**

To explore how mre11SUPsae2Δ mutations might operate, we over-expressed and purified wild-type Mre11, Mre11H37R and Mre11H37A (Fig 4A and Supplementary Fig S2F) and then subjected these to biochemical analyses. All the proteins were expressed at similar levels and fractionated with equivalent profiles, suggesting that the Mre11 mutations did not grossly affect protein structure or stability. Since Sae2 promotes Mre11 nuclease functions, we initially speculated that sae2Δ suppression would be mediated by mre11SUPsae2Δ alleles having intrinsically high, Sae2-independent nuclease activity. Surprisingly, this was not the case, with Mre11H37R actually exhibiting lower nuclease activity than the wild-type protein (Fig 4B). Furthermore, by electrophoretic mobility shift assays, we found that the H37R mutation reduced Mre11 binding to double-stranded DNA.

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**Figure 2. mre11-H37R suppresses the CPT hypersensitivity of sae2Δ cells.**

A Alignment of Mre11 region containing H37 in fungal species; secondary structure prediction is shown above.

B Western blot with anti-Mre11 antibody on protein extracts prepared from the indicated strains shows that mre11SUPsae2Δ alleles do not require Exo1, indicating that in contrast to suppression of sae2Δ nuclease activity by wild-type Mre11 (Supplementary Fig S2D).

C Alignment of Mre11 region containing H37 in fungal species; secondary structure prediction is shown above.
Conversely, mutation of H37 to alanine, which does not result in a supsae2Δ phenotype, did not negatively affect dsDNA-binding activity (Fig 4C) and only partially impaired ssDNA binding (Fig 4D).

Taken together with the fact that the lack of Sae2 only has minor effects on mitotic DSB resection (Clerici et al., 2005), the above results suggested that the sae2Δ suppressive effects of mre11SUPsae2Δ mutations were associated with weakened Mre11 DNA binding and ligase activity (Fig 4D).
Figure 4. Mre11H37R is impaired biochemically, particularly at the level of ssDNA binding.

A. Mre11 and Mre11H37R were purified to homogeneity from yeast cultures.
B. 3' exonuclease activity assay on Mre11 and Mre11H37R leading to release of a labelled single nucleotide, as indicated.
C, D. Electrophoretic mobility shift assays on Mre11, Mre11H37R and Mre11H37A with dsDNA (C) or ssDNA (D).
E. Quantification of mre11-H37R suppression of sae2Δ cell DNA damage hypersensitivity. Overnight grown cultures of the indicated strains were diluted and plated on medium containing the indicated doses of CPT. Colony growth was scored 3–6 days later. Averages and standard deviations are shown for each point.
F. Intragenic suppression of CPT hypersensitivity of mre11-nd (mre11-H125N) by mre11-H37R. Overnight grown cultures of the indicated strains were treated as in (E). Dotted lines represent data from (E). Averages and standard deviations are shown for each point.
G. Mre11 nuclease activity is not required for mre11-H37R-mediated suppression of sae2Δ CPT hypersensitivity. Overnight grown cultures of the indicated strains were treated as in (E). The dotted lines represent data from (E). Averages and standard deviations are shown for each point.
were not linked to effects on resection or Mre11 nuclease activity. In line with this idea, by combining mutations in the same Mre11 polypeptide, we established that \textit{mre11-H37R} substantially rescued camptothecin hypersensitivity caused by mutating the Mre11 active site residue His125 to Asn (Moreau et al, 2001; \textit{mre11-H125N}; Fig 4E and Supplementary Fig S2F and G), which abrogates all Mre11 nuclease activities and prevents processing of DSBs when their 5' ends are blocked (Moreau et al, 1999). Even \textit{sae2Δ mre11-H37R, H125N} cells were resistant to camptothecin and MMS, indicating that Mre11-nuclease-mediated processing of DNA ends is not required for H37R-dependent suppression, nor for DNA repair in this Sae2-deficient setting (Fig 4G and Supplementary Fig S2G). Furthermore, while \textit{sae2Δ} strains were more sensitive to camptothecin than \textit{mre11-H125N} strains, the sensitivities of the corresponding strains carrying the \textit{mre11-H37R} allele were comparable (compare curves 1 and 2 with 3 and 4 in Fig 4F) indicating that \textit{mre11-H37R} suppresses not only the \textit{sae2Δ}-induced lack of Mre11 nuclease activity, but also other nuclease-independent functions of Sae2. Nevertheless, \textit{mre11-H37R} did not rescue the camptothecin hypersensitivity of \textit{sae2Δ} cells to wild-type levels, suggesting that not all functions of Sae2 are suppressed by this \textit{MRE11} allele (Fig 4E and F).

**Identifying an Mre11 interface mediating sae2Δ suppression**

To gain further insights into how \textit{mre11\textsuperscript{SUPsae2}} alleles operate and relate this to the above functional and biochemical data, we screened for additional \textit{MRE11} mutations that could suppress camptothecin hypersensitivity caused by Sae2 loss. Thus, we propagated a plasmid carrying wild-type \textit{MRE11} in a mutagenic \textit{E. coli} strain, thereby generating libraries of plasmids carrying \textit{mre11} mutations. We then introduced these libraries into a \textit{sae2Δmre11Δ} strain and screened for transformants capable of growth in the presence of camptothecin (Fig 5A). Through plasmid retrieval, sequencing and functional verification, we identified 12 \textit{sae2Δ} suppressors, nine carrying single \textit{mre11} point mutations and three being double mutants (Supplementary Fig S3A). One single mutant was \textit{mre11-H37R}, equivalent to an initial spontaneously arising suppressor that we had identified. Among the other single mutations were \textit{mre11-L89V} and \textit{mre11-L89V}, both of which are located between Mre11 nuclease domains II and III, in a region with no strong secondary structure predictions (Fig 5B). Two of the three double mutants contained \textit{mre11-P110L} combined with another mutation that was presumably not responsible for the resistance phenotype (because \textit{mre11-P110L} acts as a suppressor on its own), whereas the third contained both \textit{mre11-Q70R} and \textit{mre11-G193S}. Subsequent studies, involving site-directed mutagenesis, demonstrated that effective \textit{sae2Δ} suppression was mediated by \textit{mre11-Q70R}, which alters a residue located in a highly conserved α-helical region (Fig 5C).

Ensuing comparisons revealed that the mutations identified did not alter Mre11 protein levels (Supplementary Fig S3B) and that \textit{mre11-Q70R} suppressed \textit{sae2Δ} camptothecin hypersensitivity to similar extents as \textit{mre11-H37R} and \textit{mre11-H37Y}, whereas \textit{mre11-L89V} and \textit{mre11-P110L} were marginally weaker suppressors (Fig 5D).

To map the locations of the various \textit{mre11\textsuperscript{SUPsae2}} mutations within the Mre11 structure, we used the dimeric tertiary structure (Schiller et al, 2012) of the \textit{Schizosaccharomyces pombe} Mre11 counterpart, Rad32, as a template to generate a molecular model of \textit{S. cerevisiae} Mre11. The resulting structure had a near-native QMEAN score (0.705 vs 0.778; Benkert et al, 2008), indicating a reliable molecular model. Strikingly, ensuing analyses indicated that the \textit{mre11\textsuperscript{SUPsae2}} mutations clustered in a region of the protein structure distal from the nuclease catalytic site and adjacent to, but distinct from, the interface defined as mediating contacts with dsDNA in the \textit{Pyrococcus furiosus} Mre11 crystal structure (Williams et al, 2008; Fig 5E; the predicted path of dsDNA is shown in black, while the \textit{mre11\textsuperscript{SUPsae2}} mutations and residues involved in nuclease catalysis are indicated in red and orange, respectively). Furthermore, this analysis indicated that H37 and Q70 are located close together, on two parallel α-helices and are both likely to be solvent exposed (Fig 5F). By contrast, the L89 side chain is predicted to be in the Mre11 hydrophobic core, although modelling suggested that the \textit{mre11-L89V} mutation might alter the stability of the α-helix containing Q70. We noted that, in the context of the Mre11 dimer, H37 and Q70 are located in a hemi-cylindrical concave area directly below the position where dsDNA is likely to bind (Fig 5E right, shown by pink hemispheres). Furthermore, by specifically mutating other nearby residues to arginine, we found that the \textit{mre11-L77R} mutation also strongly suppressed \textit{sae2Δ} camptothecin hypersensitivity (Fig 5G). As discussed further below, while it is possible that certain \textit{mre11\textsuperscript{SUPsae2}} alleles somehow influence the established dsDNA-binding interface of Mre11, we speculate that \textit{mre11-H37Y} and \textit{mre11-Q70R}, and at least some of the other suppressors, act by perturbing interactions normally mediated between the Mre11 hemi-cylindrical concave region and ssDNA (modelled in Fig 5G and discussed further below). Consistent with this idea, we found that the \textit{Mre11-Q70R} protein was markedly impaired in binding to ssDNA but not to dsDNA (Supplementary Figs S2E and S3C). However, because P110 lies in the ‘latching loop’ region of eukaryotic Mre11
Figure 5.

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that is likely to mediate contacts with Xrs2 (Schiller et al., 2012), sae2Δ suppression by this mutation might arise through altering such contacts. A recent report by L. Symington and colleagues reached similar conclusions (Chen et al., 2015).

Taken together, our findings suggested that, in addition to its established dsDNA-binding mode, Mre11 mediates distinct, additional functional contacts with DNA that, when disrupted, lead to suppression of sae2Δ phenotypes. Thus, we suggest that, during DSB processing, duplex DNA entering the Mre11 structure may become partially unwound, with the 5′ end being channelled towards the nuclease catalytic site and the resulting ssDNA—bearing the 3′ terminal OH—interacting with an adjacent Mre11 region that contains residues mutated in mre11Δsae2Δ alleles (Fig 5G and H). In this regard, we note that Mre11 was recently shown in biochemical studies to promote local DNA unwinding (Cannon et al., 2013). Such a model would explain our biochemical findings, and would also explain our biological data if persistent Mre11 binding to the nascent 3′ terminal DNA impairs HR unless counteracted by the actions of Sa2 or weakened by mre11Δsae2Δ alleles.

sae2Δ phenotypes reflect Mre11-bound DNA repair intermediates

A prediction arising from the above model is that Mre11 persistence and associated Tel1 hyperactivation in sae2Δ cells would be counteracted by mre11Δsae2Δ mutations. To test this, we constructed yeast strains expressing wild-type Mre11 or Mre11H37R fused to yellow-fluorescent protein (YFP) and then used fluorescence microscopy to examine their recruitment and retention at sites of DNA damage induced by ionising radiation. In line with published work (Lisby et al., 2004), recruitment of wild-type Mre11 to DNA damage foci was more robust and persisted longer when Sa2 was absent (Fig 6A). Moreover, such Mre11 DNA damage persistence in sae2Δ cells was largely attenuated by mre11-H37R (Fig 6A; compare red and orange curves). By contrast, mre11-H37R had little or no effect on Mre11 recruitment and dissociation kinetics when Sa2 was present (compare dark and light blue curves). Importantly, we found that HR-mediated DSB repair was not required for H37R-induced suppression of Mre11-focus persistence in sae2Δ cells, as persistence and suppression still occurred in the absence of the key HR factor, Rad51 (Fig 6B). Also, in accord with our other observations, we found that the rad50Δ allele caused Mre11 DNA damage-focus persistence in a manner that was suppressed by the mre11-H37R mutation (Fig 6C).

Previous work has established that Mre11 persistence on DSB ends, induced by lack of Sa2, leads to enhanced and prolonged DNA damage-induced Tel1 activation, associated with Rad53 hyperphosphorylation (Usui et al., 2001; Lisby et al., 2004; Clerici et al., 2006; Fukunaga et al., 2011). Supporting our data indicating that, unlike wild-type Mre11, Mre11H37R is functionally released from DNA ends even in the absence of Sa2, we found that in a mec1Δ background (in which Tel1 is the only kinase activating Rad53; Sanchez et al., 1996), DNA damage-induced Rad53 hyperphosphorylation was suppressed by mre11-H37R (Fig 7A).

While we initially considered the possibility that persistent Tel1 hyperactivation might cause the DNA damage hypersensitivity of sae2Δ cells, we concluded that this was unlikely to be the case because TEL1 inactivation did not suppress sae2Δ DNA damage hypersensitivity phenotypes (Supplementary Fig S3D). Furthermore, Tel1 loss actually reduced the ability of mre11-H37R to suppress the

Figure 6. mre11Δsae2Δ alleles bypass the need for Sa2 to remove Mre11 from DSB ends.

A IR-induced Mre11H37R foci (IRIF) persist for shorter times than Mre11-wt IRIF in exponentially growing sae2Δ cells (average and standard deviations from two or more independent experiments).

B Effects of sae2Δ and mre11-H37R on Mre11 IRIF persistence still occur when Rad51 is absent, revealing that Mre11 IRIF persistence causes defective HR (average and standard deviation from two independent experiments).

C mre11-H37R suppresses Mre11 IRIF persistence in exponentially growing rad50Δ cells (average and standard deviation from two independent experiments).
Figure 7. Tel1 participates in regulating Mre11 dynamics after DNA damage.

A mre11-H37R suppresses Tel1 hyperactivation induced by Mre11 IRIF persistence in sae2Δ cells.

B Deletion of Tel1 weakens the suppression of the sensitivity of a sae2Δ strain mediated by mre11-H37R.

C Deletion of Tel1 reduces the hyperaccumulation of Mre11 to IRIF and impairs the suppression of their persistence mediated by mre11-H37R (average and standard deviation from two independent experiments).

D mre11-H37R suppresses the sensitivity to CPT of a tel1Δ strain.

E Model for the role of MRX, Sae2 and Tel1 in response to DSBs.
camptothecin hypersensitivity of sae2Δ cells (Fig 7B). In accord with this, in the absence of Tel1, mre11-H37R no longer affected the dissociation kinetics of IR-induced Mre11 foci in sae2Δ cells (Fig 7C). Collectively, these data suggested that Tel1 functionally cooperates with Sae2 to promote the removal of Mre11 from DNA ends. In this regard, we noted that mre11-H37R suppressed the moderate camptothecin hypersensitivity of a tel1Δ strain (Fig 7D). We therefore propose that, while persistent DNA damage-induced Tel1 activation is certainly a key feature of mre11-nd, it is persistent binding of the MRX complex to nascent 3′ terminal DNA that causes toxicity in sae2Δ cells, likely through it delaying downstream HR events. Accordingly, mutations that reduce Mre11 ssDNA binding enhance the release of the Mre11 complex from DSB ends in the absence of Sae2, through events promoted by Tel1 (Fig 7E). In this model, Mre11 persistence at DNA damage sites is a cause, and not just a consequence, of impaired HR-mediated repair in sae2Δ cells.

Discussion

Our data help resolve apparent paradoxes regarding Sae2 and MRX function by suggesting a revised model for how these and associated factors function in HR (Fig 7E). In this model, after being recruited to DSB sites and promoting Tel1 activation, resection and ensuing Mec1 activation, the MRX complex disengages from processed DNA termini in a manner promoted by Sae2 and facilitated by Tel1 and Mre11 nuclease activity. Sae2 is required to stimulate Mre11 nuclease activity (Cannavo & Cejka, 2014) and subsequently to promote MRX eviction from the DSB end. However, our data suggest that Sae2 can also promote MRX eviction in the absence of DNA-end processing, as mre11-H37R suppresses the phenotypes caused by sae2Δ and mre11-nd to essentially the same extent. Thus, according to our model, when Sae2 is absent, both the nuclease activities of Mre11 and MRX eviction are impaired. Under these circumstances, despite resection taking place—albeit with somewhat slower kinetics than in wild-type cells—MRX persists on ssDNA bearing the 3′ terminal OH, thereby delaying repair by HR. In cells containing the mre11-H37R mutation, however, weakened DNA binding together with Tel1 activity promotes MRX dissociation from DNA even in the absence of Sae2, thus allowing the nascent ssDNA terminus to effectively engage in the key HR events of strand invasion and DNA synthesis (Fig 7E). Nevertheless, it is conceivable that abrogation of pathological Tel1-mediated checkpoint hyperactivation contributes to the resistance of sae2Δmre11-H37R cells to DNA-damaging agents. In this regard, we note that the site of one of the sae2Δ suppressors, P110, lies in the ‘latching loop’ region of eukaryotic Mre11 that is likely to mediate contacts with Xrs2 (Schiller et al., 2012), suggesting that, in this case, sae2Δ suppression might arise through weakening this interaction and dampening Tel1 activity.

Our results also highlight how the camptothecin hypersensitivity of strains carrying a nuclease-defective version of Mre11 does not reflect defective Mre11-dependent DNA-end processing per se, but rather stems from stalling of MRX on DNA ends. We propose that this event delays or prevents HR, possibly by impairing the removal of 3′-bound Top1 as is suggested by the fact that in S. pombe, rad50S or mre11-nd alleles are partially defective in Top1 removal from damaged DNA (Hartsuiker et al., 2009). This interpretation also offers an explanation for the higher DNA damage hypersensitivity of sae2Δ cells compared to cells carrying mre11-H125N alleles: while sae2Δ cells are impaired in both Mre11 nuclease activity and Mre11 evi- cition—leading to MRX persistence at DNA damage sites and Tel1 hyperactivation—mre11-H125N cells are only impaired in Mre11 nuclease activity. Indeed, despite having no nuclease activity, the mre11-H125N mutation does not impair NHEJ, telomere maintenance, mating type switching or Mre11 interaction with Rad50/Xrs2 or interfere with the recruitment of the Mre11–Rad50–Xrs2 complex to foci at sites of DNA damage (Moreau et al., 1999; Lisby et al., 2004; Krogh et al., 2005). In addition, our model explains why the mre11-H37R mutation does not suppress meiotic defects of sae2Δ cells, because Sae2-stimulated Mre11 nuclease activity is crucial for removing Spo11 from meiotic DSB 5′ termini. Finally, this model explains why mre11-H37R does not suppress the sae2Δ deficiency in DSB repair by SSA because the sae2Δ defect in SSA is suggested to stem from impaired bridging between the two ends of a DSB rather than from the persistence of MRX on DNA ends (Clerici et al., 2005; Andres et al., 2015; Davies et al., 2015). In this regard, we note that SSA does not require an extendable 3′-OH DNA terminus to proceed and so could ensue even in the presence of blocked 3′-OH DNA ends.

We have also found that the mre11-H37R mutation suppresses the DNA damage hypersensitivities of cells impaired in CDK- or Mec1/ Tel1-mediated Sae2 phosphorylation. This suggests that such kinase-dependent control mechanisms—which may have evolved to ensure that HR only occurs after the DNA damage checkpoint has been triggered—also operate, at least in part, at the level of promoting MRX removal from partly processed DSBs. Accordingly, we found that Tel1 deletion causes moderate hypersensitivity to camptothecin that can be rescued by the mre11-H37R allele, implying that the same type of toxic repair intermediate is formed in sae2Δ and tel1Δ cells and that in each case, this can be rescued by MRX dissociation caused by mre11-H37R (Fig 7E). Supporting this idea, it has been previously shown that resection relies mainly on Exo1 in both tel1Δ and sae2Δ cells (Clerici et al., 2006; Mantiero et al., 2007). We suggest that the comparatively mild hypersensitivity of tel1Δ strains to camptothecin is due to Tel1 loss allowing DSB repair intermediates to be channelled into a different pathway, in which Exo1-dependent resection (Mantiero et al., 2007) leads to the activation of Mec1, which can then promote Sae2 phosphorylation and subsequent MRX removal (Fig 7E). The precise role of Tel1 in these events is not yet clear, although during the course of our analyses, we found that the deletion of Tel1 reduced the suppressive effects of mre11-H37R on sae2Δ DNA damage sensitivity and Mre11-focus persistence. This suggests that, in the absence of Sae2, Tel1 facilitates MRX evasion by mre11-H37R, possibly by phosphorylating the MRX complex itself.

Given the apparent strong evolutionary conservation of Sae2, the Mre11–Rad50–Xrs2 complex and their associated control mechanisms, it seems likely that the model we have proposed will also apply to other systems, including human cells. Indeed, we speculate the profound impacts of proteins such as mammalian CtIP and BRCA1 on HR may not only relate to their effects on resection, but may also reflect them promoting access to ssDNA bearing 3′ termini so that HR can take place effectively. Finally, our data highlight the power of SVGS to identify genetic interactions—including those such that we have defined that rely on separation-of-function mutations rather than null ones—and also to inform on underlying biological and biochemical mechanisms. In addition to...
being of academic interest, such mechanisms are likely to operate in medical contexts, such as the evolution of therapy resistance in cancer.

Materials and Methods

Strain and plasmid construction

Yeast strains used in this work are derivatives of SK1 (meiotic phenotypes), YMV80 (SSA phenotypes) and haploid derivatives of W303 (all other phenotypes). All deletions were introduced by one-step gene disruption. pRS303-derived plasmids, carrying a wt or mutant MRE11 version, were integrated at the MRE11 locus in an mre11A::KanMX6 strain. Alternatively, the same strain was transformed with pRS416-derived plasmids containing wild-type or mutant MRE11 under the control of its natural promoter. Strains expressing mutated mre11-YFP were obtained in two steps: integration of a pRS306-based plasmid (pFP118.1) carrying a mutated version of Mre11 in a MRE11-YFP sae2ΔA strain, followed by selection of those ‘pop-out’ events that suppressed camptothecin hypersensitivity of the starting strain. The presence of mutations was confirmed by sequencing. Full genotypes of the strains used in this study are described in Supplementary Table S1; plasmids are described in Supplementary Table S2.

Whole-genome paired-end DNA sequencing and data analysis

DNA (1–3 μg) was sheared to 100–1,000 bp by using a Covaris E210 or LE220 (Covaris, Woburn, MA, USA) and size-selected (350–450 bp) with magnetic beads (Ampure XP; Beckman Coulter). Sheared DNA was subjected to Illumina paired-end DNA library preparation and PCR-amplified for six cycles. Amplified libraries were sequenced with the HiSeq platform (Illumina) as paired-end 100 base reads according to the manufacturer’s protocol. A single sequencing library was created for each sample, and the sequencing coverage per sample is given in Supplementary Table S3. Sequencing reads from each lane were aligned to the S. cerevisiae S288c assembly (RG4-1-1) from Saccharomyces Genome Database (obtained from the Ensembl genome browser) by using BWA (v0.5.9-r16) with the parameter ‘-q 15’. All lanes from the same library were then merged into a single BAM file with Picard tools, and PCR duplicates were marked by using Picard ‘MarkDuplicates’ library were then merged into a single BAM file with Picard tools, (v0.5.9-r16) with the parameter ‘-q 15’. All lanes from the same

MRE11 random mutagenesis

Plasmid pRS316 carrying MRE11 coding sequence under the control of its natural promoter was transformed into mutagenic XL1-Red competent E. coli cells (Agilent Technologies) and propagated following the manufacturer’s instructions. A plasmid library of ~3,000 independent random mutant clones was transformed into mre11Δsae2Δ cells, and transformants were screened for their ability to survive in the presence of camptothecin. Plasmids extracted from survivors loosing their camptothecin resistance after a passage on 5-fluoro-orotic acid (FOA) were sequenced and independently reintroduced in a mre11Δsae2Δ strain.

Molecular modelling

A monomeric molecular model of S. cerevisiae Mre11 was generated with the homology modelling program MODELLER (Sali & Blundell, 1993) v9.11, using multiple structures of Mre11 from S. pombe (PDB codes: 4FBW and 4FBK) and human (PDB code: 3T1I) as templates. A structural alignment of them was made with the program BATON (Sali & Blundell, 1990) and manually edited to remove unmatched regions. The quality of the model was found to be native-like as evaluated by MODELLER’s NDOPE (−1.2) and GA341 (1.0) metrics and the QMEAN server (Benkert et al, 2009) (http://swissmodel.expasy.org/qmean/) (0.705). The monomeric model was subsequently aligned on the dimeric assembly of the 4FBW template to generate a dimer, and the approximate position of DNA binding was determined by aligning the P. furiosus structure containing dsDNA (PDB code: 3DSC) with the dimeric model. All images were obtained using the PyMOL Molecular Graphics System.

Microscopy

Exponentially growing yeast strains carrying wild-type or mutant Mre11-YFP were treated with 40 Gy of ionising radiations with a Faxitron irradiator (CellRad). At regular intervals, samples were taken and fixed with 500 μl of Fixing Solution (4% paraformaldehyde, 3.4% sucrose). Cells were subsequently washed with wash solution (100 mM potassium phosphate pH 7.5, 1.2 M sorbitol) and mounted on glass slides. Images were taken at a DeltaVision microscope. All these experiments were carried out at 30°C.

In vitro assays

For the electrophoretic mobility shift assay (EMSA), a radiolabelled DNA substrate (5 nM) was incubated with the indicated amount of Mre11 or Mre11H37R in 10 μl buffer (25 mM Tris–HCl, pH 7.5, 1 mM DTT, 100 μg/ml BSA, 150 mM KCl) at 30°C for 10 min. The reaction mixtures were resolved in a 10% polyacrylamide gel in TBE buffer (89 mM Tris–HCl, pH 8.0, 2 mM EDTA). The gel was dried onto Whatman DE81 paper and then subjected to phosphorimaging analysis. For nuclease assay, 1 mM MnCl2 was added to the reactions and the reaction mixtures were incubated at 30°C for 20 min and deproteinised by treatment with 0.5% SDS and 0.5 mg/ml proteinase K for 5 min at 37°C before analysis in a 10% polyacrylamide gel electrophoresis in TBE buffer.

Additional Materials and Methods can be found in the Supplementary Methods.

Supplementary information for this article is available online: http://emboj.embopress.org
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Author contributions

The initial screening was conceived and designed by TO, EV, DJA and SPJ. Alignment of whole-genome sequencing data, variant calling and subsequent analysis was carried out by MH and TMK. Experiments for the in vivo characterisation of the mre11–h37r mutant were conceived by TO, IG, FP and SPJ, and were carried out by TO, FP, IG, NGJ, EV and IS. Biochemical assays were designed by SPJ, PS and HN and carried out by HN. The identification of further mre11sup2a mutants was designed by FP and SPJ and carried out by NGJ. Modelling of S. cerevisiae Mre11 was performed by BO-M, and subsequent analyses were carried out by BO-M and FP. The manuscript was largely written by SPJ and FP, and was edited by all other authors.

Conflict of interest

The authors declare that they have no conflict of interest.

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