

Mouse *RAD54* Affects DNA Double-Strand Break Repair and Sister Chromatid Exchange

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Cells can achieve error-free repair of DNA double-strand breaks (DSBs) by homologous recombination through gene conversion with or without crossover. In contrast, an alternative homology-dependent DSB repair pathway, single-strand annealing (SSA), results in deletions. In this study, we analyzed the effect of *mRAD54*, a gene involved in homologous recombination, on the repair of a site-specific I-SceI-induced DSB located in a repeated DNA sequence in the genome of mouse embryonic stem cells. We used six isogenic cell lines differing solely in the orientation of the repeats. The combination of the three recombination-test substrates used discriminated among SSA, intrachromatid gene conversion, and sister chromatid gene conversion. DSB repair was most efficient for the substrate that allowed recovery of SSA events. Gene conversion with crossover, indistinguishable from long tract gene conversion, preferentially involved the sister chromatid rather than the repeat on the same chromatid. Comparing DSB repair in *mRAD54* wild-type and knockout cells revealed direct evidence for a role of *mRAD54* in DSB repair. The substrate measuring SSA showed an increased efficiency of DSB repair in the absence of *mRAD54*. The substrate measuring sister chromatid gene conversion showed a decrease in gene conversion with and without crossover. Consistent with this observation, DNA damage-induced sister chromatid exchange was reduced in *mRAD54*-deficient cells. Our results suggest that *mRAD54* promotes gene conversion with predominant use of the sister chromatid as the repair template at the expense of error-prone SSA.

DNA double-strand breaks (DSBs) form a major threat to the integrity of chromosomes and viability of cells. Unrepaired or incorrectly repaired DSBs may lead to translocations or loss of chromosomes, which could result in cell death or uncontrolled cell growth. Eukaryotes have developed several mechanisms to repair DSBs, including nonhomologous DNA end-joining (NHEJ) and homologous recombination (HR). In *Saccharomyces cerevisiae*, DSBs are efficiently repaired through HR by the *RAD52* group genes, while a contribution of NHEJ to DSB repair is only observed in the absence of HR (32). In mammalian cells, NHEJ plays a major role in DSB repair (18). More recently, it has become clear that in addition to NHEJ, HR can play an important role in DSB repair in mammalian cells as well (22).

Several pathways of homology-dependent DSB repair have been described for *S. cerevisiae* (32). One of these pathways, single-strand annealing (SSA), specifically occurs when a DSB is made between directly repeated DNA sequences. The DSB is processed by removal of part of the 5' strand on each side of the break, exposing long 3' overhangs (25). The single-stranded DNA (ssDNA) overhangs anneal to a long complementary stretch of DNA, and nonhomologous ssDNA ends are removed. As a result, one of the repeats and the intervening sequence are deleted. In vertebrates, a similar pathway has been described (5).

An alternative homology-dependent DSB repair pathway, mediated by the *RAD52* group genes, is gene conversion (GC)

(32, 46). DSB repair through this pathway also requires the DNA around the DSB to be degraded to produce 3' ssDNA overhangs. One or both of these ends invade a homologous DNA sequence, which can be found either on the homologous chromosome or, in the S and G₂ phases of the cell cycle, on the sister chromatid. Several models for this invasion have been described, including DSB gap repair and synthesis-dependent strand annealing (11, 34). In a model for DSB gap repair, both ends invade the homologous duplex and the gap is filled by DNA synthesis. The resulting Holliday junctions are resolved either with or without crossover (CO). We will use the terms "CO" for events involving GC with CO and "GC" for GC without CO. In the simplest model for synthesis-dependent strand annealing, only one end invades the homologous sequence. After DNA synthesis primed from the invaded end, the newly synthesized strand reanneals with the other end of the DSB. Then, the second strand is synthesized, resulting in a strong bias towards non-CO (11). However, if a long tract of DNA is synthesized, the result will appear similar to CO. *RAD52* is important for almost all GC and CO pathways (32). Other genes involved include *RAD51*, *RAD54*, and *RDH54/TID1* (8, 26, 32). *RAD51* and *RAD54* are mainly required for GC. *RDH54*, a homologue of *RAD54*, is only required for GC using the homologous chromosome, while *RAD54* is involved in GC with both the sister chromatid and the homologous chromosome (2, 26, 42). In mammalian cells, similar GC and CO pathways have been found, but very little is known about the genetic requirements of the different pathways. Most of the above-mentioned genes have a homologue in mammals (22). Nevertheless, the importance of each gene can differ in mammalian and *S. cerevisiae* cells. For example, the mouse *RAD52* (*mRAD52*) gene can be mutated without a major effect on

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recombination, while it is the most important gene in *S. cerevisiae* (40).

One of the other *RAD52* group genes, *RAD54*, is clearly important in mammalian cells. The Rad54 protein belongs to the SWI2/SNF2 protein family whose members modulate protein-DNA interactions in an ATP-dependent manner (23). The *S. cerevisiae* and human Rad54 proteins are double-stranded DNA-dependent ATPases that interact with Rad51, a key player in the search for homologous template DNA (6, 14, 20, 35, 45, 48). Compared to wild-type cells, *RAD54*-deficient mouse embryonic stem (ES) cells are two- to fourfold more sensitive to ionizing radiation, methyl methanesulfonate, and mitomycin C (MMC) (10). In addition, HR in *mRAD54*-deficient cells is 5- to 10-fold reduced, as measured by targeted integration of exogenous DNA (10). This reduction in HR can explain the sensitivity of cells lacking mRad54 to DSB-inducing DNA-damaging agents, although a direct involvement of mRad54 in DSB repair has not yet been demonstrated.

Much information concerning the mechanisms of DSB repair in *S. cerevisiae* has been obtained by using a site-specific DSB induced by rare-cutting endonucleases (15). Recently, it has been shown that the *S. cerevisiae* mitochondrial enzyme *I-SceI*, which recognizes and cuts a nonpalindromic 18-bp site, leaving 4-bp 3' overhangs, works efficiently in mammalian cells, but is not toxic to these cells (17). Analysis of the repair products of the site-specific DSB allows quantitation of the relative contribution of NHEJ and different homology-dependent pathways of DSB repair in mammalian cells (7, 21, 27, 28, 47). In this study, we have investigated the relative contribution of different homology-dependent pathways to the repair of an *I-SceI*-induced chromosomal DSB in mouse ES cells that were either *mRAD54*-proficient or -deficient.

MATERIALS AND METHODS

Construction of *mRAD54* targeting vectors. Targeting vectors were constructed to integrate three different recombination-test substrates into the *mRAD54* genomic locus. The substrates were cloned into the unique *SfiI* site of exon 4, thereby disrupting *mRAD54*. The first targeting vector was made by inserting the DRneo construct (28), linearized with *XhoI*, into the *SfiI* site of a 9-kb *EcoRI* fragment from *mRAD54* encompassing exons 4, 5, and 6 (Fig. 1A) (10). The second and third targeting vectors were made by inserting the IRneo and SCneo recombination-test substrates in a similar manner (Fig. 1A) (21).

ES cell culture and electroporation. Heterozygous *mRAD54* ES cells of the genotype *mRAD54*^{+|307pur} were electroporated with the different targeting vectors and cultured on gelatinized dishes as described previously (10). The cells were split 24 h after electroporation, and hygromycin B (hygro) was added to a final concentration of 200 µg/ml. After 7 to 10 days, colonies were isolated and expanded. Genomic DNA from individual clones was digested with *StuI* and analyzed by DNA blotting using a flanking probe (Fig. 1B). The blot was rehybridized with a 700-bp 3' neomycin (*neo*) fragment to confirm a single integration of the targeting vector.

***I-SceI* transfections.** ES cells containing the recombination-test substrates were cultured in medium containing hygro at a concentration of 200 µg/ml. Transfection of 3.2×10^6 cells was done by electroporation with 6 µg of either pPGK3xnlS-I-SceI or pCBA3xnlS-I-SceI, which transiently express *I-SceI* from the phosphoglycerate kinase I (PGK) or the chicken β -actin promoter, respectively (9, 39). To determine the transfection efficiency, 6 µg of pPGKAS-*eGFP*, containing the green fluorescent protein (*GFP*) gene under the control of a PGK promoter, was cotransfected in a number of experiments. In parallel, cells were electroporated without DNA or with pBSIIKS or pPGKAS-*eGFP* alone. After electroporation, 10^3 cells were plated without selection to determine the cloning efficiency. The remaining cells were grown for 1 day without selection before they were split and cultured in medium containing G418 (200 µg/ml) or G418 (200 µg/ml)-hygro (200 µg/ml). When pPGKAS-*eGFP* had been cotransfected with the *I-SceI*-expressing plasmid, a portion of the cells was subjected to fluorescence-activated cell sorting analysis 1 day after transfection to determine the percentage of cells positive for *GFP* expression. After 8 to 11 days, cells were fixed, stained, and counted. The number of clones from the cells transfected with the *I-SceI*-expressing plasmid was corrected for the number of clones from the mock-transfected cells. To enable comparison between the number of clones from different cell lines and experiments, the absolute number of clones was divided by the cloning efficiency and transfection efficiency. The data on the number of G418- and G418-hygro-resistant clones is based on three to seven

independent experiments, using two or three independent cell lines for each genotype. In several of the experiments, colonies were isolated and expanded. Genomic DNA from individual clones was analyzed for recombination events by digestion with either *NcoI* or *EcoRI* and DNA blotting using the 700-bp 3' *neo* fragment as a probe. After analysis of DNA isolated from DRneo recombinants digested with *NcoI*, 20% of the clones showed, in addition to the banding patterns expected for SSA-CO or GC, the hybridization pattern of the original construct. These were scored as SSA-CO or GC, respectively. Colonies from all recombination substrates that had aberrations in the hybridization pattern which were difficult to interpret were not included in the analysis. Inclusion of these aberrant clones did not alter the conclusions.

SCEs. Sister chromatid exchanges (SCEs) in ES cell lines of the genotypes *mRAD54*^{+|+}, *mRAD54*^{+|/-}, and *mRAD54*^{-|/-} and a derivative of the *mRAD54*^{-|/-} line expressing the *hRAD54* cDNA were analyzed (10, 45). The *mRAD54* knockout allele in these lines was *mRAD54*^{307neo}. The cell lines were coded to prevent bias in the analysis. SCE analysis was performed according to standard procedures, with the cells either mock treated or treated with 0.2 µg of MMC/ml. At least 40 metaphases per cell line were analyzed for both the number of chromosomes and SCEs.

RESULTS

The recombination-test substrates. The three substrates that were used to measure HR frequencies in mouse ES cells are schematically depicted in Fig. 1A. They contain a hygromycin selectable marker gene (*hyg*) flanked by two inactive neomycin selectable marker genes, *S2neo* and 3' *neo*. One of the crippled *neo* genes, 3' *neo*, consists of the 3' 700 bp of the *neo* gene. The other, *S2neo*, is a full-length *neo* gene, which contains a 4-bp deletion and the 18-bp insertion of the *I-SceI* site at the position of the *NcoI* site at bp 576 of *neo* (28). Expression of the *I-SceI* enzyme can create a DSB in *S2neo*. By recombination between *S2neo* and 3' *neo*, the original *NcoI* site of *S2neo*, which is present in 3' *neo*, can be restored, creating an intact *neo* gene. The three recombination-test substrates differ solely in the relative orientation of the two crippled *neo* genes (Fig. 1A). DRneo contains both crippled *neo* genes as direct repeats. Transcription of *S2neo* occurs towards the 5' end of 3' *neo*. IRneo contains both genes as inverted repeats because 3' *neo* has been inverted relative to its orientation on DRneo. SCneo contains the genes as direct repeats, but in contrast to DRneo, transcription of *S2neo* occurs away from the 3' end of 3' *neo* (21).

Homologous integration of the recombination-test substrates in the *mRAD54* locus. We targeted the recombination-test substrates to the *mRAD54* gene of ES cells to obtain single integration of the substrates at a defined and transcriptionally active position in the genome. Consequently, the targeted cell lines are isogenic and differ only in the presence of an *mRAD54*⁺ or an *mRAD54*⁻ allele and the orientation of the crippled *neo* genes of the substrates. To achieve this, the substrates were subcloned into exon 4 of the *mRAD54* gene to create targeting vectors that would result in disruption of the gene (10). The resulting *mRAD54* alleles are referred to as *mRAD54*^{DRneo}, *mRAD54*^{IRneo}, and *mRAD54*^{SCneo}, respectively. The targeting vectors were transfected into *mRAD54*^{+|/-} ES cells of the genotype *mRAD54*^{+|307pur} (10). After selection with hygro, targeted clones were identified by DNA blotting with a unique probe outside the targeting construct (Fig. 1).

The disruption of *mRAD54* by the recombination-test substrates was confirmed by the hypersensitivity of *mRAD54*^{307pur/DRneo} ES cells to γ -irradiation (data not shown). The survival curve of the *mRAD54*^{+|DRneo} cell line after γ -irradiation was similar to that of wild-type cells, as expected, because heterozygote *mRAD54* cells show no obvious phenotype (10). Immunoblot analysis using α -hRad54 showed that mRad54 protein was present in all *mRAD54*⁺ cell lines containing the substrates but could not be detected in any of the *mRAD54* knockout cell lines (data not shown).

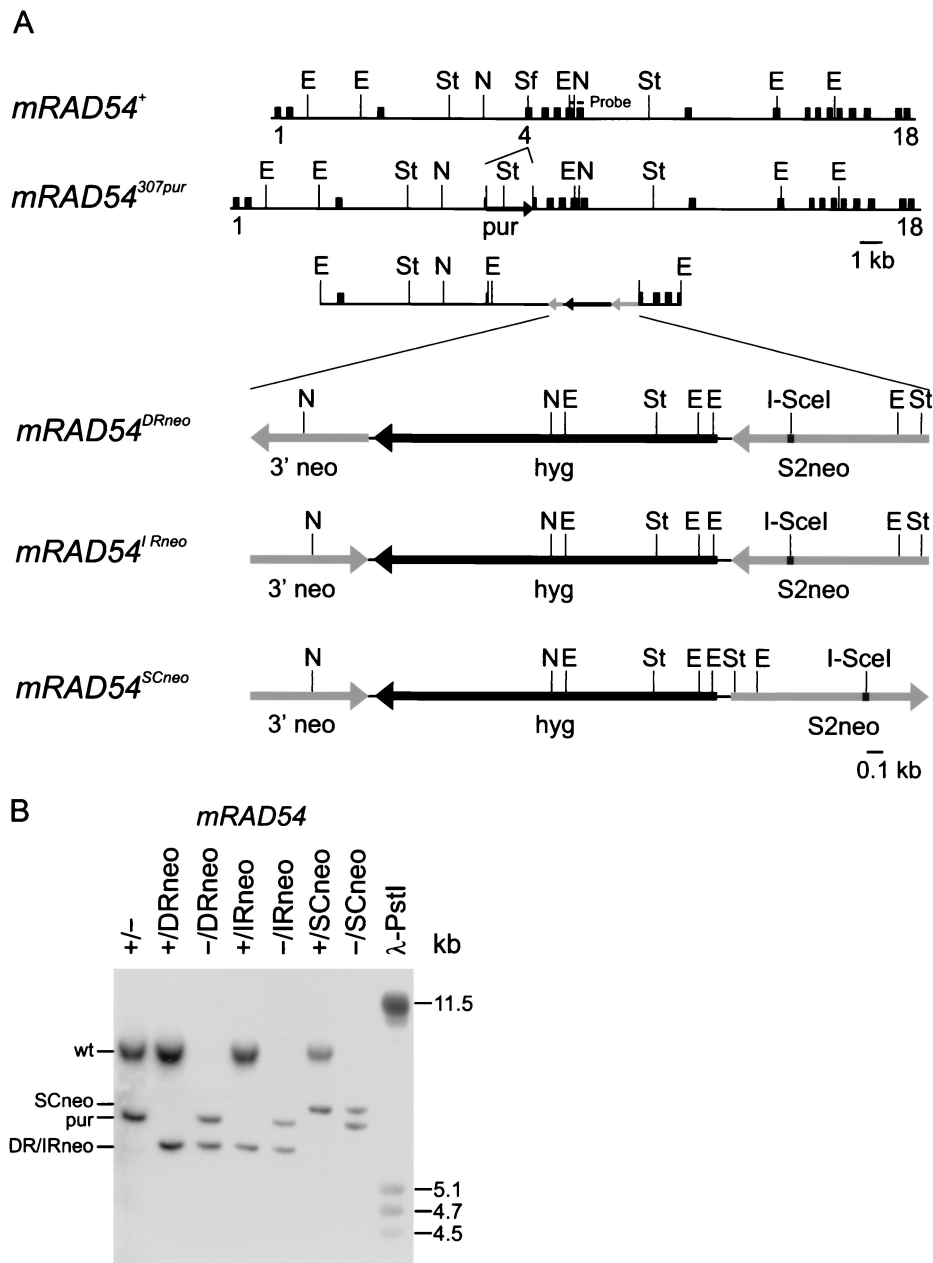


FIG. 1. Generation of *mRAD54*^{+/-} and *mRAD54*^{-/-} ES cells containing recombination-test substrates. (A) Structure of the genomic *mRAD54* locus and targeting vectors containing the substrates. The two upper lines represent the wild-type (*mRAD54*⁺) and the puromycin-targeted knockout (*mRAD54*^{307pur}) alleles, respectively. The 18 exons that encode mRad54 are indicated by boxes. The dashed line above exons 7 and 8 indicates the position of the probe used to distinguish the different *mRAD54* alleles after digestion of the genomic DNA with *StuI*. The arrow shows the position of the puromycin (*pur*) selectable marker gene. The locations of selected restriction sites are shown: E, *EcoRI*; N, *NcoI*; Sf, *SfuI*; St, *StuI*. The third line shows a generic representation of the targeting vectors. The three lower lines show the three different substrates inserted into the *mRAD54* locus in more detail. The black arrow indicates the hygromycin (*hyg*)-selectable marker gene. The gray arrow on the left represents the 700-bp 3' neomycin-selectable marker gene (3' *neo*). The gray arrow on the right represents the full-length *S2neo* gene, which contains a 4-bp deletion at the 18-bp *I-SceI* site insertion (indicated in black). (B) DNA blot of ES cells containing wild-type (+) and knockout (-) *mRAD54* alleles in addition to alleles with recombination-test substrates. Genomic DNA was digested with *StuI*. The DNA blot was hybridized with the probe indicated in panel A. Phage λ DNA digested with *PstI* was used as a size marker. The lengths of marker fragments are indicated in kilobases on the right and the positions of the different *mRAD54* alleles are shown on the left.

The DRneo substrate: DSB repair events. Transfection of an *I-SceI*-expressing plasmid in cells containing DRneo can result in a DSB in *S2neo* (Fig. 2). The DSB can be repaired by NHEJ with or without a deletion or insertion (27). NHEJ will not result in the restoration of an intact *neo* gene, and therefore NHEJ events will not be recovered. This is true for all sub-

strates. An alternative repair pathway is SSA (Fig. 2). During SSA within DRneo, complementary strands of *S2neo* and 3' *neo* will anneal, resulting in an intact *neo* gene and deletion of the intervening *hyg* gene. A third pathway to repair the DSB is HR by GC (Fig. 2). GC by recombination with *S2neo* on the sister chromatid will result in restoration of nonfunctional

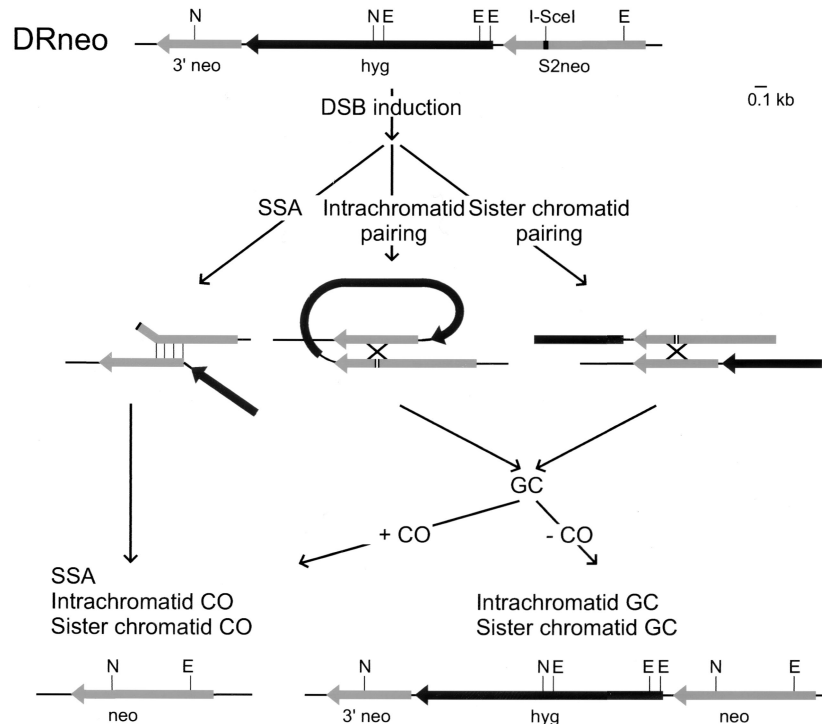


FIG. 2. Model of possible mechanisms for homology-dependent DSB repair on DRneo. The DSB induced at the I-SceI site and indicated by the gap in *S2neo* can be repaired by different repair pathways that are depicted schematically. Only repair events yielding an intact *neo* gene are shown. A summary of all possible outcomes of DSB repair is given in Table 1, and the different pathways are described in detail in the text. The annealing of the complementary ssDNA during SSA is indicated by thin vertical lines. Pairing of *S2neo* and *3' neo* (indicated by the cross) can result in GC with or without CO. Symbols are the same as those in Fig. 1.

S2neo, and therefore, these events will not be recovered. To obtain an intact *neo* gene by GC or CO, the *S2neo* containing the DSB needs to pair with either *3' neo* on the same chromatid or, in the S and G₂ phases of the cell cycle, with *3' neo* on the sister chromatid. These modes of homologous pairing are referred to as intrachromatid and sister chromatid pairing in Fig. 2. If the intermediate is resolved without a CO, the resulting clone will contain intact *neo* and *hyg* genes and a *3' neo* gene, and the cell will be resistant to G418 and hygro. On the other hand, if a single CO takes place or the GC tract continues beyond the *neo* genes, the resulting clone will contain an intact *neo* gene while the *hyg* gene and *3' neo* will be lost. The cell will only be resistant to G418. At the DNA level, the outcome of CO is therefore identical to the outcome of SSA (Table 1).

The DRneo substrate: relative efficiency of DSB repair events. The relative contribution of the different homology-dependent DSB repair pathways was investigated by transfection of an I-SceI-expressing plasmid into *mRAD54*^{+/DRneo} ES cells. As a control, a mock transfection was performed either with no DNA or with pBSIIKS or pPGKCAS-*eGFP*. Before transfection, the cells were grown by hygro selection to reduce the background due to spontaneous recombination events. After transfection, the cells were grown for 1 day without selection. Subsequently, they were divided over multiple dishes and cultured in either G418-containing medium or G418-hygro-containing medium. After 8 to 11 days, the cells were fixed and the number of colonies on each dish was counted. The frequency of spontaneously arising G418-resistant colonies varied between 10⁻⁵ and 10⁻⁶. No significant differences in the in-

TABLE 1. Possible outcomes of repair events for the different recombination-test substrates after induction of a DSB by I-SceI

DSB repair event	Possible outcome for recombination-test substrates					
	DRneo		IRneo		SCneo	
	Viability ^a	Resulting resistance gene ^b	Viability ^a	Resulting resistance gene ^b	Viability ^a	Resulting resistance gene ^b
NHEJ	+	<i>hyg</i>	+	<i>hyg</i>	+	<i>hyg</i>
SSA	+	<i>neo</i>	-		+	None
GC	+	<i>neo, hyg</i>	+	<i>neo, hyg</i>	+	<i>neo, hyg</i>
Intrachromatid CO	+	<i>neo</i>	+	<i>neo, hyg</i>	+	None
Sister chromatid CO	+	<i>neo</i>	-		+	<i>neo, hyg</i>

^a Plus and minus signs indicate whether repair through these pathways results in viable or inviable cells, respectively.

^b *neo* and *hyg* indicate the expected expression of the *neo* and *hyg* genes, respectively.

duction of G418-resistant colonies were found between transfection of a control plasmid or no DNA. The recombination frequency was increased 100- to 1,000-fold after transfection of an *I-SceI*-expressing plasmid.

G418-resistant colonies are obtained after all likely recombination events: SSA, GC, and CO. In contrast, G418-hygro-resistant colonies are only obtained after GC (Fig. 2). Therefore, the ratio of the number of G418-hygro-resistant colonies to G418-resistant colonies is an indication of the contribution of GC to all HR events. The advantage of this ratio is that it is an internal measure that can be compared directly between different cell lines and separate experiments. In addition, the ratio is not dependent on the transfection or the cloning efficiency of the cell line. In *mRAD54*^{+/DRneo} ES cells that have no defect in HR (10), this ratio of G418-hygro- to G418-resistant colonies was 0.15 ± 0.01 . Thus, around 15% of all recombination events consist of GC. The contribution of CO to the repair of a DSB is usually equal to or lower than the contribution of GC (4, 21, 32). Therefore, it is likely that SSA accounts for the majority of recombination events recovered from DRneo.

The DRneo substrate: the effect of *mRAD54* on DSB repair.

Next, we determined the effect of *mRAD54* on the repair of a DSB induced by *I-SceI* in DRneo by using *mRAD54*^{-/DRneo} ES cell lines. The ratio of G418-hygro-resistant to G418-resistant colonies shifted from 0.15 ± 0.01 , observed for *mRAD54*-proficient cells, to 0.077 ± 0.007 for *mRAD54*-deficient cells. Thus, the contribution of GC (G418-hygro-resistant clones) to the total number of recombination events (G418-resistant clones) was reduced in the absence of mRad54 protein. We conclude that the mRad54 protein is involved in repairing DSBs in vivo.

To confirm these results at the DNA level, we isolated DNA from both G418-resistant *mRAD54*^{+/DRneo} and *mRAD54*^{-/DRneo} ES cell colonies (Fig. 3). Most clones showed a hybridization pattern consistent with either GC or SSA and/or CO (Table 2). The ratio of GC to all recombination events was 0.175 for *mRAD54*^{+/DRneo} and 0.095 for *mRAD54*^{-/DRneo} ES cells (Table 2). Thus, an approximately twofold difference in the proportion of GC in the absence of *mRAD54* was again observed. However, the ratio for each genotype was slightly, but not significantly ($P > 0.2$), higher when analyzed by DNA blotting, compared to the colony formation assay.

We wished to determine whether the decrease in the ratio of GC to all recombination events observed in the absence of mRad54 was due to a decrease in the number of GCs, to an increase in SSA, or to both. Inclusion of additional controls and measuring the cloning and transfection efficiency allowed the comparison of the number of colonies obtained with different cell lines and separate experiments. The decrease in the proportion of GCs appeared to be due to both a significant increase ($P < 0.05$) in the recombination events yielding only G418 resistance, of which SSA is probably the most common, and a very slight, nonsignificant, decrease ($P > 0.10$) in the number of GCs (Fig. 4A and B). These results suggest that in the absence of mRad54 and the presence of direct repeats, ES cells shift their repair process from GC to SSA.

The IRneo and SCneo substrates: DSB repair events. The experiments with DRneo-containing cell lines yielded useful information on the frequency of GC. However, because SSA and CO result in clones that are identical at the DNA level, the relative frequencies of these repair events could not be determined. To obtain information on the usage of CO either within the same chromatid or with the sister chromatid, we constructed *mRAD54*^{+/-} and *mRAD54*^{-/-} cell lines containing IRneo and SCneo.

IRneo contains *S2neo* and *3' neo* as inverted repeats (Fig.

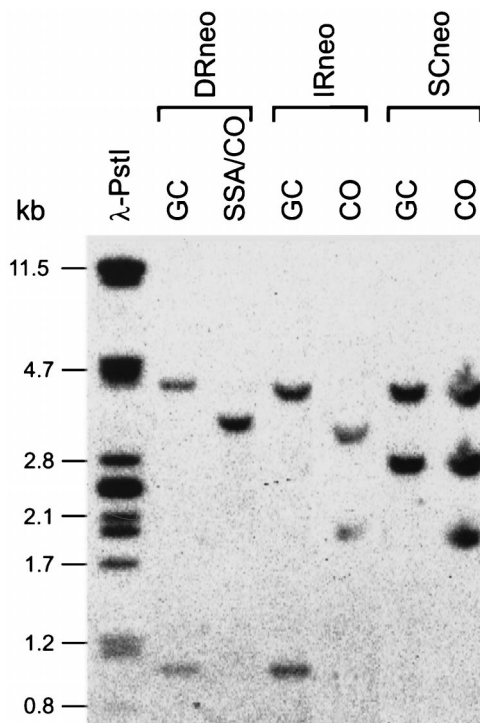


FIG. 3. DNA blot analysis of *I-SceI*-induced DSB repair events in ES cells containing the different recombination-test substrates. *mRAD54*-proficient ES cells containing either DRneo, IRneo, or SCneo were transfected with an *I-SceI*-expressing plasmid. After selection with G418 or G418-hygro, genomic DNA from individual clones was digested with *EcoRI*. The outcome of repair of the *I-SceI*-induced DSB was analyzed by DNA blotting using a 700-bp 3' *neo* probe. Only a selection of the clones listed in Table 2 is shown. As shown in Fig. 2 and 5, the sizes of the *EcoRI* fragments labeled with the *neo* probe indicate whether the DSB has been repaired by GC or CO. With DRneo, SSA results in the same molecular outcome as CO. Phage λ DNA digested with *PstI* was used as a size marker. The lengths of marker fragments are on the left.

5A). In contrast to DRneo, the *I-SceI*-induced DSB in IRneo cannot be repaired through SSA. Due to the inverse orientation of the crippled *neo* genes, nucleolytic processing of the DSB will expose identical rather than complementary ssDNA tails. However, repair of the DSB by recombination is possible through several different routes (Fig. 5A and Table 1). Both

TABLE 2. Relative contribution of different homology-dependent repair events of *I-SceI*-induced DSBs in *mRAD54*-proficient and -deficient cells containing the recombinant-test substrates

Genotype	Distribution of DSB repair events ^a			
	GC	CO (SSA)	Other	Total
<i>mRAD54</i> ^{+/DRneo}	17	80	1	98
<i>mRAD54</i> ^{-/DRneo}	6	57	3	66
<i>mRAD54</i> ^{+/IRneo}	124	2	4	130
<i>mRAD54</i> ^{-/IRneo}	112	0	8	120
<i>mRAD54</i> ^{+/SCneo}	50	54	24	128
<i>mRAD54</i> ^{-/SCneo}	55	50	37	142

^a Distribution of DSB repair events was determined by expanding colonies obtained after transfection of an *I-SceI*-expressing plasmid to ES cells of the indicated genotype and selection with either G418 (DRneo) or G418-hygro (IRneo, SCneo). DNA from the colonies was analyzed by DNA blotting after digestion with *EcoRI* or *NcoI*. The clones were classified according to their restriction pattern resulting from GC or CO. For DRneo, SSA was recovered in the same class as CO (CO-SSA). When the restriction pattern and intensity of the bands was different from the expected pattern of HR events, the clone was counted in the category "Other."

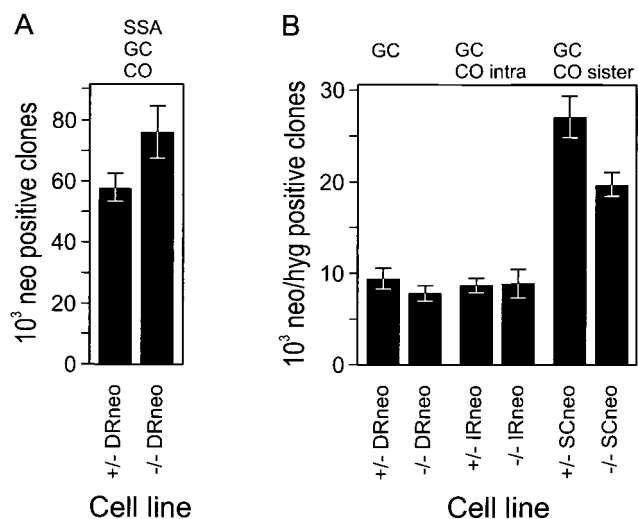


FIG. 4. Homologous recombination frequencies for the recombination-test substrates. As described in Materials and Methods, 1.6×10^6 *mRAD54*-proficient and -deficient ES cells containing the indicated substrates in the identical genomic location were transfected with pCBA3xnlS-I-SceI and processed. Shown is the normalized number of G418- or G418-hygro-resistant colonies \pm standard error of the mean for three independent experiments with two cell lines from all six genotypes. (A) HR frequency of *mRAD54*^{+/-}*DRneo* (+/-) and *mRAD54*^{-/-}*DRneo* (-/-) ES cells. Colonies containing an intact *neo* gene were obtained after repair of the I-SceI-induced DSB by SSA, GC, and CO. (B) Frequencies of GC and CO for ES cells containing the substrates. For all three substrates, *neo*- and *hyg*-containing colonies were obtained after repair of the I-SceI-induced DSB by intrachromatid GC and sister chromatid GC. The IRneo- and SCneo-containing cell lines each have one additional possibility to yield G418-hygro-resistant colonies. In the IRneo-containing lines, these clones can be formed by intrachromatid CO. For the SCneo-containing lines, they can be formed by CO after pairing with the sister chromatid.

GC and CO can occur by using 3' *neo* on either the same chromatid or the sister chromatid as a template. Both GC events will result in G418-hygro-resistant cells. In contrast, CO involving the sister chromatid results in a dicentric chromosome and an acentric chromosome, which is incompatible with cell survival. DSB repair through CO after pairing with 3' *neo* on the same chromatid results in G418-hygro-resistant cells. The orientation of the *hyg* gene will be inverted by the CO. Thus, CO can be distinguished from GC at the DNA level (Fig. 3).

SCneo contains *S2neo* and 3' *neo* as direct repeats (Fig. 5B). In contrast to DRneo, transcription of *S2neo* occurs away from the 3' end of 3' *neo*, which has implications for the outcome of DSB repair. Expression of I-SceI in a cell containing SCneo can result in a DSB in *S2neo*. SSA is a possible repair event and will pair the 5' end of 3' *neo* to the 3' end of *S2neo*. As a result, the DSB is repaired and 3' *neo* is recovered, which will not be detected because the cell will remain sensitive to G418. Similar to the other substrates, GC can occur by pairing with 3' *neo* on the same chromatid or the sister chromatid. In this way, an intact *neo* gene will be obtained and the *hyg* gene will be retained. CO after pairing with 3' *neo* on the same chromatid will yield the same outcome at the DNA level as SSA, namely a single 3' *neo* gene which will not be recovered. On the other hand, CO after unequal pairing with 3' *neo* on the sister chromatid will result in an intact *neo* gene with a partial duplication of the rest of the construct resulting in two intact *hyg* genes (Fig. 5B and Table 1). This event can be distinguished from GC by DNA blotting (Fig. 3).

The IRneo and SCneo substrates: relative efficiency of DSB repair events. To investigate the relative efficiency of the different HR repair pathways, *mRAD54*^{+/-}*IRneo* and *mRAD54*^{+/-}*SCneo* ES cells were transfected with an I-SceI-expressing plasmid, as described above for the DRneo-containing cell lines. The spontaneous recombination frequency was 10^{-5} to 10^{-6} (data not shown). The number of colonies on the pCBA3xnlS-I-SceI-transfected dishes was normalized for the number of colonies on the mock-transfected dishes, the cloning efficiency, and the transfection efficiency. The resulting recombination frequency was about 10^{-2} . In the SCneo-containing cell lines, not all recombination events are recovered, as SSA, GC using *S2neo* on the sister chromatid, and CO after pairing with 3' *neo* on the same chromatid do not result in G418 resistance (Table 1). Nevertheless, transfection of the I-SceI-expressing plasmid into *mRAD54*^{+/-}*SCneo* cells resulted in three times more colonies than transfection into *mRAD54*^{+/-}*IRneo* cells did (Fig. 4B). The number of G418-hygro-resistant colonies obtained after transfection of *mRAD54*^{+/-}*IRneo* cells with pCBA3xnlS-I-SceI was comparable to the number obtained after transfection of *mRAD54*^{+/-}*DRneo* cells (Fig. 4B).

Since both GC and CO result in G418-hygro resistance of IRneo and SCneo, we investigated the distribution of these events by DNA blotting. GC and CO can be discriminated because they result in a different restriction pattern after digestion with *Eco*RI (Fig. 3 and 5). IRneo almost exclusively showed GC, which implies that CO within the same chromatid between inverted repeats is a rare event in ES cells (Table 2). In SCneo, GC and CO contributed equally to the recovered HR events (Table 2). Thus, CO after pairing with the sister chromatid, which usually does not lead to deleterious chromosome rearrangements, is a common event. A relatively high number of SCneo-derived clones showed restriction patterns that could not be explained by GC or CO (Table 2). These clones were excluded from the analysis, but their inclusion did not alter the conclusions.

The IRneo and SCneo substrates: the effect of *mRAD54* on DSB repair. With DRneo-containing cell lines, we observed, in the absence of *mRad54* protein, a significant increase in SSA with a concomitant very slight reduction of GC. Therefore, we investigated the effect of *mRAD54* on HR in the other substrates. We transfected pCBA3xnlS-I-SceI into *mRAD54*^{-/-}*IRneo* and *mRAD54*^{-/-}*SCneo* cells and analyzed the colonies obtained as described above. DNA blot analysis revealed that there was no difference in the relative distribution of HR events between *mRAD54*-proficient and -deficient ES cell lines containing IRneo or SCneo (Table 2). *mRAD54*^{-/-}*IRneo* cells showed only GC, and *mRAD54*^{-/-}*SCneo* cells showed an equal number of GCs and COs.

The number of colonies obtained from *mRAD54*^{-/-}*IRneo* cells did not differ from the number of colonies from *mRAD54*^{+/-}*IRneo* cells (Fig. 4B). Thus, no indication was obtained for an involvement of *mRAD54* in the repair of a DSB between inverted repeats by GC. However, *mRAD54*^{-/-}*SCneo* ES cells gave rise to fewer colonies than *mRAD54*^{+/-}*SCneo* ES cells after transfection of an I-SceI-expressing plasmid (Fig. 4B). There was a consistent, statistically significant ($P < 0.05$) decrease to approximately 70% of the number of colonies obtained with *mRAD54*-proficient cell lines containing SCneo. This indicates a role for *mRAD54* in GC and CO with the sister chromatid in DSB repair in this substrate.

Influence of *mRAD54* on the induction of SCEs. To obtain independent evidence for a role of *mRAD54* in sister chromatid recombination, we measured the spontaneous and DNA damage-induced levels of SCEs in *mRAD54*-proficient and -de-

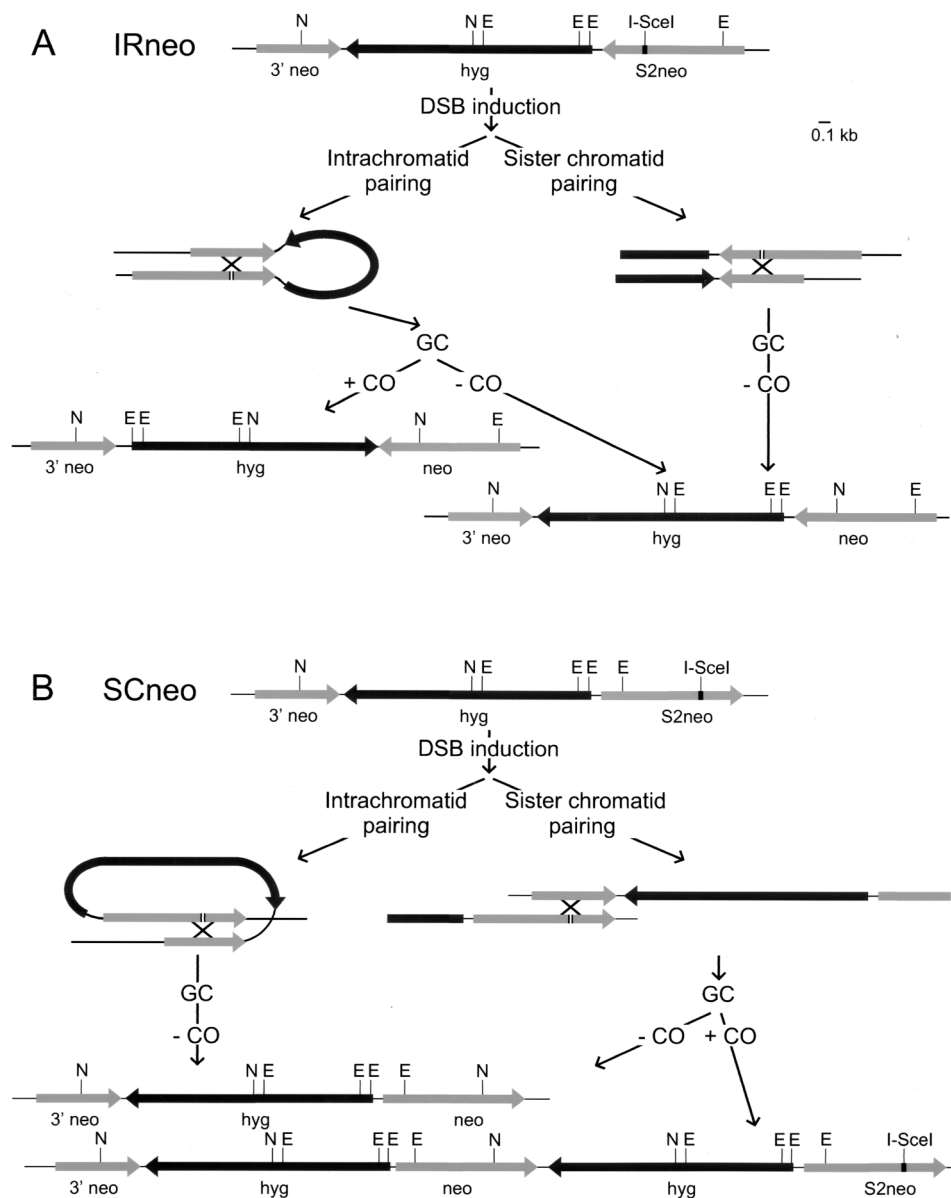


FIG. 5. Schematic representation of possible homology-dependent DSB repair pathways for IRneo and SCneo. Only repair events yielding an intact *neo* gene are depicted. A summary of all possible outcomes of DSB repair is given in Table 1, and the different pathways are described in detail in the text. Symbols are the same as in Fig. 1. The I-SceI-induced DSB is indicated by the gap in *S2neo*. Recombination between *S2neo* and *3' neo*, indicated by the cross, can lead to restoration of the original *NeoI* site resulting in an intact *neo* gene by GC with or without CO. Concerning the COs, only the product that results in an intact *neo* gene is shown. Shown are the outcomes of DSB repair events on IRneo (A) and SCneo (B).

ficient ES cells. ES cells of the genotypes *mRAD54*^{+/+}, *mRAD54*^{+/-}, and *mRAD54*^{-/-} were analyzed. The spontaneous level of SCEs found in the *mRAD54*^{-/-} cell line was slightly reduced compared to that observed in the *mRAD54*-proficient control cell lines (Fig. 6). In all cell lines, no numerical or gross structural chromosomal abnormalities were observed. DNA damage inflicted by the DNA interstrand cross-linking agent MMC increased the number of SCEs. Treatment of the cells with 0.2 μg of MMC/ml for 1 h increased the number of SCEs 2.6-fold in the *mRAD54*^{+/+} and *mRAD54*^{+/-} ES cell lines. In the *mRAD54*^{-/-} cell line, the increase in SCEs was only 1.8-fold. The difference in the average number of SCEs among *mRAD54*^{+/+}, *mRAD54*^{+/-}, and *mRAD54*^{-/-} cells was significant (Fig. 6; *P* < 0.05). In addition, we included

a derivative of the *mRAD54*^{-/-} cell line that expressed the *hRAD54* cDNA in the SCE analysis as a control. Expression of this cDNA rescues the DNA damage sensitivities of *mRAD54*^{-/-} cells (45). The expression of *hRAD54* returned the number of SCEs in the *mRAD54*^{-/-} ES cell line to wild-type levels, both spontaneously and after treatment with MMC. In all cell lines treated with MMC, no apparent chromosomal changes were observed.

DISCUSSION

The major homology-dependent DSB repair pathway for DRneo is SSA. In this study, we have analyzed HR in mouse ES cells. Using DRneo, a distinction can be made between DSB

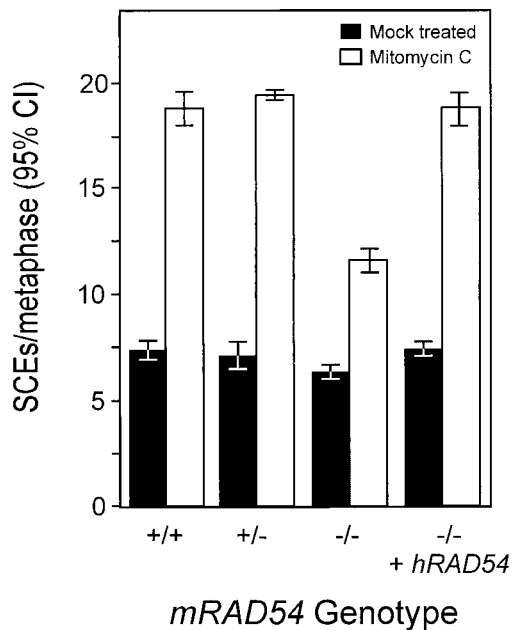


FIG. 6. Induction of SCEs by MMC in *mRAD54*-proficient and -deficient ES cells. ES cells of the indicated genotypes were either mock treated or treated with 0.2 μg of MMC/ml for 1 to 2 h, and metaphase spreads were prepared. Forty to 95 metaphases per sample were scored for the number of SCEs per cell. The frequency of spontaneous SCEs is shown in black, while the frequency of SCEs after treatment with MMC is shown in white. The error bars indicate the 95% confidence intervals.

repair through SSA and CO on one hand and GC on the other hand (Fig. 2). Our results show that approximately 15% of all homology-dependent DSB repair events within DRneo in mouse ES cells occur through recombination between the direct repeats via GC. These results appear not to be specific for mouse ES cells, because 25% of the DSB repair events in CHO cells containing integrated DRneo occur through GC (27). The separate contributions of SSA and CO to DSB repair cannot be determined with DRneo. However, a comparison to the results with SCneo, which also contains direct repeats, suggests that GC and CO occur at similar frequencies (Table 2). This implies that SSA accounts for 70% of all homology-dependent DSB repair events within DRneo in ES cells.

Consistent with our results, recombination between direct repeats through GC, when compared to SSA, accounts for the minority of detected events in a number of other assay systems, including DSB-induced events on plasmids and in chromosomes in *S. cerevisiae* and vertebrate cells (12, 19, 27, 31, 41). However, there are a number of exceptions, both in *S. cerevisiae* and mammalian cells, in which GC accounts for the majority of spontaneous and DSB-induced events (4, 13, 30, 38, 47). Variables that might contribute to observed differences among assay systems include the length and sequence context of the repeats, the distance between the repeats, the position of the DSB, and heterology in the repeats or at the ends (12, 13, 31, 33, 47). Both GC and SSA use 3' ssDNA tails as intermediates. GC requires search for homology followed by joint molecule formation actively mediated by Rad51-coated ssDNA. On the other hand, SSA involves the more passive process of annealing of complementary single-strands, although it does also, at least partially, depend on Rad52. Depending on the presence of nonhomologous ends and the length and sequence context of the repeat, these two processes

might be affected differentially. Finally, the distribution of repair events may also be dependent on the stage of the cell cycle, as GC using the sister chromatid is only possible in S and G₂.

DSB repair associated with DNA COs occurs mostly from the sister chromatid. DSB repair products of both IRneo and SCneo differ depending on whether they have been generated through GC or CO (Fig. 5). While IRneo detects intrachromatid COs, SCneo detects unequal COs between sister chromatids. It should be noted, however, that if a long tract of DNA is synthesized during GC, the result appears similar to a CO. With IRneo, DSB repair through CO occurs only in 1.5% of the analyzed repair events, while GC accounts for over 95% of the events (Table 2). This also indicates that GC tracts are generally shorter than 2.7 kb, because otherwise the outcome would have been scored as a CO. In contrast to the lack of COs with IRneo, we find that GC and CO contribute equally to DSB repair using SCneo. Thus, it appears that COs preferably arise when the sister chromatid, instead of a homologous sequence on the same chromatid, is used as the repair template. A large contribution of COs using SCneo has also been found in CHO cells, after induction of a DSB (21). A similar preference for CO using the sister chromatid has been observed in mouse L cells during spontaneous recombination between repeated sequences (4). In contrast, in *S. cerevisiae*, a preference for intrachromatid interactions has been found, as indicated by a low percentage of COs in an SCneo-like substrate (24). A high percentage of COs after intrachromatid interactions has also been found in the repair of an induced or spontaneous DSB using inverted repeats in *S. cerevisiae* (1, 37, 41, 44).

The preference for sister chromatid interactions during HR in mammalian cells could have arisen because sister chromatid recombination is, in general, less prone to the generation of chromosomal rearrangements than intrachromatid recombination. A significant fraction of mammalian genomes consists of repetitive DNA sequences. CO between these sequences will result in deleterious chromosomal rearrangements, except when the same sequence on the sister chromatid is used. Evidence thus far suggests that genome rearrangements are indeed suppressed during recombination between sequence repeats on nonhomologous chromosomes (39). Furthermore, the presence of mismatches between the repeats also prevents recombination, due to the mismatch repair system (49). *S. cerevisiae* contains hardly any repetitive sequences and will undergo less selection against allowing intrachromatid recombination. The preference for the sister chromatid in mammalian cells might occasionally result in unequal CO between sister chromatids, but if those events occur in a limited region, their potential deleterious effects could be minimized.

Comparison of the frequency of DSB repair events on the different substrates. We find that the frequency of GC is comparable among the different substrates (around 6×10^{-3} ; Table 2 and Fig. 4). It seems reasonable to assume that with all three substrates, a similar fraction of the cells receives a DSB and that a similar fraction of these DSBs is channeled into a homology-dependent repair pathway. Repair by SSA or CO using the sister chromatid will result in correct DSB repair in cells containing DRneo or SCneo. However, these events are apparently aborted in cells containing IRneo. They may cause cell death instead of resulting in GC. Otherwise, more GC events should have been recovered with IRneo. This finding of less efficient recombinational repair between inverted repeats is not unique to our assay. Both after DSB induction in *S. cerevisiae* and spontaneously in mouse cells, the frequency of recombination between direct repeats is higher than between inverted repeats (3, 41).

mRad54 influences the repair of DSBs in DRneo. A role for the mRad54 protein in the repair of DSBs has been postulated based on the ionizing radiation sensitivity and HR deficiency of *mRAD54*^{-/-} ES cells (10). The results of our study provide direct evidence that mRad54 is involved in DSB repair in vivo. The difference in DSB repair between *mRAD54*-proficient and -deficient cells is most clearly seen when the DSB is induced between direct repeats, as is the case with DRneo and SCneo (Fig. 4A and B). The absence of *mRAD54* causes a very slight reduction in GC during DSB repair of DRneo. This reduction is accompanied by a statistically significant increase in the number of COs and SSA, the latter of which is the most frequent. In *S. cerevisiae*, a similar increase in HR is seen in *rad54* mutants, both with direct repeats on plasmids and in chromosomes (16, 29, 42). The frequency of SSA (or CO) is 1.9- to 27-fold higher in *rad54* cells than in wild-type cells, while cell survival and the frequency of GC are decreased (16, 29, 42). These results suggest that there might be competition between SSA and GC (see below).

mRad54 influences recombination between sister chromatids. In cells containing SCneo, the effect of *mRAD54* on GC is more pronounced than in cells containing DRneo. Repair of the DSB through SSA is possible in SCneo, although those events are not detected. A statistically significant 27% decrease in the frequency of GC and CO is observed in the absence of *mRAD54* (Fig. 4B). Since all COs take place after pairing with the sister chromatid, *mRAD54* is clearly involved in sister chromatid recombination. This is also the case for *S. cerevisiae RAD54* (2). The contribution of GC and CO remains about equal in *mRAD54*^{-/-} cells, which indicates that *mRAD54* is involved in both GC and CO (Table 2). In contrast to these results, *S. cerevisiae RAD54* appears to be mainly involved in GC, although this has been investigated only with inverted repeats (37).

COs resulting in restoration of the *neo* gene in SCneo are the consequence of interactions with the sister chromatid and result in SCEs at the chromosomal level. Therefore, our results with SCneo predict a reduction in the level of SCEs in the absence of *mRAD54*. Indeed, we find a slightly lower level of spontaneous SCEs in *mRAD54*^{-/-} ES cells compared to that in *mRAD54*^{+/+} ES cells (Fig. 6). Because SCEs are induced by DNA-damaging agents, we have also tested whether *mRAD54*^{-/-} ES cells respond differently to MMC treatment in the SCE assay than *mRAD54*^{+/+} or *mRAD54*^{+/-} cells. Treatment of *mRAD54*^{-/-} cells with MMC yields a 1.5-fold lower induction of SCEs, compared to *mRAD54*^{+/+} cells (Fig. 6). This effect of *RAD54* on spontaneous and DNA damage-induced SCEs corresponds to results obtained with chicken-derived cells, in which a reduction in the frequency of SCEs in *RAD54*- and *RAD51*-deficient chicken B lymphocytes is observed (43). From these results, we conclude that genes required for HR are also involved in promoting SCEs. The decrease in SCEs induced by DNA damage corresponds to the similar decrease in the number of COs during DSB repair on SCneo.

The observation that *mRAD54* influences DNA damage-induced SCEs adds significantly to our results with the recombination-test substrates. The results of the SCE experiments show that *mRAD54* is involved in homology-dependent DNA repair of DNA damages that are present in naturally occurring genomic sequences. The SCE experiments overcome two restrictions of the experiments with recombination-test substrates. First, the restriction enzyme-induced DSB that initiates repair in the experiments involving the recombination-test substrates might be recognized differently from other types of DNA damages, including DSBs introduced by ionizing radia-

tion or DNA interstrand cross-linking agents. Second, in the experiments involving the recombination-test substrates, the introduction of repeated DNA sequences is necessary in order to select for successful DSB repair events. However, the presence of these repeated sequences will influence the distribution of observed repair events. SSA relies especially on the presence of repeated sequences and will be used less frequently in a more physiological situation.

The absence of mRad54 has no influence on recombination within IRneo. We find no change in the frequency of GC after induction of a DSB in IRneo in *mRAD54*-deficient cells compared to that in *mRAD54*-proficient cells (Fig. 4B). COs using the 3' *neo* on the same chromatid are rare in *mRAD54*^{+/IRneo} cells and have not been detected in *mRAD54*^{-/IRneo} cells (Table 2). Similar to our results with chromosomal substrates in ES cells, disruption of *RAD54* in *S. cerevisiae* cells has no effect on the repair of an induced DSB in inverted repeats located on a plasmid (16). In contrast, the rate of spontaneous GC between chromosomal inverted repeats is decreased 25-fold in a *rad54 S. cerevisiae* strain (37). Because *S. cerevisiae* cells display a different distribution of events, with a predominance of COs, a direct comparison between *S. cerevisiae rad54* and *mRAD54*^{-/-} ES cells is difficult (37). The lack of an effect of *mRAD54* on DSB repair between inverted repeats in ES cells also contrasts with the effects of *mRAD54* on DSB repair between direct repeats (Fig. 4). As we will discuss below, this could be due to the possibility to repair a DSB in direct repeats by SSA, which is not possible in inverted repeats.

Does mRad54 promote GC at the expense of SSA? ssDNA tails are formed as a common intermediate in SSA and GC with or without CO. Because of this common intermediate, it is likely that a certain degree of competition exists between these two pathways (12). mRad54 could have a role in promoting GC, either directly or indirectly by blocking DSBs from being processed through the SSA pathway. This would explain the increase in the number of G418-resistant colonies in *mRAD54*-deficient cells with DRneo, which results from an increase in SSA. It would also explain the decrease in G418-hygro-resistant colonies with SCneo, because an increase in SSA, which is not recovered, would cause a decrease in the recovered GC events. With IRneo, SSA is not possible, and therefore, lack of mRad54 would not have any effect on DSB repair in this substrate. Mammalian chromosomes contain a significant amount of repetitive sequences that could be used to repair a DSB by SSA, thereby resulting in deletions. Inhibition of SSA by mRad54 is therefore even more relevant in mammalian cells than in *S. cerevisiae*, where similar effects of Rad54 on SSA have been found. Direct stimulation of GC pathways by mRad54, possibly by its interaction with mRad51, would decrease the contribution of SSA to DSB repair. Alternatively, mRad54 could suppress SSA directly. It has been shown recently that the purified human and *S. cerevisiae* Rad54 proteins have ATP-dependent DNA unwinding activity (36, 48). This activity would be ideally suited for the destabilization of intermediates in SSA or the stimulation of mRad51-mediated homologous DNA pairing and strand exchange (35).

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