Identification of a novel human Rad51 variant that promotes DNA strand exchange

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ABSTRACT

Rad51 plays a key role in the repair of DNA double-strand breaks through homologous recombination, which is the central process in the maintenance of genomic integrity. Five paralogs of the human Rad51 gene (hRad51) have been identified to date, including hRad51B, hRad51C, hRad51D, Xrcc2 and Xrcc3. In searches of additional hRad51 paralogs, we identified a novel hRad51 variant that lacked the sequence corresponding to exon 9 (hRad51-ex9). The expected amino acid sequence of hRad51-ex9 showed a frame-shift at codon 259, which resulted in a truncated C-terminus. RT-PCR analysis revealed that both hRad51 and hRad51-ex9 were prominently expressed in the testis, but that there were subtle differences in tissue specificity. The hRad51-ex9 protein was detected as a 31-kDa protein in the testis and localized at the nucleus. In addition, the hRad51-ex9 protein showed a DNA-strand exchange activity comparable to that of hRad51. Taken together, these results indicate that hRad51-ex9 promotes homologous pairing and DNA strand exchange in the nucleus, suggesting that alternative pathways in hRad51- or hRad51-ex9-dependent manners exist for DNA recombination and repair.

INTRODUCTION

Homologous recombination (HR) is a fundamental process conserved in all organisms, maintaining genomic stability through the repair of exogenous and endogenous DNA double-strand breaks. HR also contributes to genomic diversity in evolution through its pivotal roles in the exchange of chromatids during meiosis (1). In addition, dysregulation of HR may lead to aberrant genetic rearrangements and genomic instability, resulting in translocations, deletions, duplications or loss of heterozygosity (2). Precise control of the HR equilibrium is therefore essential for genetic stability because both HR stimulation and repression lead to genome instability (3).

Rad51, a eukaryotic ortholog of bacterial RecA, plays a central role in the repair of double-strand DNA breaks by mediating homologous pairing and strand exchange in recombinatory structures known as Rad51 foci in the nucleus (4). Rad51 belongs to the Rad52 epistasis group in Saccharomyces cerevisiae, which is comprised of a number of the key genes (Rad50 to Rad57) involved in recombinational repair of double-strand DNA breaks (5). Among the members of the Rad52 epistasis group, Rad51 shows the highest degree of sequence conservation in evolution, with 83% amino acid sequence homology between yeast and human orthologs and 99% homology between mouse and human orthologs (6). The functional importance of Rad51 has been further emphasized by the findings that Rad51 interacts with the tumor suppressor protein, p53 (7,8), and the breast cancer-susceptibility proteins, BRCA1 and BRCA2 (9–11). Additionally, elevated levels of hRad51 have been observed in a variety of tumor cells (12–14), suggesting that strict regulation of this recombinase may be essential for maintaining genome integrity.

To date, five human Rad51 (hRad51) paralogs, Rad51B (Rad51L1), Rad51C (Rad51L2), Rad51D (Rad51L3), Xrcc2 and Xrcc3, have been identified. Each of these genes shows only a limited degree of sequence similarity to hRad51, however, they all contain the RecA domain for DNA recombination and the Walker A and B motifs for ATP binding and hydrolysis in the predicted amino acid sequences (15–18). These hRad51 paralogs have presumably arisen through a series of gene duplications in the early stages of eukaryotic evolution (19). In addition, the five hRad51 paralogs have been reported to assist the DNA strand exchange activity of hRad51, forming two distinct complexes, Rad51B-Rad51C-Rad51D-Xrcc2 and...
hRad51C-Xrc3 (20). Deficiency in any of the Rad51 paralogs has been shown to lead to increased sensitivity to DNA cross-linking agents and ionizing radiation in vertebrate cells (21–23).

In an attempt to identify additional hRad51 paralogs in humans, we searched a human testis cDNA library. We report here a novel splice variant of hRad51, hRad51-Dex9, which lacks the sequence corresponding to exon 9. This novel variant was also found in the expressed sequence tag (EST)-databases. The hRad51-Dex9 protein was localized in the nucleus and detected as an expected molecular weight of 31 kDa in the testis. The hRad51-Dex9 protein showed DNA strand exchange activity that was comparable to that of hRad51, suggesting that this novel variant also functions as a recombinase. Additionally, using site-directed mutagenesis, we found that a short basic motif located in the C-terminus of hRad51-Dex9 may play a functional role in nuclear localization of this novel variant.

MATERIALS AND METHODS

Identification of hRad51-Dex9

A human testis 5'-stretch cDNA library (Clontech) was screened using a hRad51 cDNA probe. The cDNA probe was P32-labeled by random primer labeling, and hybridization was conducted in 50% formamide, 5× SSPE (1× SSPE: 150 mM sodium chloride, 10 mM sodium phosphate, 1 mM EDTA, pH 7.4), 10× Denhardt’s solution, 2% SDS and 100 μg/ml denatured salmon sperm DNA at 42°C for 16 h. The filters were washed twice in 2× SSC (1× SSC: 150 mM sodium chloride, 15 mM sodium citrate, pH 7.0), 0.1% SDS at room temperature and then twice in 0.2× SSC, 0.1% SDS at 42°C. Next, the filters were exposed to Kodak XAR film at −70°C for varying periods of time. The positive phage clones were then sequenced using an ABI 310 automated DNA sequencer. The human EST database was also searched for identification of hRad51 paralogs using the BLASTN program (http://www.ncbi.nlm.nih.gov/entrez/BLAST). The EST A018041 clone was purchased from Open Biosystems. The nucleotide sequence reported in this paper will appear in the GenBank under accession number EU362635.

RT-PCR analysis in human tissues

Human Multiple Tissue cDNA panels (Clontech) were PCR-amplified using ExTag polymerase (Takara) with primers specific to both hRad51 and hRad51-Dex9 (forward: 5'-ttggtagaatctgctgg-3’; and reverse: 5'-aggagac agggagatctg-3’), which were derived from the flanking regions of exon 9. The reaction mixture was subjected to 30 cycles of 94°C for 30 s, 58°C for 30 s and 72°C for 40 s with a predenaturation at 94°C for 4 min and a final extension at 72°C for 7 min. The amplified PCR products were then analyzed by electrophoresis on 2.0% agarose gels.

Expression and purification of the recombinant hRad51 and hRad51-Dex9 proteins

The full-coding sequences of hRad51 and hRad51-Dex9 were PCR-amplified from recombinant phage clones using Pfu DNA polymerase (Stratagene) according to the manufacturer’s instructions. The sequences of the oligonucleotide primers are available upon request. A unique restriction site, either NotI or BamHI, was introduced into each primer to allow convenient subcloning. The PCR-amplified fragments were then gel-purified and ligated into pET28b (Novagen) or pET21c (Novagen) at the NotI and BamHI restriction sites in frame with the C-terminal hexahistidine tag. The resulting expression constructs were then confirmed to contain the desired sequences by DNA sequence analysis using the BigDye termination version 3.0 (ABI). Among the expression constructs, pET28b-hRad51 and pET21c-hRad51-Dex9 were used for expression of the hRad51 and hRad51-Dex9 proteins, respectively.

The Escherichia coli strain, BL21 (DE3) (Novagen), was used for transformation of the pET-derived expression constructs. The recombinant proteins were expressed and purified as previously described (24). However, the hRad51-Dex9 protein resulted in the formation of inclusion bodies. Denaturing and refolding of the hRad51-Dex9 protein into an enzymatically active form were done as previously published for other human proteins (25). Briefly, the inclusion bodies were precipitated by centrifugation at 8000 g for 20 min and then homogenized in 6 M urea, 10 mM K2HPO4, pH 8.2 and 3 mM β-mercaptoethanol. The solubilized recombinant proteins were then purified using Ni-NTA agarose resins (Qiagen). For refolding, the denatured hRad51-Dex9 protein was first dialyzed overnight against a buffer of 10 mM K2HPO4, pH 9.6, 200 μM CuCl2 and 2% sodium N-lauroylsarcosinate and then against a buffer of 10 mM K2HPO4, pH 9.6 and 5 μM CuCl2. Next, the proteins were further dialyzed twice against 10 mM K2HPO4, pH 7.0. The concentration of the dialyzed protein samples was then determined using a BCA Protein Assay Kit (Bio-Rad). All of the purification procedures were conducted at 4°C. The purity and size of the recombinant proteins were assessed by SDS–PAGE. The purified recombinant proteins were further confirmed by western blot analysis using a commercial hRad51 polyclonal antibody (Calbiochem).

DNA strand exchange assays

DNA strand exchange assays were done as previously described (26,27). Briefly, the recombinant hRad51 or hRad51-Dex9 protein (final concentration, 3.5 μM) was mixed with 125 ng (final concentration, 16.8 μM in nucleotides) of φX 174 viral DNA (New England Biolabs) in 20 μl buffer containing 20 mM HEPES, pH 6.5, 1 mM DTT, 6.6 mM MgCl2, 3 mM ATP, 20 mM creatine phosphate, 0.1 mg/ml creatine kinase and 50 μg/ml BSA. After 5 min of incubation at 37°C, 120 ng (final concentration, 8.4 μM in base pairs) of PstI-linearized φX 174 dsDNA (New England Biolabs) in 1 μl and 1 μl of 100 mM MgCl2 were added to the reaction mixture. Following subsequent incubation for 15, 30, 60, 120 or 240 min at 37°C, 0.5% SDS and 0.5 mg/ml proteinase K were added to stop the exchange reaction. The incubated DNA samples were then run in 0.8% agarose gels. The gels were stained with 0.1 μg/ml of syber green (Molecular Probe) for 2 h and then
distained in ddH₂O for 2 h. Images were processed using Photoshop 7.0 (Adobe).

Generation of a hRad51-Aex9-specific polyclonal antibody

A synthetic peptide (EERKRGNQNQLNRLS) was covalently conjugated to maleimide-activated keyhole limpet homocyanin. The peptide conjugate was then emulsified with an equal volume of complete Freund's adjuvant. Adult rabbits of 1.8–2.0 kg in weight were intramuscularly injected with 500 μg of the emulsified peptide conjugate four times at a 2-week interval. The rabbits were bled on Days 7 and 14 after the last injection, and the presence of antibodies was then evaluated using an ELISA assay. The antibodies were then purified using a Protein A Agarose Kit (KPL) according to the manufacturer's instructions.

Western blot analysis in human tissues

Human tissue specimens were homogenized in a lysis buffer containing 50 mM Tris, pH 7.5, 150 mM NaCl, 2% SDS, 1 mM EDTA, 1 mM PMSF, 1 mM aprotinin and 1 mM chymostatin. The protein concentrations of the tissue extracts were determined using a BCA Protein Assay (Pierce). The protein bands were visualized using an ECL detection system (Amersham-Pharmacia Biotech), or preimmune serum for 1 h at room temperature. The protein bands were then purified using a Protein A Agarose Kit (KPL) according to the manufacturer's instructions.

Subcellular localization of hRad51-Aex9

Mammalian expression constructs of hRad51 and hRad51-Aex9 were generated by PCR-amplifying their full coding sequences from recombinant phage clones using Pfu DNA polymerase (Stratagene) according to the manufacturer's instructions. The sequences of the oligonucleotide primers are available upon request. A unique restriction site, either SacI or BamHI, was introduced into the 5′ end of the primers to allow convenient subcloning. The PCR-amplified DNA fragments were then ligated into pEGFP-C1 (BD Biosciences) in frame with the N-terminal GFP tag. The resulting plasmids were then cotransfected into COS-7 cells using a Zeiss LSM510 laser-scanning microscope.

Mutagenesis of hRad51-Aex9

Site-directed mutagenesis was performed using a PCR-based DpnI-treatment method that has been previously described (28). Mutagenic primers were designed to create R264A, K265Q and Del264RK in the amino acid sequence of hRad51-Aex9. The sequences of the oligonucleotide primers are available upon request. Thermocycling was conducted using Pfu DNA polymerase (Stratagene) according to the manufacturer’s suggestions. The creation of mutations in the hRad51-Aex9 cDNA was confirmed by sequence analysis using the BigDye termination version 3.0 (BD Biosciences). To construct a C-terminal deletion mutant of hRad51, the sequence corresponding to codons 1 to 258 of hRad51 was PCR-amplified using Pfu DNA polymerase (Stratagene) with the following PCR primers: forward, 5′-ccgatgcatgccgaatgctatgcagc-3′; and reverse, 5′-cggtctcaagcaagccatgctgca-3′. A unique restriction site, either SacI or BamHI, was introduced into each primer to allow convenient subcloning. The PCR-amplified DNA fragments were then ligated into pEGFP-C1 (BD Biosciences) in frame with the N-terminal GFP tag.

RESULTS

Identification of hRad51-Aex9

The hRad51 gene is composed of 10 exons that encode a 339-amino acid polypeptide with a calculated molecular mass of 37 kDa. In an effort to identify additional hRad51 paralogs in humans, we searched a human testis cDNA library using a hRad51 cDNA probe with low stringency and obtained seven autoradiographically positive phage recombinants (data not shown). Sequence analysis of the recombinants revealed that all of the isolated clones were hRad51 cDNAs. However, one clone that contained a 1661-bp insert showed an exon–intron structure distinct from that of hRad51, specifically lacking the sequence corresponding to exon 9 of hRad51 (Figure 1). This novel splice variant of hRad51, termed hRad51-Aex9, was also identified in searches of the human EST databases (EST ID number: AI018041). We conducted complete sequencing of EST AI018041 that was obtained from a commercial source and subsequently confirmed that the hRad51-Aex9 cDNA was identical to AI018041, with the following sequence:

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1 2 3 4 5 6 7 8 9 10
1 2 3 4 5 6 7 8 9 10
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Figure 1. Schematic diagrams of the mRNA structures of hRad51 and hRad51-Aex9. Exons are shown as numbered boxes, introns as bold lines. Hatched boxes indicate the deleted exon in the hRad51-Aex9 mRNA. 'S' stands for the start codon, and 'Asterisk' for the stop codon.
exception that hRad51-Dex9 contained longer 5’- and 3’UTR sequences than the EST A1018041 clone.

The 5’-UTR of hRad51-Dex9 is at least 299 bp, the coding region is 843 bp and the 3’-UTR is 469 bp. The deletion of exon 9 causes a frame-shift at codon 259, which leads to premature termination at codon 281. The expected amino acid sequence of the hRad51-Dex9 protein consists of codons 1 to 258 of hRad51 and 22 ‘out of frame’ codons from exon 10, containing the Walker A and B ATP-binding motifs at residues 127–135 and 218–222, respectively (Figure 2). In addition, a basic motif that is composed of one lysine and two arginine residues is located at residues 303–306 of hRad51, and a similar basic motif is found at residues 264–266 in the newly created C-terminus of hRad51-Dex9 (Figure 2).

RT-PCR ANALYSIS OF HRAD51-ΔEX9 IN HUMAN TISSUES

To determine the expression of hRad51 and hRad51-Dex9 in human tissues, RT-PCR analysis was conducted using primers derived from the flanking regions of exon 9. The RT-PCR analysis was expected to generate a 467-bp fragment for hRad51-Dex9 and a 589-bp fragment for hRad51. DNA-amplicons of the expected sizes corresponding to both hRad51 and hRad51-Dex9 were most prominently detected in the testis (Figure 3). Both PCR amplicons were also detected, though to lesser extents, in the skeletal muscle, pancreas, thymus and ovary (Figure 3). Additionally, the hRad51-specific amplicon was detected in the placenta, lung, liver, kidney, spleen and colon tissues, however, the hRad51-Dex9-specific-amplicon was not

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**Figure 2.** Alignment of the amino acid sequences of hRad51 and hRad51-Dex9. The hRad51 polypeptide sequence is aligned with the predicted amino acid sequence of hRad51-Dex9. The 22 ‘out of frame’ codons are indicated with an underline in the amino acid sequence of hRad51-Dex9. Walker A and B ATP-binding motifs and basic motifs are also indicated.

**Figure 3.** RT-PCR analysis of hRad51 and hRad51-Dex9 in human tissues. A typical example of RT-PCR analysis of hRad51 and hRad51-Dex9 using poly(A)+ RNA obtained from 16 different human tissues. The analysis was repeated in triplicate, and GAPDH was used as an internal control.
detected in these tissues, suggesting that different tissue-specificities exist between hRad51 and hRad51-Δex9 (Figure 3).

The DNA strand exchange activity of the hRad51-Δex9 protein

In an effort to express and purify enzymatically active forms of the hRad51 and hRad51-Δex9 proteins, we expressed the full coding domain sequences of hRad51 and hRad51-Δex9 using an E. coli expression system. Upon induction with 1 mM IPTG at 37°C, the hexa-histidine tagged recombinant proteins of both hRad51 and hRad51-Δex9 were expressed at high levels. Fractionation of the cell lysates into different cellular compartments, such as cytoplasmic extracts, periplasmic extracts and inclusion body fractions, revealed that the recombinant hRad51 protein was present in the soluble fractions. However, the recombinant hRad51-Δex9 protein was expressed within the inclusion bodies. The insoluble hRad51-Δex9 protein was denatured by urea during purification and subsequently refolded by stepwise dialysis in the presence of N-lauroylsarcosinate and Cu²⁺. The apparent sizes of the expressed recombinant proteins were in good agreement with the deduced molecular mass, which was 38 kDa for the recombinant Rad51 protein and 32 kDa for the recombinant hRad51-Δex9 protein. The purified recombinant proteins were confirmed by western blot analysis using a commercial human Rad51 antibody (Figure 4A).

To assess the DNA strand exchange activities of hRad51 and hRad51-Δex9, we used the purified recombinant proteins with circular single-strand DNA (ssDNA) and linear double-strand DNA (dsDNA) of bacteriophage φX174. In DNA strand exchange reactions, the circular ssDNA forms joint molecules with the linear dsDNA through homologous pairing, and then the joint molecules are converted into nicked circular forms (Figure 4B). Both the recombinant hRad51 and hRad51-Δex9 proteins showed the expected joint molecules and nicked circular forms of φX174 at each of the time-intervals tested. The intensities of the bands corresponding to the nicked circular form appeared approximately the same in the either reactions with hRad51 or hRad51-Δex9 (Figure 4C), suggesting that strand exchange activity of hRad51-Δex9 is approximately similar to that of hRad51 at least in vitro. However, the hRad51-Δex9 protein showed a significantly higher activity than hRad51 in homologous DNA pairing at all the time-intervals (Figure 4C). These results are comparable with the previous findings on C-terminal deletion mutants of the E. coli RecA protein, which also showed an enhanced activity in homologous DNA pairing (29–31).

Western blot analysis of hRad51-Δex9 in human tissues

To evaluate the expression of hRad51-Δex9 at the protein level in vivo, we generated a polyclonal antibody against the peptide sequence specific to hRad51-Δex9. This hRad51-Δex9 polyclonal antibody reacted with the purified recombinant hRad51-Δex9 protein, but not with the recombinant hRad51 protein (data not shown). Human placenta, lung, testis and small intestine tissues were then tested by western blot analysis. A band with the expected molecular mass of 31 kDa for hRad51-Δex9 was prominently detected in the testis; however, this 31-kDa band was rarely detected in the other tissues tested (Figure 5A). We also investigated the expression of hRad51 and hRad51-Δex9 using a commercial antibody expected to react with both hRad51 and hRad51-Δex9. The 37-kDa hRad51 band was prominently detected in the testis, but at much lower levels in the placenta, lung and small intestine (Figure 5B). The 31-kDa band corresponding to hRad51-Δex9, however, was detected only in the testis (Figure 5A). These findings are consistent with those of the RT-PCR analysis that also showed prominent expression of hRad51-Δex9 only in the testis.

Nuclear localization of hRad51-Δex9

To investigate the cellular localization of hRad51 and hRad51-Δex9, mammalian expression constructs containing the full coding sequence of hRad51 or hRad51-Δex9 in frame with the N-terminal GFP tag were transfected into COS-7 cells. Confocal microscopic analysis of the direct fluorescence of the fusion proteins displayed subcellular signals of hRad51 and hRad51-Δex9 in the nucleus (Figure 6Aa and b). In addition, both the hRad51 and hRad51-Δex9 proteins were co-localized with nucleus-specific propidium iodide staining, further confirming the
nuclear localization of these proteins in the transfected cells (data not shown). However, the mutated hRad51 protein that did not contain the C-terminal region from codons 259 to 339 was primarily detected in the cytoplasmic area (Figure 6Ac). Taken together, these results indicate that the signal for the nuclear localization of hRad51 may reside in the C-terminus and, furthermore, that the frame-shifted region of hRad51-Δex9 may regain the residues required for nuclear localization.

A basic motif containing a stretch of lysine and arginine residues was found at residues 264–266 (RKR) in the frame-shifted C-terminal region of hRad51-Δex9. Similar types of basic motifs have been known to act as a nuclear localization signal (NLS) in a number of nuclear proteins (32,33). To determine, therefore, if this basic motif in the C-terminus of hRad51-Δex9 could function as an NLS, we generated a series of mutant constructs that harbor a del254-256RK, R264A or K265Q mutation in the basic motif. In localization studies conducted using the mutant constructs, each of the mutated hRad51-Δex9 proteins was primarily detected in the cytoplasmic areas, but rarely in the nuclei (Figure 6Ba–c). These results strongly suggest that the basic motif located in the newly created C-terminal region of hRad51-Δex9 may function as a NLS in nuclear localization of this hRad51 variant.

Figure 5. Detection of hRad51-Δex9 in human tissues by western blot analysis. Approximately 100 μg of human placenta, lung, testis and small intestine tissue extracts were subjected to western blot analysis using a hRad51-Δex9-specific antibody (A) or a commercial hRad51 antibody (B).

Figure 6. Nuclear localization of hRad51, hRad51-Δex9 and C-terminal mutants. (A) Direct fluorescence images of COS-7 cells transfected with hRad51 (a), hRad51-Δex9 (b), or hRad51-delC (c) at a magnification of ×1000. The amino acid sequences of hRad51 and hRad51-Δex9 are shown only from codons 241 to 280, and the frame-shifted region in hRad51-Δex9 is underlined. The hRad51-ΔC mutant does not contain the C-terminal sequence from codons 259 to 339. (B) Direct fluorescence images of COS-7 cells transfected with hRad51-Δex9-K265Q (a), hRad51-Δex9-R264A (b), or hRad51-Δex9-Del264_265RK (c) at a magnification of ×1000. The hRad51-Δex9-K265Q mutant harbors a substitution of Lys to Gln at codon 265, and the hRad51-Δex9-R264A mutant contains a substitution of Arg to Ala at codon 264. The residues mutated in hRad51-Δex9-K265Q and hRad51-Δex9-R264A are indicated in blue. In hRad51-Δex9-Del264_265RK, Arg-Lys residues at codons 264–265 are deleted.
DISCUSSION

Here we present a novel variant of hRad51, hRad51-Dex9, which aberrantly splices the hRad51 mRNA from exon 8 to exon 10, skipping exon 9. The predicted amino acid sequence of this novel variant contains a truncated C-terminus of hRad51, however, it retains the RecA domain for DNA recombination and the Walker A and B motifs for ATP binding and hydrolysis. With a purified recombinant hRad51-Dex9 protein, we showed that this novel variant is capable of catalyzing DNA strand exchanges in vitro, although further biochemical characterization would be required to determine the precise enzymatic properties of this hRad51 variant. In expression studies, hRad51-Dex9 was predominantly detected in the testis at both the mRNA and protein levels and, further, the hRad51-Dex9 protein was localized in the nucleus. Taken together, these findings indicate that hRad51-Dex9 catalyzes homologous pairing and DNA-strand exchange in the nucleus, suggesting that alternative pathways involving either hRad51 or hRad51-Dex9 may exist for DNA repair and recombination.

splice variants of other genes involved in DNA repair and recombination, including Rad52, Rad51D and DMC1, have been also reported (34–38). The murine and human Rad52 mRNAs undergo alternative splicing, resulting in several variants with a truncated C-terminus (34,35). Rad52 is known to catalyze the replacement of replication protein A with Rad51 on ssDNA and to promote strand exchange between complementary ssDNA and dsDNA (39,40). The human Rad52 variants interacted with both ssDNA and dsDNA; however, they did not bind to the full-length human Rad52 due to deletion of the self-interaction domain (34). Furthermore, the murine Rad52 splice variants increased the frequency of sister chromatid repair in both mammalian cells and yeast, whereas the intact murine Rad52 was more likely involved in homology-directed repair (35). Alternatively spliced forms of Rad51D and DMC1 in both humans and mice have been also identified, but their functional significance has not been evaluated (36–38). However, the presence of these variants of the proteins involved in HR further implies the presence of alternative pathways for the control of recombinational repair of dsDNA breaks.

Rad51 and its paralogs are found in the nucleus, however, it has not yet been determined if they are transported independently into the nucleus or through interactions with other proteins. BRCA2 has been known to play a critical role in the nuclear transport and foci formation of Rad51 upon exposure to exogenous damage (9–11). However, without any exogenous DNA damage, replication-associated formation of Rad51 foci occurred in a BRCA2-independent manner in CAPAN-1 cells that carry a BRCA2 truncation (41), suggesting that distinct mechanisms may be responsible for the nuclear localization and focus formation of Rad51 in the presence or absence of exogenous DNA-damaging agents. Further, several hRad51 paralogs have been shown to translocate into nucleus in a BRCA2-independent manner, using a basic motif composed of lysines and arginines as a NLS (42,43). hRad51C contains a basic motif composed of a short stretch of lysine and arginine residues at the C-terminus. Using a deletion construct of the C-terminal region, the basic motif of hRad51C was shown to function as a NLS for nuclear transport of hRad51C in mammalian cells (42). In addition, hRad51B that contains a basic motif at the N-terminus was shown to translocate into the nucleus in a BRCA2-independent manner (43). hRad51 also contains a basic motif at residues 303–306 (RKGR) in the C-terminus. This basic motif is required for the nuclear localization of hRad51-Dex9, suggesting that nuclear localization of hRad51-Dex9 may be independent of BRCA2 in the absence of any DNA-damaging agents, at least in the cultured cells tested.

Rad51 has been reported to interact with p53 and BRCA2, both of which play pivotal roles in maintaining genome integrity. In response to DNA damage, p53 modulates HR through physical interaction with several proteins implicated in recombination, including Rad51, Rad54, BLM and WRN (44,45). Using in vitro binding assays, p53 was reported to interact with the region between codons 125 and 220 of hRad51 (8). The p53-interactive region in hRad51 corresponds to the homologomerization region that is critical for formation of the functional hRad51 nucleoprotein filaments (46). The conservation of the p53-interactive region in hRad51-Dex9 suggests that this novel variant also interacts with p53, unless the absence of the C-terminal region in hRad51-Dex9 affects the physical interaction with p53. BRCA2 interacts with Rad51 through the eight conserved BRC repeats (47,48), and mutations within these repeats are associated with an increased risk of breast cancer (49,50). Electron microscopy studies showed that the BRC repeat 4 interacts with the nucleotide-binding core of Rad51, whereas the BRC repeat 3 interacts with the N-terminal region of Rad51, suggesting that the BRC repeats bind to distinct regions of Rad51 (51). The BRCA2-interactive region in hRad51 was studied using yeast two-hybrid and in vitro binding assays, which revealed that the C-terminus of hRad51 (codons 98–339) is crucial for interaction with BRCA2 (47). Our finding that the C-terminal region (codons 280–389) of hRad51 is deleted in hRad51-Dex9 suggests that this novel variant may have a different binding property from hRad51 in interaction with BRCA2. Further characterization of the interactive profile of hRad51-Dex9, particularly with p53 and BRCA2, will be necessary to determine the functional roles that this novel recombinase may play in the maintenance of genome stability and the elimination of DNA double-strand breaks.

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