Alternative end-joining is suppressed by the canonical NHEJ component Xrcc4/ligase IV during chromosomal translocation formation

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Abstract
Chromosomal translocations in hematologic and mesenchymal tumors form overwhelmingly by nonhomologous end-joining (NHEJ). Canonical NHEJ, essential for the repair of radiation-induced and some programmed double-strand breaks (DSBs), requires the Xrcc4/ligase IV complex. For other DSBs, the requirement for Xrcc4/ligase IV is less stringent, suggesting the existence of alternative end-joining (alt-NHEJ) pathways. To understand the contribution of the canonical and alt-NHEJ pathways, we examined translocation formation in Xrcc4/ligase IV-deficient cells. We find that Xrcc4/ligase IV is not required for, but rather suppresses, translocations. Translocation breakpoint junctions have similar characteristics in wild-type and Xrcc4/ligase IV-deficient cells, including an unchanged bias toward microhomology, unlike what is observed for intrachromosomal DSB repair. Complex insertions in some junctions demonstrate that joining can be iterative, encompassing successive processing steps prior to joining. Our results imply that alt-NHEJ is the primary mediator of translocation formation in mammalian cells.

Keywords
chromosomal translocation; Xrcc4/ligase IV; nonhomologous end-joining; DNA double-strand break; alternative NHEJ

Introduction
NHEJ is a major pathway for the repair of DSBs in mammalian cells¹. NHEJ is loosely defined as the joining of DNA ends without the use of extensive homology, and it is important for the correct repair of DSBs generated by ionizing radiation, as well as programmed DSBs generated in developing lymphocytes during V(D)J recombination and class switch recombination (CSR)²–⁴. Paradoxically, some type of NHEJ also appears to be responsible for generating reciprocal translocations found in human hematologic and
mesenchymal tumors, since breakpoint junctions rarely occur within homologous sequences. Rather, breakpoint junctions occur at sites of no homology or just a few bp of homology, termed microhomology. Translocation breakpoint junctions have other features of NHEJ in that they contain DNA end modifications, most often deletions, but also insertions.

A set of NHEJ factors has been identified that is required for the repair of DSBs generated by ionizing radiation and during V(D)J recombination. These canonical NHEJ factors include the DNA ligase complex, Xrcc4/ligase IV, used to seal DNA ends, and the Ku70/80 (Ku) heterodimer, which binds DNA ends and recruits a number of processing factors, including nucleases, polymerases, and the ligase complex itself. Xrcc4/ligase IV may also participate in DNA end processing through its interaction with a variety of factors. The range of activities of the recruited factors allows the NHEJ pathway to process DSBs with a variety of end structures so that they can be ligated by Xrcc4/ligase IV. End processing is complex, such that a variety of junctional sequences are possible at a single lesion. Biochemical studies support this complexity, and have even shown that the two strands of a DNA end can be independently processed and ligated in an iterative manner. These NHEJ components also suppress the other major DSB repair pathway in mammalian cells, homologous recombination (HR), presumably by protecting DNA ends from the initial resection step necessary for HR.

While critical for the repair of radiation-induced and V(D)J recombination DSBs, the canonical NHEJ factors are not absolutely required for other types of DSB repair. For example, endonuclease-generated DSBs in plasmids and chromosomes can be repaired at reduced levels in Ku and Xrcc4-deficient cells. In the immune system, Xrcc4/ligase IV promotes the efficient repair of DSBs generated during CSR, but is not essential for this process. These studies point to the existence of other pathways for the nonhomologous repair of DSBs. The poorly defined pathway(s) operating in the absence of the canonical NHEJ pathway has been termed alternative NHEJ, or alt-NHEJ.

Linked to their roles in the immune system, deficiency of several of the canonical NHEJ components in mice on a p53 background leads to pro-B cell lymphomas with Igh/Myc amplification and translocations at nonhomologous sequences. Translocations involving nonhomologous sequences at V(D)J and CSR programmed DSBs have also recently been reported in cells deficient in Xrcc4. In a more generalized system, a translocation reporter was used to demonstrate that translocations are not reduced in Ku70-deficient mouse embryonic stem (ES) cells, suggesting a role of alt-NHEJ in non-immune system translocations. However, absence of Xrcc4/ligase IV has a more severe phenotype than loss of Ku in several contexts, including telomere fusions, mouse development, and the joining of two nearby chromosome DSBs, suggesting that loss of the terminal activity of NHEJ has a more profound phenotype. Moreover, Ku is implicated in other cellular processes besides NHEJ, such as apoptosis, that may affect the recovery of translocations.

Given that Xrcc4/ligase IV does not have any other known cellular role outside of NHEJ and ligation of DNA ends is essential to NHEJ, we have now examined the role of this canonical
NHEJ component in translocation formation outside of the immune system. We find that Xrcc4/ligase IV is not required for NHEJ leading to translocations, but rather suppresses such events. Moreover, translocation breakpoint junctions have similar characteristics in wild-type and Xrcc4/ligase IV-deficient mouse cells, including a similar bias to microhomology use, as well as iterative processing of the ends to generate complex insertions. Our results imply that non-canonical NHEJ (alt-NHEJ) is the primary mediator of translocation formation in mammalian cells.

Results

Chromosomal translocations do not require Xrcc4

Ligase IV requires Xrcc4 for its stability, such that Xrcc4-deficient cells are effectively ligase IV-deficient\textsuperscript{20,30}. Xrcc4\textsuperscript{−/−} mouse cells were previously constructed by gene targeting mouse ES cells\textsuperscript{26}. The Xrcc4\textsuperscript{−/−} ES cells, like other NHEJ-deficient cells, are sensitive to ionizing radiation, defective in V(D)J recombination, and have elevated homology-directed DNA repair\textsuperscript{14,24,26}. Because a neomycin phosphotransferase (neo) gene, which would interfere with translocation selection, was still present at the targeted Xrcc4 loci in these cells, we first expressed Cre recombinase to delete the neo gene (Supplementary Fig. 1a).

Two rounds of gene targeting were then used to introduce our pCr15 translocation reporter into the neo–Xrcc4\textsuperscript{−/−} cells. This reporter consists of a neo gene split into two exons, one of which is targeted to a locus on chromosome (chr.) 17 and the other to a locus on chr. 14 (Fig. 1a)\textsuperscript{6}. The intron segment on each chromosome is demarcated by an I-SceI site, such that cleavage of both chromosomes by I-SceI endonuclease, followed by joining of DNA ends from chrs. 17 and 14 can generate a derivative chromosome 17, der(17), with a functional neo gene. Formation of a neo\textsuperscript{+} gene requires NHEJ, as the neo intron segments on chrs. 17 and 14 do not share any appreciable homology.

To determine the role of Xrcc4/ligase IV in translocation formation, we expressed I-SceI in the Xrcc4\textsuperscript{−/−} pCr15 cells and selected neo\textsuperscript{+} colonies. Surprisingly, neo\textsuperscript{+} colonies were readily obtained at a frequency of 2.7 × 10\textsuperscript{−4} (Fig. 1b, Table 1). We confirmed that the neo\textsuperscript{+} colonies arose as a result of chromosomal translocation by using fluorescence in situ hybridization (FISH; Fig. 1a, Supplementary Fig. 1c, and data not shown). All 12 translocations analyzed by FISH were reciprocal, as both the expected der(17) and der(14) chromosomes were detected. The Xrcc4 genotype of these cells was verified (Fig. 1c), as was the absence of Xrcc4 protein (Fig. 1d). Transient Xrcc4 expression in the Xrcc4\textsuperscript{−/−} cells restored wild-type levels of protein (Fig. 1d) and led to a 5-fold reduction in translocation frequency (Fig. 1c), indicating that the NHEJ complex Xrcc4/ligase IV is not required for NHEJ-mediated translocations but instead suppresses their occurrence.

Alternative NHEJ is implicated in translocation formation

Using a similar approach, we previously demonstrated that like Xrcc4, Ku70 is also not required for translocation formation\textsuperscript{24}. We compared Xrcc4/ligase IV and Ku for their role in translocation formation by directly comparing wild-type, Xrcc4\textsuperscript{−/−}, and Ku70\textsuperscript{−/−} pCr15 ES cells (Table 1). A consistent 5-fold higher translocation frequency was observed in the absence of Xrcc4 (p<0.0001) and a 3-fold higher translocation frequency in the absence of

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Ku70 (p<0.0001). The somewhat higher suppression by Xrcc4 than by Ku70 carries only a moderate level of statistical significance (p=0.043). These experiments establish that, unlike in yeast, the canonical NHEJ factors Xrcc4/ligase IV and Ku are not essential for chromosomal translocations in mammalian cells; rather, both canonical NHEJ proteins restrain translocation formation. Our results, therefore, implicate alt-NHEJ in translocation formation.

Translocation junctions are similar in wild type and NHEJ mutants

Analysis of NHEJ junctions from Xrcc4-deficient cells has previously indicated that these cells join DNA ends very differently from wild-type cells. To gain insight into the repair mechanisms leading to translocations, der(17) breakpoint junctions were analyzed from 92 Xrcc4−/−, 92 Xrcc4-complemented Xrcc4−/−, and 43 wild-type neo+ clones. For comparison, der(17) breakpoint junctions were analyzed from 42 Ku70−/− neo+ clones. Breakpoint junctions were amplified using a series of primer pairs that span the neo intron formed upon translocation (Fig. 1d). With this approach, a total of 264 of 269 breakpoint junctions could be amplified for sequencing (Supplementary Fig. 2).

The two chromosome ends that join during translocation formation have 4-base 3′ overhangs with limited complementarity, requiring processing of the ends prior to joining. A variety of DNA end modifications were observed in all genotypes (Fig. 2a and Supplementary Fig. 2). As with patient-derived translocations, deletions were the most commonly observed modification. Deletions were from one or both ends and were often short, encompassing only bases from the overhangs or the I-SceI recognition site, although longer deletions were also observed. The longest deletion was 2.1 kb and was split approximately equally between the two chromosome ends (1.0 and 1.1 kb). Longer deletions may have arisen but not survived the neo selection, although these would be expected to be a minority of events (~5%). One of the shortest deletions was missing only the terminal base of each 3′ overhang (Xrcc4−/−, Δ2; Fig. 2a); the remaining 3 bases of the 3′ overhangs were apparently filled in by a polymerase prior to joining. The lack of microhomology in this junction suggests the involvement of polymerase (pol) µ in the fill-in reaction, since this polymerase has activity even when the terminal base of the primer lacks complementarity to the template strand.

Overall, the median deletion length for wild-type, Xrcc4-complemented Xrcc4−/−, and Xrcc4−/− junctions ranged from 23 to 29 bp and was not significantly different for the three genotypes (Fig. 2b). For Ku70−/−, the median deletion length was longer (47 bp), although it also was not statistically different. The majority of deletions were <100 bp, but larger deletions of >250 bp were found in a portion of junctions for all 4 genotypes (Fig. 2b and Supplementary Fig. 3a). It is unclear what accounts for the range of deletions from the DNA ends prior to joining; nonetheless, the similarities found in the 4 genotypes indicate that loss of the canonical NHEJ components Xrcc4/ligase IV and Ku does not substantially alter the amount of nucleolytic processing prior to joining chromosome ends for translocation formation.

It has previously been reported that junctions formed during CSR in Xrcc4−/− cells are markedly biased toward those with microhomology compared with wild-type cells. We
determined the microhomology present at translocation junctions and found that the percent of total junctions containing microhomology was similar in Xrcc4−/− cells (60%) and wild-type and Xrcc4-complemented Xrcc4−/− cells (64%), although it was slightly lower in the Ku70−/− cells (54%) (Fig. 2c). Considering only the junctions without insertions, the percent of junctions containing microhomology was similar for all genotypes (wild-type, 82%; Xrcc4−/−, 76%; Xrcc4-complemented Xrcc4−/−, 78%; Ku70−/−, 76%). In all cases, the presence of microhomology was significantly greater than expected by chance, as only 44% of junctions would be expected to have microhomology if joining was random with respect to bp overlap (Fig. 2d). These results are consistent with a role for microhomology in NHEJ, but the Xrcc4−/− cells do not demonstrate a greater dependence on microhomology.

Consistent with the percent microhomology use, the distribution of microhomology lengths was similar between Xrcc4−/− and the other genotypes (Fig. 2d). Microhomology lengths were generally short, from 1 to 4 bp. The fraction of junctions with 1 bp of microhomology (16–21%) was lower than that expected by chance (28%); however, the fraction of junctions with 2 or more bp of microhomology was much greater than that expected by chance (Fig. 2d). For example, 22 to 27% of junctions were observed to have 2 bp of microhomology, whereas 10.5% of junctions would be expected by chance, a 2 to 2.6-fold higher level. The fold difference is even greater with 3 and ≥4 bp microhomology (~5 and 10-fold, respectively). Microhomologies of 5 to 7 bp were observed in a few junctions (6/207 junctions; 3%), although not in Xrcc4−/− junctions (Supplementary Fig. 2). No correlation was observed between deletion length and microhomology length for any of the genotypes (Fig. 2e and Supplementary Fig. 3b), indicating that joining of distant ends is not driven by microhomology to any greater extent than ends closer to the DSB.

Although we observed no difference in either the percent microhomology use or the microhomology length at the translocation breakpoint junctions in the various cell lines, previous analyses of NHEJ in several systems have demonstrated increased microhomology use in the absence of Xrcc4/ligase IV32,33. The bias previously observed may be sequence specific, for example, for repetitive switch regions20 and plasmids substrates with short repeats at or near the DNA ends32,33. Alternatively, the context of the DSB (intrachromosomal vs. interchromosomal; single DSB vs. two DSBs) may affect NHEJ pathway use. To determine if a single DSB is repaired with different microhomology characteristics, we analyzed intrachromosomal junctions after I-SceI expression in complemented and uncomplemented Xrcc4−/− cells. To maintain a similar sequence context for the joining events to those for translocations, we utilized a translocation clone derived from Xrcc4−/− cells that had by chance reconstituted an I-SceI site on der(17) (Supplementary Fig. 4a). I-SceI was expressed in these cells with or without Xrcc4 coexpression, and 72 h later pooled genomic DNA was analyzed by PCR and sequencing for imprecise NHEJ (I-SceI site loss; Supplementary Fig. 4a). We found that microhomology use differed in junctions from complemented and uncomplemented Xrcc4−/− cells. To maintain a similar sequence context for the joining events to those for translocations, we utilized a translocation clone derived from Xrcc4−/− cells that had by chance reconstituted an I-SceI site on der(17) (Supplementary Fig. 4a). I-SceI was expressed in these cells with or without Xrcc4 coexpression, and 72 h later pooled genomic DNA was analyzed by PCR and sequencing for imprecise NHEJ (I-SceI site loss; Supplementary Fig. 4a). We found that microhomology use differed in junctions from complemented and uncomplemented Xrcc4−/− cells (Supplementary Fig. 4b). Microhomology use in complemented Xrcc4−/− cells was not significantly different from that expected by chance (p=0.2667, one-tailed Mann-Whitney test), whereas more long microhomologies were observed in Xrcc4−/− cells, such that the distribution of microhomology use differed significantly from that expected by chance.
Thus, NHEJ of a single DSB is affected by Xrcc4 loss, unlike NHEJ of two DSBs during translocation. Still, 32% of junctions from Xrcc4<sup>−/−</sup> cells did not contain microhomology.

**Breakpoint junction insertions can be complex**

Although less frequent than deletions, insertions were also observed in a portion of the breakpoint junctions from both wild-type and mutant cells (18–29%; Fig. 2c). Approximately half of the insertions in all 4 genotypes were short (<10 bp), while the remainder were longer, extending a few hundred bp in some cases (Fig. 2a and Supplementary Figs. 2, 3c, and 5). Some of the insertions were only 1 or a few bp, and may have been added by polymerases in a template-independent manner or by template slippage<sup>34,35</sup>. Other somewhat longer insertions appear to have been templated, for example, the +5 (CAGGG) and +10 (CAGGTAATTG) insertions derived from Xrcc4<sup>−/−</sup> mutant and complemented cells, respectively (Fig. 2a). In both cases, the terminal 3 nucleotides of the chr.17 end could have primed DNA synthesis starting 11 nucleotides into the chr. 14 end (red shading, Fig. 2a); copying of at least 7 and 11 bp from chr. 14, respectively, would have provided small microhomologies for subsequent annealing to the chr. 14 end. In such a scenario, the primer for extension and the template sequences are derived from the two der(17) chromosome ends, indicating that DNA ends in these cases would have been synapsed prior to DNA synthesis.

The longest insertions for the Xrcc4<sup>−/−</sup> mutant and Xrcc4-complemented der(17) junctions, which were 231 and 396 bp, respectively, point to the complexity of breakpoint junctions with longer insertions in terms of both the derivation of the insert and the processing of each chromosome end (Fig. 2a). The 396-bp insertion was derived from sequences starting 7.7 kb upstream of the DSB on chr. 17 and was inserted after a 965 bp deletion from the chr. 17 end and a 892 bp deletion from the chr. 14 end (Supplementary Fig. 5c). The 231 bp insertion was derived from sequences starting 21 bp upstream from the DSB on chr. 14, and, hence, from a chromosome end forming the other derivative chromosome, der(14); it was inserted with no deletion from the chr. 17 end but with a 56 bp deletion from the chr. 14 end (Supplementary Fig. 5b). Because this insertion is derived from sequences forming der(14), it must have been inserted prior to synapsis of the der(17) chromosome ends. The inserted segment from chr. 14 shows an additional complexity in that it is not contiguous; it is interrupted by 2 bp (“GA”; position 161).

The complexity of the insertions led us to examine several parameters. We found that breakpoint junctions with insertions had a greater median deletion length than junctions without insertions (114 vs 25 bp, p<0.0001, two-tailed Mann-Whitney test; Fig. 3a), indicating that insertions were associated with more extensive processing of DNA ends. All 4 genotypes showed a similar trend (Supplementary Fig. 3d). Although most insertions had a unique origin, others were chimeric, being comprised of 2, 3, or even 4 inserts derived from different sources (Fig. 3b). This suggests some type of iterative processing of ends until joining is successful.

Most inserts were derived from the 2 chromosomes participating in the translocation (chr. 17, 24 inserts; chr. 14, 22 inserts), while the remaining 14 inserts were derived from other
sources, including a segment from chr. 13, a repetitive element (LINE1), and the transfected I-SceI expression vector (Supplementary Fig. 5). Microhomology was noted at one or both ends of several of the insertions, although some insertions did not have overt microhomology. Seven inserts had single bp mutations, including point mutations, deletions, and additions, indicating the involvement of an error prone polymerase. Some very short inserts of 1 to 3 bp were also observed with the longer inserts, like the “GA” described above. These may have been derived from an error prone polymerase; the alternative is template independent polymerization. The origin of the remaining 10 inserts, which ranged from 5 to 24 bp, could not be unequivocally determined.

We mapped the inserts in the der(17) breakpoint junctions on chrs. 17 and 14 (Fig. 3c). Notably, inserts were derived from all 4 chromosome ends, indicating the proximity of the ends during insertion formation. Inserts from the ~20 bp adjacent to the der(17) ends were overrepresented (15/46, 33%), making it likely that these were inserted during the der(17) joining process itself. More distant inserts did not show this bias, as der(17) and der(14) sequences were similarly represented (16 vs. 14, respectively); thus, many of these inserts were derived at an earlier step in the joining process. A distance effect was also noted in that inserts derived further from the DSB were progressively longer on average (Fig. 3c). If the inserts are derived from a copying mechanism, polymerase processivity may be affected by proximity to a DNA end. Interestingly, the most distant inserts, from 4.7 Mb away on distal chr. 17, were derived from within ~1 kb from each other from the same gene, Psmb9. Psmb9 is a known hotspot for meiotic recombination in some strains of mice36,37, although to date this gene has not been known to participate in somatic recombination or rearrangement events.

Discussion

In this report, we examined chromosomal translocations in the absence of the NHEJ ligase. In contrast to NHEJ in other systems16,17,20,27, translocations via NHEJ occur at a significantly higher frequency in the absence of Xrcc4/ligase IV. Moreover, microhomology at translocation breakpoint junctions is unaltered, also unlike other systems20,32,33 or our results with intrachromosomal joining (Supplementary Fig. 4). These results imply a suppressive role for the canonical NHEJ pathway in preventing translocations and support a central role for alt-NHEJ in translocation formation.

The question arises as to whether alt-NHEJ is a distinct pathway or a variation of the canonical pathway. The similar results we obtained in Xrcc4/ligase IV and Ku-deficient cells, both in terms of the translocation frequency and breakpoint junction characteristics, support a role for an alternative pathway of NHEJ as the major pathway in translocation formation, rather than a variation or back-up of the canonical pathway, as it seems to be in other systems. That wild-type cells have similar junction characteristics as the two mutants further supports this view. We cannot rule out, however, that other factors, for example ligase I or ligase III38–40 could slip into the canonical pathway without perturbing its outcome.
Translocation breakpoint junctions in wild-type and NHEJ-deficient cells, as well as junctions at a single DSB in the absence of Xrcc4, show biased microhomology use: an overrepresentation of 2 or more bp of microhomology and an underrepresentation of 0 or 1 bp. Microhomology would promote annealing of two DNA ends for subsequent processing, including polymerization, and ligation. For junctions with 0 or 1 bp microhomology, another mechanism may be used to stabilize ends prior to joining. Alternatively, a polymerase like pol µ may generate chance microhomology by adding nucleotides to DNA in a template-independent manner; this polymerase-generated microhomology would then act to promote annealing between chromosome ends, although it would not be observed in breakpoint junctions as microhomology. Interestingly, no correlation was observed between deletion length and microhomology use for any of the cell lines examined, implying that extensive nucleolytic digestion of DNA ends is not coupled to a search for microhomology.

Xrcc4/ligase IV and Ku are known to suppress two other DSB repair pathways, HR and single-strand annealing. Both of these pathways are initiated by DSB resection to produce 3’ single-stranded DNA tails. Since both Xrcc4/ligase IV and Ku are known to protect DNA ends from nucleolytic processing, it is tempting to speculate that canonical NHEJ components perform a similar role in suppressing translocations, by blocking access of DNA ends to alt-NHEJ components involved in end processing. A possibly related or additional role for the canonical NHEJ components is maintaining DNA ends from a DSB in close physical proximity through their end-binding activity to promote correct joining. Such a role has been suggested for Ku, since in its absence, DNA ends undergo long-range movements which would promote translocation formation. Underlying these considerations are the kinetics of NHEJ, since DSB repair occurs very quickly in wild-type cells, whereas DSBs persist in Xrcc4/ligase IV deficient cells for long periods.

It is notable in this regard that we do not see a statistically significant increase in the median deletion length at the translocation breakpoint junctions in the mutant cells compared with wild type. Thus, the increased translocation frequency in Xrcc4/ligase IV and Ku mutant cells suggests that nucleases may have access to more DNA ends for processing rather than affecting the total amount of nucleolytic processing at a particular end; alternatively, nucleases may increase resection at a DNA end, but only on one strand so as to maintain a similar deletion distribution.

Breakpoint junctions with multiple inserts provide the clearest and most complex example of iterative processing for NHEJ. For these junctions, nucleolytic processing of the DNA ends occurred, and subsequently, an insert was added to one or both ends. For junctions with three or four inserts, at least one of the ends had undergone a second round of insert addition. Since longer deletions are found in breakpoint junctions with insertions compared to those without insertions, multiple rounds of nucleolytic processing may have occurred as well. This iterative process for alt-NHEJ may facilitate joining or it may be the result of nonproductive processing events in a sort of “testing” process.

Many insertions duplicate an existing chromosomal sequence, suggesting a model whereby the insertions are added by DNA synthesis. In such a mechanism, a DNA end would prime synthesis from another chromosomal sequence acting as a template. Templated insertions...
have been termed “T nucleotides” to distinguish them from non-templated insertions\textsuperscript{46}. \textit{In vitro}, pol \( \mu \) and pol \( \lambda \) are capable of polymerizing from DNA ends that have little complementarity (pol \( \mu \) and pol \( \lambda \)) or no complementarity (pol \( \mu \)) to the template\textsuperscript{34}, and so either (or both) may be involved in these insertions. Both pol \( \mu \) and pol \( \lambda \) interact with Ku/Xrcc4/ligase IV/DNA and, where tested, this interaction stimulates polymerization\textsuperscript{10,13}. However, as we found insertions in Ku and Xrcc4-deficient cells, this interaction may not be critical for activity.

The derivation of the breakpoint junction insertions seems remarkable. Although many are derived from sequences within 10 to 100 bp from the DSB, others are derived from sequences kilobases away from the DSB. As the insertions from nearby the DSB were typically derived from the ends that form der(17), many were likely inserted during the der(17) joining process itself. Insertions from further away were nearly as likely to be derived from sequences that form der(14) as those that form der(17), suggesting that they were inserted at an earlier stage, possibly during an attempt at intrachromosomal repair. The two most distant insertions are derived from megabases away from one of the der(14) ends and from the same gene, \( \text{Psmb9} \), a meiotic hotspot of recombination in some mouse strains\textsuperscript{36,37}. The \( \text{Psmb9} \) insertions are not from the hotspot itself located at the 3′ end of the gene, but from 5′ UTR and intron sequences. Interestingly, one report has linked sequences frequently involved in translocations in somatic cells with meiotic hotspot activity, suggesting that some regions are susceptible to cleavage (breakage) in both somatic and meiotic cells\textsuperscript{47}.

In summary, alt-NHEJ appears to be the primary mediator of translocation formation in mammalian cells. Characteristics of alt-NHEJ are emerging, including a bias, but not a strict requirement, for microhomology and a role for iterative processing. Elucidating this pathway will be important for understanding mechanisms leading to oncogenic translocations.

**Materials and Methods**

**Cell culture and gene targeting**

\( \text{Xrcc4}^{-/-} \) mouse ES cells were provided by F. Alt (Harvard)\textsuperscript{26}. For construction of neo−cells (Supplementary Fig. 1a), \( \text{Xrcc4}^{-/-} \) cells were transfected with 15 \( \mu \)g pCAGGS-Cre expression vector, and G418 sensitive clones were isolated. Clones were confirmed to have undergone Cre-mediated \( \text{loxP} \) recombination to remove the neo cassette at both \( \text{Xrcc4} \) loci by PCR using primers shown in Supplementary Fig. 1b. Two sets of primers (KMO74, KMO75 and KMO77) and (KMO78 and KMO79) were used. The conditions for both PCRs, are as follows: denaturation at 95°C for 3 min, followed by 30 cycles at 95°C for 1 min, 64°C for 1 min and 72°C for 2 min.

Targeting vectors for chr.17, pC-I (\( \text{Pim1} \)), and chr. 14, rdPuroRev (\( \text{Rb} \)), were previously described\textsuperscript{6}. Targeting was confirmed at the \( \text{Pim1} \) locus using the p22 probe (a HincII-BstXI fragment) in conjunction with HincII digestion of genomic DNA\textsuperscript{48}. Targeting at the \( \text{Rb} \) locus was confirmed using probe A (a PstI-PvuII fragment) in conjunction with HindIII digestion of genomic DNA\textsuperscript{49}. The expression vector for I-SceI endonuclease, pCBASce, and

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pCAGGS were previously described\textsuperscript{50}. Human \textit{XRCC4}, a gift from J. Chaudhuri (MSKCC), was subcloned into the XhoI and XbaI sites of pCAGGS.

**Translocation analysis**

For translocation experiments, 1 × 10\textsuperscript{6} cells per well were plated in 6-well plates and transfected using Lipofectamine 2000 (Invitrogen), according to the manufacturer’s instructions. DNA for transfection was as follows: I-SceI only, 4 µg I-SceI and 4 µg GFP expression vectors; for Xrcc4 or Ku70 complementation: 4 µg SceI, 3.5 µg GFP, and 0.5 µg Xrcc4 or 4 µg SceI, and 4 µg Ku70 expression vectors, respectively. (Note: The Xrcc4 expression vector was titrated to reach wild-type levels for complementation, as analyzed by Western blotting and resistance to ionizing radiation; data not shown). 4 h after Lipofectamine addition, each well was split into two 10 cm plates; 24 h after splitting, cells were selected in 200 µg ml\textsuperscript{−1} G418. Translocation frequency was obtained by normalizing the number of \textit{neo}\textsuperscript{+} clones obtained after 7 to 10 d to the number of cells counted at 4 h post-transfection.

For translocation junction analysis, DNA was isolated from \textit{neo}\textsuperscript{+} clones using the Genelute Mammalian Genomic DNA Miniprep Kit (Sigma). PCR amplification was performed with the high fidelity Herculase II Fusion DNA Polymerase (Stratagene) under the following PCR conditions: denaturation at 95°C for 3 min, followed by 30 cycles of 95°C for 1 min, 62°C for 1 min, and 72°C for 3 min. PCR products were gel purified with Gel purification kit (Invitrogen). Purified PCR products were sequenced with the same primers that were used for PCR amplification at the DNA Sequencing Core Facility at MSKCC.

**Intrachromosomal Junction Analysis**

To analyze junctions from imprecise NHEJ at a single I-SceI site, genomic DNA was isolated from E13U18 cells 72 h after transfection of the I-SceI expression vector (with or without the Xrcc4 expression vector) and then digested \textit{in vitro} with I-SceI (NEB). (Transfection conditions were the same as that for translocation experiments.) An approximately 1.5 kb fragment was then amplified using primers surrounding the I-SceI site on der(17) by Extaq enzyme (Takara inc.) with the following primers: 17–754 TGACTCTGGCTTGTGGTTTG and 14–726 GCTGGATATGTGTCCCGTTT. PCR was done at an annealing temperature of 60°C and an extension time of 1.5 min, and products were cloned with TOPO TA Cloning kit (Invitrogen) and sequenced. For statistical analysis, a two-tailed Mann-Whitney test was used.

**Western Blotting**

Whole cell extracts were prepared with Nonidet-P40 buffer and were run on a 7.5% (w/v) Tris-HCl SDS Page and blotted using the Xrcc4 antibody clone C-20 (Santa Cruz Biotechnology), which recognizes the C terminus of the human and mouse proteins. Mouse monoclonal antibody to \textit{α}-tubulin (Sigma) was used as a loading control.

**Fluorescence \textit{in situ} hybridization**

FISH was performed by the Molecular Cytogenetics Core Facility at MSKCC. For visualization of translocations, the FITC (green) probe (Cambio #1189-14MF-01) was
hybridized to mouse chr. 14, and the Cy3 (red) probe (Cambio #1200-17MCy3-01) was
hybridized to mouse chr. 17. Images were captured using a digital FISH workstation
(Metasystems).

**Statistical Analysis**

In order to determine which statistical test to use (i.e., Kolmogorov-Smirnoff,
D’Agostino&Pearson omnibus and Shapiro-Wilk), normality tests were executed using
Prism Software (GraphPad, Inc.). Because translocation frequencies showed a normal
distribution, a two–tailed unpaired t test was applied for this analysis. Because the data do
not assume normal distribution, a one-tailed Mann-Whitney test was used for
microhomology distribution and a two-tailed Mann-Whitney test was used for other
analyses. Assuming an unbiased base composition, the probability that a junction will have
X nucleotides of microhomology expected by chance is calculated by the equation \( P(X) = \frac{(X+1)(1/4)^X(3/4)^2}{X} \) as previously described\(^{51}\).

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.
Chromosomal translocations are suppressed by XRCC4/ligase IV. (a) Translocation reporter in Xrcc4−/− pCr15 cells. DSB formation on chromosomes 17 and 14 at the I-SceI sites, followed by interchromosomal NHEJ, results in a chromosomal translocation with a neo+ gene on der(17). FISH analysis indicates that parental pCr15 cells have normal chromosomes 17 (red) and 14 (green), whereas neo+ clones have derivative chromosomes. Vertical red bars are exons 1–5 from the targeted Pim1 locus on chr.17, and the vertical green bar is exon 20 from the Rb locus. Probes are located outside the targeting arms. HII, HincII; HIII, HindIII. (b) Translocation frequency is significantly increased in Xrcc4−/− cells, but is suppressed by Xrcc4 expression. (c) Confirmation of the genotype of the endogenous Xrcc4 alleles in wild-type (WT), Xrcc4−/− and Xrcc4-complemented cells. P, parental pCr15 cells of the indicated genotypes; t, neo+ translocation clones. (d) Western blot analysis demonstrating that transient expression of Xrcc4 restores wild-type Xrcc4 protein levels to Xrcc4−/− cells.
Figure 2.
Translocation breakpoint junctions have similar characteristics in wild-type, Xrcc4−/− mutant and complemented cells, and Ku70−/− cells. (a) Representative der(17) translocation junction sequences obtained from Xrcc4−/− and Xrcc4-complemented cells. DNA ends generated by I-SceI on chrs.17 and 14 are indicated in red and blue, respectively. A summary of the various end modifications is presented to the left of each junction in bp: Δ, total deletion; µ, microhomology; +, insertion. Sequences are annotated as follows: del, deletion length from the DNA end; underline, microhomology; +, length of long insertion. The middle green sequences are short insertions from chr. 14; considering a template model for their insertion, the sequences in red shading (TAA) would be microhomology between the DNA ends that could anneal to act as a primer and the blue shading would represent microhomology for annealing after DNA synthesis between the 2 DNA ends (see text). (b) Deletion lengths for der(17) breakpoint junctions. Each value represents the combined deletion from both ends of an individual junction. The median deletion length for each genotype is indicated by a bar on the graph and the value is give below the graph. Deletion lengths do not differ significantly from each other (two-tailed Mann-Whitney test). +X4, transient complementation with Xrcc4. (c) Microhomology and insertion frequencies are similar for the four genotypes. (d) Distribution of microhomology lengths for der(17) breakpoint junctions. Only junctions with simple deletions (i.e., without an insertion) are...
(e) Lack of correlation between deletion length and microhomology use. Only junctions for *Xrec4*−/− cells are plotted.
Insertions at breakpoint junctions display complex characteristics. (a) Distribution of deletion lengths for der(17) breakpoint junctions without and with insertions. Data are combined from all four genotypes; the deletion distributions for individual genotypes are shown in Supplementary Fig. 3d. (b) Der(17) insertions are comprised of one or more distinct segments of DNA. Most insertions at the der(17) breakpoint junctions are derived from one DNA segment, although others are comprised of up to 4 segments. Most inserted segments are derived from chr. 17 (red boxes) or chr. 14 (green boxes), although some of the inserted segments are derived from other sources or are too short to be mapped precisely (white boxes). Included in this analysis are all inserted sequences >4 bp. (c) Derivation of der(17) inserts. Each arrow represents the derivation of a segment inserted at a der(17)
translocation breakpoint junction and the relative orientation of the insert relative to the centromere. Inserted segments are derived from sequences adjacent to a DSB or as far away as 4 Mb; those from nearby a DSB are generally short (≤20 bp), while more distantly-derived segments tend to be larger (up to 396 bp). Note that the inserted segments can be derived from sequences used to form either der(17) or der(14). The der(17) sequences from −1.5 kb to +1.2 kb comprise the neo intron (Fig. 1a).
Table 1

Translocation frequencies are from 4 experiments. One standard deviation from the mean is indicated. A two-tailed unpaired t test was applied, with \( p \) values derived from a comparison with \( Xrcc4^{-/-} \) cells (a) and wild-type cells (b).

| Cells     | Expression vector | Translocation frequency \((\times 10^{-4})\) | \( p \) value | 
|-----------|-------------------|---------------------------------------------|----------------|
| Wild-type | –                 | <0.001                                      |                |
|           | I-SceI            | 0.57 ± 0.11                                 | <0.001        |
|           | I-SceI+Xrcc4      | 0.62 ± 0.06                                 | <0.001 0.455  |
|           | I-SceI+Ku70       | 0.61 ± 0.10                                 | <0.001 0.610  |
| \( Xrcc4^{-/-} \) | –                 | <0.001                                      |                |
|           | I-SceI            | 2.70 ± 0.20                                 | – <0.001      |
|           | I-SceI+Xrcc4      | 0.53 ± 0.12                                 | <0.001 0.641  |
|           | I-SceI+Ku70       | 2.73 ± 0.23                                 | 0.850 <0.001  |
| \( Ku70^{-/-} \) | –                 | <0.001                                      |                |
|           | I-SceI            | 2.33 ± 0.21                                 | 0.043 <0.001  |
|           | I-SceI+Xrcc4      | 2.14 ± 0.19                                 | 0.007 <0.001  |
|           | I-SceI+Ku70       | 0.71 ± 0.02                                 | <0.001 0.046  |