Effect of the Murine Leukemia Virus Extended Packaging Signal on the Rates and Locations of Retroviral Recombination

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Reverse transcriptase (RT) switches templates frequently during DNA synthesis; the acceptor template can be the same RNA (intramolecular) or the copackaged RNA (intermolecular). Previous results indicated that intramolecular template switching occurred far more frequently than intermolecular template switching. We hypothesized that intermolecular template-switching events (recombination) occurred at a lower efficiency because the copackaged RNA was not accessible to the RT. To test our hypothesis, the murine leukemia virus (MLV) extended packaging signal (Ψ+) containing a dimer linkage structure (DLS) was relocated from the 5′ untranslated region (UTR) to between selectable markers, allowing the two viral RNAs to interact closely in this region. It was found that the overall maximum recombination rates of vectors with Ψ+ in the 5′ UTR or Ψ− between selectable markers were not drastically different. However, vectors with Ψ+ located between selectable markers reached a plateau of recombination rate at a shorter distance. This suggested a limited enhancement of recombination by Ψ+. The locations of the recombination events were also examined by using restriction enzyme markers. Recombination occurred in all four regions between the selectable markers; the region containing 5′ Ψ+ including DLS did not undergo more recombination than expected from the size of the region. These experiments indicated that although the accessibility of the copackaged RNA was important in recombination, other factors existed to limit the number of viruses that were capable of undergoing intermolecular template switching. In addition, recombinants with multiple template switches were observed at a frequency much higher than expected, indicating the presence of high negative interference in the MLV-based system. This extends our observation with the spleen necrosis virus system and suggests that high negative interference may be a common phenomenon in retroviral recombination.

Retroviruses package two copies of viral RNA into each virion (9, 28). During reverse transcription, both copackaged RNAs can be used as templates to produce a recombinant with a mixture of genetic information from each of the parental RNAs (8, 18). Retroviruses recombine at high rates (6, 15, 23, 24, 26, 29, 30, 46–48). Using murine leukemia virus (MLV)-based vectors with markers separated by 1.0, 1.9, and 7.1 kb, recombination rates of 4.7, 7.4, and 8.2%, respectively, were observed in one round of viral replication (2).

During reverse transcription of the viral genome, the virus-encoded enzyme reverse transcriptase (RT) has to perform two template-switching events (named minus- and plus-strand DNA transfer) to complete the synthesis of viral DNA (12). It has been hypothesized that RT is evolutionarily selected to have a low affinity to the template and low processivity in order to complete the two obligatory template-switching events (44). A consequence of this low processivity is that RT may also perform other nonobligatory template-switching events during reverse transcription. Because two copies of RNA are present in the virion, after dissociation from the template used for DNA synthesis, RT may reassociate with the same RNA or switch to the copackaged RNA, resulting in intramolecular or intermolecular template-switching events, respectively. Between these two events, only intermolecular template switching generates recombination.

The rates of intramolecular and intermolecular template switching were measured in a spleen necrosis virus (SNV)-based system (17). It was found that intramolecular template switching occurred far more frequently than intermolecular template switching. Furthermore, when recombination was selected in one region of the viral genome, intermolecular template switching was more likely to be observed in another region of the genome. In contrast, when recombination was not selected, intermolecular template switching was not observed in a different region of the viral genome; however, frequent intramolecular template switching was still observed. This observation led to the hypothesis that two viral populations exist. In the first population (recombinating population), RT could frequently switch templates to the same RNA or the copackaged RNA. In the second population (nonrecombinating population), RT could only switch to the same RNA frequently. Because frequent template switching occurred in both populations, the difference seemed to be the ability of the RT to interact with the other template. We hypothesized that if the barrier to intermolecular template switching is the ability to access the other RNA, then it should be possible to increase recombination by bringing the two copackaged RNAs closer together. In our experimental system, recombination is measured by the presence of two selectable markers, one from each parent, in the progeny provirus. Therefore, to test our hypothesis, the region between the two selectable markers from the two copackaged RNAs should be in close proximity to observe any effect on recombination.

Electron microscopy studies demonstrated that the two copackaged retroviral RNAs were dimerized at the 5′ end of the
RNA (27, 28, 32, 39). Genetic and biochemical studies defined a region important for dimerization through noncovalent interaction between the two viral RNAs (13, 40, 42). The region important for dimerization of MLV RNAs, the dimer linkage structure (DLS), is part of the packaging signal (Ψ) located within the 5′ untranslated region (UTR) (11, 13, 39, 40, 42). In the current report, we examined the effect of the naturally occurring dimerization between the two copackaged RNAs on recombination. The extended MLV packaging signal (Ψ′), including the DLS, was relocated from the 5′ UTR to the middle of the viral genome between two selectable markers with XhoI followed. Recombination rates between vectors with Ψ′ located in the 5′ UTR were directly compared to those from vectors with Ψ′ located between the selectable markers.

In addition, using a forced-recombination experiment, it was previously observed that the DLS was a putative recombination hot spot (31, 34, 35). To determine the general effect of the DLS on the location of template-switching events, a set of vectors that contained five sets of restriction enzyme markers was used to dissect the locations of template-switching events after one round of retroviral replication.

MATERIALS AND METHODS

Plasmid construction. Vectors pJS30 and JA32-1kb have been described previously (2); to avoid confusion, JA32-1kb is abbreviated as JA32 in this report. Vectors pJA33-1.3kb, pJa9, pJA19Neo, pJA19Hy, pJA23, pJA19Neo, and pJA19Hy were constructed by standard molecular cloning techniques (33). All of the MLV-based vectors were constructed with various derivatives of pLAEN (38) as backbones. To generate pJA33-1.3kb, pJa30 (2) was partially digested with NdeI, treated with calf intestinal phosphatase (CIP), and ligated to a linker (5′-TAACGCGTCT-3′) to create a unique NdeI site and an 8-bp insertion in the hygromycin phosphotransferase B gene (hygro) (14). To generate pJa9, pWH390-Cla was digested with EcoRI and ligated to annealed linkers PL1 (5′-AAATAAAGCTTATGGCTGCGGTT-3′) and PL2 (5′-AAATTCTCGAGGCGACATATG-3′) to generate pWBW1-pWBW1 was digested with ClaI and treated with CIP and Escherichia coli DNA polymerase I large fragment (Klenow) to fill in the recessive ends. It was treated with EcoRI and ligated to a linker (5′-AAATAAAGCTTATGGCTGCGGTT-3′) and 3′GATC to fill in the recessive ends, and self-ligated to remove the 0.6-kb sequence. Ten-fold serial dilutions of each viral stock were generated, and the viral infections were performed in the presence of 50 μg of Polybrene per ml. Viral titers were determined by the number of hygromycin-, G418-, or hygromycin plus-G418-resistant cells. Each titer shown was from one experiment.

Southern hybridization analysis. Genomic DNA purification, digestion, and Southern hybridization analysis were performed by standard molecular techniques (33). DNA transfers were performed with a vacuum blotter (Pharmacia). All blots were hybridized with probe generated by the random-priming method (5′-ACTGCAAGA CGCGTCCTATCCG-3′). DNA hybridizations were performed immediately following viral harvest. Viruses were collected from confluent monolayers and centrifuged at 400 g for 10 min to remove cellular debris.

RESULTS

Retroviral vectors used to study the effect of the MLV extended packaging signal on retroviral recombination. To examine the effect of the MLV Ψ′ on retroviral recombination, we constructed a set of vectors in which Ψ′ was located in the 5′ UTR and another set of vectors in which Ψ′ was located in the middle of the viral genome between two selectable markers. Both sets of vectors were derived from MLV and contained cis-acting elements required for viral replication and gene expression. The first set of vectors (pJS30, pJA33-1.3kb, and pJA32) contained Ψ′ in the 5′ UTR (Fig. 1). These vectors also contained hygro and neo both of which were expressed by transcripts initiated from the long terminal repeat (LTR). The translation of neo was directed by an IRES from encephalomyocarditis virus (1, 19, 20). These three vectors were highly homologous (>99%) to one another. Vector pJS30 contained functional hygro and neo (14, 22). Vector pJA33-1.3kb contained a functional neo and a nonfunctional hygro with an 8-bp frameshift insertion that destroyed an NdeI site and generated an MluI site. Vector pJA32 (2) contained a functional hygro and a nonfunctional neo with a 4-bp frameshift insertion that...
destroyed an \textit{EheI} site and generated a \textit{BssHII} site. The distance between the two inactivating mutations in \textit{hygro} and \textit{neo} was 1.3 kb. Frameshift mutations of more than 1 bp were used to inactivate genes because they exhibit a low reversion rate (2, 18).

The second set of vectors (pJA9, pJA10\^pNeo, and pJA11Hy\^p) also contained \textit{hygro} and \textit{neo} expressed from \textit{U3}-regulated transcripts (Fig. 1). However, MLV \textit{Ψ} jumped to the 5' UTR to between \textit{hygro} and \textit{neo}; in these vectors, the translation of \textit{neo} was directed by the IRES in the MLV packaging signal (4, 45). Vector pJA9 contained functional \textit{hygro} and \textit{neo} and a nonfunctional \textit{hygro} with a 2-bp frameshift deletion that destroyed a \textit{SacII} site and introduced an \textit{NdeI} site. Vector pJA10\^pNeo contained a functional \textit{hygro} and a nonfunctional \textit{neo} with a 2-bp frameshift insertion that destroyed an \textit{EheI} site and introduced a \textit{BssHII} site. The distance between the two inactivating mutations in \textit{hygro} and \textit{neo} was 1.3 kb.

**Experimental protocol used to measure the rates of recombination in one replication cycle.** The control vector pJA9 was structurally similar to the vectors (pJA10\^pNeo and pJA11Hy\^p) used to measure the rate of recombination at a 1.3-kb marker distance with \textit{Ψ} located between selectable markers (Fig. 1); however, pJA9 contained a functional \textit{hygro} and \textit{neo}. A protocol similar to that shown in (heterozygotic virions). The first two types of virions generate proviruses containing one functional drug resistance gene that confers resistance to a single drug. Recombination can occur in the heterozygotic virions to generate proviruses that contain two functional drug resistance genes that confer resistance to both drugs (18, 48). Viruses were harvested from the selected PG13 cell clones and used to infect D17 target cells. Infected D17 cells were subjected to single (hygromycin or G418) or double (hygromycin plus G418) drug selection. Virus titers were then determined by counting the numbers of drug-resistant colonies. Recombination rates were calculated from the double- and the single-drug-resistant colony titers (18). In addition, double-drug-resistant D17 cell clones were isolated, and Southern hybridization analysis was performed to confirm the recombinant genotype of the proviruses.

In this system, the recombination rates were measured in one replication cycle, which was defined by the steps that occurred from the proviral stage in PG13 helper cells to the proviral stage in D17 target cells. Viruses produced from the PG13 cells contained GaLV Env; because PG13 cells do not express the receptor for GaLV Env, reinfection during propagation of the viruses cannot occur. Additionally, viruses cannot be propagated in D17 cells, because they lack helper function. Therefore, this system allowed only a single round of replication to occur.

The control vectors generate similar single- and double-drug-resistant colony titers. In this system, recombination rates were calculated from single- and double-drug-resistant titers. Therefore, it was important to determine whether the three different drug treatments generated similar viral titers in infected cells.

The control vector pJA9 was structurally similar to the vectors (pJA10\^pNeo and pJA11Hy\^p) used to measure the rate of recombination at a 1.3-kb marker distance with \textit{Ψ} located between selectable markers (Fig. 1); however, pJA9 contained a functional \textit{hygro} and \textit{neo}. A protocol similar to that shown in
the expected bands. A unique (A2, B1, and E1) is shown in Fig. 3B; each cell clone contained representative Southern blot of three different PG13 cell clones 1.3kb and the JA32 proviruses, respectively (Fig. 3A). A representative Southern blot of three different PG13 cell clones containing a copy of JA33-1.3kb and JA32 was used to infect D17 target cells. Viral titers are shown in Table 2. The hygromycin-resistant colony titers ranged from 0.15 × 10^5 to 33.0 × 10^5 CFU/ml, the G418-resistant colony titers ranged from 0.18 × 10^5 to 31 × 10^5 CFU/ml, and the hygromycin-plus-G418-resistant colony titers ranged from 0.022 × 10^5 to 8.9 × 10^5 CFU/ml. A double-drug-resistant D17 cell clone could be generated from dual infection of the two parental viruses or a recombinant provirus that had functional hygro and neo. If the double-drug-resistant D17 cell clones contained one of each parental virus, then Southern analysis would reveal a band pattern identical to that of the PG13 cell clones with the same probe and EcoI site. This probe hybridized to the EcoI DNA fragment of JA33-1.3kb. Other abbreviations and symbols are defined in the legend to Fig. 1.

Recombination rate of MLV vectors with ψ+ located in the 5’ UTR and markers separated by 1.3 kb. Viruses harvested from eight independent PG13 cell clones that contained a copy of JA33-1.3kb and JA32 were used to infect D17 target cells. Viral titers are shown in Table 2. The hygromycin-resistant colony titers ranged from 0.15 × 10^5 to 33.0 × 10^5 CFU/ml, the G418-resistant colony titers ranged from 0.18 × 10^5 to 31 × 10^5 CFU/ml, and the hygromycin-plus-G418-resistant colony titers ranged from 0.022 × 10^5 to 8.9 × 10^5 CFU/ml. A double-drug-resistant D17 cell clone could be generated from dual infection of the two parental viruses or a recombinant provirus that had functional hygro and neo. If the double-drug-resistant D17 cell clones contained one of each parental virus, then Southern analysis would reveal a band pattern identical to that of the PG13 cell clones with the same probe and EcoI site. This probe hybridized to the EcoI DNA fragment of JA33-1.3kb.

Characterization of proviral structures in PG13 cell clones infected with JA33-1.3kb and JA32. PG13 cell clones containing JA33-1.3kb and JA32 were generated to measure the recombination rate between markers separated by 1.3 kb with ψ+ located in the 5’ UTR. The proviral structures in these cell clones were characterized by Southern hybridization analysis. Partial restriction enzyme maps of JS30, JA33-1.3kb, and JA32 are shown in Fig. 3A. The JS30 provirus had a unique NdeI site located in hygro and four EheI sites (one in each LTR, one in ψ+, and one in neo). In JA33-1.3kb, the NdeI site was destroyed to inactivate hygro, whereas in JA32, an EheI site was destroyed to inactivate neo. All three proviruses contained four EcoRV sites, two in each LTR. When genomic DNA from cell clones containing a copy of JA33-1.3kb and a copy of JA32 was digested with EheI, NdeI, and EcoRV and hybridized with probes generated from a 1.3-kb MluI-EheI fragment of JA33-1.3kb, a 1.8- and a 2.4-kb band were expected from the JA33-1.3kb and the JA32 proviruses, respectively (Fig. 3A). A representative Southern blot of three different PG13 cell clones (A2, B1, and E1) is shown in Fig. 3B; each cell clone contained the expected bands. A unique HindIII site was located in JA33-1.3kb and JA32 proviruses at the 5’ end of IRES (Fig. 3A). Therefore, each provirus was expected to generate two bands when digested with HindIII and hybridized with the aforementioned probe. Since retroviruses integrate randomly into the host genome (7), the band sizes should vary according to the site of integration. Therefore, cell clones that contain one copy of JA33-1.3kb and JA32 should produce four bands of different sizes upon HindIII digestion and Southern analysis. A representative Southern analysis with HindIII digestion of genomic DNA from the helper cell clones is shown in Fig. 3B; all cell clones (Fig. 3B and data not shown) exhibited a different band pattern, indicating that they were derived from independent infection events (Fig. 3B, lanes H).

Recombination rate of MLV vectors with ψ+ located in the 5’ UTR and markers separated by 1.3 kb. Viruses harvested from eight independent PG13 cell clones that contained a copy of JA33-1.3kb and JA32 were used to infect D17 target cells. Viral titers are shown in Table 2. The hygromycin-resistant colony titers ranged from 0.15 × 10^5 to 33.0 × 10^5 CFU/ml, the G418-resistant colony titers ranged from 0.18 × 10^5 to 31 × 10^5 CFU/ml, and the hygromycin-plus-G418-resistant colony titers ranged from 0.022 × 10^5 to 8.9 × 10^5 CFU/ml. A double-drug-resistant D17 cell clone could be generated from dual infection of the two parental viruses or a recombinant provirus that had functional hygro and neo. If the double-drug-resistant D17 cell clones contained one of each parental virus, then Southern analysis would reveal a band pattern identical to that of the PG13 cell clones with the same probe and EcoI site. This probe hybridized to the EcoI DNA fragment of JA33-1.3kb.

Characterization of proviral structures in PG13 cell clones infected with JA33-1.3kb and JA32. PG13 cell clones containing JA33-1.3kb and JA32 were generated to measure the recombination rate between markers separated by 1.3 kb with ψ+ located in the 5’ UTR. The proviral structures in these cell clones were characterized by Southern hybridization analysis. Partial restriction enzyme maps of JS30, JA33-1.3kb, and JA32 are shown in Fig. 3A. The JS30 provirus had a unique NdeI site located in hygro and four EheI sites (one in each LTR, one in ψ+, and one in neo). In JA33-1.3kb, the NdeI site was destroyed to inactivate hygro, whereas in JA32, an EheI site was destroyed to inactivate neo. All three proviruses contained four EcoRV sites, two in each LTR. When genomic DNA from cell clones containing a copy of JA33-1.3kb and a copy of JA32 was digested with EheI, NdeI, and EcoRV and hybridized with probes generated from a 1.3-kb MluI-EheI fragment of JA33-1.3kb, a 1.8- and a 2.4-kb band were expected from the JA33-1.3kb and the JA32 proviruses, respectively (Fig. 3A). A representative Southern blot of three different PG13 cell clones (A2, B1, and E1) is shown in Fig. 3B; each cell clone contained the expected bands. A unique HindIII site was located in JA33-1.3kb and JA32 proviruses at the 5’ end of IRES (Fig. 3A). Therefore, each provirus was expected to generate two bands when digested with HindIII and hybridized with the aforementioned probe. Since retroviruses integrate randomly into the host genome (7), the band sizes should vary according to the site of integration. Therefore, cell clones that contain one copy of JA33-1.3kb and JA32 should produce four bands of different sizes upon HindIII digestion and Southern analysis. A representative Southern analysis with HindIII digestion of genomic DNA from the helper cell clones is shown in Fig. 3B; all cell clones (Fig. 3B and data not shown) exhibited a different band pattern, indicating that they were derived from independent infection events (Fig. 3B, lanes H).
were expected from this digestion when hybridized with a probe containing the 3' portion of *hygro* and 5' portion of *neo*. A 1.4-kb band and a 1.9-kb band were expected to be generated from the JA10*Ψ*Neo provirus, whereas 0.8-, 1.1-, and 1.6-kb bands were expected to be generated from the JA11HyΨ provirus (Fig. 4A). A representative Southern blot of three different PG13 cell clones (C1, C2, and A4) is shown in Fig. 4B. Each cell clone contained the expected fragments consistent with the predicted structures of JA10*Ψ*Neo and JA11HyΨ. In addition, all DNAs were digested with *Hind*III to confirm that all of the PG13 cell clones used were generated through independent infection events (Fig. 4B and data not shown).

Recombination rate of MLV vectors with *Ψ<sup>+</sup>* located between two selectable markers that were 1.3 kb apart. Six PG13 cell clones coinfected with JA10*Ψ*Neo and JA11HyΨ were identified; viruses were harvested from these cell clones and were used to infect D17 target cells. The infected target cells were placed on hygromycin, G418, or hygromycin-plus-G418.

**TABLE 2. Virus titers generated from PG13 cell clones coinfected with JA33-1.3kb and JA32**

<table>
<thead>
<tr>
<th>Clone</th>
<th>Hygromycin (10&lt;sup&gt;5&lt;/sup&gt;)</th>
<th>G418 (10&lt;sup&gt;5&lt;/sup&gt;)</th>
<th>Hygromycin + G418 (10&lt;sup&gt;5&lt;/sup&gt;)</th>
<th>Recombination rate* (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>D1</td>
<td>33.0</td>
<td>22.0</td>
<td>8.5</td>
<td>7.7</td>
</tr>
<tr>
<td>L1</td>
<td>26.0</td>
<td>16.0</td>
<td>3.8</td>
<td>4.8</td>
</tr>
<tr>
<td>J2</td>
<td>31.0</td>
<td>31.0</td>
<td>8.9</td>
<td>5.7</td>
</tr>
<tr>
<td>C1</td>
<td>0.15</td>
<td>0.18</td>
<td>0.022</td>
<td>2.9</td>
</tr>
<tr>
<td>B1</td>
<td>5.6</td>
<td>5.3</td>
<td>0.93</td>
<td>3.5</td>
</tr>
<tr>
<td>A2</td>
<td>9.2</td>
<td>8.6</td>
<td>1.8</td>
<td>4.2</td>
</tr>
<tr>
<td>G1</td>
<td>9.2</td>
<td>9.4</td>
<td>2.5</td>
<td>5.4</td>
</tr>
<tr>
<td>A1</td>
<td>2.5</td>
<td>2.0</td>
<td>5.9</td>
<td>5.9</td>
</tr>
<tr>
<td>Avg ± SE</td>
<td></td>
<td></td>
<td></td>
<td>5.0 ± 0.5</td>
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</table>

* Recombination rate = (double-drug-resistant titer/lesser of the single-drug-resistant titers) × 2.

Restriction enzyme digests. In contrast, if D17 cell clones contained a recombinant provirus, Southern analyses would generate a 1.3-kb band when digested with the same three enzymes (*Eco*RI, *Nde*I, and *Ehe*I) and hybridized with the same probe (Fig. 3A). To verify that the double-drug-resistant colony titers were generated by recombinant proviruses, 16 hygromycin-plus-G418-resistant D17 cell clones were isolated, and the proviral structures were analyzed. A representative Southern analysis of three D17 cell clones (A2-1a, B1-b, and E1-b) is shown in Fig. 3B; each of the cell clones was derived from one of the three PG13 cell clones shown in the same figure. Of the 16 cell clones, 13 had a 1.3-kb band indicating that they contained a recombinant provirus (Fig. 3B and data not shown), and 3 had a genotype consistent with a double infection event. This indicated that most of the double-drug-resistant cell clones contained recombinant proviruses and the hygromycin-plus-G418 titer reflected the titer of the recombinant viruses containing functional *hygro* and *neo*.

This assay measured the formation of half of the recombinants—those containing functional *hygro* and *neo*. The other half of the recombinants, those containing inactivated *hygro* and *neo*, could not be measured in the viral titer assay. Therefore, the recombination rate for viruses generated from each cell clone was calculated by dividing the double-drug-resistant colony titers by the lesser of the two single-drug-resistant colony titers and then multiplying by 2 (Table 2). Therefore, the recombination rates of vectors JA33-1.3kb and JA32 from six PG13 cell clones ranged from 2.9 to 7.7%, with an average of 5.0% ± 0.5% (standard error [SE]).

Characterization of proviral DNA structures in PG13 cells infected with JA10ΨNeo and JA11HyΨ. PG13 cell clones containing JA10ΨNeo and JA11HyΨ were generated to examine the effect of *Ψ<sup>+</sup>* located between the selectable markers on the rate of recombination. Southern analyses were performed to examine the proviral structures in coinfected PG13 cell clones. Partial restriction enzyme maps of JA9, JA10ΨNeo, and JA11HyΨ are shown in Fig. 4A. JA9, JA10ΨNeo, and JA11HyΨ proviruses each contained a unique ClaI site between the 5' LTR and *hygro*. JA9 proviruses contained four *Ehe*I sites: one in each LTR, one in *Ψ<sup>+</sup>* and one in *neo*. JA10ΨNeo proviruses contained all four *Ehe*I sites, whereas JA11HyΨ contained only three *Ehe*I sites, because one of the sites was mutated to inactivate *neo*. A unique *Sac*II site was located in *hygro* of JA9 and JA11HyΨ; this site was mutated in JA10ΨNeo to inactivate *hygro*. Genomic DNAs from PG13 cell clones coinfected with JA10ΨNeo and JA11HyΨ were digested with three enzymes: *Ehe*I, *Sac*II, and *Cla*II. Five bands were shown in the same figure. Of the 16 cell clones, 13 had a 1.3-kb band indicating that they contained a recombinant provirus (Fig. 3B and data not shown). A unique Southern blot of three different PG13 cell clones (C1, C2, and A4) is shown in Fig. 4B. Each cell clone contained the expected fragments consistent with the predicted structures of JA10ΨNeo and JA11HyΨ. In addition, all DNAs were digested with *Hind*III to confirm that all of the PG13 cell clones used were generated through independent infection events (Fig. 4B and data not shown).
To the marker distance was increased because the observed rate had been altered, we should observe an increase in the recombination rate when the marker distance was increased. However, if the recombining population had not changed, we should not observe an increase in recombination rates when the marker distance was increased because the observed rate was already close to the maximum rate detected (8.2%) (2). To distinguish between these two possibilities, we measured recombination rates between markers 1.9 kb apart.

Vectors used to further study the effect of Ψ+ on the rate and location of the recombination events. To further characterize the effect of Ψ+ on recombination, a third set of vectors was constructed to measure the recombination rate when markers were separated by 1.9 kb. The vectors pJA23, pJA19ΨNeo, and pJA20HyΨ were very similar in sequence to the second set of vectors, pJA9, pJA10ΨNeo, and pJA11HyΨ, respectively. The structures of these viral vectors are shown in Fig. 1.

It has been proposed that the DLS region within the Ψ is a recombinational hot spot (34, 35). To examine the general effect of DLS on the locations of the intermolecular template switching events, three sets of restriction enzyme markers were placed at the 5′ end, the middle, and the 3′ end of Ψ+ of pJA19ΨNeo and pJA20HyΨ. It was shown that relocation of the packaging signal to the 3′ UTR could result in packaging of spliced RNA (21). Although it was demonstrated that the recombination rates were not affected by the packaging of spliced RNA (21), this could result in a potential bias in the analysis of the locations of the recombination events. To avoid this potential bias, the splice donor sites were destroyed by PCR mutagenesis in all three vectors used in this experiment.

Similar to pJS30 and pJA9, pJA23 is a control vector and contain functional hygro and neo. The vector pJA19ΨNeo contained a functional neo and a nonfunctional hygro with a 4-bp frameshift insertion that destroyed an NcoI site and generated an NsiI site. The vector pJA20HyΨ contained a functional hygro and a nonfunctional neo with an 8-bp frameshift insertion that destroyed an FspI site and generated an AatII site. The distance between the two inactivating mutations in hygro and neo was 1.9 kb. The presence of the NcoI-NsiI and AatII-FspI markers determined the abilities of the provirus to confer drug resistance; therefore, these two sets of markers are referred to as “selectable markers.” In addition to the selectable markers, these two vectors differed at three other restriction enzyme markers. These markers are located between hygro and neo (BamHI-ClaI), in the middle of Ψ+ (MluI-AfII), and between Ψ+ and neo (PvuII-XhoI). The MluI-AfII markers in Ψ+ were located past the minimum packaging signal and at the 5′ end of the gene containing the mutated AUG. Therefore, it was expected that this mutation would not interfere with the efficiency of RNA packaging. The other two mutations were located between Ψ+ and drug resistance genes and were not expected to interfere with the abilities of these genes to confer drug resistance. The nature of these three sets of markers was 4- to 8-bp insertions.

To determine whether the control vector, pJA23, generated similar single- and double-drug-resistant colony titers, five independent PG13 cell clones containing an intact copy of JA23 were identified by Southern hybridization analysis (data not shown). Viral titers generated from these five cell clones are shown in Table 3. Parallel to JA9 and JS30, the viral titers generated within each cell clone for JA23 were similar; thus, mutation of the SD did not adversely affect expression of either hygro or neo. The viral titers ranged from 0.4 × 10⁵ to 2.2 × 10⁶ CFU/ml for hygromycin, 0.092 × 10⁵ to 0.75 × 10⁵ CFU/ml for G418, and 0.24 × 10⁵ to 1.7 × 10⁵ CFU/ml for hygromycin-pluss-G418 drug selection. The results indicated that the titers produced by each drug selection directly reflected the amount of cells infected with the control vectors.

Proviral DNA analysis of PG13 cell clones infected with JA19ΨNeo and JA20HyΨ. PG13 cell clones containing JA19ΨNeo and JA20HyΨ were generated to further examine the effect of Ψ+ on recombination. Southern hybridization

<table>
<thead>
<tr>
<th>Clone</th>
<th>Hygromycin (10⁵)</th>
<th>G418 (10⁵)</th>
<th>Hygromycin + G418 (10⁵)</th>
<th>Recombination rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A3</td>
<td>1.1</td>
<td>0.50</td>
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<td>7.4</td>
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</tr>
<tr>
<td>Y4</td>
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<td>17.5</td>
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<tr>
<td>Avg ± SE</td>
<td></td>
<td></td>
<td></td>
<td>12.0 ± 1.5</td>
</tr>
</tbody>
</table>

*Recombination rate = (double-drug-resistant titer/lesser of the single-drug-resistant titers) × 2.
analyses were performed to determine the proviral structures. Partial restriction enzyme maps of JA23, JA19ΨNeo, and JA20HyΨ are shown in Fig. 5A. JA23, which has functional hygro and neo, has an NcoI site located in each drug resistance gene and a unique FspI site located in neo. One of the NcoI sites was destroyed to inactivate hygro in JA19ΨNeo, whereas the FspI site was destroyed to inactivate neo in JA20HyΨ. In addition, all three vectors contained an XbaI site located in each LTR.

Genomic DNAs were digested with XbaI, FspI, and NcoI and hybridized with a probe generated from a 1.3-kb NcoI-FspI fragment of JS30 (Fig. 5A). If the cell clones contained a copy of JA19ΨNeo and a copy of JA20HyΨ, 2.7- and 2.2-kb bands were expected (Fig. 5A). A representative Southern blot of three PG13 cell clones (clones 1, 7, and 15) is shown in Fig. 5B; each cell clone contained the expected bands. In addition, all DNAs were digested with HindIII and analyzed by Southern analyses as previously described to confirm that cell clones were generated through independent infection events (Fig. 5B and data not shown).

Recombination rate of MLV vectors with Ψ+ located between selectable markers that were 1.9 kb apart. Viruses were harvested from five independent PG13 cell clones that contained a copy each of JA19ΨNeo and JA20HyΨ and used to infect D17 target cells. Viral titers are shown in Table 4. The hygromycin-resistant colony titers ranged from 0.50 × 10^5 to 7.8 × 10^5 CFU/ml, the G418-resistant colony titers ranged from 0.20 × 10^5 to 20 × 10^5 CFU/ml, and the hygromycin plus-G418-resistant colony titers ranged from 1.2 × 10^6 to 5.6 × 10^6 CFU/ml.

The proviral structures in double-drug-resistant D17 cells were examined. Figure 5B shows a representative Southern blot of three D17 cell clones (1-2c, 7-8a, and 15-14b). Recombinant proviruses should generate a single 1.9-kb band when digested with XbaI, NcoI, and FspI and hybridized with the probe generated from the 1.3-kb NcoI-FspI fragment of JS30. Alternatively, the genotypes of the proviruses can be examined by amplifying a portion of the proviral sequences, using PCR and subjecting the DNA to restriction enzyme mapping (described in detail below). A total of 53 D17 cell clones were analyzed by Southern blot analysis and/or restriction enzyme mapping; 42 contained a recombinant provirus, whereas 11 were the result of a double-infection event (Fig. 5B and 6B, and data not shown).

The genotype analyses data indicated that most of the double-drug-resistant colonies contained recombinant proviruses. This indicated that the double-drug-resistant colony titers reflected the number of recombinants containing functional hygro and neo and thus could be used to calculate recombination rates. The recombination rates of JA19ΨNeo and JA20HyΨ from five independent cell clones ranged from 6.3% to 12.9%, with an average of 10.4% ± 1.2% (SE). Statistical analysis indicated that there was no significant difference between the rates of recombination at the 1.3- and 1.9-kb marker distances with Ψ+ located between hygro and neo (P = 0.30; one-way ANOVA). These results indicated that the recombination rate reached a plateau by a marker distance of 1.3 kb when Ψ+ was located between hygro and neo.

PCR amplification and restriction enzyme analysis of recombinant proviruses generated from JA19ΨNeo and JA20HyΨ. The molecular natures of the recombinant proviruses generated from JA19ΨNeo and JA20HyΨ were examined. To ensure that independent recombination events were studied, most of the cell clones were isolated from different cell culture dishes. For cell clones isolated from the same culture dishes, Southern hybridization analyses were performed to ensure that these proviruses had arisen through independent infection.

![FIG. 5. Provirals structures of JA19ΨNeo, JA20HyΨ, and JA23.](image)

**TABLE 4. Virus titers generated from PG13 cells coinfectected with JA19ΨNeo and JA20HyΨ**

<table>
<thead>
<tr>
<th>Clone</th>
<th>Hygromycin (10^3)</th>
<th>G418 (10^5)</th>
<th>Hygromycin + G418 (10^5)</th>
<th>Recombination rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.3</td>
<td>1.2</td>
<td>3.8</td>
<td>6.3</td>
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<tr>
<td>7</td>
<td>0.50</td>
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<td>12.0</td>
</tr>
<tr>
<td>15</td>
<td>1.5</td>
<td>0.44</td>
<td>2.5</td>
<td>11.4</td>
</tr>
<tr>
<td>4</td>
<td>7.8</td>
<td>20.0</td>
<td>3.7</td>
<td>9.5</td>
</tr>
<tr>
<td>25</td>
<td>1.7</td>
<td>0.87</td>
<td>5.6</td>
<td>12.9</td>
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</table>

Avg ± SE = 10.4 ± 1.2

* Recombination rate = (double-drug-resistant titer/lesser of the single-drug-resistant titers) × 2.
Of the 42 recombinant proviruses analyzed, 34 contained one template switch and 8 contained three template switches in the 2.3-kb region amplified by PCR. Among the 34 proviruses with one template switch, recombination occurred in all four regions between the five sets of restriction enzyme markers. Of these template-switching events, 8 occurred in the 5' portion of neo between FspI of JA199Neo and XhoI of JA20Hy, 1 occurred in the 3' portion of the Pvu region between PvuII of JA199Neo and AffI of JA20Hy, and 12 occurred in the 5' portion of the Psi region between MluI of JA199Neo and ClaI of JA20Hy, and 13 occurred in the 3' portion of hygro between BamHI of JA199Neo and NcoI of JA20Hy (Fig. 6B). Of the proviruses containing three template switches, four patterns were observed. Three proviruses had the first template switch between PvuII of JA199Neo and AffI of JA20Hy, the second template switch between AffI of JA20Hy and BamHI of JA199Neo, and the third template switch between BamHI of JA199Neo and NcoI of JA20Hy. Two proviruses had the first template switch between FspI of JA199Neo and XhoI of JA20Hy, the second template switch between XhoI of JA20Hy and MluI of JA199Neo, and the third template switch between BamHI of JA199Neo and NcoI of JA20Hy. One provirus had the first template switch between FspI of JA199Neo and XhoI of JA20Hy, the second template switch between AffI of JA20Hy and BamHI of JA199Neo, and the third template switch between BamHI of JA199Neo and NcoI of JA20Hy. Two proviruses had the first template switch between FspI of JA199Neo and XhoI of JA20Hy, the second template switch between AffI of JA20Hy and BamHI of JA199Neo, and the third template switch between XhoI of JA20Hy and MluI of JA199Neo, and the third template switch between MluI of JA199Neo and ClaI of JA20Hy.

Distribution of the template-switching events in the recombinant proviruses. During reverse transcription, recombination had to occur to obtain the NcoI site in hygro and the FspI site in neo to generate recombinants with functional hygro and neo. In the region between these two sets of selectable markers, JA199Neo and JA20Hy differed in three other sets of restriction enzyme sites. These five sets of markers divided the 1.9-kb region between the selected markers into four regions (Fig. 6C). Region 4 was located between NsiI-NcoI and BamHI-ClaI markers; this region was 0.66 kb in length and contained the 3' portion of hygro. Region 3 was located between the BamHI-ClaI and MluI-AffI markers; this region was 0.47 kb in length and contained the 5' portion of Psi, including the DLS. Region 2 was located between the MluI-AffI and PvuII-XhoI markers; this region was 0.43 kb in length and contained the 3' region of the Psi. Region 1 was located between the PvuII-XhoI and FspI-AflII markers; this region was 0.32 kb in length and contained the 5' portion of neo.

Of the recombinants analyzed, 34 contained a single template switch and 8 contained three template switches; therefore, a total of 58 template switches were observed (Fig. 6B). The total numbers of template switches within regions 4, 3, 2, and 1 were 20, 18, 7, and 13, respectively (Fig. 6C). If the frequencies of the recombination events were proportional to the marker distance, then the expected frequencies of template switches within regions 4, 3, 2, and 1 would be 35, 25, 23, and 17%, respectively. The observed frequencies of template switches within regions 4, 3, 2, and 1 were 35, 31, 12, and 22%, respectively. The differences between the observed and expected frequencies of template switches in regions 4, 3, 2, and 1 were not highly significant (P = 0.93, P = 0.29, P = 0.048, and P = 0.27, respectively; Pearson's chi-square test). Therefore, these results indicated that recombination events occurred in all four regions, and the presence of DLS did not cause an increased number of template-switching events in region 3.

### A.

**Parental genotypes**

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<tr>
<th>Ns</th>
<th>B</th>
<th>M</th>
<th>P</th>
<th>C</th>
<th>DLS</th>
<th>ψ</th>
<th>Neo</th>
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<td>Neo</td>
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**Template switching events:**

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<th>Distance (kb)</th>
<th>Observed (%)</th>
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<td>0.47</td>
<td>25</td>
</tr>
<tr>
<td>0.43</td>
<td>23</td>
</tr>
<tr>
<td>0.32</td>
<td>17</td>
</tr>
</tbody>
</table>

**B.**

**One template switch**

- 8
- 1
- 12
- 13

**Three template switches**

- 3
- 2
- 2
- 1

**C.**

<table>
<thead>
<tr>
<th>Ns</th>
<th>B</th>
<th>M</th>
<th>P</th>
<th>C</th>
<th>DLS</th>
<th>ψ</th>
<th>Neo</th>
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<td>Hygro</td>
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<td>ψ</td>
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</table>

**Distribution of the template-switching events in the recombinant proviruses.**

- **Fig. 6. Strategy for mapping recombinant proviruses in cell clones.** (A) Structures of the internal portions of the parental viruses. JA199Neo is shown in black, and JA20Hy is shown in white. Restriction enzyme sites are indicated above or below each parental genotype. The two selectable markers are indicated by the open circles. Arrows, hygro- and neo-specific primers for PCR; other abbreviations and symbols are the same as in Fig. 1. (B) Restriction enzyme maps of 42 recombinant proviruses containing one or three template switches. JA199Neo-derived sequences are shown in black, whereas JA20Hy-derived sequences are shown in white. The number to the right of each genotype indicates the number of recombinants with the same genotype observed in the 42 proviruses analyzed. (C) Summary of template-switching events. Restriction enzyme markers corresponding to JA199Neo and JA20Hy are shown above and below a generic recombinant structure, respectively. The distances between each set of restriction enzyme markers for four different regions are indicated below recombinant structure. In addition, the expected and observed frequencies of template-switching events are shown beneath the four regions. Expected frequencies were calculated by dividing the distance between each set of markers by the total distance between the selectable markers. Abbreviations are the same as those in Fig. 1 and 6.
Recombination during retroviral replication provides an additional mechanism besides mutation to increase variation in viral populations. Although frequent recombination has been observed in most retroviruses (6, 15, 23, 24, 26, 29, 30, 46–48), many aspects of this phenomenon are still unclear. In this report, we tested whether the rate of recombination could be altered and determined the nature of this potential alteration. Specifically, we examined the effect of the extended packaging signal on recombination and determined the general effect of the DLS on the location of the template-switching events.

Recombination rates of MLV vectors with $\Psi^+$ located in the 5′ UTR. Previously, it was determined that in one round of replication, the homologous recombination rates with markers separated by 1.0, 1.9, and 7.1 kb were 4.7, 7.4, and 8.2%, respectively (2). In all of these vectors, $\Psi^+$ was located in the 5′ UTR. It was found that the recombination rates reached a plateau when markers were separated by 1.9 kb and did not increase significantly even when the markers were further apart (7.1 kb). However, recombination rates increased as the distance between markers increased from 1.0 to 1.9 kb (4.7 and 7.4%, respectively). These data suggested that recombination rates increased in linear proportion when the distances between markers ranged from 1.0 to 1.9 kb. However, this relationship could not be determined because only two data points were obtained in the previous study. In the current study, a recombination rate of 5.0% ± 0.5% (SE) was determined with markers 1.3 kb apart. This rate is within the expected range calculated from the previous measured recombination rates when markers were separated by 1.0 and 1.9 kb (4.7% ± 1.0 kb × 1.3 kb = 6.1%; 7.4% ± 1.9 kb × 1.3 kb = 5.1%). Together with the previous published recombination rates, these data suggested that when the distances between the selectable markers were in the range of 1.0 to 1.9 kb, recombination rates increased linearly in proportion to the distance between markers. This is the first set of evidence to support that within a limited range, a linear relationship exists between recombination rates and the distances between markers.

The effect of the locations of $\Psi^+$ on the recombination rates of MLV vectors. It was previously observed that the entire viral population was capable of undergoing frequent template switching (17). However, only a subpopulation of viruses underwent recombination (intramolecular template switching). We hypothesized that the barrier to intramolecular template switching is the accessibility of the copackaged RNA. We suggested that the recombination rate may be altered by bringing the two copackaged RNAs closer together for the selectable markers by using the dimerization signal in $\Psi^+$. However, it was not clear whether the distance between the copackaged RNAs was the only factor separating the viral subpopulation that could undergo recombination from the rest of the viruses. If the distance between the copackaged RNAs was the only factor, then the size of the subpopulation would be changed by relocating $\Psi^+$. In contrast, if the distance between the two RNAs was not the only factor, then it was quite possible that the size of the recombining subpopulation would not be drastically altered. To test our hypothesis, the recombination rates of MLV vectors with $\Psi^+$ located in the 5′ UTR or between the selectable markers were compared. When selectable markers were separated by 1.3 kb, vectors with $\Psi^+$ located between hygro and neo had an approximately twofold-higher recombination rate than those from vectors with $\Psi^+$ located in the 5′ UTR. This enhancement of recombination was less pronounced when the selectable markers were separated by 1.9 kb. The recombination rates of vectors with selectable markers separated by 1.3 and 1.9 kb were not significantly different when $\Psi^+$ was located between hygro and neo (12.0 and 10.4%, respectively; $P = 0.30$; one-way ANOVA). These data also suggested that the rate of recombination reached a plateau when the markers were separated by 1.3 kb. This was in sharp contrast with the observation that when $\Psi^+$ was located in the 5′ UTR and markers were 1.0 to 1.9 kb apart, recombination rates were in linear proportion to the distances between the selectable markers. Taken together, our data indicated that with the relocation of the $\Psi^+$, the recombination rates reached a plateau at a shorter distance between the markers. However, the overall maximum recombination rates were not drastically different between vectors with $\Psi^+$ in the 5′ UTR and $\Psi^+$ between the two selectable markers: 7.4 to 8.2% and 10.4 to 12%, respectively. The maximum observed recombination rates approximated the size of the recombining subpopulation (2). Therefore, these data indicated that relocating $\Psi^+$ between the two selectable markers produced a minor alteration in the size of the recombining subpopulation. This also suggested that factors other than the distance between the two RNAs were involved in separating the recombining and nonrecombining populations.

Our study demonstrated that when markers were separated by 1.3 kb, the recombination rate was altered by placing $\Psi^+$ between the selectable markers. A previous study using an SNV-based system had shown that relocating the packaging signal to the 3′ end of the viral RNA did not alter the recombination rate when markers were separated by 1.0 kb (21). The different effects of packaging signal relocation on recombination rates in these two studies could be easily explained by the positions of the packaging signal. In the previous study, the packaging signal was moved from the 5′ UTR to the 3′ UTR; this would not affect the relative distance of the RNAs between the two selected markers. In our study, however, the packaging signal was placed between the selectable markers; this affected the relative distance of the RNAs between the two selectable markers and, as a consequence, altered the recombination rate.

Recombination in MLV exhibits high negative interference. High negative interference described a phenomenon in which a greater-than-expected probability of multiple recombination events was observed (3, 5, 17, 49). Using the observed recombination rate of 10.4% with $\Psi^+$ between markers separated by 1.9 kb, it was estimated that one in five proviruses should contain a single recombination event (see reference 17 for a detailed calculation). Since all of the proviruses analyzed in these experiments were recombinants, all proviruses contained at least one recombination event. Therefore, 1 in 5 and 1 in 25 recombinants would be expected to contain two and three recombination events, respectively. It was expected that recombinants with two template switches between the 1.9-kb region would contain a parental genotype and would not survive double-drug selection for functional hygro and neo. However, recombinants with three template-switching events could contain functional hygro and neo and should be observed. Analysis of 42 recombinant proviruses indicated that 34 contained one template switch, whereas 8 contained three template switches (Fig. 6B). Approximately 19% of the recombinant proviruses contained three template switches; this was fivefold greater than expected. Therefore, similar to the observations made with SNV (3, 17), homologous recombination in MLV also exhibited high negative interference. In addition, recombinants with multiple template switches were also frequently observed in human immunodeficiency virus type 1 (6, 50). Taken together, we propose that high negative interference is an inherent property of retroviral recombination.
The general effect of packaging signal on the locations of the template-switching events. Using a system selecting recombination in the MLV packaging signal, it was shown that template-switching events occurred frequently within a 33-nucleotide region that coincided with the DLS (31, 34, 35). These data suggested that DLS is a hot spot for recombination.

In our study, recombination can occur within any of the four regions defined by the restriction enzyme sites between the two selectable markers (Fig. 6). We found that the region containing the 5′-W′ sequences including the DLS did not experience significantly more template switching than expected. This region is 0.47 kb in length, and 25% of the recombination events were expected to occur in this region (0.47 kb/1.9 kb = 25%) if the recombination events occurred randomly. We observed that 31% of the recombinants had a template switch in this region (Fig. 6C). Therefore, the presence of DLS did not cause an increase in recombination events in this general region. This result was in agreement with our previous study using SNV-based vectors (3), in which the locations of the template-switching events throughout the entire vector genomes were determined. It was found that the 0.24-kb region containing the DLS did not experience increased template-switching events. In both the SNV and the MLV studies, we examined the frequencies of template-switching events within the general region. The exact location of the cross-overs in the recombinants could not be determined due to the lack of markers flanking the DLS. We elected not to place mutations flanking the DLS in these studies to avoid the potential impact on the frequencies of RNA heterodimer formation that could influence other aspects of our studies.

Currently, there are several series of studies pointing at different roles of DLS in recombination. It was shown that two HIV-1 clones with different DLS could undergo efficient recombination (43). Our studies indicated that the presence of DLS did not increase the recombination rate within a 0.47- or 0.24-kb region. However, forced recombination studies indicated a cluster of crossovers occurring in the DLS region (31, 0.24-kb region. However, forced recombination studies indicated a cluster of crossovers occurring in the DLS region (31, 34, 35). The data generated from these studies are not necessarily conflicting, because distinct selection pressures were applied in each system and different aspects of recombination are likely to also play important roles. Other studies are currently in progress in our laboratories to further explore the role of the DLS in recombination.

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