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Research Article  Genetics

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The genotoxic potential of retroviral vectors is strongly modulated by vector design and integration site selection in a mouse model of HSC gene therapy

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γ-Retroviral vectors (γRVs), which are commonly used in gene therapy, can trigger oncogenesis by insertional mutagenesis. Here, we have dissected the contribution of vector design and viral integration site selection (ISS) to oncogenesis using an in vivo genotoxicity assay based on transplantation of vector-transduced tumor-prone mouse hematopoietic stem/progenitor cells. By swapping genetic elements between γRV and lentiviral vectors (LVs), we have demonstrated that transcriptionally active long terminal repeats (LTRs) are major determinants of genotoxicity even when reconstituted in LVs and that self-inactivating (SIN) LTRs enhance the safety of γRVs. By comparing the genotoxicity of vectors with matched active LTRs, we were able to determine that substantially greater LV integration loads are required to approach the same oncogenic risk as γRVs. This difference in facilitating oncogenesis is likely to be explained by the observed preferential targeting of cancer genes by γRVs. This integration-site bias was intrinsic to γRVs, as it was also observed for SIN γRVs that lacked genotoxicity in our model. Our findings strongly support the use of SIN viral vector platforms and show that ISS can substantially modulate genotoxicity.

Introduction
Integrative viral vectors commonly used in gene therapy may trigger oncogenesis as a consequence of insertional mutagenesis (1, 2). Because γ-retroviral vector (γRV) gene transfer into hematopoietic stem/progenitor cells (HSPCs) caused an unexpectedly high frequency of clonal proliferation and overt leukemia in clinical trials (3, 4), safer vectors and stringent preclinical safety assays (5, 6) are urgently needed to overcome this major hurdle.

We previously tested the oncogenic potential of prototypical murine leukemia virus–derived (MLV-derived) γRV and HIV-derived lentiviral vector (LV) using an in vivo genotoxicity assay based on transduction and transplantation of tumor-prone Cdkn2a−/− murine HSPCs (7). The Cdkn2a locus has a central role in regulating senescence and preventing cell transformation caused by aberrant oncogene expression. Because Cdkn2a inactivation synergizes with several types of cancer-promoting lesions, Cdkn2a−/− mice have been invaluable in insertional mutagenesis studies for identifying cancer genes, many of which are highly relevant in human oncogenesis. The relevance of the CDKN2A pathway in human tumor suppression is well documented because of its frequent inactivation in almost all types of human cancer (8, 9). Moreover, 2 X-linked SCID (X-SCID) patients affected by γRV-induced leukemia from 2 independent clinical trials had lost expression of the CDKN2A locus as a secondary mutation (10, 11). These findings indicate that CDKN2A plays a role also in the pathophysiology of human leukemias triggered by γRV insertions in clinical trials and further validate the choice of this model to assess vector genotoxicity.

In our previous study (7), γRV treatment triggered a dose-dependent acceleration of tumor onset in transplanted mice, whereas LV did not. Because the LV tested differed from γRV in both the molecular design (12, 13) and the integration site selection (ISS) (14–17), the relative contribution of these features to the lower genotoxicity of LV remained unknown. This may be due to the self-inactivating (SIN) long terminal repeat (LTR) (18) coupled to placement of retroviral enhancer/promoter within the vector reduced genotoxicity only 2-fold in vitro (21) and was insufficient to enhance safety in an in vivo model (22). The combination

By comparing the genotoxicity of vectors with matched active LTRs, we were able to determine that substantially greater LV integration loads are required to approach the same oncogenic risk as γRVs. This difference in facilitating oncogenesis is likely to be explained by the observed preferential targeting of cancer genes by γRVs. This integration-site bias was intrinsic to γRVs, as it was also observed for SIN γRVs that lacked genotoxicity in our model. Our findings strongly support the use of SIN viral vector platforms and show that ISS can substantially modulate genotoxicity.
of strong transcriptional enhancers with the \( \gamma_RV \) preference for integrating close to promoters may imply a higher risk of altering gene expression as compared with that of LV (16, 17, 23, 24). Moreover, the putative \( \gamma_RV \) integration bias for “hot spots” or gene classes associated with cell growth and cancer may further increase the oncogenic risk of insertional mutagenesis (7, 25–29).

In order to identify the most relevant features responsible for the different genotoxicity of \( \gamma_RV \) and LV, we swapped genetic features between these vectors and tested the genotoxicity of a panel of chimeric vectors on tumor-prone HSPCs in vivo. Using this strategy, we were able to identify transcriptionally active LTR as the major determinant of genotoxicity and validate the improved safety conferred by SIN LTR design in both vector platforms. Interestingly, however, ISS, when active LTR were present, modulated genotoxicity to an unanticipated extent.

**Results**

*Vector construction and testing in tumor-prone HSC transplantation.* The panel of chimeric and parental vectors tested is shown in Figure 1A. We challenged the previously reported safety of LV by introducing the strong spleen focus–forming virus (SF) retroviral enhancer/promoter (30) in the U3 region of the LTR (LV.SF.LTR) and comparing this vector to a \( \gamma_RV \) carrying SF LTRs (RV.SF.LTR) (31). Similarly, we tested the oncogenic potential of a \( \gamma_RV \) with SIN LTRs carrying the moderately active human phosphoglycerate kinase (PGK) promoter in internal position (SIN.RV.PGK) (13). To address the position dependence of strong enhancer/promoters in genotoxicity, we placed the SF sequence in an internal position within the vector (SIN.LV.SF) and compared it with its SF.LTR counterpart. Finally, to assess the impact of promoter strength, we compared SIN LV with internal SF and PGK promoter. All vec-
Tumor-infiltrated tissues were systematically analyzed by Q-PCR to measure the relative amounts of donor-derived and recipient cells and the VCN in donor cells. Cdkn2a−/− cells constituted more than 70% of the affected tissue, indicating the donor origin of tumors. Donor cells in the tumor-infiltrated tissues, obtained from each group, had an average VCN of 9.9 ± 6.2 for LV.SF.LTR, MOI = 100, (n = 29) and 1.2 ± 1.2, MOI = 10, (n = 20); 3.0 ± 1.4 for RV.SF.LTR (n = 10); 1.14 ± 7.4 for SIN.LV.SF (n = 32); and 8.6 ± 4.1 for SIN.RV.PGK (n = 20) (Figure 1E and Supplemental Table 1).

Assessing the oncogenic risk of vector treatment. The mock-transduced control group had a median survival time corresponding to a donor cell age of 257 days, consistent with the median survival of Cdkn2a−/− mice (32, 33) and virtually identical to that in our previously published results (7), showing that cell manipulation and transplant procedures per se do not accelerate tumor onset and that our assay is highly reproducible. Survival in each experimental group was analyzed by Kaplan-Meier curves (Figures 2, A and B). Mice transplanted with HSPCs transduced at the higher dose of LV.SF.LTR (LV.SF.LTR, MOI = 100) died significantly earlier than the mock-transduced controls (median survival, 187 days vs. mock 248 days; P = 0.0001, log-rank Mantel-Cox test). Mice treated with the lower vector dose (LV.SF.LTR, MOI = 10) displayed a median survival of 211 days (Figure 2A). The median survival time was 194.5 days for RV.SF.LTR mice (Figure 2A), 227.5 days for SIN.RV.PGK mice, and 238 days for SIN.LV.SF mice (Figure 2B).

Table 1
Tumor phenotype incidence in each group of transplanted mice

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Lymphoid</th>
<th>Myeloid</th>
<th>P value vs. mock&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Guts with myeloid infiltrate</th>
<th>P value vs. mock&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>LV.SF.LTR, MOI = 10</td>
<td>20</td>
<td>7 (35%)</td>
<td>13 (65%)</td>
<td>NS</td>
<td>7/19 (36%)</td>
<td>0.0084</td>
</tr>
<tr>
<td>LV.SF.LTR, MOI = 100</td>
<td>29</td>
<td>11 (38%)</td>
<td>18 (62%)</td>
<td>NS</td>
<td>13/29 (45%)</td>
<td>0.0007</td>
</tr>
<tr>
<td>LV.SF.LTR all</td>
<td>49</td>
<td>18 (37%)</td>
<td>31 (63%)</td>
<td>0.038</td>
<td>20/48 (42%)</td>
<td>0.0007</td>
</tr>
<tr>
<td>RV.SF.LTR</td>
<td>10</td>
<td>6 (60%)</td>
<td>4 (40%)</td>
<td>NS</td>
<td>4/9 (44%)</td>
<td>0.0084</td>
</tr>
<tr>
<td>SIN.LV.SF</td>
<td>30</td>
<td>22 (73%)</td>
<td>8 (27%)</td>
<td>NS</td>
<td>2/29 (7%)</td>
<td>NS</td>
</tr>
<tr>
<td>SIN.RV.PGK</td>
<td>27</td>
<td>23 (85%)</td>
<td>4 (15%)</td>
<td>NS</td>
<td>3/25 (12%)</td>
<td>NS</td>
</tr>
<tr>
<td>Mock</td>
<td>20</td>
<td>13 (65%)</td>
<td>7 (35%)</td>
<td></td>
<td>0/17 (0%)</td>
<td>-</td>
</tr>
</tbody>
</table>

<sup>a</sup>Tumor phenotype incidence in each group of transplanted mice was compared with that of mice transplanted with mock-treated cells (mock) by Fisher’s exact test. P < 0.05 was considered significant. *Incidence of myeloid tumor infiltrate in the gut compared with that in the mock group, n, total number of mice analyzed.

Table 2
log-logistic accelerated failure time model estimates the impact on survival of vector treatment in each group

<table>
<thead>
<tr>
<th>Vector</th>
<th>n</th>
<th>log-logistic parameter ± SE&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Z</th>
<th>P value vs. mock&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>LV.SF.LTR, MOI = 100</td>
<td>29</td>
<td>-0.292 ± 0.068</td>
<td>-4.288</td>
<td>1.81 × 10&lt;sup&gt;-5&lt;/sup&gt;</td>
</tr>
<tr>
<td>LV.SF.LTR, MOI = 10</td>
<td>22</td>
<td>-0.113 ± 0.077</td>
<td>-1.455</td>
<td>1.46 × 10&lt;sup&gt;-1&lt;/sup&gt;</td>
</tr>
<tr>
<td>RV.SF.LTR</td>
<td>12</td>
<td>-0.202 ± 0.092</td>
<td>-2.191</td>
<td>2.84 × 10&lt;sup&gt;-2&lt;/sup&gt;</td>
</tr>
<tr>
<td>SIN.LV.SF</td>
<td>39</td>
<td>-0.040 ± 0.066</td>
<td>-0.598</td>
<td>5.5 × 10&lt;sup&gt;-1&lt;/sup&gt;</td>
</tr>
<tr>
<td>SIN.RV.PGK</td>
<td>34</td>
<td>-0.059 ± 0.070</td>
<td>-0.831</td>
<td>4.06 × 10&lt;sup&gt;-1&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup>Log-logistic parameters (± SE) evaluate the impact on survival of vector treatment in each group. Vectors with SF.LTR have a negative and significant impact on the survival of the LV.SF.LTR, MOI = 100, and RV.SF.LTR groups. P < 0.05 was considered significant. Z, Z score. n, total number of mice analyzed.
In order to perform a risk assessment analysis that accounted for the accelerated time of death revealed in the descriptive statistics, we used an accelerated failure time model to estimate the risk of death and the survival probability for each treatment. Within this class of models, the log-logistic distribution provided the best fit to our experimental data (Supplemental Statistical Methods). The impact of the treatment (represented by the log-logistic parameter) and the survival probability during time were estimated for each treatment group and compared with that of the mock (Table 2 and Figure 2C). For the LV.SF.LTR, MOI = 100, treatment with LV.SF.LTR, MOI = 10, did not significantly reduce the survival probability as compared with that of the mock, indicating that the genotoxicity of LV.SF.LTR at this dose was below the detection limit of our in vivo assay. Notably, mice in the LV.SF.LTR, MOI = 10, and RV.SF.LTR groups were transplanted with cells carrying similar VCN in vitro (2 and 3, respectively), indicating a higher genotoxicity of RV.SF.LTR. The same analysis performed on the SIN.RV.PGK and SIN.LV.SF groups showed no significant impact of these vectors on the survival probability in our assay, even if both groups had high VCN in vitro.

Because transduction with each vector resulted in a different integration load even when using the same MOI (MOI = 100), we adopted the VCN in tumor (VCN_tum) as a measure of dosage to perform VCN-matched comparisons between different vector treatment groups. Mice were stratified in groups having a VCN_tum ranging from 1 to 6 (VCN_tum<6) or above 6 (VCN_tum>6). The stratification criterion allowed us to compare a relevant number of mice with a similar vector load (Table 3). The survival probability of the LV.SF.LTR-VCN_tum>6 group died significantly earlier than those in the LV.SF.LTR-VCN_tum<6 group (P = 1 × 10−3) and the mock group (P < 1 × 10−4; Figure 3A). The survival of the RV.SF.LTR-VCN_tum<6 group was also significantly reduced with respect to the mock group (P = 2 × 10−4) and the VCN-matched LV.SF.LTR-VCN_tum<6 group (P = 2 × 10−2). Indeed, the LV.SF.LTR-VCN_tum<6 group, which comprised 8 mice from the MOI = 100 and 13 from the MOI = 10 transduction groups, had a survival not significantly different from that of the mock group. The SIN.RV.PGK and SIN.LV.SF groups did not show any significant acceleration with respect to the mock group when stratified according to VCN_tum>6 (Figure 3B). Similar results were obtained by applying the log-logistic model to determine the survival probability of each stratified vector treatment group compared with the mock group (Supplemental Figure 3, A–C). These results showed that vector genotoxicity is dependent on the presence of active LTRs and on the VCN of

Table 3

<table>
<thead>
<tr>
<th>Vector</th>
<th>Stratification group</th>
<th>n</th>
<th>Average VCN&lt;sup&gt;in vivo&lt;/sup&gt;</th>
<th>Median survival (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mock</td>
<td>NA</td>
<td>32</td>
<td>0</td>
<td>248</td>
</tr>
<tr>
<td>LV.SF.LTR</td>
<td>1 to 6</td>
<td>22</td>
<td>2.8</td>
<td>214.5</td>
</tr>
<tr>
<td></td>
<td>&gt;6</td>
<td>21</td>
<td>12</td>
<td>172</td>
</tr>
<tr>
<td>RV.SF.LTR</td>
<td>1 to 6</td>
<td>10</td>
<td>3.1</td>
<td>192</td>
</tr>
<tr>
<td></td>
<td>&gt;6</td>
<td>9</td>
<td>4.3</td>
<td>254</td>
</tr>
<tr>
<td>SIN.LV.SF</td>
<td>1 to 6</td>
<td>23</td>
<td>14.2</td>
<td>238</td>
</tr>
<tr>
<td>SIN.RV.PGK</td>
<td>1 to 6</td>
<td>6</td>
<td>3.7</td>
<td>225.5</td>
</tr>
<tr>
<td></td>
<td>&gt;6</td>
<td>20</td>
<td>9.5</td>
<td>238</td>
</tr>
</tbody>
</table>

n, number of mice for each stratification group.
in vitro–cultured cells or tumors. However, the negative impact on survival varies according to the type of vector and VCN.

In order to quantitatively assess the relative impact of VCN\textsuperscript{um} for each vector, we used the VCN\textsuperscript{um} of each mouse as a covariate in the accelerated failure time model. We observed that VCN\textsuperscript{um} acts linearly on the log-logistic hazard (Supplemental Statistical Methods). VCN\textsuperscript{um} affected negatively and significantly the risk of death only in interaction with the LV.SF.LTR, MOI = 100 (log-logistic parameter = −0.019; \( P = 6.51 \times 10^{-4} \)) and RV.SF.LTR (log-logistic parameter = −0.065; \( P = 6.58 \times 10^{-3} \)) but not with the LV.SF.LTR, MOI = 10, or any SIN.LTR vector (Table 4). The lack of a statistically significant impact of VCN\textsuperscript{um} for LV.SF.LTR, MOI = 10, suggests that genotoxicity was too low at this dosage to be measured in a reliable manner (Supplemental Figure 3D). This approach allowed estimation of the relative risk associated with different vectors at set dose levels. We thus plotted the risk of death at a fixed VCN\textsuperscript{um} of 1 or 10 for the genotoxic vectors (Figure 3C). RV.SF.LTR showed the highest risk, whereas LV.SF.LTR required a 10-fold higher integration load (VCN\textsuperscript{um}) to reach the same risk of \( \gamma \)RV.

**Vector integration analyses.** To gain functional evidence that LV.SF.LTR-driven oncogenesis in our Cldn2\textsuperscript{−/−} model was mediated by insertional mutagenesis and to gain more insight into the low genotoxicity profile of SIN.RV.PGK and SIN.LV.SF, we compared genes targeted by integration of these vectors in tumors and in the cells used for transplant after 2 weeks of culture (in vitro). DNA from tumor-infiltrated BM (36 LV.SF.LTR, 27 SIN.LV.SF, and 23 SIN.RV.PGK mice) was subjected to a low-sensitivity linear amplification–mediated PCR (LAM-PCR) protocol aimed at identifying provirus-genomic junctions from predominant clone(s) in mixed populations (4, 7). Standard LAM-PCR protocol was used for the in vitro–cultured cells (34). We univocally mapped a total of 529 vector integration sites on the mouse genome (UCSC Mouse Genome Browser, February 2006 release) divided into 6 data sets: LV.SF.LTR, 100 sites from tumors and 70 in vitro; SIN.LV.SF, 80 from tumors and 90 in vitro; SIN.RV.PGK, 54 from tumors and 135 in vitro. The nearest gene (known to Entrez Gene or Ensembl) was then identified by bioinformatics analysis (Supplemental Table 2). Redundant integrations were excluded from calculations (total nonredundant integrations in tumors: LV.SF.LTR = 93 and SIN.LV.SF = 78).

In the LV.SF.LTR and SIN.LV.SF data sets, the distribution of vector integrations displayed a pronounced tendency to integrate within genes (70%) without preference for transcription start sites (TSS), a pattern similar to that previously reported for other LVs (14, 16, 17, 23) (Supplemental Figure 4). On the other hand, 40% of SIN.RV.PGK integrations were located within genes, and 32% of the integrations clustered within ± 5 kb from the TSS.

Each data set was searched for matches to retroviral or Sleeping Beauty (SB) transposon common integration site (CIS) genes contained in the Retrovirus Tagged Cancer Gene Database (RTCGD) (35) and the frequency compared with the expected random frequency (Figure 4A, Supplemental Table 3A, and Supplementary Table 4). The SIN.RV.PGK had a pronounced tendency to target retroviral CIS in vitro and in tumors (\( P < 0.0001 \) vs. expected random; \( \chi^2 \) test) but not SB CIS. Several of the SIN. RV.PGK integrations targeting CIS genes mapped in the same narrow region previously targeted by retroviruses in the RTCGD (Supplemental Figure 5). On the other hand, the frequency of SIN.LV.SF integration near retroviral or SB CIS genes in vitro or in tumors was not significantly different from the expected random frequency. In contrast to the pattern observed for both SIN.LVs, LV.SF.LTR targeted both retroviral and SB CIS genes in hand, 0.004 ± 0.006 0.606 5.44
LV.SF.LTR, MOI = 10 0.054 ± 0.036 1.52 1.28
LV.SF.LTR, MOI = 100 –0.019 ± 0.005 –3.994 6.51
RV.SF.LTR –0.065 ± 0.024 –2.717 6.58
SIN.LV.SF 0.002 ± 0.003 0.591 5.54
SIN.RV.PGK 0.004 ± 0.006 0.606 5.44

A log-logistic parameters (± SE) evaluate the impact on survival of vector treatment and vector copy number on tumors in each group. \( \text{VCN}_{\text{um}} \) of LV.SF.LTR, MOI = 100, and of RV.SF.LTR has a negative impact on survival as shown by the log-logistic parameter and Z score, whereas \( \text{VCN}_{\text{um}} \) of all other vectors does not. \( P < 0.05 \) was considered significant.
vitro and in tumors at significantly high frequency ($P < 0.0001$ vs. expected random; $\chi^2$ test).

The new data sets were then evaluated with DAVID-EASE (36) and Ingenuity Pathways Analysis (IPA) software for overrepresentation of Gene Ontology (GO) classes and signaling or disease pathways (Figure 4, B and C). Analysis was performed at high stringency and the results limited to the significantly overrepresented classes with an increase of 3-fold or greater with respect to the expected random frequency (DAVID-EASE; Supplemental Tables 3 and 4) or with 3 or more genes in at least 1 data set (IPA; Supplemental Table 5) and validated by Bonferroni’s correction for multiple comparison error.

For LV.SF.LTR, the GO class Phosphorylation and Kinase Activity was overrepresented in vitro, while the GO classes Regulation of Apoptosis, Mitotic Cell Cycle, and B Cell Differentiation and the IPA classes Cancer, Cell Cycle, and Cell Death were all strongly overrepresented in tumors. As shown by $P$ value ranking, the latter were the most significant overrepresentations found among all data sets, consistently with the observed oncogenic effect of LV.SF.LTR.

For SIN.LV.SF, the only overrepresented gene classes were the GO Chromatin Modification and the IPA Molecular Transport in vitro and the GO Helicases, Protein Phosphatase, and Intracellular Protein Transport in tumors.

For SIN.RV.PGK, the IPA Cancer and Post-Translational Modification classes were strongly overrepresented in vitro together with the GO classes Chromatin Modification and GTPase Activator, whereas the only overrepresented gene classes in tumors were the GO Protein Transport and Localization and the IPA Gene Expression classes.

We then determined the targeting frequency for the overrepresented IPA classes of each data set and performed 2-tailed Fisher’s exact test for comparing vectors and conditions (Figure 4D). In tumors, LV.SF.LTR integrations at Cancer and Hematological Disease genes were significantly enriched from the in vitro data set and were more frequent than observed for the SIN.LV.SF and SIN.RV.PGK. Interestingly, SIN.RV.PGK integrations at Cell Cycle, Cell Death, and Cell Growth
and Proliferation genes were found at significantly reduced frequency in tumors compared with the in vitro data set.

**Mechanism of insertional mutagenesis by LV.SF.LTR.** In order to characterize the oncogenic mechanism of LV.SF.LTR, we measured the transcription level of genes near the vector integration site in early occurring tumors. When possible, tumors selected for analysis were transplanted into secondary mice to obtain biological replicates. For each gene near the integration site, we compared the average expression data for primary and serially transplanted tumors with an integrated vector near the tested gene (INT) and phenotype-matched tumors with integrated vector in different sites or without integrations (No INT) are plotted. Each point is the fold change relative to matched-type tumor-infiltrated BM or spleen from the mock group (control level = 1); the horizontal bar represents the average. P value of the Mann-Whitney test comparison between the samples is indicated. P < 0.05 is considered significant. Genomic region targeted by the vector (vector position and orientation are represented by arrows) is shown below each set of expression data. Genes above the thick horizontal bar (chromosome) are transcribed from left to right; those below the chromosome are transcribed in the opposite direction. (A) Tgtp, which encodes for an interferon-inducible T cell–specific GTPase and whose TSS maps 530 bp from the vector integration, was overexpressed in both tumor-infiltrated BM and spleen of 2 primary and 4 secondary transplanted mice bearing the same integration; the expression of other genes surrounding the integration was not altered (see details in Supplemental Table 6). (B) Another integration from the same groups of mice mapped within the Sos1 (37) oncogene, leading to its significant overexpression. (C) Vector integration occurred within the Eps15 (38) oncogene, leading to its overexpression in tumors of 1 primary and 2 secondary transplanted mice.

**Figure 5**
Gene expression analysis at LV.SF.LTR integration sites in tumors. (A–C) Expression of the indicated genes was measured by Q–RT-PCR on tumor-infiltrated BM or spleen cDNA (see also Supplemental Table 6). Expression data for primary and serially transplanted tumors with an integrated vector near the tested gene (INT) and phenotype-matched tumors with integrated vector in different sites or without integrations (No INT) are plotted. Each point is the fold change relative to matched-type tumor-infiltrated BM or spleen from the mock group (control level = 1); the horizontal bar represents the average. P value of the Mann-Whitney test comparison between the samples is indicated. P < 0.05 is considered significant. Genomic region targeted by the vector (vector position and orientation are represented by arrows) is shown below each set of expression data. Genes above the thick horizontal bar (chromosome) are transcribed from left to right; those below the chromosome are transcribed in the opposite direction. (A) Tgtp, which encodes for an interferon-inducible T cell–specific GTPase and whose TSS maps 530 bp from the vector integration, was overexpressed in both tumor-infiltrated BM and spleen of 2 primary and 4 secondary transplanted mice bearing the same integration; the expression of other genes surrounding the integration was not altered (see details in Supplemental Table 6). (B) Another integration from the same groups of mice mapped within the Sos1 (37) oncogene, leading to its significant overexpression. (C) Vector integration occurred within the Eps15 (38) oncogene, leading to its overexpression in tumors of 1 primary and 2 secondary transplanted mice.
sion level in tumors carrying an integrated vector near or within that gene to the expression levels in tumors of identical phenotype but with different or no integration (Figure 5 and Supplemental Table 6). Q–RT-PCR was performed on cDNA from tumor-infiltrated BM and/or spleen tissue of 12 mice to test the expression of 18 genes surrounding 11 integration sites contained in 6 different tumors. Two primary lymphoid tumors and 4 secondary transplants (2 from each primary tumor; tumors originated from different mice transplanted with the same in vitro–transduced cell populations) shared 4 integration sites. Among 9 genes tested that surrounded these integrations, Tgtp (Figure 5A) and Sos1 (Figure 5B) were strongly and significantly overexpressed with respect to the controls (24.7 ± 3-fold increase and 6.3 ± 3.8-fold increase, respectively; P < 0.001, n = 5 vs. 9). These findings were confirmed in the spleen (9.5 ± 5.4-fold increase, P < 0.001; and 5.2 ± 3.3-fold increase, P < 0.01, respectively; n = 6 vs. 12).

In another lymphoid tumor and its 2 secondary transplants, Eps15 showed a 5.4 ± 2-fold increase compared with the controls (P = 0.013, n = 3 vs. 11) among 5 genes near the vector integration (Figure 5C). All other genes tested were not significantly different from the controls. Of note, SOS1 and EPS15 have been implicated as oncogenes in human cancer development (37, 38).

In a myeloid tumor, 1 LV.SF.LTR integration mapped within intron 11 of Braf, a genomic region targeted several times by transposon integrations in sarcomas of Arf–/– mice (39) (Figure 6A). In this tumor, we detected a chimeric LV-Braf transcript that contained LV LTR and leader sequence up to the splice donor motif fused to the start of exon 13 and the remaining coding sequence of Braf (Figure 6B). This transcript must originate from the LV 5′LTR by splicing out the genomic sequence spanning from the LV splice donor to the acceptor site of exon 13 (Figure 6C). The putative protein encoded by this transcript is a truncated Braf molecule with constitutive kinase activity similar to that previously reported upon transposon integration within the same region and that has been directly implicated in cell transformation (39).

In 1 myeloid and 1 lymphoid tumor, the Nsd1 gene was targeted by 2 independent integrations 2,373-bp apart and in opposite orientation from each other (introns 5 and 6). The levels of expression of Nsd1 appeared to be reduced to about 40% in both tumors bearing the integration (n = 2 vs. 5). Interestingly, NSD1 haploinsufficiency is the major cause of Sotos syndrome and is associated with malignant tumor formation (40).

Overall, in each of the 6 tumors tested, we found at least 1 integration that resulted in either oncogene overexpression, generation...
of aberrant transcripts encoding a truncated constitutively active oncogenic protein, or putative haploinsufficiency of a tumor suppressor gene. These genotoxic events recapitulate those previously described for γ-retroviruses or transposon-driven oncogenesis.

Discussion

Here we show that LV with chimeric γRV LTR results in strong dose-dependent acceleration of tumor onset in the Cdkn2a−/− HSPC transplantation model, as observed for the γRV counterpart. On the other hand, both LV and γRV with SIN LTR appear to be neutral in our model. Although we cannot exclude that the SIN vectors tested could still display some degree of genotoxicity, their reduced oncogenic potential when compared with the vectors bearing active LTRs clearly indicates that transcriptionally active LTRs are major determinants of genotoxicity in retroviral vectors.

Beside active LTRs and vector dosage, we demonstrate that additional factors modulate vector genotoxicity to an unexpected extent. By comparing the survival of mice carrying matched copy numbers of LV or γRV with active LTRs either in the transplanted cells or in the tumors, we found that γRV was significantly more genotoxic than LV. By modeling the impact of vector and dosage on survival, we estimated that an approximately 10-fold higher integration load of LV with active LTRs is required to approach the same risk of a matched-design γRV. This does not mean that a single integration of our genotoxic LV within a cell might not trigger oncogenesis but rather that the relative oncogenic risk associated with an integration of LV and γRV, when bearing matched active LTRs, differs. This difference likely reflects the γRV integration bias for promoters (7, 14, 17, 23, 25–29) and selected gene classes involved in growth control and cancer (25–28), which may increase the probability of oncogene activation and, consequently, cancer development. Indeed, our vector integration analysis with respect to genomic features and gene classes supports a role of ISS in genotoxicity.

In the first detailed integration analysis of a SIN γRV, we observed a strong tendency to integrate not only near promoters but also near RTCGD CIS genes, which are hotspots of γ-retroviral integration retrieved from virus-induced tumors (35, 41) and are enriched in cancer genes. A bias for integration at Cancer genes was independently confirmed by IPA analysis, which showed strong overrepresentation of this gene class in the SIN RV.PGK in vitro data set. Notably, this bias was observed despite the fact that the SIN γRV did not show genotoxicity in vivo and integrations at Cancer, Cell Cycle, and Cell Death genes appeared even underselected in tumors. These findings strongly support the notion that γRVs have an intrinsic bias for integration at gene subsets enriching for cancer genes, which can be revealed even when genetic selection of cells harboring integration at oncogenes is prevented by the lack of strong transcriptional enhancers in the vector. On the contrary, SIN.LV.SF integrates near RTCGD CIS or Cancer genes at a frequency not significantly different from the expected random frequency.

LV.SF.LTR, however, showed a high frequency of integrations at RTCGD CIS and Cell Cycle, Apoptosis, and Cancer genes both in vitro and in tumors. Integrations at Cancer and Hematological Disease genes were further significantly enriched in tumors. The finding that only the genotoxic LV.SF.LTR and not the LV with SIN LTRs enriched for these types of integrations suggests that genetic selection of clones with integrations altering the expression of survival/proliferation genes occurred in vitro and in vivo. Genetic selection in vitro could occur because the cells were grown for 2 weeks before analysis (21, 42). Moreover, LV.SF.LTR targeted both retroviral and SB CIS genes, whereas the γRV tested targeted only retroviral CIS, further supporting the notion that the spectrum of targetable genes differs between the 2 vector types and that genetic selection rather than a genomic integration bias acquired by the LTR modification was the major driving force determining the LV.SF.LTR pattern of integration. Of note, because γRVs are unable to infect quiescent cells and are biased for integration at certain gene classes, the combination of effective insertion activation, typical of γRV, with the broad tropism and ISS of our LV.SF.LTR provides a mutagenic tool that may open new avenues to oncogene discovery in hematopoietic and solid tumors. We cannot exclude that the SF sequences introduced into the HIV LTR may influence the LV integration pattern by recruiting transcription factors that tether the preintegration complex to specific sites. This appears unlikely, however, because SIN.LV.SF, which bears the SF sequence in internal position, does not show an enrichment of integrations at Cancer genes as does LV.SF.LTR.

The characterization of the impact of LV.SF.LTR integration on the expression of targeted genes in tumors indicates that the mechanism of oncogenesis by LV.SF.LTR recapitulates the essential features discovered for γ-retroviruses, retrotransposons, and γ-RVs (3, 4, 7, 20, 39, 43–45). The lower risk of insertional mutagenesis by LV with active LTRs as compared with design-matched γRV can thus be explained by the lower frequency by which LVs target promoters of RTCGD CIS genes, many of which are involved in cancer.

The negative correlation between survival and VCNsum observed in mice treated with the same LV.SF.LTR dose suggests the occurrence of synergistic interaction between multiple integrations within the same cell in driving oncogenesis. This finding is in agreement with previous studies describing the cooccurrence of cooperating oncogenes in insertional mutagenesis screening performed in wild-type and tumor-prone mice (46–48) and supports the notion that both the total number of integrations administered and the average VCN per transplanted cell are relevant factors to be considered when estimating the oncogenic risk of a genotoxic vector (1, 2).

Accelerated tumor onset with an increased incidence of T cell lymphomas is a characteristic feature of MLV-mediated oncogenesis in Cdkn2a−/− mice (20, 32). In our study, the genotoxic LV.SF.LTR and RV.SF.LTR vectors induced an increased incidence of myeloid tumors that widely infiltrated peripheral organs. The reason(s) for the different phenotypic skewing induced by MLV and our vectors may be the different types of enhancers contained within the LTR and/or the different cell types targeted during chronic MLV infection in vivo as compared with a single ex vivo transduction of hematopoietic progenitors (49).

Remarkably, SIN.LV with internal PGK or SF promoter (this and our previous study; ref. 7) and SIN.RV.PGK did not accelerate tumor onset in the Cdkn2a−/− model, validating the improved safety of this configuration for both vector platforms. The improved safety of the SIN γRV design combined with a moderate internal cellular promoter (EF1α) was also recently shown by Zychlinski et al. (42) in an in vitro immortalization model.

The lack of genotoxicity of the SIN.LV.SF in our study is surprising given that (a) LV containing the same sequence duplicated within the LTR was genotoxic in our model; (b) SIN.RVs with an internal SF promoter were genotoxic both in vitro (21, 42) and upon transduction and serial transplantation of wild-type murine HSC (22); (c) a SIN LV with a strong LTR-derived murine stem
cell virus promoter in internal position was genotoxic in vitro (50); and (d) enhancers are well documented as interacting with distant promoters (20). We should consider that the background oncogenesis of Cdkn2a−/− HSPCs may hamper the detection of low-frequency or late-onset tumors induced by residual genotoxicity of the SIN.LV.SF vector and thus avoid the potential interpretation that LV can be safe even when containing strong enhancer sequences. Although our model did not reveal a difference in genotoxicity between SIN LV and SIN γRV, we show that the tendency of γRV to target cellular promoters and integrate at retroviral CIS and cancer genes remains unaffected by modifying the LTR for self inactivation. Therefore, we may speculate that SIN γRV with strong internal promoters may still pose a higher risk of genotoxicity than matched-design LV.

Nonetheless, the reduced oncogenic potential of SIN.LV.SF with respect to LV.SF.LTR is compelling and may help to unravel features of vector design contributing to genotoxicity. A possible explanation is that engagement in transgene transcription may reduce the probability and strength of interaction of the SF enhancer with neighboring genes (21, 42), while the presence of duplicated enhancer sequences increases the probability of transactivation. Another contributing factor is suggested by our finding of oncogenic LV/Braf chimeric transcripts in an LV.SF LTR tumor, which suggests that the capacity to generate fusion transcripts by splicing capture may play an important role in vector genotoxicity. The mechanism of splicing mediated activation of oncogene by LVs with active LTRs has been recently described in a cell-culture assay for insertional mutagenesis (51). The placement of enhancer/promoter sequences in the LTR upstream of a strong splice donor site increases the probability of chimeric transcript formation as compared with other configurations such as the SIN.LV.SF vector, which lacks splice donor sites downstream of the internal SF promoter. We may thus speculate that safe vector design should avoid splice donor sites downstream of strong promoters in particular if a weak polyadenylation site is present in the vector. The inclusion of strong polyadenylation sites in gene therapy vectors to reduce the probability of readthrough into cellular genes has been proposed (52).

However, the impact of this manipulation on vector genotoxicity remains to be directly tested, as it could also increase the frequency of premature transcription termination of genes targeted by vector integrations, potentially causing haploinsufficiency of tumor suppressor genes.

Other gene transfer vectors, such as those derived from spumaviruses (53), avian sarcoma leukemia virus (54), and transposons (55), display integration patterns potentially safer than those of γRVs and, for some vectors, even LVs. Further development of these platforms may establish their clinical potential.

Overall, considering our present findings together with those recently reported in other studies (21, 18, 42, 50, 51, 56), the safest design for retroviral vectors to alleviate the risk of insertional mutagenesis combines a SIN LTR and a moderately active internal promoter and should be the preferred choice at least until new technologies for site-specific gene editing or addition are fully validated for clinical applications (57, 58). The experiments described in this study were performed using a neutral transgene to investigate the oncogenic potential of several intrinsic vector features. However, both the biological activity of the transgene and the influence of the disease on target cell biology may significantly affect the oncogenic risk of vector treatment.

Methods

Mice. Wild-type FVB/N.129 mice were obtained from Charles River. FVB.129-Cdkn2a−/− mice were obtained from the National Cancer Institute Mouse Models of Human Cancers Consortium. All mice were bred and kept in a dedicated pathogen-free animal facility and were killed when they showed signs of severe sickness. All procedures were performed according to protocols approved by the Animal Care and Use Committee of the San Raffaele Institute (IACUC 225 and 320) and communicated to the Ministry of Health and local authorities according to Italian law.

Vector production. SFFV LTR enhancer/promoter sequence contained in a 415-bp EcoRV/BamHI DNA fragment from pHR55IN-CSGW17 (59) was blunt-cloned into the 7300-bp EcoRI/BamHI DNA fragment from pCCL-SIN.cPPT.hPGK.eGFP.wPRE (7) to replace the hPGK promoter and generate SIN.LV.SF. To generate the LV.SF.LTR construct, the 7300-bp EcoRI/BamHI DNA fragment from pCCL5IN.cPPT.hPGK.eGFP.wPRE was religated to generate pCCL5IN.cPPT.eGFP.wPRE (without the hPGK promoter). The same 415-bp EcoRV/BamHI DNA fragment containing the SFFV LTR enhancer/promoter sequence was blunt cloned into the BssI site in the U3 LTR region of pCCL5IN.cPPT.eGFP.wPRE, thus generating the LV.SF.LTR.

Concentrated LV stocks pseudotyped with the VSV.G envelope were produced by transient cotransfection of 4 plasmids in 293T cells and titered on HeLa cells as described (12). γRV stocks were similarly produced and titered using prkar43.3 PGK.GFP (13), pSRS.SF91 (31) (kindly provided by C. Baum, Department of Experimental Hematology, Hannover Medical School, Hannover, Germany) as transfer plasmids, pCMV-gagpol (MLV) (60), and VSV.G envelope encoding pMD2.VSV.G plasmid.

Isolation and transduction of hematopoietic progenitors. Six-week-old Cdkn2a−/− mice were killed by CO2 inhalation, and BM was harvested by flushing femurs and tibiae with PBS-2% FBS (FBS; Invitrogen). Lin− cells were purified by lineage-marker–negative selection using the Enrichment of Murine Hematopoietic Progenitors Kit (StemCell Technologies), plated at a density of 1 × 106 cells/ml, and cultured in StemSpan SFEM expansion medium (StemCell Technologies) with a cytokine cocktail composed of 100 ng/ml SCF, 100 ng/ml thrombopoietin, 100 ng/ml Flt3 ligand, and 20 ng/ml IL-3 (PeproTech).

After 24 hours prestimulation, Lin− cells were split and subjected to mock, LV, or γRV transduction (105 TU/ml or 106 TU/ml). After 12 hours, cells were washed and resuspended in the original medium and reinfected at 48 hours. Overall, cells were kept in culture for 96 hours before transplant. A sample of cells was kept for 14 days in culture to assess GFP expression by FACS analysis and genomic DNA extraction procedures.

Transplantation procedures, FACS, and histopathology. All transplantation procedures were performed as previously described (7). In brief, 6-week-old wild-type female FVB mice were lethally irradiated and injected in the tail vein with 7.5 × 106 cells/mouse. For secondary transplant, sublethally irradiated 6-week-old FVB female mice (5.75 Gy) were injected in the tail vein with marrow cells collected from tumor-infiltrated primary recipients (2 × 106 cells/mouse).

FACS analysis was performed using lineage-specific antibodies (BD Biosciences — Pharmingen) against CD11b, CD19, CD3, and the IgG isotype control on cells obtained from blood collected 6 weeks after transplant or from blood, BM, spleen, thymus, and lymph nodes of diseased mice and analyzed with FCS Express 3 software (De Novo Software) as previously described (7). Dead cells were excluded by 7-aminocanthocinomycin D staining (5 μg/ml); blood erythrocytes were lysed with 7% ammonium chloride ( StemCell Technologies).

For histological analysis, H&E staining was performed on 4-μm-thick sections of formalin-fixed, paraffin-embedded tissues. Specimens were evaluated in blinded fashion independently by 2 investigators (F. Sanvito and M. Ponzoni) as previously described. The semiquantitative scoring system was as follows: 0, no pathological infiltration; 1, mild infiltrates; 2, moderate infiltrates;
and 3, heavy infiltrates. For CD3, antigen immunolocalization was performed using rat anti-human CD3 (AbD Serotec) on formalin-fixed, paraffin-embedded 4-μm-thick sections after antigen retrieval with microwave using Tris-EDTA, pH 9. The immunoreaction was revealed by biotinylated-conjugated anti-rat antibody (Vector Laboratories) and HRP-conjugated streptavidin and using 3,3′-diaminobenzidine (DAB) as chromogen (BioGenex).

*VCN analysis.* Genomic DNA was extracted from cells (cultured in vitro for 14 days after transduction) was purified using the QIAGEN blood and cell culture DNA kit (QIAGEN). Genomic DNA from total BM or spleen or thymus of mice was purified using the QIAGEN tissue DNA kit (QIAGEN). Q-PCR analysis was performed as described (7), with probes complementary to mouse genomic β-actin (common to wild-type and Cdkn2a−/− mouse genomic DNA), GFP (common to all vectors tested), and neomycin resistance sequences (specific for Cdkn2a−/− cells) (See Supplemental Methods for information on oligonucleotides and probes used). Engraftment level was determined as the ratio between neomycin resistance and β-actin quantifications, using the DNA of a Cdkn2a−/− mouse as the standard curve. VCN was determined as the ratio between the relative amounts of GFP (common to all vector types) and the calculated engraftment levels. A GFP standard curve was made using DNA dilutions from a homozygous GFP transgenic mouse (61). Reactions were carried out according to the manufacturer’s instructions and analyzed using the ABI Prism 7900 HT Sequence Detection System (PE, Applied Biosystems).

**LAM-PCR and genomic integration site analysis.** We used 0.5–5 ng of tumor DNA and 10–100 ng of transduced pretransplant DNA, respectively, as template for LAM-PCR (62). LAM-PCR was initiated with a 25-cycle linear PCR and restriction digest using Tsp509I or HpyCH4IV and ligation of a restriction site–complementary linker cassette. The first exponential biotinylated PCR product was captured via magnetic beads and reamplified by a nested second PCR. LAM-PCR primers for LV were previously described (4, 34, 63, 64). For the yRv LAM-PCR, 2 5′-biotinylated oligonucleotides complementary to the GFP sequence (LAMGFP1, 5′-TGGACTGGCTACTTGAAGGCT-3′ and LAMGFP2, 5′-GGGATCACCTCCTGGCATGAC-3′) were used for the linear amplification step. The 2 sequential exponential amplification steps were performed with nested oligonucleotides complementary to the yRV LTR sequence (LAMRV1, 5′-GACTTGTTGCTCCTGCTTTG-3′ and LAMRV2, 5′-GGTCTCCTCGTATGTTAC-3′), each coupled with the previously described oligonucleotides complementary to the linker cassette (34). LAM-PCR amplicons were separated on Spredex gels (Elchrom Scientific) and the excised bands cloned into the TOPO TA vector (Invitrogen) or PCR-purified (QIAGEN), shotgun cloned, and sequenced (GATC Biotech). Sequences were aligned to the mouse genome (Mus musculus genome assembly February 2006, UCSC Mus musculus genome version 8) using the UCSC BLAT genome browser (http://genome.ucsc.edu). Identification of the nearest gene was performed with dedicated PERL scripts. Visualization of the RTCGD CIS integrations as a feature on the UCSC BLAT output was achieved by connecting to UCSC through the RTCGD web interface (http:// rtcgd.abc.unc.edu/), map position of each retroviral and SB CIS integrations were automatically loaded as custom tracks on the UCSC BLAT search engine. Analysis of overrepresentation of gene classes in integration data sets was performed with DAVID-EASE software 2.0 (http://david.abcc.ncifcrf.gov/home.jsp). For each of the 4 data sets, we restricted further analysis to the GO categories significantly overrepresented, identified by at least 3 genes in the Biological Process or Molecular Function systems and with a fold increase of 3 or more. The murine gene IDs in our data sets were converted to the respective human putative orthologs by searching the Ensembl database (http://www.ensembl.org/biomart/martview). Functional networks analyses were performed with IPA 6 software (Ingenuity Systems). Only significantly overrepresented classes identified by at least 3 genes were considered.

**RNA isolation and gene expression analysis.** Total RNA from tumor-infiltrating BM cells or spleen was isolated with the RNAsesy Mini Kit (QIAGEN). cDNA preparation was performed using Mo-MLV reverse transcriptase and random hexamer primers (SuperScript III First-Strand Synthesis System for RT-PCR; Invitrogen). cDNA was used as template for TaqMan Gene Expression Assays specific to each gene (Applied Biosystems) (see Supplemental Methods for gene expression assay ID and probe sequences used). Amplification reactions were performed on a 7900HT Real-Time PCR Thermal Cycler (Applied Biosystems). The relative expression level of each gene was calculated by the ΔΔCt method (65), normalized to Hprt expression (housekeeping gene control), and represented as fold change relative to mock-transduced samples (calibrator). Real-time PCR Miner software (http://www.miner.eiwindup.info) (66) was used to calculate the mean PCR amplification efficiency for each gene. The qBase software program (http://www.biogazelle.com) was used to measure the relative expression level for each gene (67). Amplification of the chimerical transcript between LVSF.LTR and Braf was obtained using oligonucleotides complementary to the lentiviral LTR (LTR 5′-CCTGCAACGACGACTAGATG-3′) and exon 22 of the Braf gene (BrafF5 5′-GACTGTCGATTCTGAAGGGC-3′). Statistics. Statistical analysis was performed with the R-statistical (version 2.1.1; http://www.r-project.org) or GraphPad Prism 4.0 software (GraphPad Software). For survival analysis, a parametric approach was used because smooth continuous estimates of the survivor function were necessary for predictive purposes as described (7). The formula describing the log-logistic distribution is as follows: \( P(\tau) = k \cdot e^{\rho \cdot t} \), where \( t \) is the time; \( \rho \) is the log-logistic parameter that determines the effect of the treatment. The log-logistic distribution provided the best fit to our experimental data (Supplemental Statistical Methods). Time of death and VCN were obtained from LV.SF.LTR, MOI = 10; LV.SF.LTR, MOI = 100; RV.SF.LTR; SIN.LV.SF; SIN.LV.PGK (7); and SIN.RV.PGK groups were used as variables in the log-logistic accelerated failure time model. The exclusion or inclusion of the SIN.LV.PGK survival data from our previous work (7) did not significantly change the results of our statistical analysis. For all statistical comparisons, \( P < 0.05 \) was considered significant.

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