Promoterless Transposon Mutagenesis Drives Solid Cancers via Tumor Suppressor Inactivation

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Abstract: A central challenge in cancer genomics is the systematic identification of single and cooperating tumor suppressor gene mutations driving cellular transformation and tumor progression in the absence of oncogenic driver mutation(s). Multiple in vitro and in vivo gene inactivation screens have enhanced our understanding of the tumor suppressor gene landscape in various cancers. However, these studies are limited to single or combination gene effects, specific organs, or require sensitizing mutations. In this study, we developed and utilized a Sleeping Beauty transposon mutagenesis system that functions only as a gene trap to exclusively inactivate tumor suppressor genes. Using whole body transposon mobilization in wild type mice, we observed that cumulative gene inactivation can drive tumorigenesis of solid cancers. We provide a quantitative landscape of the tumor suppressor genes inactivated in these cancers and show that, despite the absence of oncogenic drivers, these genes converge on key biological pathways and processes associated with cancer hallmarks.

Simple Summary: Over the past two decades, there have been many published studies reporting high-copy SB transgenic lines in which the transposon allele includes both gene-trap and internal promoter elements to drive tumorigenesis in vivo. Cancer gene discovery from end-stage solid tumors in these studies performed by our labs and others has found few proto-oncogenic insertions. However, the question remains whether these tumors are initiated by SB insertions in proto-oncogenes to promote permissible phenotypes for tumor initiation, which become dispensable for tumor maintenance. Here, our study expands upon these indirect observations to demonstrate that high-copy SB transposon alleles designed with only gene-trap elements that inactivate genes (Onc2.3) can drive tumor initiation, progression, and maintenance to end-stage tumors in the absence of sensitizing mutations.
Keywords: Sleeping Beauty transposon mutagenesis; tumor suppressor driver genes; Cul3; Rasa1; Trip12; Ras signaling; E3 ubiquitin ligase; cancer hallmarks

1. Introduction

Recent genomic analysis of physiologically and histologically normal tissues such as eyelid epidermis and esophageal squamous epithelia show that these tissues tolerate relatively high levels of mutations, typically within known tumor suppressor genes [1–3]. Sleeping Beauty (SB) insertional mutagenesis [4] is a powerful forward genetic tool used to perform genome-wide forward genetic screens in laboratory mice for cancer gene discovery [5–15]. We recently conducted and reported an SB screen to model the development of cutaneous squamous cell carcinoma in vivo and noted that approximately 30% of tumors formed without any oncogenic transposon insertions, albeit with extended latency [16,17]. This finding raises two questions: (i) Can exclusive loss of tumor suppressor genes per se drive tumorigenesis? (ii) How do the kinetics of tumorigenesis in this context compare to tumors that develop via alterations in both tumor suppressor and oncogenes?

In this study, we generated a series of new SB transposon alleles to explore the hypothesis that cumulative loss of tumor suppressor genes is sufficient to drive the initiation and progression of systemic tumorigenesis. To this end, we modified key features of an existing high copy pT2/Onc2 transposon allele [7] and engineered a new transposon construct that functions as a gene trap to inactivate gene expression. Whole body transposition of this modified high-copy, gene-trap transposon (pT2/Onc2.3 or SB-Onc2.3) via inducibly expressed SB transposase was sufficient to initiate and progress a variety of tumor types in vivo. Here, we prioritized solid tumors of the skin, lung and liver and employed high-throughput SBCapSeq approaches [9,11] to identify genome-wide SB mutations and define recurrently mutated, statistically significant candidate cancer drivers (CCDs) from bulk tumors containing more SB mutations expected by chance using the SB Driver Analysis [18] statistical framework. We provide a quantitative genetic landscape of the transposon insertions in these tissues and demonstrate that Ras signaling and members of the ubiquitin ligase complex are frequently inactivated, suggesting key roles of these biological processes and pathways during the initiation and progression of solid cancers in the absence of selected oncogenic events.

2. Results

2.1. Harnessing an SB Gene-Trap Allele for Forward Genetic Screens

2.1.1. Whole-Body Mutagenesis Using High-Copy, Gene-Trap Only SB Transposons in Wild Type Mice

We derived a novel gene-trap-only SB transposon allele, pT2/Onc2.3 (also known as pT2/SB-GT-MBM102, Figure 1a), with two alterations to the pT2/Onc2 [7] transposon vector. First, the MSCV 5′LTR promoter and splice donor sequences were removed to disable the ability of the transposon to activate gene expression upon integration into the promoter or intron regions of a gene. Second, an additional bi-directional SV40-polyA signal sequence was added to enhance the transcriptional termination ability of the transposon upon integration into genes. These changes shortened the Onc2.3 SB transposon cargo to ~1.6 kb (Figure 1a), which matches the natural size of the original fish SB [19] and optimum size to maximize transposition in mammalian cells [20]. Collectively, these modifications result in an SB transposon that can only disrupt gene expression via inactivation and facilitate the identification of inactivated tumor suppressor genes that drive tumor initiation and progression in vivo.
Figure 1. Whole-body, gene-trap exclusive SB transposon mutagenesis results in tumor formation in SB-Onc2.3 mice. (a) Comparing Sleeping Beauty transposons used in this study (T2/SB-GT and T2/Onc3) and in a similar companion study (T2/Onc2). Abbreviated allele designations for founder lines used in this work are shown to the right of each vector. (b) Kaplan–Meier survival plots comparing experimental and control SB-Onc2.3 cohorts (Mantel–Cox log-rank test, \( p < 0.0001 \), Figure S1). Mice in all SB-Onc2.3 cohorts with active SB mobilization developed solid tumors at multiple organ sites, however SB-Onc2.3-TG.14922 mice (orange line) had significantly decreased survival compared to the SB-Onc2.3-TG.14942 mice (purple line; Mantel–Cox log-rank test, \( p = 0.0015 \)). (c) Kaplan–Meier survival plots comparing the relative tumor latency between various SB cohorts with whole-body mobilization by a conditionally activated from the \( \text{Rosa26-} \text{LSL-SBase} \) allele to the \( \text{Rosa26-} \text{1lox-SBase} \) allele by whole body Beta-actin-driven Cre expression, including SB-Onc2 mice (combined data from TG.6070 and TG.6113 alleles containing an MSCV-promoter [9]), SB-Onc3 (data from TG.12740 allele containing a CAG-promoter [17]), and SB-Onc2.3 (combined data from TG.14922 and TG.14922 alleles containing no promoter) (Mantel–Cox log-rank test, \( p < 0.0001 \), Table 1).
Table 1. SB allele-specific tumor spectrum by whole-body, transposon-mediated mutagenesis in wild type mice.

| SB Cohort          | SB-Onc2.3 | SB-Onc3 | SB-Onc2 |
|--------------------|-----------|---------|---------|
| Promoter Reference | no promoter | CAG promoter | MSCV promoter |
| Tumor classification | tumors mice | tumors mice | tumors mice |
| Skin Squamous Cell Carcinoma | 5 4 (3%) | 52 20 (24%) | 0 0 (0%) |
| Skin Keratoacanthoma | 1 1 (1%) | 9 8 (10%) | 0 0 (0%) |
| Hepatocellular Carcinoma | 2 2 (2%) | 9 9 (11%) | 0 0 (0%) |
| Hepatocellular Adenoma | 4 3 (2%) | 79 42 (50%) | 0 0 (0%) |
| Adenoma, Multiple | 25 16 (12%) | 39 31 (37%) | 0 0 (0%) |
| Adenocarcinoma, Multiple | 3 3 (2%) | 9 8 (10%) | 0 0 (0%) |
| Carcinoma, Multiple | 0 0 (0%) | 6 6 (7%) | 0 0 (0%) |
| Sarcoma | 0 0 (0%) | 6 6 (7%) | 0 0 (0%) |
| Papilloma | 1 1 (1%) | 4 3 (4%) | 0 0 (0%) |
| Astrocytoma | 34 34 (26%) | 3 3 (4%) | 0 0 (0%) |
| Hemangiosarcoma | 0 0 (0%) | 3 3 (4%) | 0 0 (0%) |
| Hemangioma | 1 1 (1%) | 2 2 (2%) | 0 0 (0%) |
| Fibrosarcoma | 0 0 (0%) | 2 2 (2%) | 0 0 (0%) |
| Metastasis | 0 0 (0%) | 2 2 (2%) | 0 0 (0%) |
| Mast Cell Tumor | 0 0 (0%) | 1 1 (1%) | 0 0 (0%) |
| Trichoepithelioma | 0 0 (0%) | 1 1 (1%) | 0 0 (0%) |
| Schwannoma | 1 1 (2%) | 0 0 (0%) | 0 0 (0%) |
| Hepatoblastoma | 1 1 (1%) | 0 0 (0%) | 0 0 (0%) |
| Medulloblastoma | 0 0 (0%) | 0 0 (0%) | 6 6 (8%) |
| Lymphoma | 1 1 (2%) | 2 2 (2%) | 28 28 (36%) |
| Leukemia | 0 0 (0%) | 0 0 (0%) | 78 78 (100%) |
| Histiocytic Sarcoma | 0 0 (0%) | 8 8 (10%) | 0 0 (0%) |
| Totals | 79 107 | 237 87 | 112 78 |
| Average tumors per mouse | 0.74 | 2.72 | 1.44 |
| Average tumor latency (days) | 450 | 330 | 70 |

Distribution of cancer types between promoter-driven (SB-Onc3, SB-Onc2) and promoter-less (SB-Onc2.3) transposons [21].

After pT2/Onc2.3 plasmid linearization (Figure S1a) and microinjection into pronuclei, 56 of the resulting 101 live-born and weaned progeny were screened by Southern blot and found to carry at least one copy of the transposon transgene (Figure S1b). We reasoned that mice with the highest transgene copy number possible would allow for a maximal number of independent integration events per cell, a strategy that facilitates identification of cooperating TSGs in malignant transformation. Initially, five transgenic lines were selected to test for germ-line transmission by backcross breeding to C57BL/6J mice (Figure S1b). All five lines transmitted to offspring, but only three T2/Onc2.3 lines, TG.14913, TG.14922, and TG.14942, segregated a transposon concatemer as a discreet locus, suggesting they arose from a single event leading to high-copy concatemer integration into the respective donor genomes.

The three SB-Onc2.3 transgenic lines contain approximately the same number of copies as other T2/Onc2 high-copy lines, with estimated copy number of ~400 per transgene array (Figures S2a,b and S3a,b), and each line could be maintained as transgenic homozygotes without loss of viability, fertility, or other obvious phenotypes. However, similar to what was observed with high-copy pT2/Onc2 carrier mice [7], when combined with a constitutively active SBase allele all three T2/Onc2.3 transgenic lines had loss of viability with statistically significant reduced numbers of live-born double heterozygous carriers (Figure S4a,b). To circumvent lethality in TG.14922 and TG.14942 lines, we used a conditional LSL-SBase and Actb-Cre allele strategy [9,17] that fully restored viability in the resulting triple heterozygous carrier mice (Figure S4c–e). When combined with conditional LSL-SBase and Actb-Cre alleles, T2/Onc2.3 transgenic carrier mice (hereafter, SB-Onc2.3) ex-
hibited high rates of whole-body transposition that resulted in tumor formation (Figure 1b and Figure S5a–c).

2.1.2. Systemic, Whole Body Onc2.3 Transposition Extends Latency of Tumor Development

We generated 107 SB-Onc2.3 (n = 58 Actb-Cre | Onc2.3-TG.14942; n = 49 Actb-Cre | Onc2.3-TG.14922) and 23 control (Actb-Cre | Onc2.3-TG.14922) mice and aged them for a maximum of 550 days (1.5 years). At the predetermined endpoint, while only two control mice developed masses, many of the Onc2.3 developed tumors (Figure 1b). Tumor-free survival of both SB-Onc2.3 cohorts was significantly reduced compared to control cohorts (Kaplan–Meier log-rank test, \( p < 0.0001 \), Figure 1b). Tumor latency was significantly reduced in the TG.14922 cohort compared with the TG.14942 cohort (Kaplan–Meier log-rank test, \( p < 0.0001 \), Figure 1b) despite having indistinguishable tumor spectrums. Some of the mice censored in the survival analysis were observed to have various cancers when necropsied and/or histologically evaluated, suggesting our data represent a lower limit to the full tumor spectrum in SB-Onc2.3 mice.

The combined SB-Onc2.3 cohort exhibited a significantly longer mean tumor latency (Log rank test, \( p < 0.0001 \); Figure 1c) compared to similarly bred wild type SB-Onc2 [9] or SB-Onc3 [17] cohorts. The tumor latency and spectrum of SB-Onc2.3 mice were more similar to SB-Onc3 [17] mice, with few hematopoietic tumors and a similar wide variety of solid tumor types, compared with SB-Onc2, including cutaneous squamous cell carcinomas (cuSCC)/keratoacanthoma (cuKA) and hepatocellular adenomas (HCA)/carcinomas (HCC) (Table 1 and Table S1). Unexpectedly, SB-Onc2.3 mice exhibited a significantly larger proportion of early-stage, low-grade solid tumor types, including spinal cord and cerebellar astrocytomas (ACT) and lung alveolar adenoma (LUAA) and adenocarcinoma (LUAC) (Figure S6 and Table S1). However, with respect to transposon copy number, the average number of SB insertions observed per tumor genome in SB-Onc2.3 mice was more similar to SB-Onc2 compared with low-copy SB-Onc3 (Figures S3 and S4), suggesting that copy-number per se does not explain the differences in latency and tumor spectrum.

To better understand the genetic events driving these SB-Onc2.3 solid tumors, we generated genomic libraries from a wide variety of SB-Onc2.3-induced solid cancers and used high-throughput SBCapSeq [9,11] to identify genome-wide SB insertions. Similar to previous reported results in SB-Onc2 [9] and SB-Onc3 [17], the SB-Onc2.3 libraries were highly reproducible (Figure S7 and Table S2).

2.2. Solid Tumors Driven by TSG Inactivation in SB-Onc2.3 Mice

2.2.1. Landscape of cuSCC Driven by Transposon-Mediated Tumor Suppressor Inactivation

Skin tumors comprised 4% of the total number of tumors collected and histologically verified in SB-Onc2.3 mice, suggesting that inactivation of tumor suppressor genes is sufficient to drive keratinocyte initiation and progression to frank cuSCC in vivo. We recently reported in an SB-driven cuSCC model that activating SB insertions into Zmiz1, Zmiz2, or Mamld1 oncproteins are collectively observed in over two-thirds of cuSCC tumors sequenced using SBCapSeq [17]. This suggests that the initiation and progression of approximately one-third of skin tumors in vivo do not require positive selection for proto-oncogenes. We hypothesized that tumors without activating oncogenic insertions in the Zmiz1 or Zmiz2 genes would have higher frequencies of inactivation of the tumor suppressor genes found in the Onc2.3 system. We used SBCapSeq to quantitatively define the gene insertions in 6 SB-Onc2.3 skin tumors with confirmed cuSCC diagnosis (Tables S3 and S4) [9]. As expected, no activating SB insertions greater than background rates (\( \geq 1 \) SB-CapSeq read) were observed in the proto-oncogenes Zmiz1, Zmiz2, or Mamld1 previously reported in our SB-Onc3-driven cuSCC or cuKA cohorts [17]. In contrast, the SB-Onc2.3 cuSCC tumors had recurrent, high read-depth inactivating insertions in the Rasa1, Trip12, Cul3, Cux1, Tcf12, Nf1, Kmt2c, Ncoa2, and Crebbp genes (Figure 2). Pathway analysis using Enrichr software [22,23] and KEGG database revealed enrichment of selected SB insertions...
in genes associated with MAPK signaling pathway (Abl2, Nf1, Rasa1, Stk4; \( p = 0.003 \)) and Ras signaling pathways (Nf1, Rasa1, Stk4, Tgfb2; \( p = 0.008 \)). Based on these data, we revisited our previously reported sequencing data from the SB|Onc3 cuSCC tissues [17]. Indeed, we observed that Zmiz1/2-negative cuSCC tumors had dramatically higher inactivating SB insertions in Trip12, Kmt2c, Rasa1, Cul3, and Arid1b in ZMIZ1/2-negative cuSCC tumors relative to ZMIZ1/2-positive cuSCC genomes (Figure 2). Notably, almost all (95%, 22/23) of the Zmiz1/2-negative cuSCC contained single or cooperating mutations in those five genes (Figure 2), compared to 63% (27/43) of the ZMIZ1/2-positive cuSCC genomes. While the frequencies of individual gene hits within the Zmiz1/2-negative cuSCC genomes derived from either Onc3 or Onc2.3 SB transposon lines varied, similar cooperating relationships among the putative TSGs were conserved (Figure 2). We conclude that the Onc2.3 transposon-derived skin tumors develop via cooperating inactivation of tumor suppressor genes. This supports our prior observation that approximately one-third of SB-driven skin masses are not driven by activating SB insertions in proto-oncogenes [17].

2.2.2. Landscape of Lung Cancer Driven by Transposon-Mediated Tumor Suppressor Inactivation

Approximately one-third of lung adenocarcinomas contain oncogenic mutation in KRAS [24], and lung tumorigenesis is usually modeled in mice using oncogenic mutant Kras alleles as initiating events [25]. Using SB mutagenesis, we successfully modeled lung alveolar adenoma (LUAA) and adenocarcinoma (LUAC), demonstrating that oncogenic mutant Kras, or other strong oncogenic drivers, is not required for lung tumor initiation in the mouse. To dissect the selected SB events driving early lung LUAA/LUAC tumorigenesis, we applied the SBCapSeq methodology to analyze the insertion profiles of tumors driven by the transposons Onc2.3 (Table S5) or Onc3 (Table S6; previously reported in [17]), which can either inactivate alone or activate and/or inactivate gene expression, respectively, upon transposon insertion. In the Onc3 lung LUAA/LUAC tumors, 70% had directional, activating insertions in Rasgrf1, encoding the guanine nucleotide exchange factor RASGRF1 which functions to activate Ras signaling by exchanging GDP for GTP (Figure 3, Figure S8a,b, and Table S6). Results from 454 sequencing confirmed activating Rasgrf1 insertions among the 29 Onc3 LUAA/LUAC tumors, including 10 large tumors from flash frozen genomic DNA which were also sequenced by SBCapSeq, and 19 additional smaller tumors isolated from FFPE tumor tissues (Figure S9 and Tables S7–S9). In contrast, no insertions in Rasgrf1 were observed in SB-Onc2.3 LUAA/LUAC tumors. Several genes including Rbms3, Sik3, Cul3, Trip12, Rock1, Zfp292, and Rasa1 were more commonly inactivated in Onc2.3 compared to Onc3-driven lung tumors. Pathway analysis using the Enrichr KEGG pathway analysis and gene ontology biological processes enrichment analysis revealed statistically significant enrichment of genes with SB insertions associated with focal adhesion pathway (Rock1, Rasgrf1, Pten, and Pik3r1; \( p < 0.0001 \)), Ras signaling pathway (Rock1, Rasgrf1, and Pik3r1; \( p < 0.0001 \)), and regulation of protein ubiquitination (Cul3 and Trip12; \( p = 0.001 \)). Notably, Cul3 and Trip12—members of the E3 ubiquitin ligase complex—were exclusively inactivated in the Onc3 but not Onc3-driven lung tumors. Among the targets of TRIP12 is ASXL1, a tumor suppressor with roles in Polycomb-induced gene silencing. Two SNVs in Asxl1 were identified in mouse lung adenomas induced by MNU in the background of a KrasG12D mutation [26]. Izumchenko et al. described mutations in ASXL1 and other genes involved in DNA damage and chromatin remodeling identified in the lungs of human patients with atypical adenomatous hyperplasia, an initiating event in adenocarcinoma development [27]. RBMS3 (RNA Binding Motif Single Stranded Interacting Protein 3) is a validated TSG in human lung SCC where it acts primarily via regulation of c-Myc [28]. CUL3 is a promiscuous E3 ubiquitin ligase that partners with multiple proteins to regulate protein turnover of many targets in a context specific manner. CUL3 plays an important role in ubiquitination of KEAP1, a negative regulator of the transcription factor NRF2 which enacts a transcriptional program important in protecting lung cancer and other tumor types from oxidative stress [29]. These data suggest that, in the Onc3-driven lesions, activation of Rasgrf1 contributes to rapid lung tumorigenesis, while, in Onc2.3-driven lung
tumorigenesis, the inactivation of cooperative tumor suppressor genes drives disease after long latency, potentially through modulating protein turnover and transcriptional reprogramming. Further studies are needed to show whether these mechanisms may represent functional redundancy with key pathways initiated by oncogenic *Kras* to promote lung cancer in mice.

| Genetic Alteration | Onc2.3 | Onc3 |
|--------------------|--------|------|
| Trip12             | 83%    | 29%  |
| Tcf12              | 67%    | 60%  |
| Cul3               | 67%    | 29%  |
| Rere               | 67%    | 29%  |
| Vps13b             | 67%    | 52%  |
| Rasa1              | 67%    | 41%  |
| Zfp148             | 67%    | 12%  |
| Hnrnpm             | 56%    | 12%  |
| Ptprc              | 56%    | 24%  |
| Adgrb3             | 56%    | 0%   |
| Akap13             | 56%    | 41%  |
| Cnot1              | 56%    | 29%  |
| Cnot2              | 56%    | 29%  |
| Cux1               | 56%    | 12%  |
| Dnen1b             | 56%    | 35%  |
| Dopp10             | 56%    | 18%  |
| Uttn               | 56%    | 18%  |
| Yhdc1              | 56%    | 47%  |
| Ctc                | 56%    | 6%   |
| Fam168b            | 56%    | 18%  |
| Kanz1l             | 56%    | 0%   |
| Bmpr2              | 56%    | 29%  |
| Nf1                | 56%    | 18%  |
| Kmt2c              | 33%    | 24%  |
| Ncoa2              | 33%    | 58%  |
| Crefb              | 33%    | 18%  |
| Notch1             | 33%    | 18%  |
| Picalm             | 33%    | 18%  |
| Diap2              | 33%    | 47%  |
| Foxj3              | 33%    | 60%  |
| Fto                | 33%    | 6%   |
| R33h1              | 33%    | 24%  |
| Aida               | 33%    | 6%   |
| Arnt               | 33%    | 12%  |
| Ebf1               | 33%    | 29%  |
| Zfhx4              | 33%    | 12%  |

**Figure 2.** SB-driven cuSCC by cumulative tumor suppression. Recurrent inactivating SB insertions in Onc2.3-induced cuSCC genomes (n = 6). Comparative waterfall plots of trunk driver genes from *Zmiz1/Zmiz2*-independent SB-Onc2.3 and SB-Onc3 (n = 23 (left)) and *Zmiz1/Zmiz2*-dependent SB-Onc3 (n = 43 (right)) cuSCC tumors. (c) *Zmiz1/Zmiz2*-independent tumor segregated by SB cohort, SB-Onc3 (n = 17 (left)) and SB-Onc2.3 (n = 6 (right)).
Figure 3. SB-driven lung cancer by cumulative tumor suppression. Recurrent inactivating SB insertions in Onc2.3 (n = 6 using SBCapSeq) and Onc3 (n = 10 using SBCapSeq; n = 29 using 454T sequencing) induced LUAA genomes.

2.2.3. Landscape of Liver Cancer Driven by Transposon-Mediated Tumor Suppressor Inactivation

We previously generated hepatocellular adenomas (HCAs) and adenocarcinomas (HCC) driven by the SB-Onc3 transposon in the presence of mutant or wild type Trp53 alleles [18,30]. From 95 histologically confirmed HCAs (Table S2), we identified 43 trunk drivers, including proto-oncogenes Hras, Kras, and Rli1/Rian and tumor suppressor genes Pten, Adk, and Zbtb20 (Tables S10–S12). Most of these genes were identified in Trp53 mutant and wild type cohorts, suggesting that their role in driving HCA is Trp53-independent. Further, the majority of the putative driver genes identified in our HCA tissues were also reported in SB-driven hepatocellular carcinoma (HCC) [8,31–34], suggesting that many of the initiating genes enriched during earlier stages of hepatocyte transformation in vivo are maintained in HCC (Figure S10). SBCapSeq analysis of the four HCA and two HCC tumors from SB-Onc2.3 mice revealed no activating insertion patterns in the genomes from these tumors and the absence of hits in Hras, Kras, and Rlt1/Rian (Table S13). However, we corroborated the tumor-suppressive roles of insertions in Adk and Zbtb20 and in Nipbl, Pdlim5, Ppy1r12a, Tarc6b, Brd4, Cul3, Cttna3, Elavl1, Gphn, Nfia, Ptpn12, Taok3, and Rasa1 (Figure 4).

Using Enrichr KEGG pathway analysis and gene ontology biological processes enrichment analysis revealed statistically significant enrichment of genes with SB insertions associated with ubiquitin mediated proteolysis (Cul3, Cul5, and Trip12; \( p < 0.0001 \)), growth factor signaling (Ep300, Itgb1, and Rasa1; \( p < 0.0001 \)), scaffolding proteins involved in cell adhesion (Dlg1, Itgb1, and Pdlim5; \( p < 0.001 \)), and transcriptional regulators (Brd4, Med13l, Nfia, and Nipbl; \( p < 0.0001 \)). Notably, the inactivation of RASA1 suggests that the Onc2.3 transposon may activate Ras signaling via inactivation of the negative regulator of this pathway, in the absence of oncogenic Ras activation. Taken together, by using different transposon systems, we show that HCA in mice can be initiated by activation of oncogenes in combination with tumor suppressor inactivation or exclusively by inactivation of cooperating tumor suppressors that converge on similar mechanisms co-opted to promote tumorigenesis.
Figure 4. SB-driven liver cancer by cumulative tumor suppression. Recurrent inactivating SB insertions in Onc2.3 (n = 6 using SBCapSeq) and Onc3 (n = 52 Kras/Hras/Rtl1 oncogene negative using 454T sequencing; n = 42 Kras/Hras/Rtl1 oncogene positive using 454T sequencing) induced HCA genomes.

2.2.4. Landscape of Other Solid Tumors Driven by Transposon-Mediated Tumor Suppressor Inactivation

Finally, we conducted SBCapSeq analysis on nine other SB-Onc2.3 mouse tumors from diverse tissues and histological classifications, including three ovarian tumors (with adenoma, papillary, and granulosa cell carcinoma subtypes), a uterine adenocarcinoma, a cerebellar astrocytoma, a colon polypoid adenoma, a preputial gland carcinoma, a renal tubular cell adenoma, and a Schwannoma of the cerebellum. Individually, all but one specimen demonstrated reproducible (Table S3) and clonal SB insertions sites, as evidenced by a range of low to very high read-depth-enriched TA-dinucleotide sites within the coding
region of many known tumor suppressor genes such as *Apc, Brd4, Cul3, Cux1, Gnaq, Nf1, Pten, Rasa1, Rnf43,* and *Wac* (Figure 5 and Tables S14 and S15). Notably, the astrocytoma sample presented no evidence of clonal selection, despite exhibiting the highest total number of SBCapSeq reads. This suggests that this mass may have other non-SB driven mechanisms contributing to tumor initiation (see Note S1). Collectively, these results strongly suggest an SB-driven etiology, consistent with the studies in skin, lung, and liver described above, and in multiple previously reported studies [9–11,13,14,17,18,30].

![Figure 5. SB-driven gynecologic and other cancer histologies by cumulative tumor suppression. Recurrent inactivating SB insertions in Onc2.3-induced ovary, uterus, Schwannoma, preputial gland, and colon tumors.](image)

### 2.2.5. Common Pathways Selected in Response to Whole Genome Tumor Suppressor Gene Inactivation

We observed *Brd4, Cul3,* and *Trip12* to be commonly mutated genes across different tumor types, suggesting that there may be common mechanisms perturbed by the Onc2.3 transposon to drive tumorigenesis in the absence of oncogenic events. Therefore, we extended our analysis to statistically define recurrent tumor suppressor genes across 27 SB-Onc2.3 tumors using SB capture sequencing [9] and SB Driver analysis [18] (Figure 6 and Tables S16–S18). We reasoned that, regardless of the tumor origin, a meta-analysis of selected SB insertion events enabled by the SB Driver Analysis [18] workflow may provide a quantitative means to explore the detection of potentially novel, cooperative TSGs. We identified 1499 discovery-significant progression drivers, 476 genome-significant progression drivers (Table S17), and 53 genome-significant trunk drivers (Table S18) using the SB Driver Analysis statistical categorical framework [18] (see Materials and Methods). As expected, none of the sense-strand-oriented, recurrent SB insertions present in defined proto-oncogenes from tumors driven by SB transposons containing an internal promoter (see the SBCDDB [18,30]) were identified in any of the SB-Onc2.3 tumors. Further, meta-analysis using SB Driver Analysis, including all SB-Onc2.3 (n = 27) and SB-Onc3 (n = 10) lung tumors, showed that *Rasgyf1* was the only gene identified with an activating SB insertion pattern (Table S19).
lung tumors, showed that Rasgrf1 was the only gene identified with an activating SB insertion pattern (Table S19).

Figure 6. SB-Onc2.3-Pan-TSG driver discovery. Recurrent inactivating SB insertions into trunk drivers with >99 reads from 3 or more Onc2.3-induced cancer genomes (n = 27). SB-Onc2.3-Pan-TSG driver gene SB insertion profiles are shown on the right. Literature references summarizing studies that provide in vivo evidence for validation of the TSGs identified in this study (Table S23) [35].

The Cancer Gene Census (COSMIC v91 [36–38]) contains 313 genes with a designation of TSG and a unique mouse ortholog (Table S20). The Onc2.3-Pan-TSG dataset contains 1499 driver genes. We observed 78 genes in common between the CGC and the Onc2.3-Pan driver genes, which represent a statistically significant enrichment of shared genes than expected (Table S20; Chi-square with Yates’ correction, χ² = 104.742, two-tailed p < 0.0001). This suggests that the genes identified in our screen are enriched for known TSGs that driver tumorigenesis. Selective and statistically significant enrichment of the Onc2.3-Pan-TSG dataset was also observed with the 1004 TSGs listed in the Tumor Suppressor Gene Database [39,40] with a unique mouse orthologs. We observed 127 genes in common between the TSGdb-orthologs and the Onc2.3-Pan driver genes, which represent a statistically significant enrichment of shared genes than expected (Table S21; Chi-square with Yates’ correction, χ² = 32.662, two-tailed p < 0.0001). Overall, 167 known TSGs were identified in the Onc2.3-Pan-TSG dataset, suggesting that the additional 1333 remaining genes may represent potential novel TSG drivers (Figure 7), including Trip12, Rbm33, and Zfp292, with recurrent SB insertion sites in Onc2.3 tumors demonstrated insertion patterns predicted to inactivate target gene expression. Indeed, although not represented on either of the curated
CGC or TSGdb lists, several genes (e.g., Cep350 [11], Ncoa2 [17,33], and Usp9x [41]) have all been observed in other SB tumor screens and experimentally validated to be TSGs within mouse models and/or human cancer cells.

To gain further insights into novel pathways that lead to tumor initiation and/or progression via cooperating TSG only routes, we performed Enrichr biological pathway enrichment analysis [22,23] using the 1499 Onc2.3-Pan-TSG-Drivers. We observed over-representation of the Onc2.3-Pan-TSG-Drivers among the top 1% of all known biological processes with statistically significant enrichment in known pathways involved in cancer initiation and progression, including AR, EGF/ERGR, TGF-beta, PDGFB, mTOR, and VEGF signaling pathways, ubiquitin mediated proteolysis, chromatin modifying enzymes and organization, and RHO GTPase effectors (Table S22). Similarly, Enrichr gene ontology enrichment analysis reaffirmed our manual curation, namely that genes with roles in diverse processes, including protein ubiquitination, cadherin binding, focal adhesion, and chromatin, are over-represented among the top 1% of all known biological processes (Table S22). We conclude that this set of known and suspected TSGs represents a rich resource for prioritizing future efforts to catalogue the genes in the cancer genome that can drive tumor initiation and/or progression without oncogenic mutations.

3. Discussion

In this study, we investigated whether in vivo tumor initiation and progression was possible solely by inactivation of tumor suppressor genes. We generated a new SB transposon allele that lacks the internal promoter present in all SB transposons used to-date in mouse models of cancer [6–8,18,30,42]. Here, we demonstrate that the Onc2.3 transposon restricts SB mutagenesis to act as a gene trap, selectively inactivating genes. While SB genetic screening studies have reported tumor suppressor genes in mouse models of cancer, these screens have often been conducted using strong oncogenic initiating mutations,
including \textit{Kras}^{G12D} \cite{10,41}, \textit{Braf}^{V600E} \cite{11,43}, \textit{Apc} loss \cite{14,44}, \textit{Pten} loss combined with \textit{Blm} loss \cite{45}, or HBV expression \cite{31}. Similar studies using the \textit{piggyBac} transposon have been combined with \textit{Blm} loss \cite{46}. Importantly, we demonstrate that our Onc2.3 transposon system enables the evaluation of tumorigenesis in vivo by exclusive loss of tumor suppressor genes without introduction of an initiating oncogene \cite{47,48}. Notably, whole-body mutagenesis using the Onc2.3 transposon resulted in distinct cell types that were more susceptible to tumorigenesis than others. This may be due to the fact that exclusive tumor suppressor gene inactivation leads to a longer tumor latency in some tissues, or that some cell types in the mouse have robust mechanisms to withstand tumorigenesis without an exogenous or induced oncogenic hit. This latter phenomenon has been well-documented in the literature, particularly for mouse models of human cancers driven by oncogenic sensitizing events, including melanoma, pancreatic and colon cancers \cite{10,11,14,18,30}, notably absent from the observed tumor types with the Onc2.3 promoterless transposon.

Mutagenesis methods to study skin tumorigenesis typically rely on inducing point mutations that target both tumor suppressors and oncogenes \cite{49–51} or insertional mutations without point mutations \cite{9,11}. In cuSCC tumors driven by the SB-Onc2.3 transposon, we noted a high frequency of gene inactivation in chromatin remodelers. This suggests that in the absence of an oncogenic hit, inactivation of chromatin remodeler genes can drive cuSCCs, presumably due to their dynamic ability to regulate expression of genes, miRNAs and lncRNAs involved in different biological processes and pathways intimately involved in tumorigenesis. Indeed, inactivation of chromatin remodeler genes has been shown to drive the initiation and progression of many different cancers \cite{52–57}. We recently reported functional evidence that shRNA-mediated knockdown of \textit{KMT2C} or \textit{CREBBP} in immortalized primary human keratinocytes was sufficient for cell transformation in vitro but was insufficient to promote tumor growth in vivo, suggesting that cuSCC development requires additional cooperating oncogenic events \cite{17}. This is supported by the recent findings of Ichesi et al. who reported that heterozygous loss of \textit{Crebbp} alone, or together with heterozygous loss of its paralog \textit{Ep300}, cooperates with mutant Ras to drive keratinocyte hyper-proliferation in vivo \cite{58}. Furthermore, a recent genetic screen demonstrated that inactivation of \textit{Fbxo7} or \textit{Sic9a3} could transform mouse embryonic fibroblasts cells in vitro within the context of oncogenic \textit{Kras}^{G12D} but was not sufficient to form tumors in vivo \cite{59}. Our SB-Onc2.3 is also of relevance to human cuSCC, as the majority of patients are older individuals, and recent studies indicate that cancer driver mutations are tolerated in physiologically normal squamous tissues, even in relatively younger individuals \cite{1,2}. This is consistent with the observation in both this study (using SB-Onc2.3) and our previous study (using SB-Onc3) \cite{17} that cumulative tumor suppressor inactivation over time may be an important and under recognized route to keratinocyte transformation initiation and cuSCC progression.

In both cuSCC and liver tumors, inactivation of the \textit{Rasa1} gene, encoding a RAS-inhibiting GTPase, was a frequent, recurrent event. \textit{Rasa1} inactivation was observed in a previous SB study investigating resistance mechanisms to FGFR inhibition in invasive lobular breast cancer \cite{60}. Our data suggests that Ras-mediated MAPK-ERK signaling may play a role in driving both cuSCC and hepatocellular adenocarcinoma. In support of this, the Onc2.3-driven cuSCCs also exhibited inactivation of \textit{Nf1}, encoding a negative regulator of Ras signaling important in many cancer types. Mutations in \textit{RASAI} and \textit{NF1} co-occur in human non-small cell lung cancer \cite{61}, which may reflect independent mechanisms by which \textit{RASAI} and \textit{NF1} inhibit Ras or the clonal diversity within these tumors that preferentially selects one of these mechanisms.

The majority of the putative driver genes identified in HCA driven by Onc2.3 were also reported in SB-driven hepatocellular carcinoma (HCC) \cite{8,31–34}, supporting the role for these events in HCC initiation. While we did not identify SB inactivation of \textit{Ncoa2} and \textit{Sav1} in our tumors, these genes were previously identified in liver cancer mouse models driven by SB mutagenesis in the presence of Myc activation or \textit{Pten} loss \cite{33,62}. It is likely that initiating oncogenic events influence the evolutionary routes of tumor development,
dictating the selection and maintenance of cooperating drivers. The extent to which context-dependent SB insertion events impact tumor development remains to be determined. Our HCA/HCC data suggests that while tumors initiated by an oncogenic mutation may exhibit shorter tumor latencies than tumors lacking exogenous initiating oncogenic events and may have distinct subsets of inactivated genes, both populations converge on similar signaling pathways. Importantly, the frequent development of cuSCC and HCA using the Onc2.3 SB transposon demonstrates that tumor suppressor gene inactivation is sufficient for de novo transformation of both keratinocytes or hepatocytes in vivo.

When all the Onc2.3 tumors were analyzed together, we noted that the negative regulators of the Ras signaling pathway such as Rasa1 and Nf1 were commonly inactivated in liver and cuSCC, while genes encoding members of the E3 ubiquitin ligase such as Trip12 and Cul3 were frequently inactivated in all tumor types analyzed. Additionally, in cuSCC tumors without activation of the Zmiz proto-oncogenes, the frequency of Trip12 and Cul3 inactivation were markedly increased, relative to tumors with Zmiz oncogene activation. The E3 ubiquitin ligase-proteasome have diverse roles, including regulating the Ras, MAPK, and PI3K/Akt signaling pathways, gene expression and cell death [63]. This range of biological processes may explain why Trip12 and Cul3 inactivation is frequently observed in the Onc2.3 tumors. Indeed, a previous Sleeping Beauty screen by Dorr et al. using the Onc2 transposon system also found inactivation in Cul3 [64]. The authors functionally validated the tumor suppressive role of CUL3 in A549 and H522 human lung cancer cell lines and showed that shRNA knockdown of this gene increased proliferation rates, relative to control. The second E3 ligase associated gene Trip12 observed in our screen has been shown to be essential in the regulation of the cell cycle [65]. Furthermore, we note that the genes discussed here were also observed in an earlier in vivo recessive screen that modeled systemic tumor development in the context of Pten and Blm loss sensitized background [46]. Since our screen was conducted in the absence of any sensitizing mutations, this observation suggests that there are core pathways that are necessary for tumor development in various anatomic sites.

Our Onc2.3 model is not without limitations. First, in cuSCC tumors, it is not clear why we did not observe inactivation of tumor suppressors frequently described in human squamous epithelium such as TP53 and NOTCH1. Inactivation of NOTCH1 has been postulated to drive the evolutionary dead-end of nascent pre-cancerous population of cells [66]. This is based on the observation that the frequency of NOTCH1 mutations decreases in esophageal and skin cancers, relative to physiologically normal epithelial tissues [1,3,66]. Consistent with this, in our previously reported Onc3 screen, we noted that the frequency of Notch1 inactivation was decreased in cuSCC versus keratoacanthoma tissues, suggesting that Notch1 loss is not essential for cuSCC development [17]. While we did not observe inactivation of Trp53, our previous Onc3 screen suggests that there were no obvious differences in the insertion profiles of tumors with either mutant or wild type Trp53 background, albeit the rate of tumorigenesis was significantly reduced in Trp53 mutant mice. Additionally, as noted in the SB Cancer Driver Database, the low frequency of transposon-driven Trp53 inactivation is not unique to our study, and has been observed in other SB screens reported to date [30]. Second, we noted that genes involved in chromatin remodeling were inactivated in all of the tumor types sequenced, albeit at different frequencies. It is possible that loss of these tumor suppressors may activate oncogene expression, thereby promoting tumorigenesis. However, given the significantly extended latency of tumor development in the Onc2.3 mice, it is unlikely that such genes confer strong advantage in clonal expansion, and the chromatin remodeler loss likely affects tumor suppressive processes. Similarly, without genomic sequencing analysis, we cannot exclude the possibility that spontaneous driver mutations that confer cooperating oncogenic and/or tumor suppressive alterations, are also contributing to the Onc2.3-driven tumors analyzed in this study. However, extensive genomic analyses of whole genome, exome or transcriptome from SB-driven melanoma [11,43], myeloid leukemia [9] and cuSCC [17] have revealed very few non-transposon derived alterations, including any
known or suspected oncogenic events. Last, in this study, we focused only on determining the transposon insertion profiles across various tumor types. The inactivation of genes involved in chromatin remodeling and protein ubiquitination suggests the necessity to profile the transposon mutagenesis-driven tumors on other platforms (e.g., transcriptome, proteome) to gain a more comprehensive understanding of the molecular events driving tumorigenesis.

Collectively, our data strongly implicate routes to tumorigenesis via cooperative inactivation of tumor suppressor driver genes alone and preclude the necessity of oncogenic events. The dramatic reduced penetrance and increased latency of skin, lung, and liver tumors when subjected to systemic gene-trap only transposon mutagenesis highlights three important points: (i) gene inactivation per se is sufficient to drive de novo transformation in vivo of keratinocytes, lung alveolar epithelium, and hepatocytes and tumor initiation in vivo; (ii) transformation by gene inactivation alone is likely to be the exception rather than rule as penetrance of proto-oncogenic driven cuSCC, LUAC, and HCA is substantially higher; and (iii) cuSCC development has no absolute requirement for oncogenic drivers (e.g., activating ZMIZ or RAS oncoproteins) and can occur via cumulative inactivation of tumor suppressors, which, in either instance, converge on similar altered cancer hallmark signaling pathways. Thus, we conclude that the SB-Once2.3 model delivers a novel in vivo platform that allows systemic gene inactivation in a non-sensitized background and provides a heterogeneous background that promotes evolutionary routes to tumorigenesis.

4. Materials and Methods

4.1. Generation of pT2/Onc2.3 Vector

SB transposon vector pT2/Onc2 [7] (a gift from A. Dupuy) was sequentially digested with restriction enzymes Xmal (NEB, Boston, MA, USA), at position 717 bp, and EcoRI (NEB) at position 729 bp; at 37 °C for 30 m, each, per manufacturer’s suggested protocol, to linearize the plasmid. Linear plasmid was visually confirmed as an ~4.9 kb on a 1% agarose TAE gel stained with EtBr viewed under UV light. A small piece of agarose gel containing the linear pT2/Onc2 was excised, purified using GFX columns (Amersham Biosciences, Marlborough, MA, USA, #414581), and saved for ligation. A PCR cloning strategy was used to insert an extra bi-directional SV40-polyA termination sequence into the pT2/Onc2 vector. Using 100 µM each of two complimentary overlapping (bold) oligos, Xmal $\rightarrow$ GAGA TGG CTT, EcoRI $\rightarrow$ TGG AAT, T2/Onc2 vector was sequenced with the manufacturer’s suggested protocol. Ligation products were electroporated into MAX efficiency DH10B competent cells (Invitrogen #18297010), plated on LB agar plates with ampicillin, and incubated 37 °C for 18 h. Miniprep DNA from selected trans­ductants was sequentially digested with Xmal (NEB) and EcoRI (NEB), at 37 °C for 30 m each, per manufacturer’s suggested protocol. The ligation product was run on a 2% agarose TBE gel stained with EtBr viewed under UV light, and a small piece of gel containing the 108 bp fragment was excised, purified using GFX columns (Amersham Biosciences #414581) prior to ligation. Finally, a ligation reaction using a 1:5 ratio of pT2/Onc2:Xmal-EcoRI

Commented [MBM5R4]: Yes, they denote the restriction cut sites for the enzymes listed, respectively

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Commented [MBM5R4]: Yes, they denote the restriction cut sites for the enzymes listed, respectively
linear vector and SV40-polyA: Xmal-EcoRI insert, respectively, was performed using T4 DNA Ligase at room temperature for 30 min, followed by heat inactivation at 65 °C for 20 min. Ligation products were electroporated into MAX efficiency DH10B competent cells (Invitrogen #18297010), plated on LB agar plates with ampicillin, and incubated at 37 °C for 18 h. Miniprep DNA from selected transformants was digested with HindIII (NEB), at 37 °C for 60 min each, per manufacturer’s suggested protocol. Recombinant pT2/Onc2.2 vector was identified by the presence of three fragments at 3.658 kb, 0.8 kb, and 0.6 kb fragments. pT2/Onc2.2 vector DNA was cut with restriction enzyme Ascl (NEB #R0558S), at 37 °C for 60 min, to linearize the plasmid. Linear plasmid was visually confirmed on a 1% agarose TAE gel stained with EtBr viewed under UV light. A small piece of agarose Hin dIII (NEB), at 37 °C for 60 min, was digested with HindIII (NEB), at 37 °C for 60 min, using the manufacturer’s suggested protocol. Recombinant pT2/Onc2.2 vector was identified by the presence of three fragments at 3.658 kb, 0.8 kb, and 0.6 kb fragments.

4.2. Generation of T2/Onc2.3 Transgenic Mice

pT2/Onc2.3 DNA was linearized with restriction enzyme with Scal (NEB #R3122) at position 3,502 bp (Figure S1a) at 37 °C for 90 min, followed by heat inactivation at 80 °C for 20 min. pT2/Onc2.3:Scal linear plasmid, at 2, 4, and 10 ng/μL, was prepared for microinjection into (B6C3)F2 hybrid embryos using standard techniques [7]. Tail biopsy genomic DNA from founder animals was digested with Drai (NEB) and BamHI (NEB), run on a 0.8% TAE agarose gel at 30 v for 16 h, transferred to membrane, and screened by Southern blotting using a 32P-labeled En2-SA (splice acceptor) probe, a 500 bp En2-SA PCR product with pT2/Onc2.3 template and primers T2.3-En2.5Probe, 5′-GCTGCAATAAACAAAGTTGGGCGC-3′ and T2.3-En2.3Probe, 5′-CTTGAGGTCAACATTTAAGGTAGCC-3′, and standard meth-
ods. Individual transgenic lines (n = 5) were established by backcrossing founders to C57BL/6J (JAX, Bar Harbor, ME, USA, #000664). Germ line transmission was confirmed by Southern blotting using standard techniques. All five lines transmitted to offspring, but only three lines, T2/Onc2.3(TG.14913) (TgTn(sb-T2/Onc2.3)14913Mbm), T2/Onc2.3(TG.14922) (TgTn(sb-T2/Onc2.3)14922Mbm), and T2/Onc2.3(TG.14942) (TgTn(sb-T2/Onc2.3)14942Mbm), segregated a transposon concatemer as a single, discreet locus by Southern blot genotyping. Subsequent offspring were genotyped by multi-plex PCR using primers specific to the SB transposon T2.3-sense, 5′-CTG TCA GGT ACC TGT TGG TCT GAA AC-3′ and T2.3-antisense, 5′-CCT CAA GCT TGG GTG CGT G-3′, which yields a 370 bp PCR product, and control primers Actb-sense, 5′-ACA AGG TCA AAA CTC AGC AAC AAG T-3′ and Actb-antisense, 5′-GCT GAG AGG GAA ATT GTT CAT TAC A-3′, which yields a 700 bp PCR product. Multiplex PCR genotyping protocol for SB-Onc2.3 mice may be downloaded from figshare http://dx.doi.org/10.6084/m9.figshare.12811301 [68].

The following alleles were used to construct the SB-driven mouse model of multiple solid tumor histologies: Actb-Cre (FVB/N-Tg(ACTB-cre)2Mrt/J [69]); Trp53flox/+ (FVB.129P2-Trp53tm1Brn/Nci [70]); Trp53LSL-R172H/+ (129S4-Trp53tm2Tyj/Nci [71]); T2/Onc2 (TG.12740) (TgTn(sb-T2/Onc3)12740Njen [8]); and Rosa26-LSL SBase or SBaseLSL (Gt(ROSA)26Sor(tm2(sb11)Njen [44]). The resulting cohorts of mice were on mixed genetic backgrounds consisting of C57BL/6J, 129, C3H and FVB. Genotyping by PCR assays with primers specific to the alleles was performed. No sample size estimate was used. The production and characterization of all SB and SBase mouse strains was supported by the Department of Health and Human Services, National Institutes of Health, and the National Cancer Institute. Mice were bred and maintained in accordance with approved procedures directed by the respective Institutional Animal Care and Use Committees in the National Cancer Institute Frederick National Lab, A*STAR Biological Resource Centre, and H. Lee Moffitt Cancer Center and Research Institute/University of South Florida (IACUC studies: IS00008185, IS00007898, IS00006747, IS00003681, IS00002945). Gross necropsies were performed and all masses were documented and prepared for subsequent analysis. Both sexes were used for experiments, including reporting of cohort data survival analysis. No randomization or blinding was performed, all mice were assigned to groups based on genotype.

4.3. Histological Analysis

Histological analysis of necropsy tissues was performed on 5-µm sections of formalin-fixed, paraffin-embedded (FFPE) samples stained with hematoxylin and eosin (H&E). Nuclear staining of SB transposase (SBase) was confirmed by immunohistochemistry on FFPE tissues after antigen retrieval (pH 9) and endogenous peroxidase inhibition followed by overnight incubation with mouse antibody to SBase (anti-SBase; R&D Systems, Minneapolis, MN, USA; pH 9; 1:200 dilution), and then by incubation with primary antibody; chromogen detection (with HRP polymer, anti-rabbit or anti-mouse, with Envision System from Dako, Carpinteria, CA, USA) and hematoxylin counterstaining were performed per manufacturer’s instructions. Genomic DNA (gDNA) was isolated from flash frozen or FFPE necropsy specimens using Gentra Puregene DNA isolation kit (Qiagen, Germantown, MD, USA) protocol for tissue using the manufacturer’s protocol (see Note S2). Source histology image files may be downloaded from figshare https://doi.org/10.6084/m9.figshare.12816305 [72].

4.4. Mapping SB Insertion Sites

Full details for the SBCapSeq protocol, an enhanced SBCapSeq method optimized for sequencing from solid tumors [73], will be published elsewhere (Mann, Mann et al., in preparation; for general protocol and concept, see also [9,74]). Briefly, for selective SB insertion site sequencing by liquid hybridization capture, gDNA (0.5 µg per sample) of bulk tumor genomes was used for library construction using the AB Library Builder™ System (Thermo Fisher Scientific), including random fragmentation and ligation of bar-coded Ion Xpress™ sequencing adapters (Thermo Fisher Scientific). Adapter-ligated
templates were purified by Agencourt® AMPure® beads (Beckman Coulter, Indianapolis, IN, USA) and fragments with insert size of ~200 bp were excised, purified by Agencourt® AMPure® beads, amplified by 8 cycles of adapter-ligation-mediated polymerase chain reaction (aLM-PCR), and purified by Agencourt® AMPure® beads with elution in 50 µL of TE (1× Tris-Ethylenediaminetetraacetic acid [EDTA], pH8). Capture hybridization of single or multiplexed up to 12 barcoded libraries (60 ng per sample) was performed using custom xGen® Lockdown® Probes (IDT, Coralville, Iowa, UAS; full details available at https://doi.org/10.35092/yhjc.11441001) [73]. All 120-mer capture and blocking oligonucleotide probes were purchased from IDT as UltraD N A Oligos with Standard Desalting. Captured library fragments were further amplified by 12 cycles of aLM-PCR and run using an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, UAS) or TapeStation (Agilent Technologies) to estimate enrichment. Captured libraries were quantified by Qubit® Fluorometer (Thermo Fisher Scientific) and quantitative Real-Time-PCR (qRT-PCR) were used to dilute libraries for template preparation and Ion SphereTM Particle (ISP, Thermo Fisher Scientific) loading using Ion Chef™ System (Thermo Fisher Scientific) and sequencing on the Ion Proton™ platform (Thermo Fisher Scientific) with PL13 semiconductor wafer chips per manufacturers recommended instructions. High-throughput sequencing of up to 39 multiplex captured libraries was carried out per PL13 chip to achieve an average of 4.5 million reads per barcode. Reads containing the transposon IRDR element were processed using the SBCapSeq bioinformatic workflow as described [9]. For splink_454 sequencing (see Note S3), SB insertion reads were generated by 454 GS Titanium sequencing (Roche) of pooled splinkerette PCR reactions with nested, barcoded primers was performed as described previously [10,11,75]. Pre- and post-processing of 454 reads to assign sample DNA barcodes, filter out local hopping events from donor chromosomes, and map and orient the SB insertion sites across the entire nuclear genome of the mouse was performed. Donor chromosomes SB insertions were filtered away prior to identification of common insertion sites using SB Driver Analysis [18]. Using SB insertion site density to infer the likely donor integration site, we identified chromosome 1 for both TG.14922 and TG.14942 (data may be downloaded from figshare https://doi.org/10.6084/m9.figshare.12816191 [76].

4.5. SB Driver Analysis Discovery, Progression and Trunk Drivers

Using the SB Driver Analysis framework [18], Discovery significant progression drivers are defined as statistically significant recurrently altered genes using False Discovery Rate (FDR; Benjamini–Hochberg procedure) multiple hypothesis testing p-value correction and Genome significant progression drivers are defined as statistically significant recurrently altered genes using Family Wise Error Rate (FWER; Holm–Bonferroni procedure) multiple hypothesis testing p-value correction from all SB insertion events within a dataset (no read depth filtering) to capture the widest possible list of potential driver genes. In contrast, Genome significant trunk drivers are defined as statistically significant recurrently altered genes using FWER multiple hypothesis testing p-value correction from SB insertion events with a pre-determined minimum read depth filtering of the dataset to report a statistically stringent prioritized driver gene list. Trunk drivers have the highest probability to be represented recurrently across tumors with the highest read depths, suggesting early and/or highly selected insertion events during bulk tumor growth. As a rule, Progression Dare a subset of Discovery Drivers, and Trunk Drivers are often also Progression and/or Discovery Drivers [18]. The BED files containing all SB insertions from histologically verified SB-Onc2.3 tumor specimens were filtered to consider only insertions represented by 100 or more reads in three or more tumors. When genes from a specimen genome were found to contain >1 SB insertion in the same RefSeq gene, only the SB insertion with the highest sequence read count was used for SB Driver Analysis calculations. SB Driver Analysis [18] was performed to identify the genes significantly enriched for insertions at high read counts with a FWER-corrected p < 0.05 were termed trunk driver genes based on the inferred high clonality across the tumor population [18].
4.6. qRT-PCR

Total RNA was purified and DNase treated using the RNeasy Mini Kit (Qiagen). Synthesis of cDNA was performed using SuperScript VILO Master Mix (Life Technologies). Quantitative PCR analysis was performed on the QuantStudio 12K Flex System (Life Technologies) or 7900HT Sequence Detection System (Applied Biosystem). All signals were normalized to the levels of GAPDH TaqMan probes. TaqMan probes were obtained from Life Technologies.

4.7. Software

Unless otherwise noted, bioinformatic analysis pipelines, report generation, and figure visualization were performed using bash, R, Python scripts, Vector NTI, and GraphPad Prism 8 software (Version 8.1.1).

4.8. URLs

TgTn(sb-T2/Onc2.3) 14942Mbm, http://www.informatics.jax.org/allele/MGI:644175
5; TgTn(sb-T2/Onc2.3)14922Mbm, http://www.informatics.jax.org/allele/MGI:644175
7; TgTn(sb-T2/Onc2.3)14913Mbm, http://www.informatics.jax.org/allele/MGI:6441760;
Mouse Genome Informatics (MGI) database, http://www.informatics.jax.org/; Git(ROSA)26
Sortm2(sb11)Njen, http://www.informatics.jax.org/allele/MGI:3839796; FVB/N-Tg(ActB-
cre)2Mrt/J, https://www.jax.org/strain/003376; FVB/129P2-Trp53tm1Brn, https://www.
jax.org/strain/008462; 12954-Trp53tm2tyj, https://www.jax.org/strain/002101; Enrichr,
http://amp.pharm.mssm.edu/Enrichr/; Tumor Suppressor Gene Database, https://bioinfo.
uth.edu/TSGene/; UniProt, http://www.uniprot.org/; COSMIC Cancer Gene Census,
https://cancer.sanger.ac.uk/census/; VENNY 2.1, https://bioinfogp.cnb.csic.es/tools/
venny/; Oncoprinter, http://www.cbioportal.org/oncoprinter.

5. Conclusions

Based on our study, high-copy SB transposon alleles designed to only contain elements that inactivate genes (Onc2.3) can drive tumor initiation, progression, and maintenance to end stage tumors in the absence of sensitizing mutations.

Supplementary Materials: The following are available at https://www.mdpi.com/2072-6694/13
2/225/s1, Notes S1–S3: Promoterless Transposon Mutagenesis Drives Solid Cancers via Tumor
Suppressor Inactivation [77], Tables: Promoterless Transposon Mutagenesis Drives Solid Cancers via Tumor Suppressor Inactivation [78], Table S1: Tumor incidence and subgroup classifications by cohort groups, Table S2: Sequencing projects summary by cohort, Table S3: Specimen metafile data for projects sequenced using SBCapSeq protocol with Ion Torrent Proton sequencer, Table S4: BED file of SB insertions for Skin-Onc2.3_SBC cohort, Table S5: BED file of SB insertions for Lung-Onc2.3_SBC cohort, Table S6: BED file of SB insertions for Lung-Onc3_SBC cohort, Table S7: Specimen metafile data for projects sequenced using Splink_PCR protocol with 454 GS-FLX Titanium pyrosequencer, Table S8: BED file of SB insertions for Lung29_454, Table S9: SB Driver Analysis with discovery (FDR_hit = TRUE) and progression (FWER_hit = TRUE) for the Lung-Onc3_454 dataset (n = 29), Table S10: BED file of SB insertions for HCA95_454, Table S11: SB Driver Analysis with discovery (FDR_hit = TRUE) and progression (FWER_hit = TRUE) for the HAC-Onc3_454 dataset (n = 95), Table S12: Trunk SB Driver Analysis with discovery (FDR_hit = TRUE) and progression (FWER_hit = TRUE) for the HAC-Onc3_454 dataset (n = 95), Table S13: BED file of SB insertions for Liver-
Onc2.3_SBC cohort, Table S14: BED file of SB insertions for Other-Onc2.3_SBC cohort, Table S15: BED file of SB insertions for Ovary-Onc2.3_SBC cohort, Table S16: BED file of SB insertions for Onc2.3-Pan-Tumor-SBC27 cohort, Table S17: Discovery and progression SB Driver Analysis for Onc2.3-Pan-Tumor-SBC27 cohort, Table S18: Trunk SB Driver Analysis for Onc2.3-Pan-Tumor-SBC27. (Note: Six trunk drivers—Aox2, Crlf3, Dgkd, Fgfr2, Ilf3, and Zfp292—were manually removed from the dataset because supporting evidence from three related lung masses from a single animal provided insufficient evidence for tumor recurrence.), Table S19: Discovery and progression SB Driver Analysis for SB1 Lung combined Onc2.3 and Onc3 cohorts, Table S20: COSMIC v91 Cancer Gene Census TSG enrichment with SB-Onc2.3 Pan-tumor drivers, Table S21: TSGdb enrichment with SB-Onc2.3
Pan-tumor drivers, Table S22: Enrichr biological pathway and process analysis with SB-Onco2.3 Pan-tumor drivers, Table S23: Literature references summarizing studies that provide in vivo evidence for validation of the TSGs identified, Table S24: Rasgrf1 gene expression analysis of from SB-Onco3 Lung tumor genomes, Figure S1: Transgenic SB-T2/Onc2.3 founder mice, Figure S2: Estimating SB transposon concatamer copy number from different SB strains, Figure S3: Estimating SB transposon concatamer copy number from different SB strains, Figure S4: Embryonic lethality in double heterozygous mice, Figure S5: Overview of SB-induced tumor studies, Figure S6: Necropsy and histological analysis of SB-Onco2.3 tumors, Figure S7: Evaluating the reproducibility of SBCapSeq from bulk SB-Onco2.3 tumors, Figure S8: Activating SB insertions into the Rasgrf1 locus drive overexpression an oncogenic delta-N isoform, Figure S9: Landscape of candidate trunk drivers mutated in SB-induced LUA with Spltink_45T sequencing, and Figure S10: Landscape of candidate trunk drivers mutated in SB-induced HCA with Spltink_45T sequencing.

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