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Recombinant Adeno-Associated Viral Integration and Genotoxicity: Insights from Animal Models

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Currently, clinical gene therapy is experiencing a renaissance, with new products for clinical use approved in Europe and clinical trials for multiple diseases reporting positive results, especially those using recombinant adeno-associated viral (rAAV) vectors. Amid this new success, it is prudent to recall that the field of gene therapy experienced tragic setbacks in 1999 and 2002 because of the serious adverse events related to retroviral and adenoviral gene delivery in two clinical trials that resulted in the death of two patients. In both cases, the toxicity observed in humans had been documented to occur in animal models. However, these toxicities were either undetected or underappreciated before they arose in humans. rAAVs have been tested extensively in animals and animal models of disease, largely without adverse events, except for transient elevation in liver enzymes in some patients. However, a small but growing number of murine studies have documented that adeno-associated viral gene delivery can result in insertional mutagenesis. Herein, the aggregate data are reviewed from multiple murine studies where genotoxicity associated with rAAV treatment has been observed. The data emphasize the need for a proactive position to evaluate the potential risks and possible solutions associated with AAV-mediated gene therapy.

Keywords: AAV, gene therapy, insertional mutagenesis, genotoxicity, cancer, HCC.

INTRODUCTION

The current level of enthusiasm for gene therapy as a viable therapeutic approach to treat a variety of human diseases is unprecedented. No longer is gene therapy a largely academic undertaking as biotechs, pharma companies, venture capitalists, and investors become increasingly more involved in the field.1–3 However, after two highly publicized deaths occurring in two separate gene therapy clinical trials in the late 1990s and early 2000s, the future of gene therapy appeared bleak.4,5 The toxicity observed in these clinical trials was caused by immunogenicity to an adenoviral vector and then to insertional mutagenesis by a retroviral vector. These types of toxicities had been observed in animals,6,7 but the risk in humans was deemed to be minimal. Current viral vectors are designed to be less mutagenetic, as in the case of SIN lentiviruses, or are intrinsically less immunogenic than the those that caused severe adverse events in earlier clinical trials.

One such vector, recombinant adeno-associated virus (rAAV), is inherently non-pathogenic and less likely to result in insertional mutagenesis because the majority of vector genomes remain episomal after transduction.8,9 This viral vector is suitable for both localized and systemic gene delivery and is capable of transducing a variety of cell types. To date, numerous naturally occurring AAV serotypes have been isolated from a number of different species, and novel capsids have been genetically engineered.10 A number of rAAVs have been shown to exhibit tropisms for hepatocytes, myocytes, and neuronal cells.

AAV has several advantages over other commonly used viral vectors that have been adapted for gene delivery, such as retroviral, lentiviral, and...
DNA encapsulation. During the latent stage, the are integral to AAV genome replication and proper are DNA elements that are 145 bp in length and that flanked by inverted terminal repeats (ITR). The ITRs and assembly-activating protein (AAP), which are replication proteins (Rep), the capsid proteins (Cap), and multiple open reading frames coding for the genome is 4.7 kb in size and consists of three promoters included both latent and lytic phases. The AAV ge- location on human chromosome 19. AAV2 provirus preferential integrates into a specific does not integrate, vector genomes and transgene non-replicating episomes. Because rAAV largely integrates also means that the therapeutic transgene to the wild-type AAV precursor (reviewed by Daya and Berns). The life cycle of wild-type AAV includes both latent and lytic phases. The AAV genome is 4.7 kb in size and consists of three promoters and multiple open reading frames coding for the replication proteins (Rep), the capsid proteins (Cap), and assembly-activating protein (AAP), which are flanked by inverted terminal repeats (ITR). The ITRs are DNA elements that are 145 bp in length and that are integral to AAV genome replication and proper DNA encapsulation. During the latent stage, the AAV2 provirus preferential integrates into a specific location on human chromosome 19.

The wild-type AAV genome is substantially altered to create the rAAV genome used in gene therapy, and these alterations result in dramatic differences in physical characteristics between the wild-type AAV and rAAV. When rAAVs are genetically engineered to carry exogenous sequences, such as enhancers, promoters, transgenes, and polyA signals, most of the wild-type AAV genome is removed because AAVs have a packaging constraint of approximately 4.7 kb. While the Rep and Cap proteins are necessary for rAAV production, the AAV ITRs are the only sequences necessary for rAAV packaging of the regulatory and transgene sequences. Hence, rAAV integration into the host genome is predicted to be greatly reduced because the sequences coding for the native AAV proteins needed for integration are not present in the recombinant AAV. However, a low number of random integrations do occur following gene delivery using rAAV. The rAAV vectors’ reduced ability to integrate also means that the therapeutic transgene exists predominantly in the host cell nucleus as non-replicating episomes. Because rAAV largely does not integrate, vector genomes and transgene expression are lost over time because of the inability to replicate with the host cell’s DNA.

Unlike adenoviral and gamma-retroviral vec- tors, rAAV has been used in numerous clinical trials with no vector- or viral-related severe adverse events, other than transient transaminitis. A gene therapy pilot safety study using an adeno- virus to treat a patient with ornithine transcarba- mylase deficiency resulted in a fatality thought to be the result of a systemic inflammatory response to the adenoviral capsid. This type of severe adverse immune response has not been observed in clinical trials that utilize rAAV gene therapy, presumably because AAV is less immunogenic than adenovirus. In fact, only mild elevations of liver enzymes have been reported after systemic rAAV delivery in patients. Clinical trials that utilized gamma-retroviral mediated ex vivo gene transfer to treat X-linked severe combined immunodeficiency and Wiskott–Aldrich syndrome have reported insertional mutagenesis by the gene therapy vector, which caused leukemia in patients. Because the rAAV transgene exists in a predominately episomal state and integrates at a very low frequency following rAAV transduction, the chances of insertional mutagenesis by a rAAV transgene are greatly reduced in comparison with retroviral and lentiviral vectors, which usually integrate into the host genome. However, an increased incidence in hepatic carcinoma (HCC) in mice after rAAV gene delivery has been reported, and some of these studies have shown that inser- tional mutagenesis by rAAV was the most likely cause of the HCC. Herein, some of the studies, listed in Table 1, that observed or interrogated HCC formation after rAAV gene delivery are discussed. The key observations from each study are highlighted, and unresolved issues are discussed.

AAV AND HCC: AN OVERVIEW OF PUBLISHED STUDIES

Lysosomal storage diseases (LSDs) are a relatively large class of inherited metabolic disease that are typically caused by a deficiency in a soluble lysosomal enzyme. LSDs are ideal candidates for systemic gene therapy. They are simple, monogenic diseases, and some proportion of lysosomal enzymes can be secreted from corrected cells and taken up by distant cells through a receptor-mediated process known as cross-correction. Mucopolysaccharidosis type VII (MPSVII) is an invariably fatal LSD caused by a deficiency in the lysosomal enzyme, β-glucuronidase (GUSB). β-Glucuronidase is ubiquitously expressed, and its deficiency leads to the accumulation of undegraded glycosaminoglycans in most tissues, including the liver spleen, heart, kidney, bones, eyes, and brain. A spontaneously arising mouse mutant was discovered...
Table 1. Studies that observed or investigated HCC formation after rAAV gene delivery

<table>
<thead>
<tr>
<th>Loci with integration(s)</th>
<th>Genome</th>
<th>Enhancer/promoter</th>
<th>Time of associated with HCC</th>
<th>Publication AAV serotype configuration Transgene Route of delivery Dose</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>GUSB</td>
<td></td>
<td></td>
<td></td>
<td>Donsante et al. [2001] AAV serotype 1 ss CMV/CBA</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Donsante et al. [2007] ss CMV/CBA</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Bell et al. [2005] ss Alb, CBA, SV40, TBG, Z12*</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td>A1AT, EGFP</td>
<td></td>
<td></td>
<td></td>
<td>Lacer et al. [2006] ss AMPB/TBG</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Embury et al. [2008] ss CBA* (WPRE)</td>
<td>Neonatal 7.3 x 1013 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Li et al. [2011] ss APOE/hAAT</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Wang et al. [2012] ss APOE/hAAT</td>
<td>Neonatal 7.3 x 1013 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Rian et al. [2013] ss CMV/CMV</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Embury et al. [2008] ss CBA* (WPRE)</td>
<td>Neonatal 7.3 x 1013 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Li et al. [2011] ss APOE/hAAT</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Li et al. [2011] ss APOE/hAAT</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Walia et al. [2015] ss APOE/hAAT</td>
<td>Neonatal 1014 vg/kg</td>
</tr>
</tbody>
</table>

HCC, hepatic carcinoma; rAAV, recombinant adeno-associated virus.

at The Jackson Laboratory that has a single base pair deletion in the Gusb gene, is completely deficient in GUSB activity, and mimics most of the biochemical, histological, and clinical features of human MPSVII [34,35]. The MPSVII mouse has been used extensively in preclinical studies to determine the efficacy of enzyme replacement therapy, stem cell-mediated therapy, and gene replacement therapy.

One of the first systemic AAV-mediated gene therapy experiments was performed using MPSVII mice. [36,37] Intravenous injection of a first-generation AAV2 vector into newborn MPSVII mice resulted in widespread transduction, relatively high levels of human GUSB activity, and prevention of lysosomal storage material in most tissues of the body, including the brain. A follow-up study in the MPSVII mice showed that this approach led to persistent GUSB expression and resulted in dramatic improvements in bone development, retinal function, auditory function, body weight, and life-span. [38] However, it was discovered that a number of long-lived (>1 year) rAAV-treated MPSVII mice from this study developed HCC. [39] It was virtually impossible to determine the cause of the tumors in that study, since the experiment was not designed as a toxicity study, and the number of animals with HCC was relatively small. In addition, the tumor samples were negative for GUSB activity, and the rAAV genome could not be detected by polymerase chain reaction (PCR) using primers specific for the GUSB cDNA.

A larger follow-up study in newborn MPSVII and normal mice was performed in an attempt to replicate the findings of Donsante et al. and to determine the cause of the HCC. Intravenous injection of the same AAV2 vector resulted in 30–60% of both normal and MPSVII mice developing HCC at ≥13 months of age. [40] It was also shown that the incidence of HCC in long-lived MPSVII mice receiving bone-marrow transplantation and a transgenic mouse expressing ~20-fold greater than normal levels of GUSB was nearly zero. The most striking and unexpected finding was that four of the six tumors evaluated had the recombinant AAV vector integrated within a 6,000 bp region of the Rian locus on mouse chromosome 12. This region contains several genes and is rich in micro- and snoRNAs. Integration of the rAAV vector dysregulated this locus. Another interesting finding was that all of the integrants were rearranged such that the GUSB cDNA was deleted. This explains the lack of a PCR product and absent GUSB activity in the tumors examined in the original report. [39] Although rAAV integration into the Rian locus was associated with the development of HCC, a...
cause and effect relationship was suggested but not proven. However, a subsequent study showed that there was a high frequency of integration into the same locus following intravenous injection of an rAAV vector containing a promoter and enhancer combination (no transgene) flanked by sequences that are homologous to the Rian locus.\(^41\) Integration of the promoter/enhancer elements and subsequent dysregulation of this locus led to nearly 100% penetrance of the HCC phenotype. These data strongly suggest that disruption of this locus by integration of an rAAV vector containing a strong promoter/enhancer combination causes HCC in mice.

The development of HCC following intravenous injection of an AAV vector was still generally considered an isolated event, even though there were several additional reports of tumorigenesis following rAAV administration. Portal-vein injection of an rAAV vector containing the woodchuck hepatitis virus post-transcriptional regulatory element and the phenylalanine hydroxylase cDNA into the mouse model of phenylketonuria also resulted in a high frequency of HCC.\(^42\) The authors of this report concluded that the HCC was caused by expression of the hepatitis virus protein X-phenylalanine hydroxylase fusion protein. Although immunohistochemical analysis showed that the hepatitis virus protein X-phenylalanine hydroxylase fusion protein was expressed in the liver of experimental animals, the expression appeared to be limited to the normal liver tissue and not the tumor. In retrospect, these data suggest that the tumors were not caused by expression of the hepatitis virus protein X-phenylalanine hydroxylase fusion protein, rather a potential integration of a rearranged rAAV vector that did not express the transgene, but manifested genotoxicity similar to what was seen with the MPSVII mice and vectors.

A high incidence of HCC was also observed in ornithine transcarbamolase (OTC)-deficient mice following portal-vein injection of rAAV vectors encoding either OTC or LacZ.\(^43\) OTC-deficient mice are known to have a high incidence of spontaneous liver tumors. Statistical analyses of the various groups suggested that mice treated with the LacZ vector had an increased risk of HCC. A retrospective analysis of some of the samples from this study discovered numerous integration sites in the tumor tissue, including integrations into the Rian locus.\(^44\)

There is also evidence suggesting that systemic delivery of rAAV vectors does not lead to the development of HCC. In 2005, a large retrospective study of mice treated with rAAV did not find an association between rAAV treatment and HCC in mice.\(^45\) A second, relatively large study, involving 132 mice, specifically designed to address the issue of HCC following intravenous administration of rAAV vectors, showed no statistically significant increase in HCC in mice injected with an rAAV vector encoding clotting factor IX.\(^46\) However, the small number of HCCs observed in this study, five in total, four of which occurred in the rAAV-treated group, may not have been sufficient to detect a significant association. Nevertheless, this study also used integration profiling and gene expression analysis to demonstrate that the HCCs appeared free of clonal integration events, or gene expression changes related to rAAV integration.\(^46\) A separate study showed that hepatic tumors were observed in young adult mice following intravenous injection of a high dose (\(2 \times 10^{10}\) vp/mouse) of an rAAV vector expressing a factor IX cDNA containing the Padua mutation, but occurred at an insignificant rate of \(\sim 15\%\). Genomic characterization was not performed on the HCC in this study. One significant difference between studies has been the developmental timing of the injections. Donsante et al. injected the mice on postnatal day (PND) 1–2; Bell et al. and Li et al. injected young adult animals.\(^39,45,46\) These data suggest that mice injected during the newborn period may be more susceptible to the development of rAAV-associated HCC and, furthermore, that the HCC association was not mouse model dependent.

In a complex study using varied scAAV vectors, applied with and without the agent camptothecin, and/or partial hepatectomy, Rosas et al. assayed HCC formation in cancer-prone and immunodeficient mice, with the intention that genotoxic effects of rAAV may be magnified.\(^48\) While the authors did observe HCC formation, and even integration events into cellular oncogenes, the very high background rate of HCC formation and relatively low number of recovered integration events failed to resolve whether rAAV was carcinogenic.

The next set of studies confirmed what was initially observed by Donsante et al., specifically that mice treated in the neonatal period with high doses of rAAV had a tendency to develop HCCs later in life. As with earlier studies that have observed HCC after rAAV administration, the most recent studies were designed to assess efficacy not toxicity in mouse models of either Sandhoff disease (SD) or methylmalonic acidemia (MMA). In the case of SD, neonatal mice were injected via the superficial temporal vein with a relatively high dose (\(2.5 \times 10^{14}\) vg/kg) of an rAAV9 vector that used the CMV promoter to drive the expression of the HEXB cDNA,
Figure 1. (A) A genomic alignment showing an alignment of 31 syntenic vertebrate sequences around the *miR341* locus of the mouse, and the position of a small simple repeat within the gene. (B) A base pair resolution map shows the CGGT repeat with mapped AAV integrations indicated as colored dots, with the corresponding study. The relative conservation of each base was measured and depicted with a logo plot where the height is proportional to conservation and boldness indicates more sequences.
and followed for beneficial effects. At the experimental end point of 43 weeks, gross examination during necropsy revealed tumors in 8/10 neonatally rAAV9-HexB–injected control and SD mice; seven animals had liver tumors, and one had multiple lung tumors. Tumors were not observed in untreated mice, but it should be noted that untreated SD mice uniformly perished, typically by 15 weeks of age. In 2/7 of the liver tumors, integrations into the Rian locus were observed, while the single mouse that developed lung cancer was found to have an rAAV integration in fibroblast growth factor receptor 2 (Fgfr2), a gene known to have altered expression in cancer. Although this study lacked large numbers of control animals, the penetrance of HCC in the mice treated as neonates clearly implicates rAAV therapy as a predisposing factor to carcinogenesis.

The recent report by Chandler et al. is the largest mouse study to report rAAV genotoxicity to date. It used large numbers of mice that were treated with a variety of rAAV vectors and serotypes, or that were untreated. In this study, the colony-background incidence of HCC measured at <10% (n = 51 mice), whereas control (n = 24) and MMA mice (n = 24) treated in the neonatal period with a dose of $1 \times 10^{11}$ vg/pup of therapeutic or reporter rAAVs developed HCC at a rate of 50–75% by 2 years of age. This study confirmed the findings of several previous reports that identified clonal rAAV integrations in the Rian locus, with upregulation of proximally located micro-RNAs and genes as the likely cause of the HCC. Identifying insertional mutagenesis of the Rian locus as the cause of HCC was not a surprising finding because previous studies to identify genes involved in carcinogenesis have identified mutations in the Rian locus as a cause of HCC in mice.

This study also reported several unique findings with regard to rAAV toxicity and integration patterns. First, the risk of the HCC was found to be rAAV dose-dependent. Second, a majority of the causative HCC-rAAV integrations were mapped to Mir341, a locus within Rian. As depicted in Fig. 1A, a genomic alignment centered on Mir341 revealed several unusual features. The locus appears only to be present in rodents, and contains an embedded repeat. Upon closer inspection, this repeat contains a unique array of two simple quadranucleotide repeats (CCGT and GGCT), some of which are internally repeated, as well as 11 CpG dinucleotides, yet is flanked by conserved sequences (Fig. 1B). Most rAAV...
HCC integrations mapped into this tiny microRNA, and perhaps most importantly, one rAAV, which utilized an hAAT promoter and synthetic MUT allele (designated synMUT in the Fig. 1B), did not cause upregulation of proximally located miRNAs and genes or HCC, suggesting that rAAV vector design could influence genotoxicity.

The use of a sensitive integration-capture method that relied on high throughput sequencing (HTS) allowed for the detection of thousands of rAAV integration events from approximately 40 samples has afforded a limited assessment of integration preferences. To query integration preferences in a very direct fashion, the number of integrations in a locus was divided by the size of the locus as measured in base pairs in order to normalize for frequency. Approximately 149 of a total of 1,205 loci with an rAAV integration had multiple rAAV integrations (Supplementary Table S1; Supplementary Data are available online at www.liebertpub.com/hum). Table 2 shows the loci with the highest rAAV integration frequency, which ranges from $2.19 \times 10^{-3}$ to $4.40 \times 10^{-7}$ integrations per base pair. Of importance, and in contrast to previous studies that have suggested the rAAV integration is largely random, it is noted that rAAV integration frequencies varied significantly between different loci, suggesting an integration preference by rAAV (Table 2) not yet observed in smaller rAAV data sets. This analysis also suggests that the characterization of many more integration events will be needed to further define sites preferred for integration by rAAV.

## CONCLUDING REMARKS AND FUTURE DIRECTIONS

Given the recent observation that AAV integrations appear to be present in some human HCCs and, in subset, may be pathologically relevant, a careful reconsideration of possible genotoxicity mediated by rAAV in preclinical models is warranted as vectors are studied before entering the clinic. What is clear from a number of reports is that mice treated in the neonatal period with high doses of rAAV that contains strong promoter-enhancer elements, or vectors that target integration into Rian, have a very high chance of developing HCC, and in the HCC, remnants of the vector will be likely be present in Mir341. However, it needs to be emphasized that distinct, non-Mir341 rAAV integrations within Rian do occur and appear pathogenic. The results of the limited reanalysis of available integration data presented in Table 2 confirms that site preferences for rAAV do exist, especially when one considers that some view a common integration site (CIS) as a region of $\leq 30$ kb that harbors two or more integrations (the equivalent of $\geq 6.7 \times 10^{-5}$ integration per base pair). Many more rAAV integrations must be captured and analyzed in order to define rAAV CISs and to determine whether sequence motifs, chromatin states/marks, and vector characteristics influence integration preferences. A recent publication describing a hybrid AAV-piggyBac transposase delivery method might also add important information about whether AAV integration events can cause HCC.

Another area of future experimental need surrounds the precise characterization of rAAV integration events. A particularly relevant observation is that both sides of the integration junction have never been captured in any rAAV HCC study. Thus, it remains possible that rAAV integration might be associated with a complex rearrangement or even translocation, in addition to the possibility of vector concatamerization. Classical genomic cloning, tiling, and/or new methods may need to be

### Table 2. Genes that are frequent targets of rAAV integration from Chandler et al.

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Unique HCC integrations</th>
<th>Unique normal liver integrations</th>
<th>Total number of integrations in gene</th>
<th>Locus size (bp)</th>
<th>Integration frequency normalized to gene size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mir341 (Rian)</td>
<td>18</td>
<td>3</td>
<td>21</td>
<td>96</td>
<td>$2.19 \times 10^{-3}$</td>
</tr>
<tr>
<td>AC152083.1 (Rian)</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>87</td>
<td>$1.15 \times 10^{-2}$</td>
</tr>
<tr>
<td>Mir212</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>91</td>
<td>$1.0 \times 10^{-2}$</td>
</tr>
<tr>
<td>Alb</td>
<td>14</td>
<td>9</td>
<td>23</td>
<td>753</td>
<td>$4.33 \times 10^{-3}$</td>
</tr>
<tr>
<td>Igfbp1</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>284</td>
<td>$3.52 \times 10^{-3}$</td>
</tr>
<tr>
<td>Gm14719</td>
<td>1</td>
<td>0</td>
<td>11</td>
<td>1,845</td>
<td>$1.08 \times 10^{-3}$</td>
</tr>
<tr>
<td>7SK</td>
<td>2</td>
<td>0</td>
<td>2</td>
<td>981</td>
<td>$1.02 \times 10^{-3}$</td>
</tr>
<tr>
<td>Afp</td>
<td>23</td>
<td>28</td>
<td>51</td>
<td>18,194</td>
<td>$2.8 \times 10^{-3}$</td>
</tr>
<tr>
<td>Gm16326</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>414</td>
<td>$2.42 \times 10^{-3}$</td>
</tr>
<tr>
<td>AV036118</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>725</td>
<td>$1.38 \times 10^{-3}$</td>
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<tr>
<td>Gm10800</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>755</td>
<td>$1.32 \times 10^{-3}$</td>
</tr>
<tr>
<td>Gm11620</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>1,845</td>
<td>$1.08 \times 10^{-3}$</td>
</tr>
<tr>
<td>Dnajb8</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>989</td>
<td>$1.01 \times 10^{-3}$</td>
</tr>
</tbody>
</table>
applied to elucidate fully the genomic features of rAAV integration and whether they might contribute to carcinogenesis.

A set of parallel studies should also be undertaken, allowing the growing genomic resources associated with human HCCs to be queried to ascertain whether there is a recurrent set of genes, RNAs, and/or regulatory elements commonly dysregulated in HCC. By integrating HTS-based analyses of integrations in liver tissue from rodents (and other species) treated with rAAV, an assessment of genotoxic risk might be approximated by simply mapping rAAV integrations, especially if they fall into a CIS, onto a framework of genetic elements that are associated with human HCC. A pattern of vector integration into loci associated with human HCC, if seen in numerous samples from different animals, might indicate a potential hepatic toxicity, and suggest further monitoring, such as the assessment of circulating alpha-fetoprotein and/or abdominal imaging, in clinical applications.

While the current safety profile of rAAV in human clinical applications remains impeccable, the preclinical animal model data, coupled with an incomplete understanding of rAAV integration, should continue to motivate vector safety studies in preclinical models using modern genomic methods.

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AUTHOR DISCLOSURE

No competing financial interests exist.

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