Adeno-associated viral (AAV) vectors packaging the CRISPR-Cas9 system (AAV-CRISPR) can efficiently modify disease-relevant genes in somatic tissues with high efficiency. AAV vectors are a preferred delivery vehicle for tissue-directed gene therapy because of their ability to achieve sustained expression from largely non-integrating episomal genomes. However, for genome editing applications, permanent expression of non-human proteins such as the bacterially derived Cas9 nuclease is undesirable. Methods are needed to achieve efficient genome editing in vivo, with controlled transient expression of CRISPR-Cas9. Here, we report a self-deleting AAV-CRISPR system that introduces insertion and deletion mutations into AAV episomes. We demonstrate that this system dramatically reduces the levels of *Staphylococcus aureus* Cas9 protein, often greater than 79%, while achieving high rates of on-target editing in the liver. Off-target mutagenesis was not observed for the self-deleting Cas9 guide RNA at any of the predicted potential off-target sites examined. This system is efficient and versatile, as demonstrated by robust knockdown of liver-expressed proteins in vivo. This self-deleting AAV-CRISPR system is an important proof of concept that will help enable translation of liver-directed genome editing in humans.

**RESULTS**

**AAV-CRISPR Is Capable of Disrupting Episomal AAV Genomes In Vivo**

The liver is one of the principal sites where AAV-based gene therapy is being applied successfully in humans. We tested whether AAV-CRISPR could disrupt a co-expressed AAV-GFP transgene in mouse liver. Fifteen possible GFP-targeting gRNAs (Table S1) were first screened in 293-GFP cells, and the most efficient one (Sa_G-13) was selected for in vivo studies (Figure S1). An AAV8 vector encoding *Staphylococcus aureus* Cas9 (SaCas9) and the GFP-targeting gRNA was co-delivered with an AAV expressing emerald GFP (EmGFP) in an equimolar ratio, and mouse livers were harvested 3, 7, 14, 21, and 28 days after delivery (Figure 1A). Cas9 cutting of the GFP transgene did not significantly reduce the genome copy number of AAV-GFP or AAV-SaCas9 episomes (Figures 1B and 1C). AAV-GFP editing rates increased in a linear fashion over time, reaching...
Figure 1. AAV-CRISPR Removal of an Episomal AAV Transgene

(A) AAV8 vectors encoding emerald GFP (EmGFP) under the control of a small liver-specific promoter, HLP (top), and SaCas9 driven by a synthetic liver-specific HLP promoter with either a control guide RNA (gRNA) or a GFP-targeted gRNA (bottom). Experimental timeline with treatment groups is shown at right. (B) AAV-GFP genome copy numbers. (C) AAV-SaCas9 genome copy numbers. (D) GFP indel rates calculated via next-generation sequencing (NGS). (E) Western blots for GFP and beta-tubulin in mouse livers treated with a control gRNA or GFP-gRNA vector. #Failed AAV injection based on the absence of detectable vector genomes by qPCR. Data are indicated as mean ± SD. *p < 0.05.
an average of 90% insertion or deletion (indel) rate after 4 weeks (Figure 1D). The GFP protein level was significantly reduced after 2 weeks and was virtually undetectable by 3 weeks (Figure 1E), establishing the proof of concept that CRISPR-Cas9 can eliminate AAV-expressed transgenes in vivo.

**Screening the Activity of Self-Deleting gRNAs**

To develop a self-deleting AAV-CRISPR platform, we first designed gRNAs targeting different functional domains of the SaCas9 coding sequence (Figure S2; Table S2). We then tested the activity of 19 potential self-deleting SaCas9 gRNAs in vitro using a firefly-luciferase-based single-strand annealing assay. Briefly, short oligos containing the gRNA target sites and corresponding protospacer-adjacent motif (PAM) were cloned between two direct repeats of a truncated firefly luciferase gene containing two intervening stop codons after the SaCas9, and each candidate gRNA. The ratio of firefly luciferase activity is proportional to cutting efficiency by CRISPR-Cas9, and the relative efficiencies of different gRNAs can be quantitatively measured. The firefly luciferase reporter plasmids were co-transfected into HEK293FT cells along with a renilla luciferase control, and the direct repeats anneal, forming a full-length functional firefly luciferase in a subset of DNA-repair events (Figure S3A). Firefly luciferase activity is proportional to cutting efficiency by CRISPR-Cas9, and the relative efficiencies of different gRNAs can be quantitatively measured. The firefly luciferase reporter plasmids were co-transfected into HEK293FT cells along with a renilla luciferase control, SaCas9, and each candidate gRNA. The ratio of firefly:renilla luciferase activity of each self-deleting gRNA was normalized to that of the most efficient GFP-targeting gRNA. Seventeen of the 19 gRNA had detectable activity against SaCas9 target sites (Figure S3B). The “Self-5” gRNA had the highest activity, and was selected for further testing in vivo along with “Self-1,” which disrupts the enzymatic RuvC-1 nucleosome domain closest to the start codon.

**Testing Self-Deleting SaCas9 gRNAs In Vivo**

We next sought to determine whether our self-deleting CRISPR system could reduce SaCas9 protein in vivo. To do this, we compared the Self-1 and Self-5 gRNAs for their ability to eliminate AAV-expressed SaCas9 protein in the liver. Male C57BL/6J mice were injected with an AAV-SaCas9 vector targeting an endogenous gene, either alone or in combination with a second AAV vector expressing either Self-1 or Self-5 (Figure S4A). SaCas9 protein was readily detectable by western blotting in livers at 1 month after injection. Interestingly, the protein levels of SaCas9 were similarly decreased by Self-1 and Self-5 relative to the mice receiving the SaCas9 vector alone (Figure S4B). Based on this, we selected Self-1 for further development, as indels near the N terminus would reduce the likelihood of creating immunogenic SaCas9 peptides. Additionally, Self-1 is located at the nuclelease RuvC-like portion of SaCas9, which is potentially more critical to the activity of the nuclease than the REC domain, where Self-5 targets.

**Dose-Response of SaCas9 Removal with the Self-1 gRNA**

We next tested whether increasing the molar ratio of Self-1 to SaCas9 vector would more efficiently remove SaCas9 protein. Male C57BL/6J mice were injected with an AAV vector encoding SaCas9 and a gRNA targeting an endogenous gene (Mttp). A second AAV vector expressing the self-deleting gRNA (Self-1) was co-delivered at a 1:1, 1:2, or 1:3 molar ratio (Figure 2A). To ensure that editing could reach its maximal value, these animals were followed for 6 weeks prior to liver harvest. Self-deletion of SaCas9 did not significantly reduce AAV genome copy number, as seen previously with AAV-GFP (Figures 2B and 2C). Without the self-deleting gRNA, indels at the endogenous target site (Mttp) were present at a frequency of 28% via next-generation sequencing (NGS). With the self-deleting gRNA co-delivered, the indel rates at the endogenous Mttp target site were significantly lower, at 15%, 9%, and 7%, respectively, with increasing Self-1 doses (Figure 2D). The 1:3 ratio of Cas9:Self-1 resulted in slightly lower endogenous editing efficiency, as compared to a 1:1 ratio (7% versus 15%; p < 0.05). Disruption of the SaCas9 transgene ranged from 18% to 23% via Tracking of Indels by Decomposition (TIDE), with minimal variation between the self-deleting groups (Figure 2E). SaCas9 protein level in liver tissue was significantly reduced in all three groups receiving the self-deleting gRNA vector (down 70%, 84%, and 84%) (Figures 2F and 2G) but with no significant differences among the three ratios tested.

**Endogenous and Self-Deleting SaCas9 gRNAs Exhibit No Measurable Off-Target Activity**

A critical concern regarding current CRISPR-Cas9-based gene editing strategies is the cutting activity at potential off-target sites. These off-target events may cause unwanted tumorigenic mutations or large chromosomal rearrangements. Potential off-target sites were first identified in silico using the bioinformatics tool COSMID and subsequently examined using targeted NGS. Of the 8 potential off-target sites associated with the Mttp-targeting gRNA, only off-target site 3 (OT3) showed indel formation above background levels at a low rate of 0.13%–0.22% (Figure 3A). However, on closer inspection of the reads, this site occupies an area of five direct Gs and most likely represents PCR and sequencing error due to the chemistry of the Illumina platform. The self-deleting gRNA at the highest dose did not display any detectable off-target cutting activity at the 22 predicted potential off-target sites (Figure 3B). NGS revealed 38% indel rates at the SaCas9 locus at the highest self-deleting gRNA dose, indicating a robust on-target cutting activity.

**Timing of AAV-CRISPR Self-Deletion**

We next examined the time course of SaCas9 self-deletion in relation to an endogenous target. Mice were injected with an AAV-SaCas9 vector targeting Mttp alone or in combination with a 1:1 molar ratio of Self-1. Livers were harvested at 1, 2, and 4 weeks from the mice receiving Self-1 and compared to those from animals with SaCas9 alone at 4 weeks (Figure 5A). Mttp and SaCas9 self-editing rates increased in a linear fashion between 1 and 4 weeks (Figures S5B and S5C). At the protein level, SaCas9 was lower at all time points with the self-deleting gRNA (Figure S5D). The lower level of SaCas9 protein at 1 and 2 weeks is likely a result of the normal gradual increase in AAV expression in this tissue (which typically requires 10–14 days to reach its peak) rather than self-deletion. Since self-deletion occurs simultaneously with on-target cutting, we reasoned that higher rates of on-target editing might be achieved by slightly delaying the delivery of Self-1. To test this,
we compared mice injected with SaCas9 targeting Mttp alone, mice that received a co-injection of Self-1, and a third group that received the injection of Self-1 5 days later (Figure S6A). Robust editing of the endogenous target was observed with 24% for AAV-SaCas9 alone. In this experiment, on-target editing decreased dramatically with co-injection of Self-1. Interestingly, endogenous editing was preserved in the group receiving Self-1 after the 5-day delay (26%) (Figure S6B). However, this group had no detectable self-deletion and a complete absence of GFP protein, indicating that Self-1 did not transduce the liver in these mice (Figures S6C and S6D). Thus, it appears that even as early as 5 days after AAV8 administration, additional AAV8 vectors are blocked from entering the murine liver by the host immune system.

Versatility for Editing Different Genomic Targets

We next sought to determine whether our self-deleting gRNA system could achieve comparable editing efficiency at other endogenous targets. To test this, we selected the low-density lipoprotein receptor (Ldlr) and the apolipoprotein E (ApoE) genes. LDLR and ApoE are primarily liver-expressed proteins involved in the clearance of ApoB-containing lipoprotein particles, and both are potential targets for therapeutic genome editing. Mice were injected with SaCas9 vectors with gRNAs targeting either Ldlr or Apoe, with or without co-injection of Self-1 at a 1:1 ratio, and followed for 28 days (Figure 4A). High levels of on-target editing were observed for both Ldlr and Apoe, and this was not significantly reduced with co-injection of Self-1 (Ldlr: 41.9% versus 37.5%; Apoe: 33.8% versus 30.4%) (Figures 4B and 4C). Co-injection of Self-1 introduced indels in the AAV-SaCas9 vector targeting Ldlr (25.9%) as well as Apoe (50.1%) at high frequency (Figure 4D). In the liver, LDLR protein was decreased by 80% with AAV-SaCas9 alone, and a similar 79% reduction was achieved with the addition of Self-1 (Figures 4E and 4F). Although SaCas9 protein was not completely eliminated by Self-1, it was decreased by 73%, relative to AAV-SaCas9 alone (Figure 4G). Likewise, ApoE protein was reduced in plasma from mice treated with or without Self-1 (70% versus 62%), indicating highly efficient disruption of this primarily liver-expressed secreted protein (Figures 4H and 4I). SaCas9 protein levels dropped by 79% with Self-1 co-injection, showing a second example with efficient on-target editing despite SaCas9 self-deletion (Figure 4J).
Self-Deletion Generates Novel AAV Integration Patterns

Recombinant AAV are believed to be largely non-integrating in the absence of Rep, and integration events in human gene therapy studies appear to be rare and randomly distributed. However, we and others have previously reported insertion of fragments of inverted terminal repeats (ITRs) from AAV vectors at CRISPR-Cas9-generated double-strand breaks, as well as whole vector genome insertions. Importantly, we also uncover an unexpected safety concern due to the insertion of whole and truncated AAV vector genomes at endogenous loci cut with AAV gene therapy in humans. We show that a self-deleting gRNA targeting the coding sequence of SaCas9 can introduce inactivating mutations in AAV episomes in vivo. This results in efficient, although not complete, elimination of the SaCas9 protein over several weeks. For most targets, SaCas9 can be dramatically reduced without significantly compromising on-target editing. Off-target mutagenesis was not detected with the self-deleting gRNA or with the on-target gRNA at predicted sites in the mouse genome, consistent with the reported high specificity of SaCas9. Importantly, we also uncover an unexpected safety concern due to the insertion of whole and truncated AAV vector genomes at endogenous loci cut with AAV-CRISPR.

AAV-CRISPR has been used in recent years to modify endogenous genes in the liver, heart, skeletal muscle, retina, and brain. However, to our knowledge, this is the first demonstration that AAV-CRISPR can disrupt recombinant AAV episomes with high efficiency in vivo. In designing this approach, we intended to cut AAV episomes with Cas9 to promote their degradation. Interestingly, this did not occur for either AAV-GFP or AAV-SaCas9 and suggests that the liver favors non-homologous end joining (NHEJ) repair of blunt-ended extrachromosomal DNA over degradation. Despite this, inactivating mutations introduced into AAV episomes can efficiently silence transgene expression. In the case of our self-deleting SaCas9 system, the levels of this nuclease are reduced 79% or more at the protein level. For some targets, SaCas9 can be dramatically reduced without significantly compromising on-target editing. Off-target mutagenesis was not detected with the self-deleting gRNA or with the on-target gRNA at predicted sites in the mouse genome, consistent with the reported high specificity of SaCas9. Importantly, we also uncover an unexpected safety concern due to the insertion of whole and truncated AAV vector genomes at endogenous loci cut with AAV-CRISPR.

FIGURE 3. Endogenous Mttp and SaCas9 gRNAs Exhibit No Detectable Off-Target Activity

(A) NGS revealed no detectable off-target activity at 8 potential sites of the Mttp gRNA. The asterisk (*) indicates significance against saline-injected group, and the pound symbol (#) indicates significance against Self/Cas9 1:1 via one-way ANOVA and Tukey’s post hoc multi-comparisons test. (B) Self-deleting gRNA revealed no detectable off-target activity in vivo at the highest dose of gRNA (1:3). OT1’s higher indel rate is due to direct “G” repeats in the sequence. Data are indicated as mean ± SD. *p < 0.05.
in on-target efficiency was seen for Mttp. The relative molar excess of AAV episomes within the cell may serve as a natural delay switch, allowing for the editing of endogenous targets before a complete dropoff in SaCas9 activity occurs.

Our self-deleting AAV-CRISPR system relies on efficient co-transduction with two different AAV vectors, and variations in co-delivery may explain some differences in editing efficiency for Mttp versus Ldlr and Apoe. Ideally, a single vector system would be preferred,
which would guarantee simultaneous delivery of on-target and self-deleting gRNA to the same cells. Our attempts to develop a single vector system in which the self-deleting gRNA was co-expressed in cis with SaCas9 were unsuccessful. Even with the use of liver-specific promoters, indel mutations were introduced into the AAV genomes during viral packaging and did not express SaCas9 in vivo (data not shown). In addition, accommodating an on-target gRNA in the same vector would be a significant engineering challenge, due to the packaging limits of AAV vectors. However, given that we have established proof of concept for AAV-SaCas9 self-removal, and recent successes in AAV gene therapy in humans, this merits further exploration.

Our self-deleting AAV-CRISPR system generally reduces SaCas9 protein in the range of 70% to 84%. It is interesting to note that there is always residual SaCas9 in the liver with this approach, even at 4–6 weeks after AAV administration. One possible explanation is that there is imperfect overlap in the hepatocytes transduced with AAV-SaCas9 and Self-1. In our experience, the doses we used will deliver to 98%–99% of hepatocytes, and very efficient removal of LDLR and ApoE protein were achieved. Therefore, this probably does not solely account for the residual SaCas9 protein. A second possibility is that there may be a negative-feedback loop present. Since the enzymatic activity of SaCas9 is critical for its own disruption, the system could reach a steady state corresponding to a theoretical minimum, below which the remaining SaCas9 protein cannot effectively edit the AAV episomes encoding it. While it would be ideal to completely remove SaCas9 protein for clinical applications, this may not be feasible or absolutely essential. Rather, substantial decreases in SaCas9 protein may be acceptable in the liver, which is a highly regenerative tissue. Even if a handful of SaCas9-positive cells persist, these might be eliminated by the host immune system without significant pathological consequences, in much the same way as AAV capsid-presenting hepatocytes are purged by cytotoxic T cells. Since genome editing is a permanent modification, durable therapeutic benefit could still be achieved, provided most of the edited hepatocytes either survive or expand to repopulate the liver.

The integration of full as well as truncated AAV genomes at CRISPR-Cas9-generated cut sites is a novel and unexpected finding. The risk of insertional mutagenesis is an important safety concern for any gene therapy, which certainly has significant implications for the clinical use of AAV-based genome editing. It is well known that treatment of neonatal mice with high doses of recombinant AAV vectors can result in tumor formation in the liver through integration into the Rian locus, via cis-acting effects of strong promoter elements. Likewise, a recent report identified wild-type AAV2 integrations in known tumor suppressors and proto-oncogenes in human hepatocellular carcinoma biopsies. While the relevance of these findings to humans receiving recombinant AAV gene therapy is a subject of debate, it is clear that insertional mutagenesis should be avoided as much as possible. In the context of delivery of CRISPR-Cas9 with AAV vectors, insertion of whole vector genomes could be particularly problematic, as this event creates an artificial “hot spot” for integration, which will occur far more frequently than other homology-driven events. The risk of tumorigenesis with whole vector genome insertions is unknown but would likely depend on the transgene promoter activity, the gene being edited, and the genomic context of the target site.

Immunity to Cas9 has been found in humans (https://www.biorxiv.org/content/biorxiv/early/2018/01/05/243345.full.pdf), and mice, and there is concern that prolonged Cas9 expression could provoke elimination of edited cells and severe tissue damage. Our self-deleting system would significantly limit the window of SaCas9 protein expression in the majority of hepatocytes, though not completely eliminating it from all cells. Although cytotoxic T cells respond to AAV capsids in humans receiving liver-directed gene therapy with AAV, this can be effectively managed with prednisone. Short-term immunosuppression is now becoming standard practice for AAV trials in humans, and it is believed that prednisone preserves transgene expression by delaying T cell responses until AAV capsids are cleared. The risks of genome editing with CRISPR-Cas9 in human liver are currently unknown, but these could likely also be mitigated by immunosuppression prior to Cas9 elimination. Our self-deleting AAV-CRISPR system establishes critical proof of concept that transient expression of Cas9 with AAV vectors can be achieved, capitalizing on the unparalleled delivery efficiency of this vector system. Refinements to the current self-deleting AAV-CRISPR system are needed to provide complete Cas9 removal while avoiding insertional mutagenesis at CRISPR-Cas9 on-target cut sites.

MATERIALS AND METHODS

gRNA Design

gRNAs were designed targeting S. aureus Cas9, GFP, Mttp (exon 2), and Apoe (exon 2) by manual inspection based on the presence of a canonical NNGRRT PAM at the target site. Potential off-target sites were identified using the web-based bioinformatics program CRISPR Off-target Sites with Mismatches, Insertions and/or Deletions (COSMID). Search criteria for off-target sites included a NNGRR PAM (instead of NNGRRT), and a maximum of 3 mismatches and 2 base insertions or deletions relative to the target in the M. musculus (Mm10 build) genome. gRNAs with closely matching off-target sites were excluded and redesigned. gRNA sequences are shown in Table S2.

Figure 5. AAV Whole Genome Insertions at a CRISPR-Generated Cut Site

(A) Primers were designed to detect full-length AAV vector genome integrations from the 5′ and 3′ sides of the cut site. The endogenous gene-specific primers A and C are unique to the Mttp, Ldr, and Apoe loci, while the AAV genome primer B is common. (B) AAV-Self-1 genome insertions in the Mttp locus are present in both forward and reverse orientations and do not increase from 1–4 weeks. (C) AAV-Self-1 insertions at the Mttp locus are not affected by increasing Self-1 dose. (D) AAV-Self-1 insertions in both orientations in the Ldr locus. (E) AAV-Self-1 insertions in both orientations in the Apoe locus.
Figure 6. HITI Insertions of Truncated AAV-SaCas9 Vector Genomes at Endogenous Target Sites
(A) Partial AAV genomes are capable of integrating into endogenous gene targets after self-editing via homology-independent targeted integration (HITI). Primers were designed to detect self-edited vector integration from the 5' and 3' side of the target site. Primers A and C are specific to the target locus, while primer B is common within AAV-SaCas9 near the breakpoint. (B) Truncated AAV genome insertions with self-editing detected at the Mttp locus at 1–4 weeks after AAV. (C) Dosage of self-deleting gRNA does not change the frequency of truncated AAV vector insertion at the Mttp locus. (D) Truncated AAV genome insertions at the Ldr locus with self-editing. (E) Truncated AAV genome insertions at the Apoe locus with self-editing. For all targets, nonspecific bands in lanes without Self-1 likely arise from aberrant priming of AAV-SaCas9 episomal DNA and concatamers.
Plasmid Construction
Luciferase SaCas9 targets were cloned into the pLV45.1-SSA-Luciferase backbone. AAV plasmids containing the ITRs from AAV2 were used to construct CRISPR plasmids using gene synthesis and standard molecular biology approaches. Plasmid 1162_pAAV-HLP-EmGFP-spA encodes Emerald GFP driven by a small synthetic liver-specific promoter, HLP.42 Plasmid 1313_pAAV-U6-BbsI-MluI-gRNA-SACas9-HA-OLLAS-spA encodes the SaCas9 transgene derived from px602 (Addgene plasmid #61593, a gift from Feng Zhang) for liver-specific expression, with an upstream gRNA cloning cassette. gRNAs targeting GFP or Mttp were constructed using the plasmid 1518_pAAV-U6-SA-eGFP-gRNA-HLP-SACas9-HA-OLLAS-spA. A self-deleting gRNA targeting the RuvC domain of the S. aureus Cas9 gene was separately cloned into a similar backbone upstream of EmGFP (1530_pAAV-U6-BbsI-gRNA-SA-EmGFP), generating 1531_pAAV-U6-Self1-gRNA-SA-HLP-EmGFP. Targeting of Ldrl was accomplished using the previously described 1375_pAAV8-U6-SA-WtmLdrlEx14-gRNA-N22-CB-SACas9-HA-OLLAS-spA vector.43 A gRNA targeting Apoe was cloned into 1255_pAAV-U6-SaBsi-MluI-gRNA-CB-SACas9-HA-OLLAS-spA,43 generating 1377_pAAV8-U6-SA-mApoe-Exon2-gRNA2-N22-CBSACas9-HA-OLLAS-spA. All clones were verified by sequencing, as well as individual digestion with XmaI, SnaBI, and PvuII to confirm intact ITRs. Complete sequences are listed in the Supplemental Information, and plasmids are publicly available on Addgene or upon request.

Single-Strand Annealing assay for gRNA Screening
24 hr prior to transfection, 10,000 HEK293FT cells were seeded into 96-well plates. Transfections were performed in triplicate, with each mix containing 100 ng DNA measured by Qubit and 0.4 μL Lipofectamine 2000. A typical reaction contained 20 ng Firefly target, 40 ng px601 (Addgene plasmid #61591, a gift from Feng Zhang), 24 ng self-editing plasmid, 4 ng renilla, and 11 ng pUC19 plasmid. Luciferase expression was measured 48 hr post-transfection using the Dual-Glo Luciferase Assay System (Promega, E2920).

AAV Production
AAVs were generated as previously described,1,3 with several modifications. The adeno viral helper plasmid pAdDeltaF6 (PL-F-PVAdF6) and AAV packaging plasmid pAAV2/8 (PL-T-PV0007) were obtained from the University of Pennsylvania Vector Core. These plasmids were co-transfected with the AAV transgene construct into 293T cells using polyethylenimine (PEI). Cell pellets were harvested and purified using a single CsCl density gradient centrifugation. Fractions containing AAV vector genomes were pooled and then dialyzed in 100,000 molecular weight cut off (MWCO) cassettes against three washes of PBS at 4°C overnight to remove CsCl. Purified AAVs were then concentrated using an Amicon 100-kDa MWCO centrifugal filtration device (UFC510024) before storage at −80°C until use. AAV titers were calculated after DNase digestion using qPCR against a standard curve and primers specific to S. aureus Cas9 and GFP (Table S3).

Animals
Male C57BL/6J mice, 6–8 weeks of age, were obtained from Jackson Laboratories and kept with a light cycle from 7 a.m. to 9 p.m. Animals were allowed free access to food and water and maintained on a standard chow diet. Individual AAV vectors were injected at a dose of 5 × 1011 genome copies (GCs) per animal, with exception of the self-deleting vector, which ranged from 5 × 1011 to 1.5 × 1012 GCs per mouse. AAVs were diluted in 300 μL sterile PBS and delivered via intraperitoneal injection. All treatment conditions were randomly allocated within each cage of mice at the time of injection. Mice were fasted 5 hr prior to injection and all subsequent blood collection. Blood was collected via retro-orbital bleeding using heparinized Na-telson collection tubes, and plasma was isolated by centrifugation at 10,000 × g for 20 min at 4°C. All experiments were approved by the Baylor College of Medicine Institutional Animal Care and Use Committee (IACUC) and performed in accordance with institutional guidelines under protocol numbers AN-6243 and AN-7243.

TIDE Analysis
Primers amplifying the target regions were designed flanking the cut site, approximately 350 bp away on each side (Table S2). The gene of interest was then amplified via PCR, and the products were separated with electrophoresis on agarose gels and extracted using the QIAquick Gel Extraction Kit (QIAGEN, catalog no. 28704). Primers designed for PCR amplification were then used for Sanger sequencing of the targeted regions. Indel percentages were calculated via TIDE,24 using a control chromatogram for comparison. Decomposition windows, left boundaries, and indel ranges were optimized to have the highest alignment possible. The significance cutoff was maintained at p < 0.001 for all analyses. Primers are provided in Table S3.

Western Blotting
Liver lysates were prepared by homogenizing liver pieces in 10 volumes of RIPA buffer (50 mM Tris [pH 8.0], 150 mM NaCl, 1 mM EDTA, 1.0% Triton X-100, 0.1% SDS, 0.5% sodium deoxycholate) supplemented with Complete Mini Protease Inhibitor Cocktail (Roche, reference #11836153001) at a frequency of 2.5 Hz, four times, in a benchtop bead mill homogenizer. Samples were cleared by centrifugation at 15,000 × g, and the supernatant was collected. Protein concentrations were determined using a bicinchoninic acid (BCA) protein assay (Pierce, catalog no. 23225) according to the manufacturer’s instructions. Liver lysates (80 μg protein) were diluted in 4× NuPAGE MOPS SDS Running Buffer, NP0002 (Life Technologies, reference no. NP0007) supplemented with 5% beta-mercaptoethanol to a 20-μL final volume. Samples were denatured by heating to 95°C for 5 min and cooled on ice until gels were loaded. Proteins were resolved by SDS-PAGE on 4%–12% gradient gels (Invitrogen, reference nos. NP0322BOX and WGF1402BX10) using 3-morpholinopropan-1-sulfonic acid (MOPS) running buffer (Life Technologies, reference no. NP0002) and transferred to polyvinylidene fluoride (PVDF) membranes. Blocking was carried out for 1-hr rocking at 50 rpm on shaking platform with a 2:1 ratio of Odyssey Blocking Buffer (LI-COR Biosciences, P/N 927-40000) to PBS with 0.05% Tween 20 (PBS-T). Primary antibodies were diluted in a solution of
PBS-T supplemented with 0.1% BSA. Primary antibodies were then detected using goat anti-rabbit (680 nm; Rockland Immunocroohe, RL6111440020.5) and goat anti-mouse (800 nm; Rockland Immunocroohe, RL6111450020.5) secondary antibodies diluted in PBS-T + 0.1% BSA for 2 hr. Fluorescent imaging was performed on an Odyssey Classic Imager (LI-COR Biosciences). All western blots were performed in a similar manner, and antibody catalog numbers and dilutions are provided in Table S4.

**Deep Sequencing**

Genomic DNA extracted from mouse livers was amplified using locus-specific primers containing common adaptor sequences, and a second round of PCR amplification was used to add sample indexes as previously described. Amplicons for all target regions were purified using magnetic beads, pooled in equimolar amounts, and sequenced using the Illumina MiSeq platform. Alignment of sequence reads to reference sequences and indel quantification were carried out as previously described. All deep sequencing primers and indel analyses are provided in Tables S6–S10.

**Statistics**

All data are shown as the mean ± SD. Comparisons involving two groups were evaluated by a two-tailed Student’s t-test. For comparisons involving three or more groups, a one-way ANOVA was applied, with Tukey’s post-test used to test for significant differences among groups. In all cases, significance was assigned at p < 0.05.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes six figures, ten tables, a list of plasmids used, and FASTA sequences and can be found with this article online at https://doi.org/10.1016/j.omtm.2018.11.009.

**AUTHOR CONTRIBUTIONS**

W.R.L., G.B., C.M.L., and A.L. conceived and designed the studies; A.L., C.M.I., A.E.H., K.E.J., M.D.G., W.L., K.S.B., A.M.D., H.D., and A.R. performed the experiments and analyzed data; A.L., W.R.L., C.M.I. and G.B. wrote the manuscript.

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