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A molecular toolbox for rapid generation of viral vectors to up- or down-regulate neuronal gene expression in vivo

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We introduce a molecular toolbox for manipulation of neuronal gene expression in vivo. The toolbox includes promoters, ion channels, optogenetic tools, fluorescent proteins, and intrinsic artificial microRNAs. The components are easily assembled into adeno-associated virus (AAV) or lentivirus vectors using recombination cloning. We demonstrate assembly of toolbox components into lentivirus and AAV vectors and use these vectors for in vivo expression of inwardly rectifying potassium channels (Kir2.1, Kir3.1, and Kir3.2) and an artificial microRNA targeted against the ion channel HCN1 (HCN1 miRNA). We show that AAV assembled to express HCN1 miRNA produces efficacious and specific in vivo knockdown of HCN1 channels. Comparison of in vivo viral transduction using HCN1 miRNA with mice containing a germ line deletion of HCN1 reveals similar physiological phenotypes in cerebellar Purkinje cells. The easy assembly and re-usability of the toolbox components, together with the ability to up- or down-regulate neuronal gene expression in vivo, may be useful for applications in many areas of neuroscience.

Keywords: intronic miRNA, ion channel, lentivirus, AAV, RNAi, hippocampus, cerebellum, virus

INTRODUCTION

Tools for manipulation of gene expression in neurons in vivo are important for investigation of nervous system function and may be of therapeutic benefit. Using viruses to manipulate neuronal gene expression in vivo is increasingly popular due to the flexibility and economy of generating viruses compared with transgenic animals. Nevertheless, it can be time consuming to assemble new viral constructs containing different promoter, gene of interest, and reporter elements. Furthermore, many of these elements have not been widely validated for use in the central nervous system and their performance may vary between studies. A toolbox of standardized modular components that can be rapidly assembled into new viral constructs is therefore desirable. A critical component of such a toolbox will be methods to disrupt expression of targeted genes. One promising approach is gene knockdown by RNA interference (RNAi). However, current methods using RNAi in neurons are associated with off target effects and toxicity (Grimm et al., 2006; McBride et al., 2008; Ehler et al., 2010; Martin et al., 2011). In non-neuronal cells intrinsic expression of miRNAs has been suggested to alleviate these problems, while simultaneously enabling stable, high level co-expression of reporter genes (Du et al., 2006). However, it is not clear if this approach will produce specific or selective gene knockdown when introduced into neurons in vivo using viruses.

To address the goal of establishing a molecular toolbox for in vivo up- and down-regulation of neuronal gene expression, we generated and validated a set of vectors containing cassettes encoding promoters, genes of interest, intronic miRNA constructs, and reporter proteins flanked by recombination sites. These cassettes can be assembled into various combinations and inserted into a viral backbone through a single recombination reaction. The system is extremely flexible, allowing rapid assembly of gene-reporter fusion constructs, bicistronic expression of genes, and expression of intrinsic interfering RNAs for gene knockdown. We show that viruses produced from these components give stable over-expression or efficient knockdown of genes in neurons in vivo, while simultaneously expressing reporter genes detectable by native fluorescence. By focusing on manipulation of the HCN1 gene, which encodes an ion channel that has been relatively well characterized using knockout mice (e.g., Nolan et al., 2003, 2004, 2007; Chen et al., 2009, 2010; Huang et al., 2009), we go on to provide evidence that intrinsic expression of artificial miRNAs causes effective knockdown of gene expression and does not appear to be associated with off-target effects on physiological properties of neurons. Because this molecular toolbox enables identification and investigation of transduced cells by a variety of techniques, it may be useful for addressing a broad range of questions within neuroscience.

METHODS

VECTOR CONSTRUCTION

The principle behind the molecular toolbox is to allow rapid generation of new constructs through recombination cloning. To create the initial att-flanked PCR fragments for recombination, traditional digestion, and ligation methods were used for some constructs, whereas other were generated by PCR amplification or synthesized directly.

Restriction enzyme-based cloning

pEYFP-HCN1 was constructed by amplifying the HCN1 sequence from pEGFP-HCN1 (gift from Bina Santoro, Columbia University, New York) and cloning it into the EcoRI and BamHI sites.
of pEYFP-C1 (Clontech). pFCK-ChR2-mCherry was constructed by replacing the EGFP sequence in pFCK-ChR2-EGFP (from Ed Boyden, MIT, obtained via Addgene) with mCherry digested from pmCherry-N1 (Clontech) using AgeI and BsrGI. To enhance transgene expression we used a version of the woodchuck hepatitis virus post-transcriptional regulatory element optimized for safety (oPRE; Schambach et al., 2006). A lentivirus with oPRE cloned into the backbone (pLenti-oPRE) was generated by PCR amplifying the oPRE sequence from pMP71-GFP-opre (provided by Axel Schambach, Hannover Medical School, Hannover) and cloning it into the Xhol and XbaI sites of pLenti6/BLOCK-it-DEST backbone (Invitrogen).

To create a Gateway-compatible adeno-associated virus (AAV) vector, pCAGW-ChR2-Venus (from Karel Svoboda, obtained via Addgene; Petreanu et al., 2009) was digested with AgeI and EcoRV to remove the CAGW promoter and the ChR2-Venus insert. Reading Frame B of the Gateway Vector Conversion System (Invitrogen) was ligated into the AAV backbone to create pAAV-Gateway.

Recombination cloning

While several technologies are available for recombination cloning (e.g., Ma et al., 1987; Bubeck et al., 1993; Oliner et al., 1993) here we use the Gateway system (Invitrogen). This recombination cloning system uses a three-step strategy to generate expression constructs (Hartley et al., 2000; Sasaki et al., 2004; Yahata et al., 2005). Cassettes encoding individual components of the constructs are amplified by PCR using primers containing att sites that allow recombination. The PCR cassettes are then recombined into a donor vector with matching att sites to generate an entry vector. By using alternative matching pairs of att sites different arrangements of entry vectors can be recombined into a destination vector to generate new expression constructs in a single reaction.

Generation of att-flanked cassettes. Cassettes encoding the toolbox components were generated by PCR amplification using primers containing the required att recombination site sequences in addition to 18–25 base pairs of template-specific sequence. Two hundred nanomolars of each primer was added to 1–2 μl of template DNA and 22.5 μl Platinum PCR SuperMix High Fidelity (Invitrogen). Thirty-five cycles of amplification were performed at 94°C for 30 s, 55°C for 30 s, and 68°C for 60 s.

Generation of entry vectors. To generate an att-flanked PCR cassette into a vector, 50 fmol of purified att-flanked PCR products were added to 50 fmol of the appropriate pDONR vector (Invitrogen), 2 μl of BP Clonase II enzyme (Invitrogen), and TE Buffer pH 8.0 to a total of 10 μl. The recombination reactions were incubated at 25°C overnight generating entry vectors. One microliter of proteinase K solution (Invitrogen) was added and the reactions were incubated at 37°C for 10 min. Three microliters of each reaction was transferred into TOP10 cells (Invitrogen) and positive transformants were selected with kanamycin (Sigma-Aldrich).

Generation of viral destination vectors. To generate viral vectors, 10 fmol of each of the required entry vectors were added to 20 fmol of the destination vector (pLenti6/BLOCK-it/DEST, pLenti-oPRE, or AAV-Gateway), 2 μl LR Clonase II Plus enzyme mix (Invitrogen), and TE Buffer pH 8.0 to a total volume of 10 μl. The reactions were incubated overnight at 25°C and treated with 1 μl proteinase K solution (Invitrogen) at 37°C for 10 min. The entire reaction was transformed into SblI3 cells (Invitrogen) and positive transformants were selected using ampicillin (Invitrogen) and blasticidin (Autogen Bioclear).

General purpose cassettes. CAMKIIa short (0.4 kb) and long (1.3 kb) promoter variants (Dittgen et al., 2004) were amplified from mouse genomic DNA. Synapsin and enhanced synapsin promoters were amplified from pBSIISK-SYN-GFP-WPRE and pBSIISK-E/SYN-GFP-WPRE respectively (vectors provided by Hiroyuki Hioki, Kyoto University, Kyoto). The Netrin G1 putative promoter was amplified from mouse genomic DNA. A 1.1-kb sequence corresponding to the first 345 bp + 754 bp immediately upstream of the transcription start site of the Netrin G1 (Ntng1) transcript variant a, gene (NM_030699.2) was cloned. Fluorescent reporter cassettes were amplified from pEYFP-C1, pEYFP-C1, pECFP-C1, and pmCherry-C1 (Clontech). An IRES-EYFP cassette was amplified from pPRIG (provided by Patrick Martin, UNICE, Nice). A cassette containing oPRE was generated by flanking the oPRE sequence with the appropriate recombination sites to ensure it would always insert 3’ to any coding cassettes during recombination into a viral vector. EYFP-HCN1 cassettes were amplified from pEYFP-HCN1. Channelrhodopsin-mCherry was amplified from pFCK-ChR2-mCherry. CAMKII(0.4)-Halorhodopsin-EYFP was amplified from pFCK-Halo-EYFP (from Ed Boyden, MIT, obtained via Addgene; Han and Boyden, 2007). Following PCR, the amplicons were gel purified using a QIAquick gel extraction kit (Qiagen). To generate entry vectors, all cassettes were recombined into the appropriate donor vector.

Reporter constructs. To generate a reporter construct for visualization of neurons in vivo, pENTR-L1-CAMKII(0.4)-L4, pENTR-R4-EYFP-R3, and pENTR-L3-oPRE-L2 were recombined with pLenti6/BLOCK-it-DEST to create pLenti6-CAMKII(0.4)-EYFP. To test the Netrin G1 promoter a reporter lentivirus was made by recombining pENTR-L1-NetrinG1-L4, pENTR-R4-EYFP-R3, and pENTR-L3-oPRE-L2 with pLenti6/BLOCK-it-DEST to create pLenti6-NetrinG1-EYFP.

Ion channel constructs. The coding sequences of Kir2.1 (M. musculus potassium inwardly rectifying channel, subfamily J, member 2 (Kcnj2) NM_008425) fused to a C-terminal mCherry sequence was synthesized by Mr Gene. The sequence was flanked by L5-L2 att recombination sites to enable recombination cloning, pENTR-L1-CAMKII(0.4)-R5 and pENTR-L5-Kir2.1-mCherry-R2 were recombined with the pLenti-oPRE backbone to create the lentivector pLenti–CAMKII(0.4)–Kir2.1-mCherry. The coding sequences of Kir3.1 [M. musculus potassium inwardly rectifying channel, subfamily J, member 3 (Kcnj3) NM_008426.1] and Kir3.2 [M. musculus potassium inwardly rectifying channel, subfamily J, member 6 (Kcnj6), transcript variant Girk2A-1, NM_010606] fused to a C-terminal EGFP sequence and flanked by R4-R3

1http://mrgene.com
att sites were synthesized by Mr. Gene. Entry vectors encoding the Kir3-EGFP constructs were recombined with pENTR-L1-CAMKII(0.4)-L4, pENTR-L3-oPRE-L2, and pLenti6/BLOCK-IT-DEST to create the lentivectors pLenti6-CAMKII(0.4)-Kir3.1-EGFP and pLenti6-CAMKII(0.4)-Kir3.2-EGFP.

**miRNA constructs.** BLOCK-IT miR RNAi sequences targeting mouse HCN1 (Mm4510951, Mm4510952, and Mm4510953; Invitrogen) were cloned into the synthetic intron of pSM155 (provided by Guangwei Du, University of Texas Health Science Center, Houston) using procedures described in detail elsewhere (Du et al., 2006). Briefly, RNAi sequences were designed so that once annealed, the duplexes had four nucleotide overhangs compatible with the cohesive ends produced by BsmBI digestion of pSM155. The RNAi duplexes were annealed and ligated into pSM155. A cassette containing the intronic miRNA upstream of EGFP was then amplified with B4r and B3r PCR primers and recombined to generate the entry vector pENTR-R4-HCN1miR-EGFP-R3. To create a negative control miRNA construct, a cassette containing a miRNA targeting luciferase upstream of an EGFP sequence was amplified from pSM155-Luc (also from Guangwei Du) with B4r and B3r primers and recombined to create the entry vector pLenti6-ESYN-HCN1miR-EGFP. To generate AA V expressing miRNAs for experiments in Figures 5–7, pENTRL1-ESYN-L4 and pENTR-L3-oPRE-L2 were recombined with pAA V-Gateway and pENTR-R4-HCN1miR-EGFP-R3 or pENTR-R4-LucmiR-EGFP-R3 to create pAA V-ESYN-HCN1miR-EGFP and pAA V-ESYN-LucmiR-EGFP.

**PCR PRIMERS**

CAMKII(1.3) attB1: GGGGACAAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
CAMKII(0.4) attB1: GGGGACAAGTTTGTACAAAAAAGCAGGCTACTTGTGGACTAAGTTTGTTC
CAMKII attB5r: GGGGACAACTTTTGTATACAAAGTTGTTGCCCGCAGAACTAGGGGCCACT
CAMKII attB4: GGGGACAACTTTTGTATAGAAAAGTTGGTGCCCGCAGAACTAGGGGCCACT
SYN attB1: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
SYN attB5r: GGGGACAACTTTTGTATACAAAGTTGTTGCCCGCAGAACTAGGGGCCACT
SYN attB4: GGGGACAACTTTTGTATACAAAGTTGTTGCCCGCAGAACTAGGGGCCACT
IRES-EGFP attB3: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
ChR2 attB5: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
ChR2 attB4r: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
HCN1 attB5: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
HCN1 attB4r: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
K2P9.1 attB4r: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
K2P9.1 attB3r: GGGGACAACAGTTTGTACAAAAAAGCAGGCTCATTATGGCCTTAGGTCACTT
HCN1F EcoRI: AATTCTAATGGAAGGCGGCG
HCN1R BamHI: GATCTAAATTCGAAGCAAAACGGGG

**CELL LINE CULTURE AND TRANSFECTIONS**

HEK293FT cells (Invitrogen) were maintained at 37°C, 5% CO2 in DMEM with GlutaMAX I, 4,500 mg/l glucose and 110 mg/l sodium pyruvate (Gibco) supplemented with 50 U/ml each of penicillin and streptomycin (Sigma) and 10% v/v FCS (Invitrogen). DNA transfections of miRNA constructs were performed using FuGENE6 (Roche) according to manufacturer’s instructions.

**VIRUS PRODUCTION**

Lentiviruses were generated by triple transfection of HEK293FT cells (Invitrogen) with the modified lentivector, psPAX2 (courtesy of Didier Trono, EPFL Lausanne, obtained via Addgene) and the pLP/VSVG envelope plasmid (Invitrogen) using calcium phosphate transfection. The lentiviruses were harvested in serum-free medium after 3 days, filtered, and concentrated in primed Centriplus-80 filter devices (Millipore). Functional titer was determined by EGFP fluorescence after serial dilution on HEK293FT cells and was 0.5–2 × 10^8 TU/ml. Hybrid AAVs containing the ITR
from AA V2 and Capsid from AA V1 were produced by Vectorbiolabs\textsuperscript{2}. Titer was determined by measurement of vector genome copies and was $5 \times 10^{12}$ GC/ml.

**STEREOTAXIC INJECTION**

All animal experiments were carried out according to the guidelines laid down by the University of Edinburgh’s Animal Welfare Committee and in accordance with the UK Animals (Scientific Procedures) Act 1986. C57BL6/J/OlaHsd mice aged 4–6 weeks were anesthetized with isoflurane and mounted into a stereotaxic frame. 2, -1.5, and 0.5 either with the injecting arm angled back 10˚ and positioned over the midline, or angled back 30˚ and rotated 45˚ counter-clockwise to target more lobules. Following surgery, animals were returned to their home cages for at least 2 weeks before being used for analysis.

**IMAGING**

Mice were terminally anesthetized with sodium pentobarbital and perfused with 4% PFA. The brains were post-fixed and immersed in 30% sucrose overnight before 30 μm coronal sections were cut on a freezing microtome. Native fluorescence of reporter genes was acquired at Nyquist sampling rates, were deconvolved using Huygens software (Scientific Volume Imaging, Netherlands), and the resulting three-dimensional images were analyzed using NIH ImageJ software.

**FACS SORTING**

Lentivirus expressing Kir2.1-mCherry was stereotaxically targeted to the dentate gyrus of 4-week-old C57BL6/OlaHsd mice. Two weeks after injections the mice were culled and the brains were sectioned on a freezing microtome. The slices were transferred to oxygenated ACSF and then incubated at 37˚C in 5% CO\textsubscript{2} for 45 min to dissociate the cells. The digestion reaction was stopped by the addition of 100 μl BSA (10 mg/ml; Sigma-Aldrich), 11 μl FCS (Invitrogen), 10 μl DNase (1,000 U/ml; Sigma-Aldrich), and 10 μl 5 mM Leupeptin (Sigma-Aldrich). The samples were triturated through a fire-polished Pasteur pipette until no clumps were visible. Two milliliters of cold ACSF + HEPES was added to each sample and they were passed through a 70-μm sieve (Becton Dickinson) to remove any remaining cell clumps. The cells were counted with a hemocytometer and resuspended in cold ACSF + HEPES to a density of $1 \times 10^6$ cells/ml. One microliter per milliliter of Live/Dead stain with a green fluorescent dye (Invitrogen) was added to each sample and they were incubated on ice for 30 min in the dark. The samples were centrifuged at 800 rpm and washed twice in 1 ml PBS + 1% FCS to remove unbound stain. Samples from an un.injected brain were prepared alongside the experimental brains to use as controls for the fluorescence-activated cell sorting (FACS) sorting and subsequent RT-PCR. A sample from the uninjected brain was heat-treated at 50˚C to provide a positive control for calibration of the Live/Dead stain. The samples were sorted on a FACS Aria II (BD Biosciences, USA) with gating parameters set to collect live mCherry-fluorescent cells. The yield of positive cells isolated from the dentate gyrus of a single brain ranged from 5,249 to 14,624 cells.

**REAL TIME PCR**

Total mRNA was extracted using the Arcturus PicoPureAR RNA Isolation Kit (Applied Biosystems) in combination with the RNase-Free DNase Set (Qiagen) according to manufacturer’s instructions. Taqman Gene expression assays were used for amplification of target genes. Specific assays used were Mm00484848_m1 (Mcm6), Mm01326464_m1 (NeuroD6), Mm01253033_m1 (Gfap), and Mm00446968_m1 (Hprt1). RT-PCR reactions were set up using the TaqmanAR RNA-to-C\textsubscript{T} 1-Step Kit (Applied Biosystems). One microliter of mRNA template was added to 1 μl of the gene-specific probe and primers mix, 10 μl 2× TaqMan RT-PCR mix, 0.5 μl 40× TaqMan RT Enzyme mix, and 7.5 μl H\textsubscript{2}O. Reverse transcription was performed at 48˚C for 15 min followed by activation of AmpliTaq Gold DNA Polymerase at 95˚C for 10 min and 40 cycles of amplification at 95˚C for 15 s and 60˚C for 1 min on an Mx3000P QPCR system (Stratagene). Quantification was performed using the comparative C\textsubscript{T} method with normalization to Hprt1. Results were expressed as mean ± standard error of the mean and statistical analyses (ANOVA and Student’s t-test) were performed using Smith’s Statistical Package, version 2.80 or R (The R Foundation).

**ELECTROPHYSIOLOGY**

Cerebellar brain slices were prepared and whole-cell patch-recordings made from visually identified cerebellar Purkinje cells following previously described procedures (Nolan et al., 2003; Zonta et al., 2011). Briefly, sagittal cerebellar slices were cut in cold (4–6˚C) modified oxygenated ACSF of the following composition (mM): NaCl 86, Na\textsubscript{2}H\textsubscript{2}PO\textsubscript{4} 1.2, KCl 2.5, NaHCO\textsubscript{3} 25, CaCl\textsubscript{2} 0.2–1 μM, Mg\textsubscript{2}+ 1 mM, HEPES 10, Na\textsubscript{2}ATP 2.5, Na\textsubscript{2}GTP 0.3, and glucose 25, sucrose 75, at 6˚C for 10 min. Four hundred micrometers coronal slices were cut through the hippocampus on a Vibratome 3000 sectioning system. The slices were transferred to oxygenated ACSF + 10 μM HEPES at 37˚C and incubated for 10 min. The dorsal dentate gyrus from each slice was microdissected out under a dissecting microscope. All samples from a single brain were pooled together in 1 well of a 12-well plate with 900 μl oxygenated ACSF + HEPES. One hundred microliters papain (200 U/ml, Sigma-Aldrich) was added to each well and the samples were incubated at 37˚C in 5% CO\textsubscript{2} for 45 min to dissociate the cells. The digestion reaction was stopped by the addition of 100 μl BSA (10 mg/ml; Sigma-Aldrich), 11 μl FCS (Invitrogen), 10 μl DNase (1,000 U/ml; Sigma-Aldrich), and 10 μl 5 mM Leupeptin (Sigma-Aldrich). The samples were triturated through a fire-polished Pasteur pipette until no clumps were visible. Two milliliters of cold ACSF + HEPES was added to each sample and they were passed through a 70-μm sieve (Becton Dickinson) to remove any remaining cell clumps. The cells were counted with a hemocytometer and resuspended in cold ACSF + HEPES to a density of $1 \times 10^6$ cells/ml. One microliter per milliliter of Live/Dead stain with a green fluorescent dye (Invitrogen) was added to each sample and they were incubated on ice for 30 min in the dark. The samples were centrifuged at 800 rpm and washed twice in 1 ml PBS + 1% FCS to remove unbound stain. Samples from an un-injected brain were prepared alongside the experimental brains to use as controls for the fluorescence-activated cell sorting (FACS) sorting and subsequent RT-PCR. A sample from the uninjected brain was heat-treated at 50˚C to provide a positive control for calibration of the Live/Dead stain. The samples were sorted on a FACS Aria II (BD Biosciences, USA) with gating parameters set to collect live mCherry-fluorescent cells. The yield of positive cells isolated from the dentate gyrus of a single brain ranged from 5,249 to 14,624 cells.

\textsuperscript{2}www.vectorbiolabs.com
White et al.  

**RESULTS**

A MOLECULAR TOOLBOX FOR RAPID GENERATION OF VIRAL CONSTRUCTS

To enable rapid generation of new viral constructs we constructed a library of vectors containing cassettes compatible with the Gateway cloning system (Figure 1). The Gateway system uses site-specific recombination for directional cloning of PCR products and their subsequent subcloning into destination vectors. DNA fragments are amplified by PCR using primers that contain specific recombination sites (att sites). Entry vectors are generated using the BP Clonase enzyme to catalyze the insertion of the DNA fragment into a donor vector containing matching att sites. To generate the final construct the entry vector can then be recombined with a destination vector through the addition of LR Clonase. By including different att sites in each vector, att-flanked cassettes from up to four entry vectors can be directionally recombined into a destination vector in a single reaction (Figures 1A,C). Compared with conventional cloning approaches this reduces the number of

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**Figure 1** | Schematic illustrating modular cloning system. (A) Schematic illustrating recombination of entry vectors carrying cassettes flanked by att sites with a viral destination vector. In a single reaction recombination between matching att sites inserts the cassettes into the vector in the correct order, generating a viral expression construct. (B) Components of the molecular toolbox flanked with att sites ready for recombination. For simplicity, only the att-flanked regions of the entry vectors are illustrated in (B,C). (C) Examples of recombination reactions to generate the lentiviral expression construct plenti-CAMKII(0.4)-Kir2.1-mCherry and the AAV expression construct pAAV-ESYN-HCN1mR-EGFP.
steps and time required to generate new constructs and allows the same components to be easily rearranged to form new viral vectors. The att-flanked cassettes that we describe here encode neuronal promoters, genes of interest, reporter genes, or intronic miRNAs (Figure 1B). The promoter cassettes are flanked with the appropriate att sites to ensure they insert as the first cassette in the viral backbone and therefore drive expression of downstream cassettes. The reporter cassettes are flanked with att sites permitting their expression either alone or as N- or C-terminal fusion proteins. Because we wanted to transduce neurons with either lentivirus or AAV and as there are currently no AAV vectors available that are compatible with the Gateway cloning system, we converted pACAGW-ChR2-Venus-AAV (Petreanu et al., 2009) into a Gateway-compatible destination vector by removing the CAG promoter and the ChR2-Venus insert and replacing them with a recombination cassette. Functionality of all the modular components of the toolbox was confirmed in vitro before using a subset to generate viruses for further in vivo validation (Table 1).

To evaluate expression of ion channels tagged with fluorescent reporter molecules, we first produced lentiviruses assembled from constructs encoding either an EYFP reporter alone, or Kir potassium channel-reporter fusion proteins, with expression driven by the short variant of the CAMKIIα promoter (CAMKII(0.4)-EYFP, CAMKII(0.4)-Kir2.1-mCherry, CAMKII(0.4)-Kir3.1-EGFP, and CAMKII(0.4)-Kir3.2-EGFP). To examine their expression in hippocampal neurons in vivo, we stereotaxically injected the viruses into CA1. Two weeks later, we sectioned the brains and imaged the native fluorescence in virally transduced cells. All four viruses expressed their transgenes at high levels within neurons. The subcellular localization of the reporter protein differed between

<table>
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<th>Component</th>
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<th>Level of validation</th>
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<tr>
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</table>

The components of the toolbox that are in entry vectors ready for recombination to generate new viral vectors. The level of validation to date is indicated with observations regarding in vivo expression where appropriate.
the viruses, indicating differential trafficking of each ion channel (Figure 2A). The Kir2.1 channel was localized to the soma and proximal dendrites of CA1 pyramidal neurons, whereas Kir3.1 and Kir3.2 were present throughout the proximal and distal dendrites. Importantly, the differences in distribution of each ion channel reflect expression patterns for native channels in vivo (Miyashita and Kubo, 1997; Koyrakh et al., 2005) suggesting that this approach will be useful to study ion channel localization (cf. Piskorowski et al., 2011). Expression of Kir2.1-mCherry results in an inwardly rectifying potassium current confirmed by electrophysiological recordings from acute slices (data not shown) in agreement with previous findings using a similar construct in cultured neurons (Burrone et al., 2002). We are yet to assess the functional consequences of expression of Kir3.1-EGFP and Kir3.2-EGFP, but similar constructs yield functional channels in expression systems (Stevens et al., 1997).

Promoters other than CAMKIIα can also be successfully used to drive neuronal gene expression (Dittgen et al., 2004). To determine if a fragment of the putative Netrin G1 promoter can drive gene expression in vivo, we assembled the lentivirus NetrinG1-EYFP and injected it into the dentate gyrus. Consistent with in situ hybridization data from the Allen Brain Atlas which suggests the presence of Netrin G1 mRNA in the dentate gyrus, we found EYFP fluorescence in dentate gyrus granule cells transduced with NetrinG1-EYFP (Figure 2B). However, expression was not limited to granule cells, as we also observed occasional fluorescence in interneurons and glia (Table 1). The enhanced synapsin (ESYN) promoter also gave strong transgene expression in dentate gyrus granule cells transduced with the virus ESYN-LucmiR-EGFP (Figure 2C). The native fluorescence from this virus (and the other reporter viruses) was bright enough to allow imaging of dendritic spines (Figure 2C). Transgene expression was visible from 4 days post-injection, reached maximal intensity by 7–10 days post-injection and remained stable for at least 9–12 weeks (the longest time points we have analyzed for AAV and lentivirus respectively, data not shown).

**ISOLATION OF NEURONS TRANSDUCED WITH VIRUSES IN VIVO FOR DOWNSTREAM ANALYSIS OF GENE EXPRESSION**

Expression of virally encoded fluorescent proteins in addition to interfering RNAs or transgenes enables identification of transduced neurons and in principle might allow their isolation from non-transduced neurons. We therefore assessed the amenability to FACS of neurons transduced using viruses generated from the toolbox (Figure 3).

We stereotaxically injected CAMKII(0.4)-Kir2.1-mCherry into the dentate gyrus of mice aged 4–6 weeks (Figure 3B). Two weeks later the dentate gyrus was manually dissected out and dissociated (see Methods and Figure 3A). The cells were stained with a cell death marker and sorted by FACS to isolate a live Kir2.1-mCherry expressing population. We were able to harvest up to 14,624 positive cells from a single brain. RNA was extracted from these cells and screened by RT-PCR for markers of different hippocampal cell types to examine the purity of the sorted population. Samples of RNA from cells dissociated from either the entire dorsal hippocampus or just the dentate gyrus dissected from uninjected brains were used as controls.

**FIGURE 2 | Lentiviral expression of transgenes in hippocampal neurons.** Examples of gene expression from lentiviruses injected into the hippocampus. (A) Expression of EYFP or Kir-reporter fusion proteins in CA1 pyramidal neurons of the hippocampus driven by the CAMKII short promoter. (B) Expression of EYFP in dentate gyrus granule cells driven by the Netrin G1 promoter. (C) EGFP expression in dentate gyrus granule cells driven by the ESYN promoter. Reporter protein expression is bright enough to enable imaging of dendritic spines (upper and lower right panels). Higher magnification spine images (lower right panel) correspond to different depths from a Z-stack of the region indicated by the white box (upper right panel). Scale bars = 100 μm for all images except the spine image, where the scale bar = 5 μm.
To evaluate the accuracy of cell isolation and the specificity of neuronal transduction we used RT-PCR of mRNA isolated from sorted cells to compare expression of anatomical and cell type markers. Expression of Neurod6, which is a marker for the CA1 and CA3 regions of the hippocampus (Lein et al., 2004), was over 20-fold higher in the entire hippocampus compared with the dissected dentate gyrus. Neurod6 was undetectable in three out of four samples of sorted Kir2.1-mCherry positive cells and the remaining sample had extremely low expression of Neurod6 mRNA that was significantly less than the entire hippocampus (Figure 3C). This indicates minimal contamination of the samples with cells from CA1 or CA3. Expression of Mcm6, which is a marker for dentate gyrus positive cells and the remaining sample had extremely low expression of Neurod6 mRNA that was significantly less than the entire hippocampus (Figure 3C). 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REDUCTION OF NEURONAL GENE EXPRESSION BY VIRAL EXPRESSION OF INTRONIC miRNAs

Whilst our previous constructs altered neuronal gene expression through over-expression of transgenes, it is also useful to be able to reduce expression of endogenous genes. This can be achieved through the use of RNAi. However, early approaches for RNAi that used RNA Pol-III promoters to express high levels of short hairpin RNAs (shRNA) can be toxic in vivo (McBride et al., 2008; Ehlert et al., 2010). More recently, Pol-II promoters have been used to express artificial miRNA sequences. This reduces the off-target effects (McBride et al., 2008) but can result in poor expression levels of cocistronic reporter genes (Du et al., 2006). To address this issue, vectors were developed in which the miRNA is contained within a synthetic intron upstream of the reporter gene (Du et al., 2006). This allows for efficient expression of both the reporter and the miRNA from a Pol-II promoter (Figure 4A). We speculated that viruses expressing artificial intronic miRNAs and reporter genes would have improved specificity of gene knockdown in neuronal populations in vivo.

To test this possibility, we constructed AAVs targeting expression of the HCN1 ion channel and evaluated their efficacy in cerebellar Purkinje neurons. Oligonucleotide sequences designed against HCN1 were cloned between the miRNA arms in the artificial intron of the pSM155 plasmid vector as previously described.
To check their effectiveness, the HCN1 miRNA vectors were cotransfected with an HCN1-m-Cherry plasmid into HEK293FT cells. The most effective HCN1 miRNA (MmiS10953) successfully reduced expression of HCN1-mCherry to almost undetectable levels whereas a negative control miRNA targeting luciferase (Luc miR) had no effect (Figure 4B). We therefore generated AAV vectors in which a cassette encoding the intronic HCN1 miRNA and EGFP reporter was inserted downstream of the ESYN promoter in a modified AAV backbone (AAV-Gateway). A negative control vector encoding Luc miR-EGFP was also constructed and viruses produced.

Injection of the AAVs into the cerebellum demonstrated widespread, high level expression of the EGFP reporter in Purkinje neurons within 2 weeks (Figure 5A). We evaluated the efficacy of the knockdown approach by measuring the hyperpolarization-activated current ($I_{\text{f}}$) recorded from cerebellar Purkinje cells in brain slices. Previous experiments with knockout mice suggest that HCN1 channels are the major ion channel type responsible for $I_{\text{f}}$ recorded from cerebellar Purkinje cells (Nolan et al., 2003). Consistent with this idea, and with the efficacy of the artificial miRNA approach, $I_{\text{f}}$ was abolished in Purkinje cells transduced with AAV expressing the HCN1 miRNA (Figure 5B-C). In contrast, Purkinje cells transduced with AAV expressing the miRNA targeted against luciferase had $I_{\text{f}}$ with amplitude and kinetics comparable to non-transduced neurons (Figure 5C-D). Thus, efficacious suppression of HCN1 function by the viral expression of HCN1 miRNA is not due to non-specific effects of viral transduction or expression of miRNAs.

To further address the specificity of the miRNA approach, we took advantage of the ability of electrophysiological techniques to quantify key physiological properties of neurons. We compared measures of spontaneous action potential firing, action potential waveform, and basic integrative properties, in control and HCN1 miRNA transduced neurons from mice with global deletion of HCN1 and wild-type littermate controls (Figures 6 and 7). We reasoned that if knockdown with HCN1 miRNA is specific, then the physiological properties of Purkinje cells transduced with AAV expressing HCN1 miRNA should be similar to Purkinje cells from mice with global deletion of HCN1. Differences between physiological properties due to viral transduction or off-target effects of the HCN1 miRNA should be apparent by a similar effect of transduction with the HCN1 miRNA AAV on Purkinje cells from wild-type mice and mice with global deletion of HCN1.

We first examined spontaneous action potentials generated by cerebellar Purkinje cells. In agreement with previous studies of global HCN1 knockout mice (Nolan et al., 2003), we find that transduction with the HCN1 miRNA AAV does not affect the frequency of spontaneous action potentials recorded from Purkinje cells using either cell-attached or whole-cell configurations (Figures 6A-C). Properties of the action potential waveform were also similar in both uninfected and infected Purkinje cells from control mice and mice with global deletion of HCN1 (Figures 6A,D,E). These data are consistent with the notion that HCN1 channels do not play a pacemaker role in cerebellar Purkinje cells (Raman and Bean, 1999; Nolan et al., 2003; Carter and Bean, 2011) and also indicate that transduction with AAV expressing intronic microRNAs and EGFP does not have off-target effects on ion channels important for generation of action potentials.
FIGURE 6 | Expression of HCN1 miR does not affect spontaneous firing of cerebellar Purkinje cells. (A–D). Examples of spontaneous action potentials recorded from cerebellar Purkinje cells in brain slices obtained from control mice (A,B) or global HCN1 knockout mice (C,D). Purkinje cells were either positive for EGFP indicating expression of HCN1 miRNA (B,D) or were EGFP negative indicating they did not express HCN1 miRNA (A,C). (E) Frequency of spontaneous action potentials recorded from Purkinje cells in the cell-attached configuration. There was no significant effect of genotype (p = 0.59) or AAV infection (p = 0.14). (F) Frequency (upper panel), half-width (middle panel), and peak after-hyperpolarization (lower panel) for spontaneous action potentials recorded from Purkinje cells in the current-clamp configuration. There was no significant effect of AAV on any parameter (p > 0.14) and no significant effect of genotype on spike frequency or AHP duration. There was a significant effect of genotype on the AHP peak (p = 0.034). This effect is small and is inconsistent with previous comparisons suggesting it may simply reflect a chance outcome expected from repeated statistical testing. Statistical analysis uses ANOVA. Group sizes were as follows: HCN1+/+, HCN1 miR–ve (n = 7 cells, 3 mice); HCN1+/+, HCN1 miR+ve (n = 9 cells, 3 mice); HCN1+/-, HCN1 miR–ve (n = 9 cells, 3 mice); HCN1+/-, HCN1 miR+ve (n = 9 cells, 3 mice).

We next examined responses of Purkinje cells to injection of current steps though the recording electrode. In all experimental groups negative current steps switched off action potential firing. The relationship between action potential frequency and the amplitude of the current step was similar in Purkinje cells from wild-type and global HCN1 knockout mice that were uninfected or that were infected with the HCN1 miRNA AAV (Figures 7A–C). In contrast, the relationship between the amplitude of current steps and the modal membrane potential was significantly modified by deletion of HCN1 and also by knockdown of HCN1 following transduction with the HCN1 miRNA AAV (Figures 7A,B,D). Thus, whereas in Purkinje cells from wild-type mice the membrane potential remained close to spike threshold even during current steps that switched off spike firing, in Purkinje cells expressing HCN1 miRNA or from mice with deletion of HCN1, the membrane potential reached more hyperpolarized potentials. Importantly, there was no significant difference between any of the HCN1 knockdown or HCN1 deletion groups in the relationship between current and membrane potential. These results are consistent with the idea that HCN1 channels are primarily activated in Purkinje cells by stimuli that inhibit spontaneous action potential firing and that they act to stabilize the membrane potential close to spike threshold (Nolan et al., 2003). They also indicate that the effects of AAV mediated expression of HCN1 miRNA are specific to HCN1 channels and do not involve changes in the properties of other membrane ion channels, or general pathological effects.

DISCUSSION

We have described a molecular toolbox of components to engineer lentiviral and AAV vectors for up- or down-regulation of neuronal
gene expression in vivo. The system is extremely flexible and allows rapid generation of viral constructs. Viruses produced from these vectors yield stable expression of transgenes at levels high enough to detect native fluorescence in neurons. This allows easy identification of virally transduced cells for downstream analysis of the effects of gene manipulation.

**A MODULAR APPROACH TO VIRUS ASSEMBLY**

Constructing viral vectors using traditional cloning methods is time consuming and inefficient as it requires restriction enzyme sites that must be compatible with insert sequences and the vector. Altering or rearranging elements within a vector often involves a new cloning strategy leading to unnecessary duplication of work. We have used a recombination cloning system to build a library of validated interchangeable components for the generation of viral vectors. New vectors can be quickly assembled using a single recombination reaction reducing both the time and expertise required. Entry vectors encoding the desired components must be generated initially, but they can then be utilized repeatedly in various combinations to create new expression constructs. By creating a Gateway-compatible AAV vector, the same entry vectors can now be used to produce both lentiviruses and AAV. Additional viral vectors, such as rabies or HSV could also be adapted to this system, further increasing the utility of the toolbox. In this way, the same components can be easily swapped between different viral systems, whilst minimizing the variability between them.

The toolbox could potentially be applied to diverse neuroscientific questions. Using different promoters we have expressed various reporter proteins, ion channels, and miRNA cassettes in neurons in vivo. The high level of native reporter fluorescence detectable from these viruses facilitates both imaging and electrophysiology-based techniques as well as ex vivo isolation of
transduced cells. Using FACS to detect the reporter, we isolated relatively pure populations of transduced granule cells from the dentate gyrus. These cells could then be subjected to further ex vivo analysis of their DNA, RNA, or protein profiles to examine the consequences of the gene manipulation in more detail. This approach has the advantage of performing the manipulation in an in vivo context whilst allowing the use of a wide range of sensitive in vitro assays for the subsequent analysis of transduced cells. It is also possible to isolate many more transduced cells than is practical using other current techniques such as laser capture microscopy. It is important to note however, that the purity of the cell population isolated will depend on both the specificity of the transgene promoter and the accuracy of virus injection and tissue dissection.

In contrast to generation and maintenance of knockout and transgenic mice, which can be time consuming and expensive, the use of viruses to alter gene expression in vivo has several advantages. In particular, viruses can be rapidly produced and are easily targeted to well-defined brain regions at specific time points. However, there are potential disadvantages to consider when planning viral experiments. Transgene expression levels may vary unpredictably depending on how many viruses transduce each cell and where lentivirally expressed DNA integrates into the cell’s genome. In addition, while careful injection of lentiviruses and AAV directly into the brain parenchyma results in little or no immune response (Chamberlin et al., 1998; Mazarakis et al., 2001; Abordo-Adesida et al., 2005; Lowenstein et al., 2007) any surgical procedure still has the potential to induce local inflammatory responses. This can be counteracted through the use of careful surgical techniques, viral production and purification methods (Scherr et al., 2002; Baekelandt et al., 2003) and injection of lower viral titers (Chamberlin et al., 1998; Abordo-Adesida et al., 2005; Lowenstein et al., 2007). Nevertheless, where appropriate residual effects must still be excluded through the use of experimental controls.

EFFICIENT AND SPECIFIC GENE KNOCKDOWN USING INTRONIC ARTIFICIAL microRNAs

Traditionally vectors for RNAi express shRNAs at high levels from strong Pol-III promoters. The shRNAs can result in toxicity by disrupting the biogenesis of natural miRNAs (Grimm et al., 2006), causing off-target silencing through accumulation of antisense RNAs (Birmingham et al., 2006) and stimulating innate immune responses to double stranded RNA (Samuel, 2001; Bridge et al., 2003; Fish and Kruthof, 2004). In vivo, these off-target effects can cause cell death and lethality (Grimm et al., 2006; Martin et al., 2011). Therefore, subsequent approaches have mimicked endogenous miRNAs by expressing artificial pri-miRNAs from Pol-II promoters. The pri-miRNAs are expressed at lower levels, but are more efficiently processed, minimizing their side effects (McBride et al., 2008). The use of Pol-II promoters also enables co-cistronic expression of a reporter gene in a cell-specific and regulated manner. A weakness of this approach is that processing of the pri-miRNA can block translation of the co-cistronic reporter gene as the resultant mRNA is unstable due to the lack of either a 5′ CAP or polyA tail (Du et al., 2006). Placing the miRNA within an artificial intron upstream of the reporter gene addresses this problem by allowing cellular splicing mechanisms to excise the miRNA whilst leaving an intact mRNA for co-expression of the reporter gene (Du et al., 2006). This approach has been used recently to generate a BAC transgenic mouse (Garbett et al., 2010), but viral delivery of artificial intronic miRNA to neurons has not been reported. Our results show that an AAV can be used to express an intronic miRNA targeting the HCN1 ion channel resulting in specific knockdown of Ih in cerebellar Purkinje neurons without apparent off-target effects.

The molecular toolbox we have developed allows for the rapid generation of lentiviral and AAV vectors. Nevertheless, it is still important to rigorously test each component in order to generate useful viruses. When expressed within a viral context, promoter sequences may not always yield the anticipated pattern of activity. Although our neuronal promoters gave the expected high levels of expression within principle neurons (Dittgen et al., 2004; Hioki et al., 2007; Nathanson et al., 2009a,b; van Hooijdonk et al., 2009), we also observed occasional expression in interneurons as has been reported previously (Nathanson et al., 2009a). Transgene expression driven by the Netrin G1 promoter sequence did not recapitulate endogenous Netrin G1 expression patterns and a Netrin G2 promoter sequence failed to express in vivo despite high activity in vitro (data not shown). Ensuring viral transgene expression is restricted to very specific subpopulations of neurons may require additional strategies, such as using viral constructs incorporating FLEX switches in combination with Cre-expressing transgenic animals (Schnütgen et al., 2003; Atasoy et al., 2008). As new technologies for the control of gene expression become available, they can also be incorporated into the molecular toolbox. We anticipate these tools will be of widespread use for investigation of nervous system function and pathology.

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