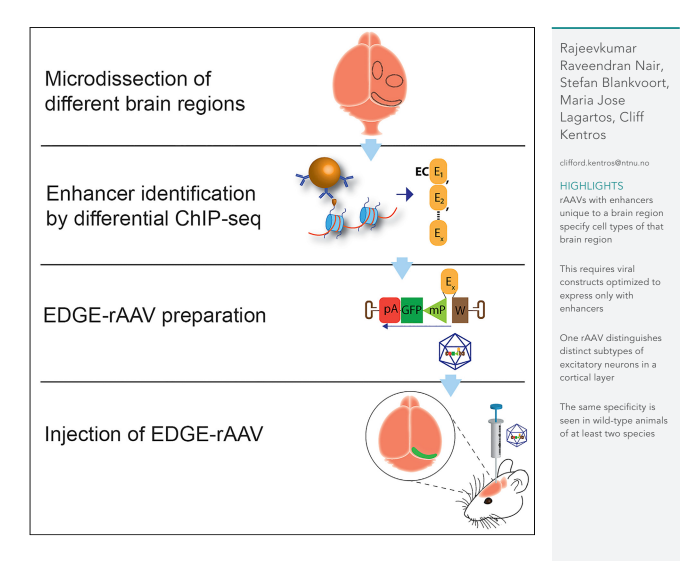
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Article

Enhancer-Driven Gene Expression (EDGE) Enables the Generation of Viral Vectors Specific to Neuronal Subtypes

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SUMMARY

Although a variety of remarkable molecular tools for studying neural circuits have recently been developed, the ability to deploy them in particular neuronal subtypes is limited by the fact that native promoters are almost never specific enough. We recently showed that one can generate transgenic mice with anatomical specificity surpassing that of native promoters by combining enhancers uniquely active in particular brain regions with a heterologous minimal promoter, an approach we call EDGE (Enhancer-Driven Gene Expression). Here we extend this strategy to the generation of viral (rAAV) vectors, showing that some EDGE rAAVs can recapitulate the specificity of the corresponding transgenic lines in wild-type animals, even of another species. This approach thus holds the promise of enabling circuit-specific manipulations in wild-type animals, not only enhancing our understanding of brain function, but perhaps one day even providing novel therapeutic avenues to approach disorders of the brain.

INTRODUCTION

The mammalian brain is the most complex biological structure known, with innumerable distinct cell types differing in cytoarchitecture, electrophysiological properties, gene expression, and connectivity (Luo et al., 2008; Zeng and Sanes, 2017). Understanding brain function requires understanding neural circuits at the level of specificity at which they operate. Recent years have seen the development of truly revolutionary molecular tools that allow neuroscientists to elucidate precise neural connectivity (Callaway and Luo, 2015) and monitor (Chen et al., 2013) and manipulate (Boyden et al., 2005; Roth, 2016; Sternson and Roth, 2014) neural activity. However, optimal use of these tools to examine the functional circuitry of the brain requires the ability to deliver them specifically to particular elements of neural circuits (i.e., neuronal cell types), rather than as a nonspecific bolus affecting all of the neurons in a brain area. The use of molecular genetics is the only method by which one can perform truly cell-type specific manipulations, as evidenced by a variety of studies using transgenic animals expressing transgenes from neuronal promoters (genomic regions just upstream of the transcriptional start site) (Kanter et al., 2017; Miao et al., 2017). However, such approaches are limited by the fact that, because individual genes are expressed in a variety of cell types in the brain, promoters are not specific to a single neuronal cell type. Although estimates vary (ENCODE Project Consortium, 2012), there are at least an order of magnitude more cis-regulatory elements (i.e., enhancers and repressors, distal genomic regions that help regulate where and when promoters transcribe DNA) than promoters, suggesting that enhancers may be more specific. This led us to take an approach to the generation of molecular genetic tools that we call Enhancer-Driven Gene Expression (EDGE), based on identifying the cis-regulatory elements uniquely active in particular brain regions and combining them with a heterologous minimal promoter. When we used this strategy to make transgenic mice, they were indeed significantly more specific than the presumed parent gene, often driving expression primarily in particular sets of neurons in the brain region they were derived from (Blankvoort et al., 2018).

However, although transgenic animals are powerful tools for the analysis of neural circuits, they have limitations. They are costly in both time and resources, can be subject to insertional effects (Matthaei, 2007; Feng et al., 2000), and are most practical in a limited number of species. Moreover, although they are often excellent models of disease, transgenic technologies are far from therapeutic applications. Recombinant adeno-associated viral vectors (rAAVs) can overcome many of the above issues. They can be made relatively quickly, generally do not insert into the genome or replicate, and can be used in a variety of species (Watakabe et al., 2015) including humans and therefore have clinical potential as well (Bouard et al., 2009; ¹Kavli Institute for Systems Neuroscience and Centre for Neural Computation, NTNU, Norway

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Dias et al., 2018; Kotterman and Schaffer, 2014; Mendell et al., 2017). However, efforts to generate cellspecific viral vectors by capsid modifications (Koerber et al., 2008, 2009, Klimczak et al., 2009) or using promoters (Delzor et al., 2012; Kugler et al., 2003; Shevtsova et al., 2005) have been largely unsuccessful to date to address a particular cell type, with a few notable exceptions (Dimidschstein et al., 2016; Hartl et al., 2017), and even those are likely to have multiple subclasses. This is in large part because the relatively small payload size of rAAVs puts most native promoters out of reach. However, most enhancers are much smaller than promoters, raising the intriguing possibility of targeting specific neuronal cell types in any species by adapting EDGE to viral vectors, provided the background expression of the viral backbone and promoter can be minimized. Toward this end, we present results demonstrating enhancer-based viral vectors that specifically express in particular neurons of the entorhinal cortex (EC) in two different species of wild-type animals.

RESULTS

Optimization of rAAV Design for Enhancer-Driven Gene Expression

Because one can obtain some degree of apparent specificity with rAAVs by means other than transcriptional regulation, we took steps to ensure that any observed specificity comes from the enhancer element used. Most notably, AAV serotypes exhibit distinct tropisms for different cell types: for instance, AAV8 is most efficient for oligodendrocytes and astrocytes (Aschauer et al., 2013; Hutson et al., 2012) and AAV 1, 2, 5, 7, 8, 9 prefer neurons (Aschauer et al., 2013; Castle et al., 2016; Davidson et al., 2000; During et al., 2003) (although they are by no means exclusive to them), whereas rAAV9 appears well suited for cortical neurons (Aschauer et al., 2013) and a variety of AAVs with engineered capsids show specific tropisms (Deverman et al., 2016; Tervo et al., 2016). We therefore used a single serotype (AAV2/1) with a wide tropism for neurons (Hauck et al., 2003) for the vast majority of our efforts toward engineering rAAVs transcriptionally specific to particular subtypes of neurons. We selected AAV 2/1, a chimera between capsid-1 (less efficient neuronal transduction [Castle et al., 2016]) and capsid-2 (vast tropism [Wang et al., 2003]) because of its broad transduction efficiency (Hauck et al., 2003) and to prepare viruses with high purity (During et al., 2003; Mcclure et al., 2011) via heparin columns (see Transparent Methods).

Because injections of small volumes of rAAVs can appear specific because of the specific parcellation around the injection site, we used a medial entorhinal cortex (MEC) enhancer (MEC13-53) known to be specific to a particular subset of neurons in the entorhinal cortex (Blankvoort et al., 2018) in transgenic animals so we knew what to look for. Figure 1A shows the expression pattern obtained from crossing one of the MEC13-53 tTA driver lines to a payload line expressing the helper transgenes for the Δ G-rabies monosynaptic tracing system (Blankvoort et al., 2018). Expression in this cross was limited to Reelin-positive (RE+), Calbindin-negative (CB-) excitatory projection neurons in layer (LII) of the EC (Kitamura et al., 2014; Varga et al., 2010; Witter et al., 2017). Finally, we injected the same large (400 nL in mice, as opposed to the ~50-nL injections typically used with nonspecific rAAVs) volume of each virus into multiple animals using the same EC coordinates and compared only green fluorescent protein (GFP)-expressing rAAVs of similar titer (see Table S1 and Transparent Methods). For the purposes of comparison, Figure 1B shows the widespread strong expression throughout the various layers of the entorhinal cortex (as well as subiculum and parasubiculum) resulting from injecting a control AAV with a relatively (it has been shown to prefer neurons) nonspecific cytomegalovirus promoter (CMV-rAAV) of the same serotype and similar titer.

The initial step in obtaining viruses capable of driving expression as specific as the EDGE transgenic animals in wild-type brains is to find a minimal viral promoter that is capable of robust expression *only* when paired with a heterologous enhancer. This is complicated by the fact that the viral inverted terminal repeats (ITRs) themselves have transcriptional activity (Carter et al., 1993; Flotte et al., 1993; Haberman et al., 2000), as can be seen by the very weak (but still above autofluorescence) nonspecific expression obtained from a viral construct with neither a promoter nor an enhancer (Figure 1C). Note that the expression levels in Figure 1C are far below those seen with the other viruses: each panel in Figure 1 has been differentially post-acquisition processed to aid visualization, the "background" expression seen in Figure 1C would otherwise be imperceptible (see Figure S1 for comparison of each image with the same processing). To minimize this issue, we reversed the orientation of the expression cassette relative to the ITRs such that the sense strand was under the influence of the 3' ITR, which we attenuated by putting WPRE (Zufferey et al., 1999) between the 3'ITR and the enhancer (see schematics in 1C, D). The substantial reduction in background expression enabled us to recapitulate MEC LII-specific expression in a wild-type mouse (Figure 1D) with a mutated minimal CMV promoter (CMV*) (Loew et al., 2010). Roughly similar results varying in amount

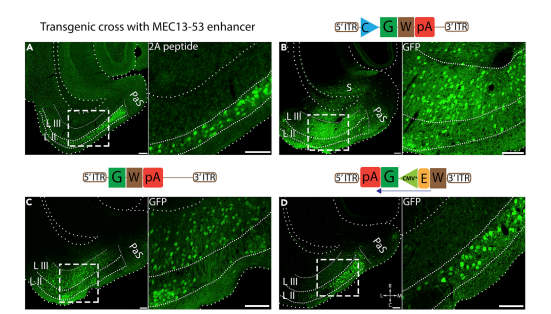


Figure 1. Optimization of rAAV Constructs for Enhancer-Dependent Gene Expression

(A) Transgene expression in a MEC13-53 tTA X tetO-TVAG transgenic cross visualized by anti-2A immunostaining is restricted to RE + LII projection neurons in EC (Blankvoort et al., 2018). Since this is a different antibody, this is purely a qualitative comparison.

(B) Injection of a nonspecific (CMV-rAAV) virus into the EC shows broad label throughout the entire region, including all layers of EC, as well as subiculum (S) and parasubiculum (PaS).

(C) The same construct without a minimal promoter shows weak nonspecific expression throughout the region that would not be visible at normal image settings (see Figure S1).

(D) Changing the orientation of the expression cassette leads to a marked reduction in nonspecific expression of MEC13-53 rAAV (see inset in [C] and [D], note that most of the LIII label in [D] is not cellular, unlike in [C], and when it is, it is very light, i.e., from baseline transcription). All murine injections were 400 nL. NB: images were differentially modified to best visualize the GFP expression pattern in each panel; comparisons of these images with the same post-acquisition settings are shown in Figure S1 (see Transparent Methods). Note that all label above background auto-fluorescence was treated as positive, even though there were two markedly distinct intensities of label. See also related Figures S1 and S2. Schematics of the viral designs are depicted on top of the corresponding image. ITR, inverted terminal repeat; W, woodchuck

hepatitis virus post-transcriptional regulatory element; pA, human growth hormone polyadenylation signal; E, enhancer; G, Green fluorescent protein; C, cytomegalovirus promoter; CMV*, mutated minimal cytomegalovirus promoter. Scale bar, 100 μ m.

and specificity were obtained with other minimal promoters (Figure S2), but we selected CMV* for all subsequent experiments (and hereafter simply refer to the enhancer) as it was the smallest one that worked well. The specificity of the expression of this virus as compared with a nonspecific CMV-rAAV virus is quantified in Figure 2. Although still clearly far more specific than the CMV-rAAV, the quantification of MEC13-53 rAAV does not seem as specific as it looks in the figure panels because in our counts we did not distinguish between weak "background" label (such as that seen in Figures 1C and S1 without a promoter) and the strong specific labeling (see below).

MEC13-53 EDGE rAAVs Express Specifically in Layer II Stellate Cells in Wild-Type Mice and Rats

The neuron-specific stain NeuN (Boccara et al., 2015) confirms the robust LII-specific expression of the MEC13-53 rAAV (Figures S3A and S3C) in neurons (100% of labeled cells were NeuN+, data not shown). Weak, "background" GFP expression was observed in other layers as well in both this virus (Figure S3A, inset) and in the rAAV backbone (i.e., the same virus lacking the enhancer, Figure S3B, inset), which in contrast did not strongly label any cells. Within LII of MEC there are two major classes of excitatory principal neurons, RE + stellate cells and CB + pyramidal cells (Rowland et al., 2018; Witter et al., 2017), with RE label providing a sharp boundary between MEC and parasubiculum (Varga et al., 2010; Witter et al., 2017) (see arrows in Figures 2A and 2E inset). We therefore performed immunohistochemical analysis comparing

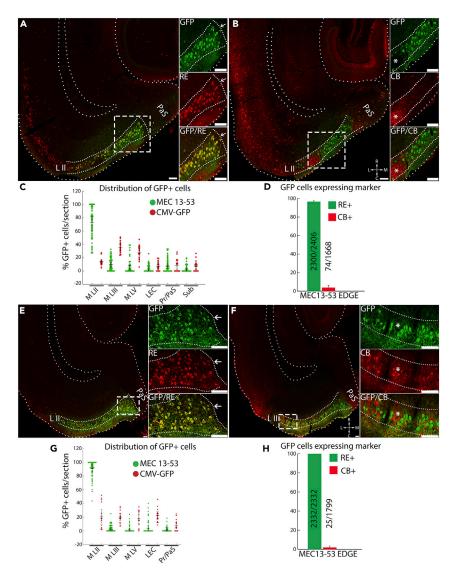


Figure 2. MEC13-53 EDGE rAAVs Recapitulates the Cell-Type Specificity Seen in the MEC13-53 EDGE Transgenic Crosses in WT Mice and Rats

Equal amount of MEC13-53 rAAV was injected into MEC of wild-type mice (A–D) and rats (E–H). Insets show anti-GFP (top), marker (middle), and overlay (bottom) of box in main panel. Sections of MEC13-53 rAAV injections counterstained with anti-RE antibody (red, [A] and [E]) and anti-CB antibody (red, [B] and [F]); with a CB + cluster (asterisks) in the insets in (B) and (F). Note the extensive co-localization of the RE stain with the GFP, the sharp delineation of the entorhinal/ parasubicular boundary by both labels (arrows, [A] and [E]), and the exclusion of viral label from the CB clusters (asterisks, [B] and [F]).

(C) Proportion of GFP-expressing cells in different parahippocampal regions for both MEC13-53 and nonspecific CMV-rAAV. Each point is a section, note the large number of sections where 100% of the cells are in LII and 0% in other regions exclusively in the MEC13-53 rAAVs compared with the controls (for pictures of control injections see Figures 1B, S4, and S5). A total of 13,096 and 8,540 GFP + cells were counted from three mice injected with CMV-rAAV and seven mice with MEC13-53 rAAV, respectively; data represented as mean \pm SEM. In (G), 7,191 and 2,831 GFP + cells from sections were counted from MEC13-53 rAAV and CMV-rAAV, respectively, from three rats. Quantitation of results are shown in (D) (for mice) and (H) (for rats), showing overlap of GFP with cell-marker Reelin stain (green) in LII MEC of mice (96%) and rats (complete overlap). About 4% overlap of GFP with Calbindin (red) was observed in mice and <2% overlap in rats, with number of cells counted in MECLII region. MEC-LII GFP + cells were counted from separate RE and CB immunostained sections from seven mice and three rats injected with MEC13-53 rAAV; data represented as mean \pm SEM. See also related Figures S3–S6. Scale bar, 100 µm; all images were processed identically.

these markers with viral GFP and found that, for the MEC13-53 rAAV, 96% (2,300/2,406) of GFP + cells in layer II were RE+ (Figures 2A and 2D), whereas 4% (74/1,668) were CB+ (Figures 2B and 2D). In contrast, for injections of roughly equal amounts of the ubiquitous CMV-rAAV, only 34% (319/929) of GFP + LII cells were RE +, whereas 10.5% (142/1,353) were CB+. Thus, the MEC13-53 rAAV drives transgene expression specifically in a particular subset of excitatory neurons in EC of wild-type mice, i.e., RE + EC LII neurons (stellate cells in MEC), avoiding the adjacent CB + pyramidal cells, like the transgenic lines based on the same enhancer.

Although this nicely illustrates the specificity of this EDGE rAAV, perhaps the greatest utility of EDGE rAAVs is that, because enhancers are highly conserved (Cotney et al., 2013) and can be obtained from any tissue sample, they have the potential to work across species. As seen in Figures 2E-2G, S3C, and S4C, the MEC13-53 rAAV derived from mouse EC is, if anything, more specific in the rat. Figures 2E and S3C shows GFP expression almost exclusively in MEC LII (as quantified in Figures 2G and S6B), whereas the few labeled neurons in the virus with no enhancer have no layer specificity (Figure S3D), just as in mouse (Figure S3B). Similarly, 100% (2,332/2,332) of MEC LII GFP + neurons in rats injected with MEC13-53 rAAVs were RE+ (Figures 2E and 2H), whereas only 1.4% (25/1,799) were CB+ (Figures 2F and 2H), even though the two excitatory subtypes are intermingled (Witter et al., 2017). This, and the presence of LII-specific label throughout the dorsoventral and medio-lateral axes of the MEC (Figure S4C), provides compelling evidence for cellular specificity. Note that, with the nonspecific CMV-rAAV, 35% (189/518) of GFP + LII cells were RE +, whereas 46% (285/613) were CB+ (Figure S5). It is interesting to note that, although these two markers are largely mutually exclusive, there are reports of a very small subpopulation of RE + neurons that are also CB+ (Fuchs et al., 2016; Varga et al., 2010), so the single-digits label with the MEC13-53 virus may be those cells. Clearly, though, the two rAAVs with the same serotype have very different expression patterns, both in terms of layer and cellular specificity.

Systemic Administration of Blood-Brain Barrier Crossing MEC13-53 EDGE Recapitulates MEC Layer II Stellate Cell Expression

Although we are mainly interested in developing tools to be used in analysis of the EC, it is interesting to ask whether this enhancer would express in other brain regions if it were systemically administered. We therefore packaged the MEC13-53 EDGE enhancer (shown with the 2/1 serotype in Figures 1 and 2) into the blood-brain barrier crossing PHP (Deverman et al., 2016) serotype and performed noninvasive intravenous injections via the tail vein. Systemic injections of MEC13-53 EDGE PHP resulted in much sparser GFP + cells overall, but they are also mostly confined to layer II of MEC throughout the caudal forebrain (Figures 3A and S7). However, we also noticed sparse expression of the transgene in regions other than MEC, typically also in brain regions we would sometimes see transgene expression in MEC13-53 transgenic lines (Figures 3B and 3C, Table S2). Curiously, we did not see expression in LII of the piriform cortex, the major site of non-EC expression in the MEC13-53 transgenic lines, possibly due to the particular tropism of the PHP capsid. Furthermore, we confirmed that these GFP + cells in MEC are RE+ (Figure 3D). These results suggest that EDGE rAAV can retain its particular cell-type specificity, even when assembled in a serotype with a different innate tropism.

EDGE rAAVs Recapitulate the Expression Pattern of Their Respective Transgenic Lines

To examine whether this is a general strategy, we created EDGE rAAVs with several other enhancers with known specificity (Blankvoort et al., 2018). Although not all enhancers that worked as transgenic lines worked in rAAVs, roughly half (Figure 4, left column) did indeed appear to recapitulate the specificity (or relative lack thereof, 4A, B) of the corresponding EDGE lines (Figure 4, right column). The MEC13-104 rAAV (Figure 4A) recapitulates the relatively sparse labeling of a subset of LIII neurons (arrows) seen in the MEC13-104 line (Figure 4B), whereas the converse is true for the mainly LIII-specific LEC13-8 (compare 4C with 4D) line. Thus, the relative densities of the layer-specific label appear to be enhancer specific, suggesting that the minority of cells that strongly express outside of their primary layer may not be "noise." Ongoing experiments explore the functional distinctions between the cells labeled by the various enhancers, which may label distinct subsets of what has been considered a single neuronal cell type, e.g., stellate cells.

DISCUSSION

Our prior work showed that identification of cis-regulatory elements uniquely active in finely dissected cortical subregions allows one to generate genetic tools specific to cells in that subregion, an approach

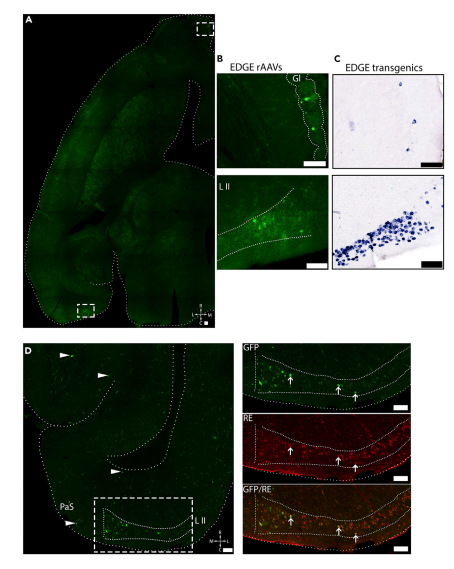


Figure 3. Recapitulation of LII MEC Specificity of MEC13-53 Using a BBB-Crossing rAAV Serotype (A and B) (A) Representative image of the GFP + neurons in horizontal brain section from a mouse injected with 10¹² particles of MEC13-53 rAAV PHP, intravenously into tail vein. The boxes in (A) are zoomed in (B). (C) MEC13-53 transgene expression in same regions as in (B) are in the MEC13-53 tTA X tetO-TVAG transgenic cross. (D) Sections of MEC13-53 rAAV PHP injected brain counterstained with anti-RE antibody. Insets show anti-GFP (top), Reelin (middle), and overlay (bottom) of box in main panel. Label is throughout the layers of EC and sparsely in other regions (arrow heads, [D]). Note the extensive co-localization of the RE stain with the GFP (arrows).

See also Figure S7. Scale bar, 100 μ m. See also related Table S2.

we call EDGE (Blankvoort et al., 2018). Here we show that one can use the same approach to make rAAVs with similar specificity in both mouse and rat, provided the vector and minimal promoter's innate transcriptional activity is minimized. This clearly cross-validates the initial identification of enhancers in our prior work (Blankvoort et al., 2018): although transgenic lines might show highly specific expression patterns purely due to insertional effects (although not the same pattern in multiple founders, as we saw), rAAVs typically do not insert into the genome (Mccarty et al., 2004), so cannot show such effects. In other words, although the precise functional significance of the enhancers presented here remains unknown, they clearly are "true" enhancers, reflecting some genetic subgroup of excitatory neurons in the entorhinal cortex of wild-type mice and rats. Taken together, these data lead to two very interesting conclusions: (1) given that the numbers of enhancers may run into the millions (as opposed to ~44,000 promoters) (ENCODE Project Consortium, 2012), they may provide access to the ever-growing number of neuronal cell types

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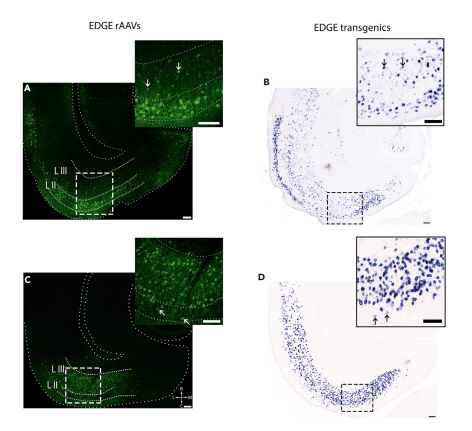


Figure 4. EDGE rAAVs Recapitulate the Distinct Layer-Specific Expression Patterns Seen in EDGE Transgenic Mice

Comparison of expression patterns obtained by injection of EDGE rAAVs (left column) with those seen in transgenic crosses made with the same enhancers (right column). Wild-type mice were injected with 400 nL of EDGE rAAVs (A) MEC13-104 and (C) LEC13-8. Transgene expression in the corresponding EDGE transgenic crosses ([B], MEC13-104 tTA X tetO-TVAG) and ([D], LEC13-8 tTA X tetO-HM3) visualized by ISH on horizontal sections using the respective transgene probes. The sparse expression of the transgene in minor layers is indicated by arrows both in EDGE transgenics and viruses. Scale bar, 100 μ m; all sections are horizontal, and all rAAV figures underwent the same image processing. See also related Table S1 and Data S1.

than promoters, which may be far greater than generally assumed (Zeisel et al., 2015; Cembrowski et al., 2016; Tasic et al., 2018; Saunders et al., 2018); and (2) although we do not do so here, one could conceivably take this approach toward generating neuronal subtype-specific transgene expression in species other than the traditional genetic models of mouse, zebrafish, fly, and worm, because one can do the required epigenomic analyses on any tissue sample.

There has been a lot of effort over the years toward making cell-type-specific viral vectors, but even in those cases when a minimal native promoter is useful (i.e., when a single marker defines the cells, e.g., TH-AAV [Gompf et al., 2015] and CaMKII-AAV [Nathanson et al., 2009]) the AAVs are not fully restricted to cells expressing the gene. We have previously shown that using single, uniquely active enhancers can lead to far greater specificity than that of native parent promoters (Blankvoort et al., 2018), at least in transgenesis. That enhancers drive expression similarly in both transgenic lines and viruses is not a particularly surprising result. It has been known for decades that enhancers drive cell-specific expression (Grosveld et al., 1987; Noonan and Mccallion, 2010; Shen et al., 2016) in a variety of species. Enhancers for the six homeobox genes related to the fly *distal-less* gene (Cohen and Jurgens, 1989) (*Dll* in fly, Dlx in vertebrates) have been shown to play a crucial role in morphogenesis in many species (Anderson et al., 2000). One such enhancer in the Dlx 5/6 gene cluster has been shown to be critical to the development of interneurons in particular (Stenman et al., 2003), and a recent paper (Dimidschstein et al., 2016) used this enhancer element in a viral vector to obtain interneuron-specific expression in a variety of species, nicely showing

that enhancers can be used to drive expression in viral vectors. However, as is true for most genetically defined enhancers active early in development, Dlx5/6 drives expression across broad classes of neurons (e.g., interneurons in general) throughout the brain, rather than to particular interneuronal subclasses and/or subregions.

More recently, several groups have begun to incorporate cis-regulatory elements into their strategies for creating viral vectors specific to neuronal subtypes. Such efforts are likely furthest along in the retina, where Juttner and colleagues (2019) created a broad rAAV resource targeting subtypes of retinal neurons using strategies based on genes of interest (GOIs) identified in a priori transcriptomal analysis (Siegert et al., 2012) and epigenetic analysis (Hartl et al., 2017) of known retinal cell types. Although most of these constructs are simply the minimal promoters of the GOIs, some also are based on the local epigenetic landscape, using strategies such as conservation, methylation patterns, and transcription factor binding sites to identify likely cis-elements for GOIs. Although the results in retina can be quite impressive, little is known how specific such vectors would be in the rest of the brain. As for the brain, Hrvatin et al., 2019 recently published an interesting screening strategy called PESCA (Paralleled Enhancer Single Cell Assay), in which multiple rAAVs containing barcoded putative enhancers (they use the term Gene Regulatory Elements, or GREs) are screened via single-cell transcriptomics (scRNAseg) rather than the more traditional one-at-a-time anatomical techniques shown here. Although scRNAseq does not always reflect actual viral expression, this technique nevertheless promises to greatly increase the throughput involved in first-pass screening of rAAVs. In a very interesting study, Graybuck and colleagues compare scRNAseq data to the epigenetic single-cell Assay for Transposase-Accessible Chromatin with Sequencing (scATACseq) data from layer-specific transgenic mice. Hits that co-register in both the transcriptomic and epigenetic clusters are then cloned into PHP.B Cre-rAAVs and systemically (retro-orbitally) injected into a Cre-reporter mouse for anatomical characterization

The overwhelming similarity of these various approaches is the idea that individual cis-regulatory elements may be more specific than promoters. Each strategy has two stages: identifying likely cis-regulatory elements and then making and screening the resulting rAAVs. The major difference clearly comes at the identification stage: each of these other GRE (as opposed to promoter) -based approaches has been based on a priori knowledge of the transcriptomics of whatever cell type one is looking for, often even taking advantage of transgenic animals, whereas EDGE simply looks for regionally specific chromatin marks in reproducibly dissected bulk tissue. The advantage of the former is resolution: by a "deep dive" into subtypes of what we had originally thought were cell types, one both gets at the scale of neuronal diversity and immediately puts the cell types in context, whereas with pure differential screens of bulk tissue such as EDGE you really do not know what cell types you will get, you just know that they are more or less specific to your tissue of interest. However, the flexibility (one simply needs ChIP of an ROI), ease of doing EDGE in other species, and ability to discover truly new cell types counterbalance this disadvantage. A more purely technical difference is between bulk ChIPseq and ATACseq. Although the latter can be done with much less tissue (even single cells), the former's use of particular histone marks may provide greater specificity for active enhancers rather than other forms of open chromatin. At the screening level, systemic viral injections (Graybuck, 2019) with AAV serotypes that cross the blood-brain barrier clearly give you the best idea of where a particular enhancer can express throughout the brain. We regret that we are as of yet unable to obtain permission to perform retro-orbital AAV injections from our local regulators, so our systemic injections were with a less effective technique (tail vein), lowering the effective titer. If PESCA (Hrvatin et al., 2019) can reliably be done on bulk tissue, however, it may end up as a better screen for our purposes. All in all, there are advantages to each approach that make them largely complementary, suggesting that combinations of these techniques and comparisons between the resulting datasets (ChIP versus ATAC, bulk versus single cell) may well end up being the best overall approach.

Thus, the most important aspect of these and other papers is not that enhancers can work in viral vectors, it is illustrating the promise of applying modern genomic techniques to the study of the precise neural circuitry of the vertebrate brain. The striking diversity of enhancers found in these tiny subregions of cortex (numbers comparable with those found for entire organs) may indicate a similar diversity of neuronal cell types in the brain. However, the relationship between enhancers and cell types remains unclear. Indeed, the expression patterns we obtain are arguably more specific than our current understanding of neuronal cell type (Luo et al., 2008; Zeng and Sanes, 2017). For instance, stellate cells are a generally accepted excitatory neuronal cell type of the medial entorhinal cortex (Rowland et al., 2016; Varga et al., 2010; Witter et al.,

2017). However, we show that distinct enhancers drive expression in EC LII stellate cells to different degrees in both transgenics and rAAVs. The question becomes whether these enhancer-driven expression patterns reflect functionally distinct stellate cells, or states of stellate cells, or just random subsets of the same indivisible cell type. In the specific case of stellate cells, a recent paper used optogenetic tagging to show that stellate cells of the MEC exhibit a variety of quite distinct receptive field properties (i.e., they can be grid cells or spatial cells or border cells), suggesting that there are many functional subtypes of stellate cells (Rowland et al., 2018). More generally, the relationship between differential enhancer usage and neuronal cell types is a highly non-trivial question, not least because there is not even complete agreement even as to how to define neuronal cell types (although there are notable exceptions) (Cembrowski et al., 2016; Tasic et al., 2018; Tremblay et al., 2016), let alone how many there are. There are several other interesting explanations for differential enhancer usage beyond cell type; for instance, it could dictate distinct states of a single cell type. In support of this, neural activity drastically changes the chromatin landscape of the brain, including which enhancers are active (Gallegos et al., 2018; Malik et al., 2014). It will likely take years of anatomical, molecular, and physiological characterization of these tools to disentangle such questions, so for our current purposes the most important consideration is that these enhancer-based molecular genetic tools remain true to type, as appears to largely be the case, comparing the virus to the transgenic.

It should be noted, however, that specificity is almost never absolute, especially with viral vectors. Although we obtain neuronal subtype-specific results with large injections into the entorhinal cortex (Figures 2 and S4), it is likely that any cell type in other brain regions that express the transcription factor(s) appropriate for a particular enhancer would be labeled as well, as can be seen with the systemic injections shown in Figure 3. Thus, we do not claim that the rAAVs shown here are necessarily 100% regionally specific; indeed, it is hard to imagine that a particular enhancer is only used once in development. Rather, we demonstrate clear cell-type specificity when the MEC13-53 rAAV is injected into a particular brain region, which is nevertheless good enough for the study of neural circuitry. Moreover, many more cells are infected than show strong GFP label, and there is a baseline level of transcription from other elements in the viral construct (i.e., the minimal promoter and the ITRs). This implies that superfection of enough rAAVs could lead to discernible nonspecific transgene expression in any cell regardless of the promoter, something that is shown most clearly by making viruses containing no exogenous promoter whatsoever (Figure 1C). Viral expression is thus not all-or-nothing, but the difference between background and enhancer-driven expression levels can be quite marked (Figure S1). This background expression inherent to rAAVs can be quite problematic when a little bit of expression can have a large effect. This is true when expressing enzymes such as recombinases or when complementing replication-competent viruses (e.g., pseudotyped ΔG rabies [Weible et al., 2010]) but is likely not an issue with transgenes whose effects vary roughly linearly with their expression levels, such as the chemogenetic (Sternson and Roth, 2014) and/or optogenetic tools (Boyden et al., 2005) used to study neural circuits.

Thus, identification of the active enhancers of a mere four cortical subregions of the mouse brain has led to a variety of transgenic, and now viral tools for circuit analysis that appear to work across species, at least in rodents. Since, in principle, one can do this on any reasonably well-annotated genome, one could conceivably develop tools for anatomically specific "circuit-breaking" tools in any species, even our own. Thus, not only will circuit-specific tools greatly facilitate our understanding of normal and pathological brain function, but they could also in time possibly provide circuit-specific therapeutic avenues. For example, it has been known for decades that preclinical stages of Alzheimer's disease (AD) are characterized by neuronal loss and accumulation of neurofibrillary tangles in the superficial layers of trans-entorhinal cortex (Braak and Braak, 1991), a region roughly equivalent to rodent MEC layer II. In addition, intracellular amyloid- β is found specifically in MEC layer II RE + neurons in human AD pathology and rodent disease models (Kobro-Flatmoen et al., 2016). Given the emerging consensus that AD may progress trans-synaptically (De Calignon et al., 2012; Spires-Jones and Hyman, 2014), it is conceivable that one could use something like a MEC13-53 rAAV to deliver therapeutic agents directly to the presumed pre- α cells, thereby stopping AD before it starts. More generally, it is possible that the reason that many neurological and neuropsychiatric disorders are resistant to drug therapy is that they are imbalances in particular neural circuits, not diseases of the entire brain. A drug having tropism for multiple circuits (as most do) would then by definition produce unwanted side effects: it may do the right thing in the right circuit, but it does the wrong thing to normal circuits. Results like those presented here allow the hope that investigators may one day be able to design interventions with the specificity required to treat the complex diseases of the brain.

Limitations of the Study

Although we think that we have made a substantive contribution toward the generation of circuit-specific tools that could be used outside of traditional genetic models, we freely acknowledge the limitations of our data. Although it is indeed true that active enhancers can be identified in any tissue sample of reasonable size from any species and used to make EDGE-rAAVs in ways similar to that presented here, we have only showed the same specificity for stellate cells in two rodent species—larger animals such as primates pose significant challenges with viral vectors. In addition, although we can see remarkable cellular specificity when EDGE rAAVs are injected into the region they were designed for, systemic administration suggests that the enhancer may also express in other cell types if injected in other regions. Regardless, we feel that these are quite useful tools for the analysis of neural circuits.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.100888.

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AUTHOR CONTRIBUTIONS

C.K., R.R.N., and S.B. conceptualized the study. R.R.N did the data curation. Formal analyses were done by R.R.N., M.J.L., and S.B. C.K. did the funding acquisition. Investigation was done by R.R.N., M.J.L., and S.B. Methodology was by R.R.N., S.B., and C.K. Project administration and supervision were done by C.K. and R.R.N. Resources for the study were from R.R.N., M.J.L., and S.B. Validation was by R.R.N., S.B., M.J.L., and C.K. Original draft was prepared by R.R.N. and C.K. Review and editing was by C.K., R.R.N., S.B., and M.J.L.

DECLARATION OF INTERESTS

C.K., S.B., and R.R.N. are inventors on US Patent Application no. 62/584,282, Appl. Norwegian University of Science and Technology (NTNU), which is related to this work. The authors have no other competing interests to declare.

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Supplemental Information

Enhancer-Driven Gene Expression (EDGE)

Enables the Generation of Viral Vectors

Specific to Neuronal Subtypes

Rajeevkumar Raveendran Nair, Stefan Blankvoort, Maria Jose Lagartos, and Cliff Kentros

1 Supplemental information



Image settings to visualize all in similar intensity

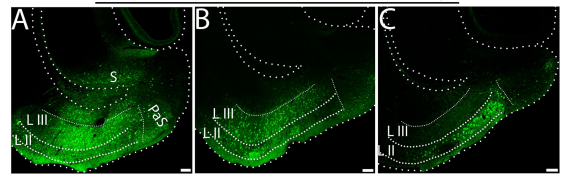


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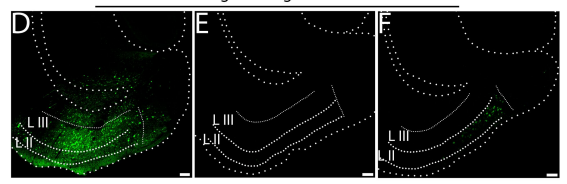


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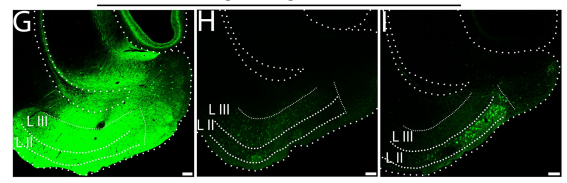


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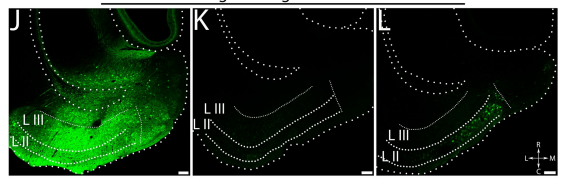
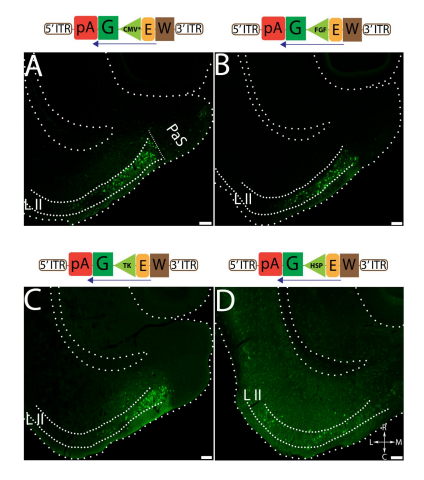
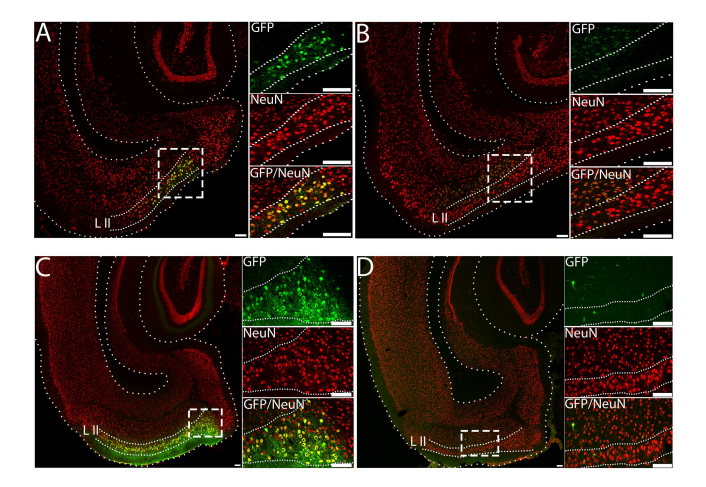


Figure S1. Visualization of GFP signal depends critically on post-processing 4 (related to Figure 1). The same images shown optimally in Figure 1B-D are shown 5 with the same post-acquisition processing for the purposes of comparison of strong 6 versus weak viral GFP expression seen with the different viral constructs. Left column 7 is CMV rAAV, middle column is the promoterless-rAAV, and the right column is the 8 MEC13-53 rAAV. Top row (S1A-C) shows images at optimized settings as shown in 9 Figure 1B-D; second row (S1D-F) shows images at optimization settings for CMV-10 rAAV applied to all images; third row (S1G-I) shows optimization settings for 11 promoterless-rAAV; fourth row (S1J-L) shows optimization settings for MEC13-53 12 rAAV. Note that at the settings for both CMV-rAAV and MEC13-53 the background 13 GFP seen in the promoterless-rAAV (S1E, K) is not visible, while at the settings 14 optimized for promoterless-rAAV the label in the parenchyma makes it impossible to 15 visualize individual cells in the CMV-rAAV, and makes it look like there is (much lighter) 16 label in LIII in the MEC13-53 rAAV (see Figure 1D inset). Schematics of the viral 17 designs are depicted on top of the corresponding image. Scale bar = $100 \mu m$. 18

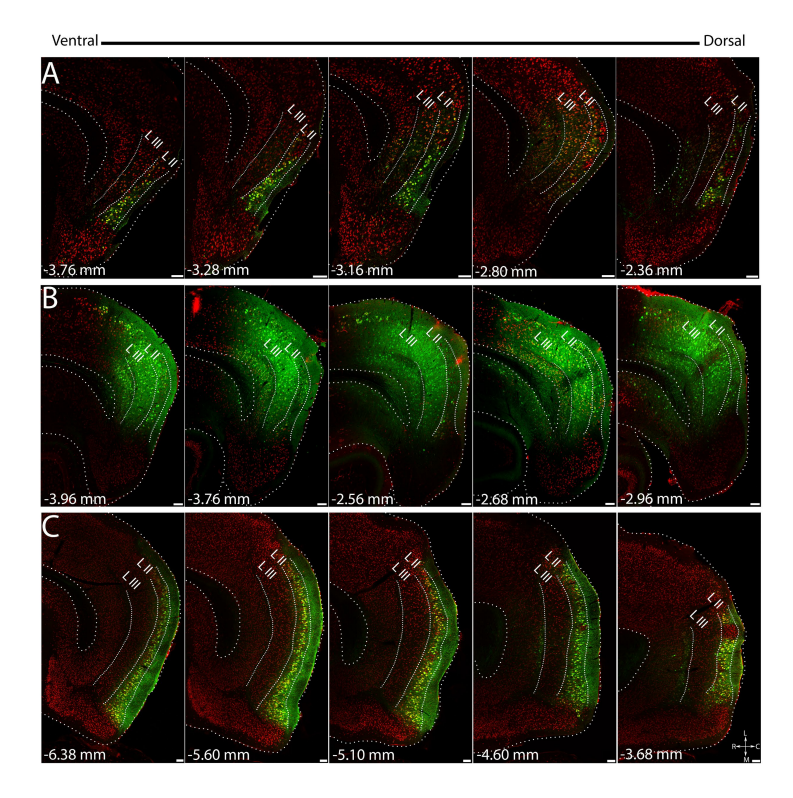


21 Figure S2. Optimization of minimal promoter for EDGE-rAAV constructs (related

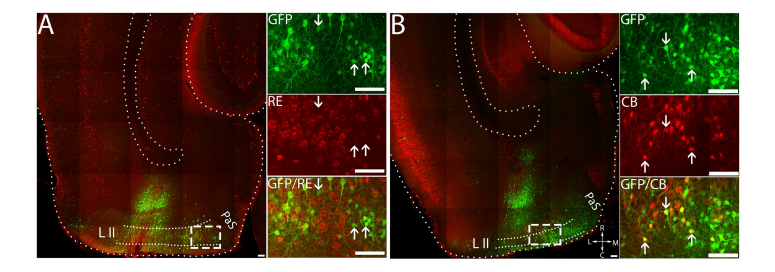
to Figure 1). 400 nl of MEC13-53 EDGE rAAVs with the various minimal promoters (A) CMV*, (B) FGF4, (C) TK or (D) HSP68 (see Methods for details) were injected into MEC in wild-type mice. While each minimal promoter led to layer-II specificity when combined with the MEC13-53 enhancer, we chose to use minimal CMV* because of its smaller size and limited nonspecific expression in other layers. Schematics of the viral designs are depicted on top of the corresponding image. See related supplemental Data S1 for sequences. Scale bar = 100 μ m.



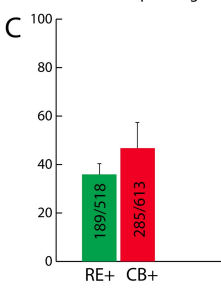
- 30 Figure S3. MEC13-53 drives transgene expression in LII MEC neurons (related
- 31 to Figure 2). Equal amounts of MEC13-53 rAAV (A, C) or CMV*-rAAV (B, D, i.e.
- 32 *identical to A except without an enhancer) were injected into MEC of wild-type mice*
- (*A*, *B*) and rats (*C*, *D*). Insets show anti-GFP (top); NeuN marker (middle); and overlay
- 34 (bottom) of box in main panel. Note the extensive co-localization of the NeuN stain
- 35 with the GFP. Scale bar = $100 \ \mu m$.



37 Figure S4. EC LII specificity of MEC13-53 throughout the caudal forebrain (related to Figures 1 and 2). Representative images of the GFP+ and NeuN+ 38 neurons in horizontal sections at multiple dorso-ventral levels from, (A) a mouse brain 39 injected with 400 nl and (C) a rat brain injected with 1000 nl of MEC13-53 rAAV. 40 MEC13-53 drives transgene expression preferentially in MEC LII throughout dorso-41 ventral axis. (B) Representative images of the GFP+ and NeuN+ neurons in multiple 42 horizontal sections in dorso-ventral axis from a mouse brain injected with CMV-rAAV. 43 Label is throughout the layers of EC and also in subiculum. Sterotaxic coordinates 44 were identified based on anatomical features using Paxinos G & Franklin K (for mouse 45 brain) and Paxinos G & Watson C (for rat brain). Scale bar =100 µm. 46







48 Figure S5. CMV-rAAV is not specific to any cell-type in rat (related to Figure 2).

1000 nl CMV-rAAV was injected into MEC of wild-type rats. Insets show anti-GFP 49 (top); marker (middle); and overlay (bottom) of box in main panel, marker is RE in A 50 and CB in B. The arrows in A represent GFP+ cells that non-overlap with RE, while 51 arrows in B show cells that are GFP+ and CB+. (C) Quantitation of results shown in 52 A, B, showing 35 % overlap of GFP with cell-marker RE stain (green) in LII MEC, and 53 46 % overlap of GFP with CB (red), with number of cells counted. MEC-LII GFP+ cells 54 were counted from separate RE and CB immunostained sections from 3 rats injected 55 with CMV-rAAV, data represented as mean \pm SEM. Scale bar = 100 μ m. 56

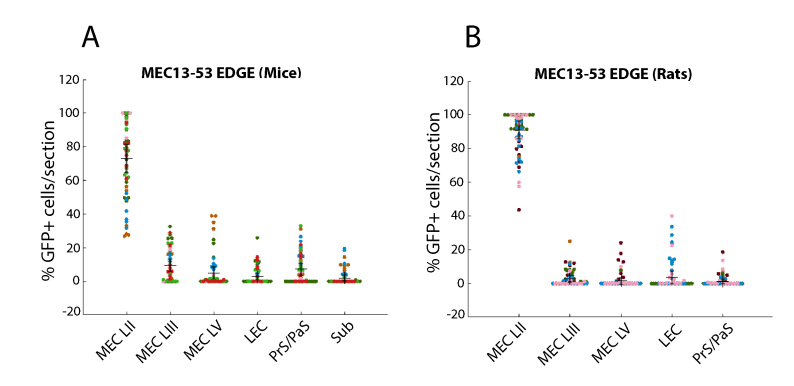


Figure S6. Layer II specific expression of MEC13-53 rAAV in multiple injections 58 in two species (related to Figures 1 and 2). MEC13-53 rAAV was injected into MEC 59 of wild-type mice or rats. Sections were stained with anti-GFP, and GFP+ cells were 60 counted in multiple sections per animal across various subdivisions of the 61 parahippocampal area (see Transparent methods). The percentage of GFP+ 62 cells/section analysed for different groups were plotted. (A, B) Distribution of 63 percentage GFP+ cells/section across the different parahippocampal regions in all 64 MEC13-53 EDGE injected mice (A) and rats (B). MEC13-53 rAAV LII sections with 65 relatively few GFP+ cells are distal to the injection site). For (A) and (B) each colour 66 represents the percentage GFP+ cells/section that belong to the same animal. The 67 horizontal lines depict the mean percentage ± SEM. 68

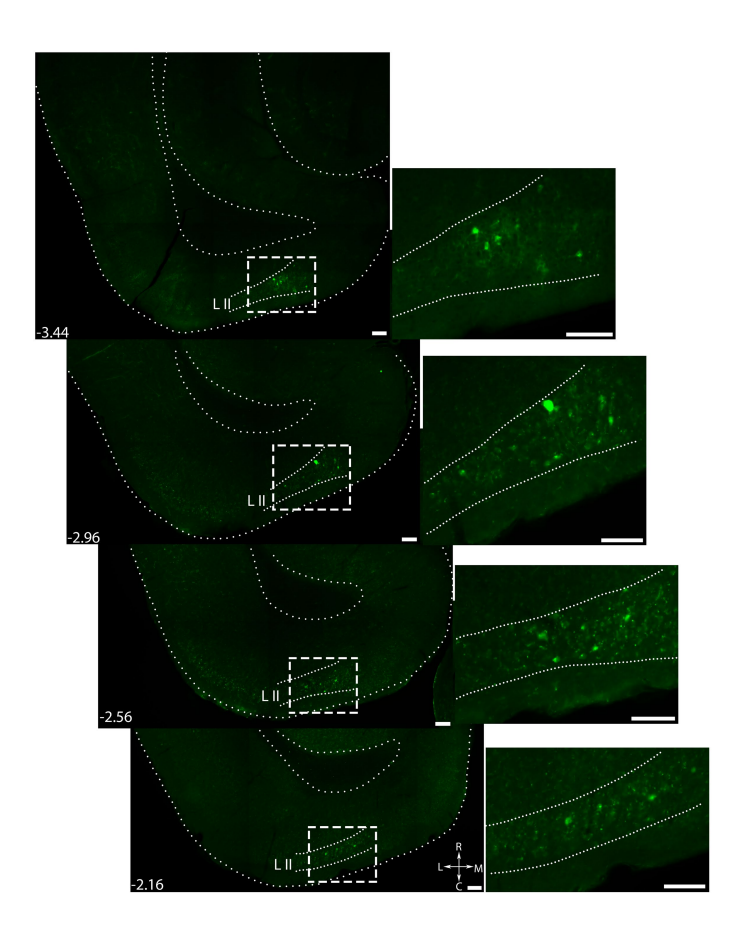


Figure S7. EC LII specificity of MEC13-53 rAAV PHP throughout the caudal
forebrain (related to Figure 3). Multiple dorso-ventral levels from the same mouse
brain injected with MEC13-53 EDGE PHP into tail vein shows MEC13-53 drives
transgene expression preferentially in MEC LII throughout dorso-ventral axis.
Sterotaxic coordinates were identified based on anatomical features using Paxinos G
& Franklin K (for mouse brain). See supplemental Table S2. Scale bar = 100 µm.

77 Transparent Methods

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit anti-GFP	ThermoFisher	Cat# A-11122,RRID:AB_221569
	Scientific	
Mouse anti-Reelin	Merck Millipore	Cat# MAB5364, RRID:AB_2179313
Mouse anti-Calbindin	Swant	Cat# 300, RRID:AB_10000347
Mouse anti-NeuN	Merck Millipore	Cat# MAB377, RRID:AB_2298772
Rabbit anti-2A peptide	Merck Millipore	Cat# ABS31, RRID:AB_10615498
Experimental models: Organ	nisms/Strains	
C57BL/6J mice	Jackson laboratory	IMSR Cat# JAX:000664,
		RRID:IMSR_JAX:000664
Long Evans rats	Charles River	RGD Cat# 2308852, RRID:RGD_2308852
MEC13-53 tTA X tetO-TVAG	Blankvoort <i>et al</i>	
	2018	
MEC13-104 tTA X tetO-	Blankvoort <i>et al,</i>	
TVAG	2018	
LEC13-8 tTA X tetO-	Blankvoort <i>et al,</i>	
GCaMP6	2018	
AAV-293 cell line	Agilent	Cat# 240073 , CVCL_6871
Stbl3 <i>E. coli</i> strain	ThermoFisher	Cat# C737303
	Scientific	

Reagents, chemicals, kits				
AAV helper free system	Agilent	Cat# 240071		
pHelper	Agilent	Cat# 240071		
pRC	Agilent	Cat# 240071		
pXR1	NGVB, IU, USA			
pAAV PHP.B	Deverman <i>et al,</i>			
	2016			
Maxiprep kit	Qiagen	Cat# 12663		
Fetal bovine serum	ThermoFisher	Cat# 16000-044		
	Scientific			
Benzonase nuclease HC	Merck Millipore	Cat# 71206-3		
HiTrap® Heparin columns	GE	Cat# 17-0406-01		
Amicon Ultra centrifugal	Merck Millipore	Cat# Z648043		
filters				
Power SYBR™ Green PCR	ThermoFisher	Cat# 4368577		
Master Mix	Scientific			
DNase I	ThermoFisher	Cat# EN0521		
	Scientific			
DIG-labelled riboprobe	Roche	Cat# 11277073910		
ISH blocking reagent	Roche	Cat# 11 096 176 001		
Nitroblue tetrazolium chloride	Roche	Cat# 11 383 213 001		
5-Bromo- 4-chloro- 3-indolyl-	Roche	Cat# 11 383 221 001		
phosphate, 4-toluidene salt				
Levamisole	Vector	Cat# SP-5000		

Instruments, softwares				
Confocal microscope, Zeiss	Zeiss			
LSM 880				
Zen 2012 software	Zeiss			
Hamilton needle	Hamilton	Cat# HAMI7762-06		
Nanoliter injector,	World Precision			
Nanoliter2010	Instruments			
Microsyringe pump controller,	World Precision			
Micro4 pump	Instruments			
Axio Scan. Z1 scanner	Zeiss			
Zen 2.3 software	Zeiss			

82

83 CONTACT FOR REAGENT AND RESOURCE SHARING.

84 Further information and requests for resources and reagents should be directed to

and will be fulfilled by the Lead Contact, Cliff Kentros (<u>clifford.kentros@ntnu.no</u>).

86

87 EXPERIMENTAL MODEL AND SUBJECT DETAILS

88 Rodent Details

Experiments were carried out using C57BL/6J mice (IMSR Cat# JAX:000664, RRID:IMSR_JAX:000664) obtained from Jackson laboratory, USA and Long Evans rats (RGD Cat# 2308852, RRID:RGD_2308852, Charles River, USA). EDGE transgenic crosses were created at the Kavli Institute for Systems Neuroscience and Centre for Neural Computation as described (Blankvoort et al., 2018). All experiments were conducted in compliance with protocols approved by the Norwegian Food Safety
Authorities and European Directive 2010/63/EU (FOTS ID 6269). All mice and rats
were housed in enriched environment cages in a 12 hr light/dark cycle with food and
water ad libitum.

98 METHODS DETAILS

99 Molecular cloning of constructs

All rAAV constructs were generated on backbone plasmid pAAV-CMV-MCS-WPRE-100 hGH PolyA (modified by cloning WPRE after the MCS in pAAV-MCS #240071, Agilent, 101 USA). Control rAAV constructs were generated as follows: for ubiquitously expressing 102 rAAV construct (CMV-rAAV) without a region-specific enhancer was generated by 103 cloning the enhanced GFP (GFP) into the MCS of the pAAV-CMV-MCS-WPRE-hGH 104 PolyA. For the promoterless construct for testing the transcriptional activity of ITRs, 105 CMV promoter was removed from CMV-rAAV. To synthesize EDGE rAAV constructs, 106 the CMV promoter, MCS and hGH PolyA sequences except WPRE were removed 107 from pAAV-CMV-MCS-WPRE hGH PolyA. An expression cassette consisting of a 108 109 hybrid promoter (composed of a region specific enhancer and minimal promoter), GFP and PolyA sequence were then subcloned into the plasmid in reverse orientation 110 relative to the ITRs, to circumvent any promoter activity from the 5'ITR. The expression 111 cassette in the reverse orientation was cloned into the plasmid upstream of the WPRE 112 which thus minimized the promoter activity from the 3'ITR. Various EDGE rAAV 113 constructs with the revised design were generated by cloning murine enhancers 114 115 obtained from our initial enhancer screen such as MEC-13-53, MEC-13-104 or LEC-13-8 and different minimal core promoters: a variant of CMV (CMV*, derived from the 116 sequence of pTRE3G, Clontech, USA) (Loew et al., 2010), fibroblast growth factor 4 117

(FGF4) (Murtha et al., 2014), HSV-TK (sequence from NEB, USA) or HSP68 118 (Blankvoort et al., 2018, Cotney et al., 2013) into the expression cassette in the reverse 119 orientation. Sequences of the EDGE rAAVs, the region specific enhancers and the 120 minimal promoters used in the study are given below. Plasmids were maintained in 121 the Stbl3 E. coli strain (#C737303, ThermoFisher, USA) to avoid ITR-mediated 122 recombination. Enhanced GFP, WPRE, LEC-13-8 and minimal promoters were 123 synthesized by Genscript, USA. Positive clones were confirmed by restriction 124 digestion analyses and subsequently by DNA sequencing. Endotoxin-free plasmid 125 126 maxipreps (#12663, Qiagen) were made for rAAV preparations.

127 rAAV preparations

128 EDGE rAAVs were packaged in AAV serotype 2/1 (having a mosaic of capsid 1 and 2) (Hauck et al., 2003) using Heparin column affinity purification (McClure et al., 2011). 129 Specifically, a pAAV construct generated as described above with AAV helper 130 plasmids encoding the structural elements, were transfected into the AAV-293 cell line 131 (CVCL 6871, Agilent, USA). The day before transfection, 7 x 10⁶ AAV-293 cells were 132 seeded into 150 mm cell culture plates in DMEM containing 10% fetal bovine serum 133 (#16000-044, ThermoFisher, USA) and penicillin/streptomycin. Co-transfection of 134 plasmids such as pAAV-containing the transgene, pHelper, pRC (#240071, Agilent, 135 USA) and pXR1 (NGVB, IU, USA) was carried out next day. After 7 hours, the medium 136 was replaced with fresh 10% FBS-containing DMEM. The AAV-293 cells were cultured 137 for two days following transfection to allow rAAV synthesis to occur. The AAV-293 cells 138 filled with virus particles were scraped from the cell culture plates, then isolated by 139 centrifugation at 200 x g. The cell pellet was then subjected to lysis using 150 mM 140 NaCl-20 mM Tris pH 8 buffer containing 10 % sodium deoxycholate. The lysate was 141 treated with benzonase nuclease HC (#71206-3, Millipore) for 45 minutes at 37°C. 142

Benzonase-treated lysate was centrifuged at 3000 x g for 15 mins and the clear supernatant then subjected to HiTrap® Heparin High Performance (#17-0406-01, GE) affinity column chromatography using a peristaltic pump (McClure et al., 2011). The elute from the Heparin column was concentrated using Amicon Ultra centrifugal filters (#Z648043, Millipore). The titer of the resultant viral stock was determined by quantitative PCR as approximately 10¹¹ infectious particles/ml.

EDGE rAAVs were packaged in AAV PHP.B and purified using iodixanol density 149 gradient method. Specifically, pAAV construct was transfected into the AAV-293 cell 150 line (CVCL 6871, Agilent, USA) along with AAV helper plasmids encoding the 151 structural elements. The day before transfection, 7 x 10⁶ AAV-293 cells were seeded 152 into 150 mm cell culture plates in DMEM containing 10% fetal bovine serum (#16000-153 044, ThermoFisher, USA) and penicillin/streptomycin. PEI mediated co-transfection of 154 plasmids such as pAAV-containing the transgene, pHelper, pAAV PHP.B (Deverman 155 et al., 2016) was carried out next day. After 24 hours, the medium was replaced with 156 fresh 10% FBS-containing DMEM. The AAV-293 cells were cultured for two days 157 following transfection to allow rAAV synthesis to occur. The medium and AAV-293 158 cells filled with virus particles were scraped from the cell culture plates, then isolated 159 by centrifugation at 200 x g. The cell pellet was then subjected to lysis using 20 mM 160 Tris, 300 mM NaCl and 20 mM MgCl₂, pH 7.6 buffer. The supernatant was mixed with 161 40% PEG for 2 hours in ice for precipitation of virus particles. Centrifuge the PEG 162 treated medium at 4000 x g for 15 minutes. The lysate and the PEG-precipitate was 163 treated with benzonase nuclease HC (#71206-3, Millipore) for 45 minutes at 37°C. 164 Benzonase-treated lysate was centrifuged at 3000 x g for 15 mins and the clear 165 supernatant then subjected to iodixanol gradient ultracentrifugation. 4 different layers 166 of the gradient, i.e. 15%, 25%, 40%, and 58% of iodixanol was built in a Beckman and 167

Coulter quick-seal centrifuge tube. Phenol red was added to the 25% and 58% layers to aid visualization of the layers within the tube. The virus containing supernatant was layered above the 15% iodixanol by slowly dripping the solution onto the top layer of the gradient. Seal the tip of the tube using a heating device (Beckman and Coulter). The 40% iodixanol layer, after ultracentrifugation at 200,000 g for 2 hours at 18°C, was collected and buffer exchanged with DPBS using Amicon Ultra centrifugal filters (#Z648043, Millipore) (modified protocol from Addgene, USA).

175 Titration of the rAAVs

The titration of the rAAVs prepared for the study was carried out by guantitative PCR 176 (Aurnhammer et al., 2012, Gray et al., 2011) using Power SYBR™ Green PCR Master 177 Mix (#4368577, ThermoFisher, USA), the following primers were used for GFP; 178 forward primer- 5'-AGCAGCACGACTTCTTCAAGTCC and reverse primer 5'-179 TGTAGTTGTACTCCAGCTTGTGC (modified protocol from Addgene, USA). A known 180 concentration (2 x 10⁹ molecules/µl) of a pAAV construct containing the GFP 181 sequence was used for generating the standard curve. 5 serial dilutions of plasmid 182 from 2×10^8 to 2×10^5 were made in PCR grade water for creating the standard curve. 183 The purified rAAVs were treated with DNase I (#EN0521, ThermoFisher, USA) at 37°C 184 for 30 minutes, to eliminate any contaminating plasmid DNA carried over from the 185 rAAV production process. DNAse-treated rAAVs were serially diluted for the qPCR 186 titration (from 1:20 to 1:2500) in PCR grade water. A mastermix of the reagents for 187 the qPCR was prepared consisting of the SYBR Green PCR Master Mix, the primers 188 and PCR-grade water. 5µl each from the standards and the rAAV dilutions along with 189 15µl of mastermix were subjected to qPCR at 95°C 10 min / 95°C 15 sec / 60°C 1 min/ 190 repeat 40x/ melt curve using StepOne machine (Applied Biosystems, USA). Data 191 analyses were performed by StepOne2.3 software and by Microsoft excel. 192

193 Stereotaxic Injections

For rat experiments, the rAAVs were stereotactically injected into three-four months old male Long-Evans rats. Injections were performed with 1 μ l rAAV at a titer of ~1 x10¹¹ infectious particles/ml, into the MEC of the rats. The rats were deeply anaesthetized with isoflurane gas (induction with 5 % isoflurane (v/v), maintenance at 1 % isoflurane (v/v), airflow of 1200 ml/min). To maintain the body temperature of the animal, a heating pad at 37°C was used.

Rats were injected subcutaneously with buprenorphine hydrochloride (Temgesic®, 200 Indivior) and Metacam® (Boehringer Ingelheim Vetmedica) at the prescribed dosage. 201 Local anaesthetic Bupivacaine hydrochloride (Marcain[™], AstraZeneca) was applied 202 203 at the place of incision. The head was fixed to the stereotaxic frame with ear bars, and 204 the skin at the incision site was disinfected with 70 % ethanol and iodine before the incision was made using a sterile surgical scalpel blade. After incision, the mouthpiece 205 206 and ear bars were adjusted so that bregma and lambda were aligned horizontally. Mediolateral coordinates were measured from the mid-sagittal sinus, anterior-207 posterior coordinates were measured from posterior transverse sinus, and dorso-208 ventral coordinates were measured from the surface of the brain. A craniotomy was 209 made around the approximate coordinate, and precise measurements were made with 210 the glass capillary/Hamilton needle (HAMI7762-06) used for virus injection. 211 Coordinates for rat injections were 4.6 mm lateral, 0.2 mm anterior to the posterior 212 transverse sinus and 2.6 mm deep, with the glass capillary/needle lowered at 10° 213 pointing towards the nose. A single injection of 1 µl virus was conducted at a speed of 214 100 nl/min using a nanoliter injector (Nanoliter2010, World Precision Instruments, 215 Sarasota, FL, USA), controlled by a microsyringe pump controller (Micro4 pump, 216 World Precision Instruments). After completion of the injection, the capillary was 217

retracted after a 10 minutes delay, to give the virus time to diffuse. Finally, the wound
was rinsed with saline and the skin was sutured. The animals were left to recover in a
heating chamber, before being returned to their home cage. Next day Metacam was
administered orally and their health was checked daily.

For mouse experiments, 10-15 weeks-old adult C57BL/6J mice (male or female) were 222 anaesthetized with isoflurane (induction with 5 % isoflurane (v/v), maintenance with 1 223 % isoflurane (v/v), airflow of 1200 ml/min). After applying the local analgesic Marcain 224 (40 µl, 0.25 mg/ml, SC), the global analgesic Temgesic (0.03 mg/ml, 100-150 µl per 225 226 mouse dependent on bodyweight, SC), and Metacam (2.5 mg/ml, 100-150 µl per mouse dependent on bodyweight, SC) the head was fixed in a stereotaxic frame. 227 Subsequently the skull was exposed by a single incision of the scalp, craniotomies 228 229 were made approximately 5 mm posterior and 3.3 mm lateral of the bregma. Then, the virus solution was injected at a location 0.3-0.5 mm anterior to the transverse sinus 230 and at a depth of 1.8-2.0 mm from the brain surface. Unless otherwise stated, all 231 injections were bilateral injections of 400 nl rAAV injected at a rate of 50 nl/min. Mice 232 were given a second post-operative injection of Metacam the next day, and their 233 weight was monitored until stable. 234

235 Tail vein injections

For tail vein injections, 10-15-weeks-old adult C57BL/6J mice (male or female) were anaesthetized with isoflurane (induction with 5 % isoflurane (v/v), maintenance with 1 % isoflurane (v/v), airflow of 1200 ml/min). Anaesthetized mouse was placed on a heat pad maintained at 37 °C during the procedure. The tail was wiped using 70% alcohol and warmed using water pad to cause vasodilation of the vein. Mice were injected with EDGE rAAV PHP virus preparation into lateral tail vein to a total volume of 100 µl in PBS (approx. 1 x 10¹² infectious particles/mouse) using a 30-gauge insulin syringe (#720-2555, Omnican, B.Braun). Remove the needle from the vein and apply slight
pressure to the puncture site with a dry piece of sterile cotton until the bleeding has
stopped.

246 **Perfusions**

After 4 weeks, the rodents were sacrificed. Rodents were anaesthetized with pentobarbital and perfused transcardially with freshly prepared 0.9 % saline followed by 4 % paraformaldehyde in 0.1 M Phosphate buffer (pH 7.4) with 0.9 % saline. The brains were stored in 4% PFA overnight before being transferred to 30% sucrose solution for approximately two days.

252 Immunostaining

Horizontal rat brain sections of 50 µm were prepared using a sliding microtome at 253 -30°C. Brain sections were stored at -20°C in 0.1 M phosphate buffer containing 25 % 254 glycerin and 30 % ethylene glycol. Multiple labelling of free-floating sections was 255 carried out as briefly described. Usually, every sixth section in the series was selected 256 for immunostaining and washed in phosphate-buffered saline (PBS). Sections were 257 permeabilized and blocked for 1 hour at room temperature using PBS containing 0.1 % 258 Triton X-100 and 3 % normal donkey serum, or, when staining for reelin and calbindin 259 0.5 % Triton X-100 and 5 % goat serum and when staining for NeuN 0.3 % Triton X-260 100 and 3 % BSA (PBS++). Sections were subsequently incubated with primary 261 antibodies in PBS++ at 4°C for two days with mild shaking. PBS-washed sections were 262 incubated for 2 hours at room temperature with secondary antibodies diluted in PBS++ 263 (or PBS containing Triton X-100 without serum/BSA). 264

265 Solution containing 2.5 % 1,4-diazabicylo[2.2.2]octane/polyvinyl alcohol 266 (DABCO/PVA) was used to mount the sections in Polysine slides (Menzel-Glaser, 267 ThermoFisher, USA). Antibodies used were rabbit anti-GFP (Thermo Fisher Scientific Cat# A-11122, RRID:AB_221569, 1:500), mouse anti-Reelin (Merck Millipore Cat#
MAB5364, RRID:AB_2179313, 1:1000), mouse anti-Calbindin (Swant Cat# 300,
RRID:AB_10000347, 1:5000), mouse anti-NeuN (Millipore Cat# MAB377,
RRID:AB_2298772, 1:1000) and rabbit anti-2A peptide (Millipore Cat# ABS31,
RRID:AB_10615498, 1:2000). All corresponding secondary antibodies were from
ThermoFisher/Life technologies or Jackson ImmunoResearch laboratories, USA,
used at a dilution of 1:400.

275 In situ hybridization

Mice were perfused transcardially with 4% paraformaldehyde (PFA) in RNAse free 276 PBS. The brain was extracted and stored in 4% PFA overnight before being 277 transferred to 30% RNAse free sucrose solution for approximately two days. The brain 278 was then sectioned horizontally in 30 µm thick sections and divided into a set of 279 approximately 6 series and stored in a -80°C freezer. A series was then thawed before 280 use. To stain transgenes TVAG or HM3, 30µm thaw-mounted sections were 281 hybridized overnight at 62°C with a DIG-labelled riboprobe for TVAG or HM3 282 (approximately 1:500; Roche, Cat. 11277073910) and then incubated in Blocking 283 solution (600µl MABT, 200µl sheep serum, 200µl 10% blocking reagent (Roche, Cat. 284 No. 11 096 176 001) for 2-3 hours at room temperature. Slides were drained of the 285 blocking solution, and antibody solution (1:5000 dilution of sheep anti-dig alkaline-286 phosphatase (AP) in blocking solution) was added to the slides. The slides were 287 transferred back to the Perspex box and incubated at room temperature overnight. 4g 288 of polyvinyl alcohol (Mol. Wt. 70000 - 100000) were transferred to a 50ml 289 290 polypropylene centrifuge tube, and AP staining buffer (100mM NaCl, 50mM MgCl₂, 100mM Tris pH 9.5, 0.1% Tween-20) was added till the total volume of the solution 291 was 40 ml. The solution was shaken to dissolve the solid material, and further heated 292

in a water bath. When the solution was clear, it was cooled down to 37°C. The slides 293 were washed 5 times in MABT in room temperature for 4 min each wash. Further, the 294 slides were washed 2 times in AP staining buffer for 10 min in room temperature while 295 the slides were shaken. NBT (Nitroblue tetrazolium chloride, Roche. Cat. No. 11 383 296 213 001, 140 µl/40ml), BCIP (5-Bromo- 4-chloro- 3-indolyl- phosphate, 4-toluidene 297 salt, Roche, Cat. No. 11 383 221 001, 105 µl/40 ml) and Levamisole (Vector, Cat. No. 298 SP-5000, 3.2ml/40ml) were added to the polyvinyl alcohol solution, and mixed well 299 before transferring it to a Coplin jar together with the slides. The slides were incubated 300 301 at 37°C for 5 hours. After incubation, the slides were washed 2 times in PBS + 0.1% Tween-20 to stop the staining reaction. The slides were further washed 2 times in 302 ddH₂O and left air dry at room temperature for overnight. The slides were cleared in 303 xylene and coverslipped. The stained sections were imaged using automated Axio 304 Scan. Z1 scanner (Zeiss), Zen 2.3 software with transmitted white light as the light 305 source. 306

307 QUANTIFICATION AND STATISTICAL ANALYSIS

308 Confocal Imaging and Analysis

Brain sections were imaged using a confocal microscope (Zeiss LSM 880, Zen 2012 309 software) with either Plan-Apochromat 40x/1.4 Oil DIC M27 oil immersion or Plan-310 Apochromat 20x/0.8 air immersion objectives. Image acquisition was done at identical 311 capture settings in confocal microscope. Captured images were processed identically 312 using Zen 2012 software and figures were prepared using Adobe photoshop. Confocal 313 314 images of GFP expression for different viral constructs were captured using 488nm laser. For figure S1, identical post-acquisition processing for the purposes of 315 comparison of strong versus weak viral GFP expression seen with the different viral 316

constructs, was carried out as explained below. Confocal images (czi.) were opened 317 in Zen 2012 software. Intensity range indicator tool was used to visualize, and display 318 tab was used modify the brightness intensity level of the acquired image. The 319 brightness intensity levels for the green channel was modified until the optimum 320 intensity was attained, as displayed by the range indicator. Once the parameters for 321 the optimal brightness intensity for green channel for one viral construct was identified, 322 identical changes in the intensity levels were applied to the other identically captured 323 confocal images for strong versus weak viral GFP expression comparison (Figure S1). 324 For e.g. in 2nd row (D-F) in Figure S1, the optimal intensity parameters for visualizing 325 D was determined using range indicator and display tab and subsequently applied 326 identically to the E and F. 327

328 The quantification of GFP+, NeuN+, RE+ or CB+ cells was performed out manually using Zen 2012 software. Approximately ten 50 µm thick horizontal sections were 329 selected from the dorso-ventral axis (-1.5 mm to -4 mm from Bregma) per brain. 330 Counts were carried out on the confocal images (.czi format) of rAAV injected brain 331 sections immunostained for GFP or respective markers (NeuN, RE, CB). For the layer 332 specificity analyses shown in Figure 2C, 13096 (3 mice) and 8540 GFP+ cells (7 mice) 333 were counted from mice, injected with CMV-rAAV and MEC13-53 rAAV respectively. 334 335 Data about MEC13-53 rAAV injections in mice includes counts from two different batches of the MEC13-53 rAAV. Any fluorescent signal greater than background auto-336 fluorescence was considered as positive, even though there was often a baseline 337 transcription from the rAAV promoter construct versus the enhancer-assisted (EDGE) 338 signal, which was typically orders of magnitude greater. For analyses in rat brains 339 (Figure 2G), 7191 GFP+ cells from sections were counted from 76 horizontal sections 340 from 5 separate hemispherical injections of MEC13-53 rAAV in 3 rats (-2 mm to -6 mm 341

342 from Bregma) and 2831 GFP+ cells were counted for CMV-GFP. For the quantification of MEC-LII EDGE GFP cells expressing cell-markers, 2406 (RE co-immunostained) 343 and 1668 (CB co-immunostained) GFP+ cells were counted from 7 separate MEC13-344 53 rAAV injected mice (Figure 2D). For similar analyses in rats (Figure 2H), 2332 and 345 1799 MEC-LII GFP+ cells were counted from separate RE and CB immunostained 346 sections from 5 different hemispherical injections of MEC13-53 rAAV (we analysed 31 347 sections for RE and 27 sections for CB) from 3 rats. Analyses were done in Microsoft 348 excel and graphs were made in Adobe illustrator. For the supplemental figure S6, 349 350 plotspread toolbox of Matlab was used to represent individual measurements of each tissue section analysed per animal and per group. For the statistical analysis of the 351 percentage of GFP positive cells, we used SPSS software. For the comparison of the 352 means, One-way ANOVA test was used and LSD Post Hoc test was conducted with 353 a significant level of 0.01. 354

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