Flexible, AAV-equipped Genetic Modules for Inducible Control of Gene Expression in Mammalian Brain

Godwin K Dogbevia1,2, Martin Roßmanith1, Rolf Sprengel3 and Mazahir T Hasan1,4

Controlling gene expression in mammalian brain is of utmost importance to causally link the role of gene function to cell circuit dynamics under normal conditions and disease states. We have developed recombinant adeno-associated viruses equipped with tetracycline-controlled genetic switches for inducible and reversible control of gene expression in a cell type specific and brain subregion selective manner. Here, we characterize a two-virus approach to efficiently and reliably switch gene expression on and off, repetitively, both in vitro and in vivo. Our recombinant adeno-associated virus (rAAV)-Tet approach is highly flexible and it has great potential for application in basic and biomedical neuroscience research and gene therapy.

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Introduction

The recombinant adeno-associated viruses (rAAVs) are powerful gene delivery vehicles for long-term, cell typespecific gene expression in living animals.1,2 Previously, we deployed the rAAV technology to deliver genetically encoded activity sensors to optically record neuronal activity in selected brain regions.3,4 Further, we have applied the rAAVs to inducibly delete a specific gene in a selective brain region,5,6 thereby linking gene function(s) with learning-dependent synaptic plasticity8 and memory formation.9 With AAVs, after virus infection in cells, maximum transgene expression can be observed in a matter of 2 weeks, both in vitro8 and in vivo.6 The flexibility to engineer designer rAAVs with different capsid proteins10 is an added advantage for selective targeting of diverse cell types. The rAAV DNA exit in the cell nucleus as episomal concatemers,8 and in some cases, it can integrate at a specific DNA site in the host genome,9 without any observable side effects. Importantly, rAAVs are largely nonpathogenic,10,11 which make them very attractive gene delivery vehicles for human gene therapy. Indeed, promising results have been reported with rAAVs to treat a number of diseases, including Leber’s Congenital Amaurosis,12 Hemophilia,13 congestive heart failure,14 and Parkinson’s disease.15

Our rAAVs are equipped with tetracycline (tet)-controlled genetic switches for inducible and reversible control of gene expression8 in any brain region(s) and cell type of choice, and gene expression can be controlled by doxycycline (Dox) at any time point chosen by an experimentalist. The use of rAAVs equipped with the tet inducible genetic switches thus paves the way to link gene activity to neuronal circuit function(s) in complex biological processes such as learning and memory,16 emotion,17 addiction,18 stress/anxiety,19 depression,20 pain,21 and aging.22 Here, we have characterized the different parameters for reliable control gene expression in the mammalian brain using a two-virus approach equipped with the ITA (Tet-off) and rITA (Tet-on) systems. Importantly, to combat various neurological disorders and malfunctions, our two rAAV approach is suitable for gene therapy.

Results

Tet-controlled gene expression in vitro and in vivo

There are two complementary tetracycline (tet) inducible systems,23,24 which are based on three key components (Figure 1a,b): (i) a tet transactivator (tTA)25 or a reverse tTA (rtTA),26,27 which are potent synthetic transcriptional factors, (ii) a unidirectional minimal tet promoter (Ptet)28 or a bidirectional minimal tet promoter (Ptetbi)29 to express multiple gene(s) of interest, when either tTA or rtTA binds to the tet operators (tetO)s located in Ptet/Ptetbi, and (iii) an inducer, doxycycline (Dox), a hydrophobic derivative of tet that controls gene expression and can rapidly distribute in different tissues in vivo, and easily crosses the blood–brain barrier.29 The operating principle of the tet systems (Figure 1c) is as follows: in the absence of Dox, tTA binds to the tetOs located in Ptet/Ptetbi to activate gene transcription, and in the presence of Dox, gene transcription can be switched-off, when the tTA/Dox complex unbinds the tet promoters (Ptet/Ptetbi). The reverse tTA (rtTA) variants binds to Ptetbi only in the presence of Dox (rtTA+/Dox) to activate Ptetbi mediated gene transcription, and without Dox, rtTA is unable to bind Ptetbi, terminating gene transcription.

We have developed recombinant rAAVs to inducibly and reversibly control gene expression in the mammalian brain. With our two-virus approach, we placed tTA or rtTA under control of a pan-neuronal specific promoter (human synapsin or hSYN) (Figure 1a,b). For quantifying gene expression, we placed the firefly luciferase (fLUC)30 and the tdTomato (tdTOM)31 genes under control of a Ptes bi (Figure 1a,c). With Ptetbi, these two genes were coexpressed for (1) live fluorescence imaging of neurons with tdTOM and (2) quantitative recording of luciferase (LUC) activity (Figure 1a,b). To normalize for the variability in gene expression, which is largely due to variability in virus infection of cells, we included a “tracer” virus

1Department of Neurobiology, Max Planck Institute for Medical Research, Heidelberg, Germany; 2Institute of Experimental and Clinical Pharmacology and Toxicology, University of Lübeck, Lübeck, Germany; 3Max Planck Research Group at the Institute for Anatomy and Cell Biology, Heidelberg University, Heidelberg, Germany; 4Charité-Universitätsmedizin, NeuroCure Cluster of Excellence, Berlin, Germany. Correspondence: Mazahir T Hasan, Max Planck Institute for Medical Research, Heidelberg, Germany. E-mail: mazahir.t.hasan@gmail.com

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to express the renilla luciferase (rLUC) in neurons. The use of specific light-emitting substrates for fLUC and rLUC, D-luciferin and coelenterazine, respectively, enabled us to quantify gene expression in a ratiometric manner (fLUC/rLUC) (Figure 2b), which are displayed here as relative light units (RLU) ± SEM.

First, we infected hippocampal organotypic slices with a cocktail of two rAAVs: rtTA2 system (rAAV-hSYN-tTA2 + rAAV-Ptetbi-fLUC/tdTOM) and rtTA system (rAAV-hSYN-rtTA2-nM2 + rAAV-Ptetbi-fLUC/tdTOM). Two weeks after virus infection, we detected robust Dox-induced, rtTA-dependent and tTA-dependent gene expression activation by Dox and removal of Dox (Figure 2a, middle and right panels). Expression of tdTOM was undetectable without Dox (Figure 2a, left panel). Because the half-life time of the tdTOM protein is about 24 hours, it takes several days to deplete the tdTOM signal, when using the rtTA system after Dox treatment (data not shown).

The fLUC on the other hand has a half-life time of only a few hours, enabling determination of Dox-controlled, rtTA/rtTA2-dependent gene expression time course, both in vitro and in vivo. To quantify gene expression, we performed the LUC assays using a cocktail of three viruses; the rtTA system (rAAV-hSYN-tTA2 + rAAV-Ptetbi-fLUC/tdTOM + rAAV-hSYN-rLUC) and the rtTA system (rAAV-hSYN-rtTA2-nM2 + rAAV-Ptetbi-fLUC/tdTOM + rAAV-hSYN-rLUC). With the rtTA system, RLU activity was about 18-fold higher over the baseline (without rtTA or with rtTA/+Dox). In the case of the rtTA system, the RLU activity without Dox was near the baseline levels, and RLU activity increased by about 20-fold by Dox (Figure 2c). We further determined that the maximum eLUC activity from brain samples infected with the rAAV-hSYN-eLUC was quite similar to brain samples infected with the Tet-inducible systems (rAAV-hSYN-tTA2 + rAAV-Ptetbi-fLUC/tdTOM, -Dox and rAAV-hSYN-rtTA2-nM2 + rAAV-Ptetbi-fLUC/tdTOM, +Dox) (Supplementary Figure S3). In the case of the inducible systems, the low virus titers (rAAV-Ptetbi-fLUC/Venus) hindered signal detection from the intact skull. However, the eLUC signal was detected, when the skull was removed (data not shown). We propose that in future studies long-term regulated gene expression in mouse brain should be feasible by noninvasive in vivo bioluminescence imaging. We further suggest that noninvasive in vivo bioluminescence imaging using the rAAV-hSYN-eLUC can serve as a reliable tool to control for the variability in virus injection and will help to identify properly rAAV-injected animals.

Repetitive ON/OFF gene expression cycles in vitro
Because access and removal of Dox occurs rapidly in cultured cells, we used dissociated neurons as a model system to investigate the time course of rtTA-dependent gene expression inactivation by Dox and re-activation upon Dox removal and rtTA-dependent gene expression activation by Dox and virus infection (Supplementary Figure S2). The observed basal fLUC activity in the uninduced state is thus most likely due to the intrinsic leakiness by the tet responder virus (rAAV-Ptetbi-fLUC/tdTOM), and leakiness increases with increasing number of rAAV-Ptetbi virus particles in infected cells. Consistent with our previous observations,6 intrinsic leakiness by rAAV-Ptetbi module is likely mediated by the rAAV inverted terminal repeat (ITR) enhancer-like activity onto the Ptetbi.32 To reduce leakiness, we had to lower the rAAV-Ptetbi virus titer.

We also performed noninvasive bioluminescence imaging with the tet inducible systems using the enhanced firefly luciferase (eLUC), but the eLUC activity was not detected in the mouse brain through an intact skull (data not shown). However, strong bioluminescence signal was detected through an intact skull, skin and fur with the mouse cortex injected with rAAV-hSYN-eLUC (Figure 2c). We further determined that the maximum eLUC activity from brain samples infected with the rAAV-hSYN-eLUC was quite similar to brain samples infected with the Tet-inducible systems (rAAV-hSYN-tTA2 + rAAV-Ptetbi-fLUC/tdTOM, -Dox and rAAV-hSYN-rtTA2-nM2 + rAAV-Ptetbi-fLUC/tdTOM, +Dox) (Supplementary Figure S3).
inactivation upon Dox removal. In these experiments, we applied a three virus cocktail: the tTA system (rAAV-hSYN-tTA2 + rAAV-Ptetbi-fLUC/tdTOM + rAAV-hSYN-rLUC) and the rtTA system (rAAV-hSYN-rtTA2-nM2 + rAAV-Ptetbi-fLUC/tdTOM + hSYN-rLUC). Two weeks after virus infection, neurons were treated with Dox in the medium and RLU activity was calculated at various time points. After Dox treatment for 24 hours, RLU activity with the tTA and rtTA systems reached a baseline and a maximum, respectively, and it remained nearly unchanged after an additional 24 hours (Figure 3a,b). Further, Dox-controlled on/off gene expression cycles with the tTA and rtTA systems were possible with an interval of 48 hours. These experiments were performed by separating the virus-treated primary dissociated neurons into four groups (n = 4 per group, each group) lasting six days and RLU activity was measured from cell lysates. With the tTA and rtTA systems over a period of 6 days, the magnitude of RLU activity regulation was 10- and 12-fold, respectively (Figure 3c,d).

Repetitive ON/OFF gene expression cycles in vivo
To determine an optimal Dox concentration required to switch-off and switch-on gene expression in tTA and rtTA-dependent manner, respectively, we injected a three-virus cocktail bilaterally into cortex and hippocampus of 10 wild-type mice. Two weeks after virus injection, mice were treated with different concentrations of Dox (3, 10, 30, and 100 μg/g body weight, n = 2 mice per group). Two days after intraperitoneal (i.p.) Dox injection, RLU activity was measured from virus injected brain lysates (four samples per mouse, i.e., cortices and hippocampi of both brain hemispheres). With the tTA system, 10 and 100 μg/g body weight of Dox reduced gene expression levels by 7- and 10-fold, respectively (Figure 4a). With the rtTA system, however, 30 and 100 μg/g body weight of Dox increased gene expression by 13- and 16-fold, respectively (Figure 4b).

To determine the number of days required for gene reactivation after a single i.p. Dox injection (with tTA) and gene

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**Figure 2 Functional gene expression in vitro and in vivo.** (a) Dox-controlled, rtTA and tTA dependent gene activation in rat organotypic hippocampal slices infected with two viruses as depicted in the images above (scale bar 500 μm). With the rtTA system, tdTOM expression was undetectable in the absence of Dox (left panel, but it was robust with Dox (middle panel). With the tTA system, however, there was very strong tdTOM expression in the absence of Dox (right panel). (b) Firefly luciferase activity measurements of lysates from infected organotypic hippocampal slices treated with or without Dox. The mean of the luciferase activity in the absence of Dox was equated to 1. The data shown are mean of 4 independent infected organotypic slices and expressed as relative light units (RLU; fLUC/rLUC ± SEM). (c) In vivo bioluminescence in the somatosensory cortex in two mice infected with eLUC gene under control of a human hSYN promoter. Bioluminescence signal was detectable only after intraperitoneal injection of D-luciferin.
Tetracycline-controlled Inducible Gene Expression in Mammalian Brain

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Discussion

We have generated two recombinant rAAVs with tet controlled genetic switches for long-term, inducible and reversible regulation of gene expression in neurons. The fLUC and the tdTOM genes were placed under a bidirectional tet promoter (P_{tetbi}), allowing for Dox-controlled, tTA- and rtTA-dependent regulated gene expression. The fLUC protein has a short half-life of roughly 2–4 hours and a large dynamic range that is five orders of magnitude. The LUC activity assay is highly sensitive, with minimal background activity, and is ideally suited for quantitative measurement of gene expression.

We characterized our two-virus approach to determine the time course of gene activation and inactivation in the mammalian brain. Our results show that both tTA and rtTA systems provide about 12-fold change in gene regulation. With the tTA system, gene expression (without Dox) is enabled from the very beginning. Although Dox treatment terminates gene transcription within minutes, the preexisting ILUC protein, for example, with its intrinsic half-life time of a few hours, will reach undetectable background levels in about 48 hours. With the tTA system, when it is desired that to keep gene expression switched-off, Dox must be constantly administered. However, upon Dox removal, gene expression can be switched-on in about a week. This slow time course is due to slow Dox clearance from the brain. With the rtTA system, on the other hand, gene expression can be rapidly and fully switch-on within 24 hours, but it takes about a week to switch-off gene expression, again due to slow Dox clearance from the brain. These results clearly show that the tTA and rtTA systems are highly complementary and can be flexibly applied to investigate specific biological processes, and they are also highly suited for gene therapy applications. Previous studies have shown that controlling the amounts of Dox levels allow for graded control of gene expression, demonstrating that the tet systems operate as a rheostat. However, the rheostat can be converted to on/off switches, when transactivator is combined with a repressor. These features allow for flexible application of the tet systems for biological studies.

The P_{tetbi} vectors are intrinsically leaky. Although this leakiness is minimal, it can produce enough gene-product to interfere with cell physiology. The internal terminal repeat (ITR) of rAAVs have potential enhancer-like activity. Previously, we discovered that two oppositely oriented DNA sequences flanking the P_{tetbi} insulate the P_{tetbi} from the ITR. However, removal of one such DNA sequence (a gene, for example) brings the ITR very close to the P_{tetbi}, thereby increasing Dox-independent expression or leakiness. In our current study, we have observed that the minimal leakiness of a
With the rtTA system, gene activation (single Dox i.p.) and in vivo time courses, and Dox-controlled regulated gene expression inactivation (after single Dox i.p.) was also 2 and 6 days, respectively.

**P** after a single intraperitoneal Dox injection (30 μg/g, < 0.0001).

tTA-dependent gene re-activation with a single intraperitoneal Dox were used as baselines, respectively. (tTA treated (100 μg/g) and Dox-untreated mean luciferase/renilla ratios (ranging from 0 to 100 μg/g). For the tTA and rtTA systems, Dox-treated with different concentrations of Dox per gram body weight and displayed here as relative light units (RLU). (tTA system rtTA system

Figure 4 Dox doses, inactivation (tTA) and activation (rtTA) time courses, and Dox-controlled regulated gene expression in vivo. Wild-type mice were injected with the tTA system (three viruses; AAV-hSYN-tTA2, AAV-P_{bi}-fLUC/tdTOM, and AAV-hSYN-rLUC) or the rtTA system (three viruses; AAV-hSYN-rtTA2-nM2, AAV-P_{bi}-fLUC/tdTOM, and AAV-hSYN-rLUC). For each time point, measurements were made from 12 cortical and hippocampal extracts from three independently injected mice, and fLUC/rLUC activity ratios were measured and data calculated as mean ± SEM and displayed here as relative light units (RLU). (a,b) Mice were treated with different concentrations of Dox per gram body weight (ranging from 0 to 100 μg/g). For the tTA and rtTA systems, Dox-treated (100 μg/g) and Dox-untreated mean luciferase/renilla ratios were used as baselines, respectively. (c,d) Time course (days) for tTA-dependent gene reactivation with a single intraperitoneal Dox injection (10 μg/g, \( P < 0.0001 \)) and rtTA-dependent gene inactivation after a single intraperitoneal Dox injection (30 μg/g, \( P < 0.0001 \)). (e,f) With the tTA system, gene inactivation (single Dox i.p.) and reactivation (after single Dox i.p.) was 2 and 6 days, respectively. With the rtTA system, gene activation (single Dox i.p.) and inactivation (after single Dox i.p.) was also 2 and 6 days, respectively.

rAAV-P_{bi} can be substantially decreased by reducing the responder virus (rAAV-P_{bi}) titer.

We have demonstrated here that our rAAV-based approach is highly suitable for long-term regulated gene expression in living mammals. It should be pointed out that a previous study elegantly demonstrated the use of a single rAAV system, harboring a transactivator (tTA or rtTA) together with a \( \text{Ptet} \) for regulated gene expression.\(^{36} \) However, due to the limited cloning capacity of the rAAVs, a single rAAV approach can accommodate small DNA sequence lengths, which substantially limits the utility of a single virus approach for many applications. Our two rAAV approach on the other hand provides a flexible solution to simultaneously control the expression of multiple genes.

**Materials and methods**

**Animal welfare.** All experiments were performed in accordance with the animal welfare guidelines of the Max Planck Society and approved by the local authorities (Regierungspräsidium Karlsruhe). We housed 6–10-week-old C57BL/6 wild-type and NMRI albino mice under standard conditions in a 12-hour light/dark cycle in Makroline cages type 2A with food and water.

**Plasmid constructs.** In the current study, we generated new rAAV plasmids, (i) pAAV-P_{bi}-fLUC/tdTOM and (ii) pAAV-hSYN-eLUC (enhanced luciferase)\(^{36} \) and (iii) pAAV-hSYN-rLUC. The other plasmids (pAAV-hSYN-tTA, pAAV-hSYN-rtTA, and pAAV-hSYN-EGFP) were described previously.\(^{6,37} \) To generate the plasmid pAAV-P_{bi}-fLUC/tdTOM, firefly luciferase gene (1.6kb fragment) was excised from the plasmid pBl1-5 (ref. \(^{28} \)) with Pst1/Xba1, blunted and subsequently ligated into a HindIII/Sph1 blunt-end ligated to the plasmid pAAV-P_{tetbi-iCre/tdTOM, replacing iCre with fLUC. Similarly, the plasmid pAAV-hSYN-rLUC was generated by blunt-end ligation by replacing the EGFP from pAAV-hSYN-EGFP (BamHI/HindIII) with a 947 bp rLUC fragment from the plasmid pRL-CMV (Nhe/Xba1). To generate the plasmid pAAV-hSYN-eLUC, the enhanced luciferase gene (eLUC) including SV40pA was excised from the plasmid pEULc-test (a gift from Dr. Nakajima) using EcoRI/BamH1 and a 1.9 kb fragment was ligated into the vector pAAV-hSYN-vector (without tTA) using EcoR1/BamH1. To generate the plasmid pAAV-P_{bi}-eLUC/Venus, the 1.6 kb eLUC fragment was amplified by PCR from the plasmid pEULc with the forward primer EcoRV 5’-AAATTCGATCTGCTATCCGACCCAC-3’ and the reverse primer SpeI 5’TGCTCATGACTGGCCGCGCTT-3’. The PCR generated fragment was cloned into the vector pAAV-P_{bi}-iKK2DN/Venus (unpublished plasmid).
by removing IKK2DN with EcoRV and SpeI and replacing it with eLUC to generate the plasmid pAAV-P_{tetbi}-eLUC/Venus.

Preparation and purification of rAAVs via Heparin column. Recombinant adeno-associated viruses (rAAVs) serotype 1 and 2 were generated by cotransfection of HEK293 cells and purified by modified heparin column purification as described. In brief, plasmids corresponding to the various rAAVs used in this study were individually cotransfected with pDP1 and pDP2 (ratio 2:1:1) helper plasmids in HEK293 cells by the calcium-phosphate method. Forty-eight hours after transfection, cells were harvested into a 50 ml falcon tube followed by centrifugation at 1,000 rpm for 15 minutes. The cell pellet was resuspended in 45 ml of 150 mmol/l NaCl, 20 mmol/l Tris.HCl (pH 8.0), frozen in liquid nitrogen and stored at −70 °C. Cells were thawed at room temperature (RT) and incubated at 37 °C with 40 U/ml of benzonase and 0.5% NaDOC for 60 minutes with frequent mixing. Lysed cells were centrifuged and frozen at −70 °C for at least a few hours and subsequently thawed and centrifuged at 3,900 rpm for 15 minutes, supernatant collected and frozen at −70 °C overnight. Frozen supernatant was thawed and spin for 15 minutes at 4,000 rpm. The cleared supernatant was run through a pair-equilibrated 1 ml heparin column. The HiTrap Heparin HP column (GE Healthcare Bio-Sciences AB, Uppsala, Sweden) was serially washed with 20 ml of 100 mmol/l NaCl, 20 mmol/l Tris.HCl (pH 8), 1 ml of 200 mmol/l NaCl, 20 mmol/l Tris.HCl (pH 8) and 1 ml of 300 mmol/l NaCl, 20 mmol/l Tris.HCl (pH 8). The virus was eluted from the heparin column with 1.5 ml of 400 mmol/l NaCl, 20 mmol/l Tris.HCl, 3 ml of 450 mmol/l NaCl, 20 mmol/l Tris.HCl (pH 8) and 1.5 ml of 500 mmol/l NaCl, 20 mmol/l Tris.HCl (pH 8). The eluate was concentrated in a 15 ml Amicon Ultra (100,000 cut-off) concentrator (Merck Millipore, Darmstadt, Germany). The virus was concentrated further into a final volume of 250 µl and filtered through 0.2 µm Acrodisc filter (Merck Millipore, Darmstadt, Germany).

Virus infection of hippocampal organotypic slices and dissociated neurons. Organotypic slices were prepared as described previously and slices were infected with 500 nl of virus cocktails by local application on individual slices. The different virus cocktails are as follows: the TTA system (rAAV-P_{tetbi}-eLUC/tdTom + rAAV-hSYN-rLUC + rAAV-hSYN-tTA) and the rTA system (rAAV-P_{tetbi}-eLUC/tdtOM + rAAV-hSYN-rLUC + rAAV-hSYN-rLUC + rAAV-hSYN-rLUC). Media were changed 5 days after infection, and subsequently replaced every 3 days. Dissociated neurons on the other hand, were infected by adding 1 µl of the virus cocktail into 1 ml of medium. Two weeks of virus infection, brain slices were lysed for quantitative luciferase activity measurements or fixed for fluorescent imaging of tdTom.

Quantifying luciferase gene activity in dissociated neurons. Dissociated primary neurons were infected with a TTA virus cocktail or RTA virus cocktails. Two weeks later, cells infected with the TTA and rTA systems were incubated for 48 hours either without or 1 µg/ml of Dox, respectively. Subsequently, cells were lysed with 100 µl of passive lysis buffer and 20 µl of the lysates were used for measuring firefly and renilla luciferase activities. For the repeated cycle of gene activation and inactivation, dissociated primary neurons in 24-well plates were infected with TTA and rTA systems. Two weeks after infection, cells were put into five groups each consisting of four wells. Each group was incubated with or without Dox over 48 hours. The Dox-treated medium was replaced with a conditioned medium without Dox also for another 48 hours. This was repeated for five cycles of Dox addition or withdrawal. The removal of Dox was done by washing three times with conditioned medium without Dox. At the end of two cycles, cells were harvested for luciferase assay. The firefly luciferase (fLUC) and renilla luciferase (rLUC) assays were performed sequentially in one reaction tube according to the manufacturer’s instruction (Promega Corporation, Wisconsin). To normalize gene activity, fLUC/rLUC mean ratios were calculated as relative light units (RLU) ± SEM, and for each time point, measurements were performed from four independent cell lysates. For detailed dual luciferase reporter assay protocol, refer to Dual-Luciferase Reporter Assay System protocol instructions for use of products e1910 and e1960 (Promega Corporation).

Stereotactic virus injection into the mouse brain. Six- to 10-week-old C57BL/6 and NMRI albino mice were deeply anesthetized (ketamine 100 mg/kg, and xylazine 5 mg/kg) and secured in a Kopf stereotactic setup (Kopf Instruments, California). The surgical procedure was started only after animals no longer responded to tail or paw pinch. Throughout the surgery, animals were kept on a heating pad to prevent hypothermia. Cream (Bepanthen, Bayer, Leverkusen, Germany) was applied to the eyes of mice to prevent dehydration. The foreskin on the skull was cut open to expose the skull. With the help of a drill, small holes (about 50–100 µm) were drilled into the head of the mice at different coordinates. Approximately 300 nl of viral cocktail was injected via glass pipette (tip diameter 10–20 µm) into each brain region. For in vivo bioluminescence imaging, mice were injected in the motor cortex. For in vivo luciferase activity measurement, mice were injected in the somatosensory cortex and in the hippocampus of both brain hemispheres. After injection, the skin was sutured and the wound disinfected. Newly injected mice were kept on heating blocks at 37 °C until they woke up and were fed wet food. The coordinates used for the different injections with reference to the bregma are as follows: for hippocampus (−1.70 mm bregma, 1.5 mm lateral, 1.5 mm deep), for motor cortex (0.5 mm bregma, 1 mm lateral, 500 µm deep) and for somatosensory cortex (−1.70 mm bregma, 1.5 mm lateral, 500 µm deep). To avoid postoperative dehydration, mice were injected with physiological saline solution. To reduce moderate acute pain, buprenorphine (Buprenex, 0.1 µg/g body weight) was injected after the surgical procedure. Mice were placed on a heating-pad while recovering from surgery.

Quantifying luciferase gene activity in brain tissues. To measure luciferase activities from brain lysates, wild-type mice were injected with the TTA system (rAAV-P_{tetbi}-eLUC/tdTom + rAAV-hSYN-rLUC + rAAV-hSYN-tTA) and the rTA system (rAAV-P_{tetbi}-eLUC/tdTom + rAAV-hSYN-rLUC + rAAV-hSYN-rLUC). Two weeks after virus injection, mice were either...
injected with Dox (rTTA injected mice) or without (tTA injected mice), and 48 hours later mice were anaesthetized with isoflurane and decapitated or virus infected brain regions (somatosensory cortex and hippocampus) were carefully dissected and frozen in liquid nitrogen. Brain extracts were prepared and later sonicated in 200 μl PBS and 20 μl of the extract was used to measure firefly and renilla luciferase activities as described above.

**Bioluminescence imaging in vivo.** Bioluminescence imaging was performed with an EM-CCD Digital camera C9100-13 (Hamamatsu Photonics Deutschland GmbH, Herrsching am Ammersee, Germany) in combination with a dark box (AEQUORIA MDS) (Hamamatsu Photonics Deutschland GmbH, Herrsching am Ammersee). At the beginning of the experiment, BALB/c mice were stereotactically injected with rAAV-hSYN-eLUC into the cortex of the right brain hemisphere. After recovery from surgery the animals were anesthetized with a mixture of ketamine (65 μg/g) and xylazine (14 μg/g) by i.p. injection. To avoid dehydration, all mice were injected with 500 μl of physiological saline (0.9 % NaCl). Furthermore, the animals were i.p. injected with an aqueous solution of D-luciferin (300 μg/g).

Right after the last injection, mice were placed on a stage in the Hamamatsu dark box. At first, a bright field image was taken for later analysis followed by bioluminescence recording using the streaming mode (six images possible with an exposure time of 10 minutes each). All data was recorded with the aid of Hokawo imaging software version 2.1, also provided by Hamamatsu.

**Immunostaining with NeuN antibody.** Immunostaining was carried out on wild-type mice or rat organotypic slices infected with a TTA system (rAAV-P, bi-ILUC/tetTOM and rAAV-hSYN-tTA). In brief, mice were perfused with warm PBS and fixed in 4% paraformaldehyde in 1× phosphate buffered saline (PBS) prior to decapitation. Brains were removed and postfixed in 4% paraformaldehyde at 4 °C overnight, followed by embedding in 2.5% agarose. Coronal sections with 60–100 μm in thickness were prepared by vibratome slicing and stored in PBS 4 °C. Tissues were blocked in 4% normal goat serum supplemented with 1% bovine serum albumin (BSA), 0.3% Triton X-100, and incubated overnight at room temperature in anti-NeuN (mouse monoclonal, Chemicon, California) primary antibody diluted in 1× PBS, 1% (BSA), and 1% normal goat serum/0.3% Triton X-100. The next day, tissues were washed twice in 1× PBS/0.3% BSA/0.1% Triton X-100 followed by anti-mouse FITC secondary antibody (1:200, Jackson Immuno Research, Pennsylvania) for 1 hour at room temperature. Slices were washed 2× in PBS and mounted with Aqua Poly/Mount on glass slides with cover slips. Immunostained sections were analyzed using an Axiosvert 200 M confocal microscope with LSM 5 PASCAL (Zeiss, Jena, Germany) coupled to 543 nm HeNe and 450–530 nm Argon lasers (Lasos Lasertechnik GmbH, Jena, Germany).

**Statistical analysis.** All data are presented as mean ± standard error of the mean. Figure 2b was analyzed with t-tests \(P < 0.0001\). Multiple group comparisons were performed with one-way analysis of variance with \(P < 0.05\) is considered significant.

**Supplementary material**

**Figure S1.** Neuron-specific gene expression.

**Figure S2.** Firefly luciferase activity in relative light units.

**Figure S3.** Enhanced firefly luciferase expression using different rAAV expression systems.

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