A Transposon-Mediated System for Flexible Control of Transgene Expression in Stem and Progenitor-Derived Lineages

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SUMMARY

Precise methods for transgene regulation are important to study signaling pathways and cell lineages in biological systems where gene function is often recycled within and across lineages. We engineered a genetic toolset for flexible transgene regulation in these diverse cellular contexts. Specifically, we created an optimized piggyBac transposon-based system, allowing for the facile generation of stably transduced cell lineages in vivo and in vitro. The system, termed pB-Tet-GOI (piggyBac-transposable tetracycline transactivator-mediated flexible expression of a genetic element of interest), incorporates the latest generation of tetracycline (Tet) transactivator and reverse Tet transactivator variants—along with engineered mutants—in order to provide regulated transgene expression upon addition or removal of doxycycline (dox). Altogether, the flexibility of the system allows for dox-induced, dox-suppressed, dox-resistant (i.e., constitutive), and dox-induced/constitutive regulation of transgenes. This versatile strategy provides reversible temporal regulation of transgenes with robust inducibility and minimal leakiness.

INTRODUCTION

Complex multicellular organisms contain a wide spatial and temporal diversity of cell types. Gene expression and signaling mechanisms are tightly coordinated processes that are recycled throughout development, maturation, and aging, thus maintaining a smaller overall gene number. This makes the investigation of gene function dependent on the ability to precisely manipulate gene expression in each spatial and temporal context. In the CNS, one such example is the Notch signaling pathway, which is involved in a multitude of cell types from development to late neurodegeneration (Ables et al., 2011). The dynamic cellular contexts and diverse signaling interactions create the need to temporally regulate gene activity in order to precisely study gene function. Moreover, transgene manipulation has emerged as an important technique to study and treat disease. Specifically, directing neuronal subtype differentiation is important for disease modeling, and regulated growth factor secretion is a promising therapy for several neurodegenerative diseases (Behrstock et al., 2006; Marchetto et al., 2011).

Tetracycline (Tet)-regulated systems have been used to temporally and spatially regulate gene expression in various methodologies (Furth et al., 1994; Gossen and Bujard, 1992). Specifically, the bacterial Tet transactivator (tTA) has been optimized to silence gene expression downstream of a Tet-regulated promoter in the presence of doxycycline (dox), a Tet analog. In addition to this “Tet-Off” system, a “Tet-On” system uses a reverse tTA (rtTA) in order to activate transgene expression in the presence of dox (Mansuy and Bujard, 2000). Tet-inducible systems have suffered from “leaky” gene expression and low inducibility of transgene expression over baseline (Mansuy and Bujard, 2000). These drawbacks, along with epigenetic silencing of these elements through endogenous methylation, contribute to the limitations related to CNS transgene regulation (Zhu et al., 2007). While new tTA and rtTA variants help overcome some of these limitations (Zhou et al., 2006), their use in neural stem cell populations is unexplored.

Plasmid electroporation (EP) to the brain has become a commonly used method of in vivo transgenesis in development (Breunig et al., 2007; Saito and Nakatsuji, 2001). However, like most episomal methods of transgenesis, EP suffers from plasmid dilution (Chen and LoTurco, 2012). Thus, studying complex lineage trees or postnatal neuro- and gliogenesis has been limited (Chen and LoTurco, 2012). To mitigate this, the recently characterized piggyBac transposon system permits stable integration of transgenes and can be used with various standard in vitro transfection methods to generate stable cell lines (Chen and LoTurco, 2012; García-Marqués and López-Mascaraque, 2013; Kahlig et al., 2010).

However, the robustness of the new tTA and rtTA variants and the ability of pBASE to promote stable genetic integration of such complex inducible/reversible genetic systems in stem and progenitor-derived cells have not yet been described. Our results indicate that this genetic system...
RESULTS

We constructed a custom genetic system for stable Tet-regulated flexible transgene expression using piggyBac transposition, termed pB-Tet-GOI (piggyBac-transposable Tet-mediated flexible expression of a genetic element of interest). pB-Tet-GOI consists of three plasmids: (1) a pBase-expressing plasmid for transposon integration, (2) a reverse Tet transactivator (rTTA) plasmid (or transactivator variants [rTA2 and rTA2-CA] discussed below) to allow for dox-regulation of the GOI, and (3) a Tet response GOI plasmid (Figure 1A). This tripartite system provides flexible, “mix and match” use of the different rTAA/rTAA variants with different GOI response plasmids. pBASE itself recognizes the piggyBac terminal repeats (TRs) that flank the rTAA and response plasmids and thereby catalyzes stable genomic insertion of these plasmids to avoid plasmid dilution with cell division (Chen and LoTurco, 2012). Importantly, the pBASE plasmid is episomal and dilutes with cell division, preventing the stably inserted rTAA and response plasmids from continuously “hopping” in and out of the genome. The rTAA plasmid yields constitutive expression of rTAA-V10 protein. With dox, the rTAA-V10 binds to the Tet-Bi promoter of the response plasmid and induces bidirectional transcription of two genes of interest (i.e., “Tet-On”). Finally, the Tet response plasmid features a constitutively expressed TagBFP2 reporter with a nuclear localization signal (NLS) and a V5 epitope tag (Figure 1A) for fluorescent and immunocytochemical (ICC) identification of transgenic cells. In addition, this design was employed to minimize spurious Tet-responsive gene expression by using the TagBFP2 cistron as a genetic insulator, minimizing possible upstream promoter/enhancer activity due to insertion in active chromatin and preventing transcriptional interference—including enhancer activity attributed to the upstream piggyBac TR (Shi et al., 2007).

We first generated a Tet response plasmid termed pB-TRE-Bi-EGFP (Figure 1B) expressing enhanced green fluorescent protein (EGFP). Next, we incorporated the transcription factor NEUROGENIN 2 (NEUROG2) that promotes glutamatergic-like neuron generation during cortical development (Zhang et al., 2013) and the DLX2 homeobox protein that promotes interneuron generation (Petryniak et al., 2007) into separate response plasmids to functionally test the ability of our system to indcuously direct differentiation. These two plasmids were defined as pB-TRE-Bi-EGFP/Ngn2 (i.e., Neurog2) and pB-TRE-Bi-EGFP/HA-Dlx2 (Dlx2 with a hemagglutinin [HA] epitope tag) (Figures 1C and 1D). For initial tests of the system, we transfected mouse N2a cells and a human cortical progenitor cell line (HuNPC) (Svendsen et al., 1998) with the three different response plasmids, along with rTAA-V10 and pBase. After culture for 4 days with or without dox, western blot analysis revealed robust GFP protein levels only in N2a cells grown with dox and no substantive GFP production without dox (Figure 1E). These collective results highlight both the inducibility and non-leakiness of the system. V5 epitope levels (from the TagBFP2-NLS in the response plasmid) demonstrated equivalent transfection and protein loading (Figure 1E). Moreover, Ngn2 and HA (from the Dlx2 construct) were detected only in their respective groups when ICC analysis showed that HuNPCs nucleofected with pB-TRE-Bi-EGFP/HA-Dlx2 and grown with dox produced TagBFP2, GFP, and HA. In contrast, cells grown without dox produced only TagBFP2 (Figures 1F–1G1). Similarly, HuNPCs nucleofected with pB-TRE-Bi-EGFP constitutively expressed nuclear TagBFP2 but expressed GFP only with dox (Figures 1H and 1H1). Finally, Ngn2 and GFP were induced only with dox in the pB-TRE-Bi-EGFP/Ngn2 group (Figures 1I–1J1). Interestingly, even though HuNPCs nucleofected with pB-TRE-Bi-EGFP are cortically derived, they did not express detectable Ngn2 (Figures 1I1 and 1K1). Taken together, pB-Tet-GOI was readily inducible, non-leaky, and could be used to bidirectionally express a reporter and GOI simultaneously in vitro.

To analyze whether the pB-Tet-GOI system expresses proteins that are biologically active and at sufficient levels to direct specific neuronal differentiation, HuNPCs were nucleofected with each response plasmid and differentiated for up to 6 weeks. Live cells were imaged daily for GFP and TagBFP2 autofluorescence. Notably, each group displayed divergent morphological changes and growth patterns over time (Figures S1A–S1C). Sibling cells from these three groups were fixed at 4 and 14 days for ICC examination of differentiation into TUJ1+ neurons and glial fibrillary acidic protein (GFAP)+ astroglia using confocal microscopy (Figures S1D–S1F2). Induced expression markedly increased neurogenesis at the expense of astrocytogenesis as early as 4 days and to a lesser extent at 14 days in both the pB-TRE-Bi-EGFP/HA-Dlx2 and pB-TRE-Bi-EGFP/Ngn2 groups (Figures 1L and 1M). The divergent morphologies were most clearly seen by computer-assisted cell tracing of the GFP+ cells at this time point (Figures S1G–S1H).

To assess the ability of our system to generate pure populations of stable undifferentiated human cell lines, we nucleofected three groups of HuNPCs as mentioned above, expanded them in culture without dox, and then sorted cells to isolate BFP+ cells (Figures S1J–S1L). All groups, expanded as undifferentiated neurospheres without dox, did not express GFP but did express the neural progenitor...
marker, SOX2 (Figures S1M–S1M2 and data not shown). Following dox addition, pB-TRE-Bi-EGFP/Ngn2 gained GFP expression while SOX2 expression was lost, suggesting that the immature cells can differentiate on demand by dox addition (Figures S1N–S1N2 and S1O–S1O2).

In addition to rtTA-V10 (Tet-On), we engineered tTA2 ("Tet-Off") and tTA2-CA ("Tet-insensitive") plasmids (Figure 2A). tTA2-CA has a H100Y mutation, making it dox insensitive (Hecht et al., 1993) to provide ubiquitous GOI expression without dox. Finally, we incorporated tTA2-CA into the response plasmid (instead of the transactivator plasmid) to permit temporally inducible but subsequently permanent expression of genes under the Tet-Bi promoter, yielding pB-TRE-Bi-EGFP/i-tTA2-CA). More precisely, dox

Figure 1. Validating the pB-Tet-GOI System for Transgene Manipulation

(A) Schematic of pB-Tet-GOI system for transposon-mediated integration along with inducible and reversible transgene expression.

(B–D) Response plasmids utilized for directed differentiation experiments.

(E) Western blot analysis of response groups grown with and without dox.

(F–K) Immunocytochemical staining for HA (DLX2 epitope tag), GFP (dox reporter), TagBFP2 (constitutive reporter), and NGN2 in HuNPCs nucleofected with indicated response plasmids.

Scale bar, 100 µm.

(L and M) Quantification of HuNPCs harboring indicated response plasmids and differentiated for 4 days (I) or 2 weeks (J) after dox.

Error bars represent mean ± SEM. *p < 0.05, **p < 0.01; n = 3 biological replicates per condition per time point.

See also Figure S1.
Figure 2. Transactivator Variants Allow for Flexible Manipulation of Transgene Expression

(A) Chart of transactivator variants and expected properties.

(B) Schematic of plasmid encoding pB-TRE-Bi-mpClover-P2A-Luc2P.

(C) Design of in vitro assay for transactivator variant validation.

(D–H) Luciferase assay reveals changes in firefly luminescence in the presence and absence of dox (as noted in panels) for the respective transactivator variants. n = 4 biological replicates per condition per time point. Samples normalized to Renilla.

Error bars represent mean ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.

See also Figure S2.
induces rtTA-V10 activation, leading to i-tTA2-CA expression and continual TRE-Bi self-activation (Figure 2A).

In order to rapidly and quantitatively assay these transactivator variants, we incorporated a destabilized firefly luciferase (Luc2P) gene (Figures 2B and 2C). Further, Luc2P was co-expressed using a P2A element with myristilated/palmitololated-Clover (mpClover; Figure 2B), which is an EGFP variant (Lam et al., 2012) tagged to membranes. pB-TRE-Bi-mpClover-P2A-Luc2P-transduced cells did not exhibit mpCLOVER expression or LUC2P activity without dox (Figures 2D–2H; Figures S2A–S2E and SA). With dox, rtTA-V10 induced mpCLOVER at 8 hr, which was detectable up to 16 hr after dox removal (Figures S2B1–S2C1). However, mpCLOVER displayed more rapid kinetics in that luminescence was noticeably diminished at 16 hr (Figure 2F). This may reflect the longer half-life of mpCLOVER when compared with destabilized Luc2P. Five days after dox removal, neither mpCLOVER expression nor LUC2P activity was detectable (Figure 2G; Figure S2D1). Dox was subsequently re-added to the culture, and mpCLOVER expression and LUC2P activity was detected (Figure 2H; Figure S2E1). This demonstrates that the system can be induced, reversed, and subsequently induced (i.e., On → Off → On).

As expected, cells expressing the pB-TRE-Bi-EGFP/rtTA2-CA (Tet-insensitive) plasmid displayed constitutive mpCLOVER expression and luminescence without dox (Figures 2D–2H; Figures S2A2–S2E2). In contrast, cells expressing the pB-TRE-Bi-EGFP/i-tTA2-CA combined plasmid did not exhibit basal mpCLOVER expression or luminescence without dox (Figure 2D; Figure S2A3). However, after a single dox administration, both were constitutively expressed—even after dox removal (Figures 2E–2H; Figures S2B3–S2E3).

tTA2 (Tet-Off) exhibited high basal luciferase activity and mpCLOVER expression, similar to that of dox-induced rtTA-V10 (Figure 2D; Figure S2A4). mpCLOVER expression decreased with dox addition until it was undetectable at 5 days, and it rebounded 3 days after dox removal (Figures S2B4–S2D4). Luminescence decreased more rapidly (again, presumably due to the attached PEST destabilization domain), approaching baseline at 16 hr after dox addition, though rebounding (along with mpCLOVER fluorescence) after dox removal (Figures 2E–2H; Figure S2E4).

To confirm transposition, we nucleofected neural stem cells with our pB-Tet-GOI system, alternate piggyBac expression vectors encoding fluorescent proteins and episomal expression vectors expressing analogous fluorescent proteins in various combinations—with and without pBase. In the presence of pBase (and dox), pB-Tet-Bi-EGFP and pB-mCherry demonstrated negligible loss of co-expression and retained high levels of fluorescence when assayed by vital imaging at 3 and 6 days post-transduction (group 1; Figures S2F and S2G). Without pBase, the identical combination of plasmids (with dox) displayed relatively similar co-expression at 3 days, but both GFP and mChERRY expression were almost undetectable by 6 days (group 2; Figures S2F and S2G). Mixing different piggyBac or episomal vectors in the presence of pBase demonstrated that the episomal plasmid expression was lost at 3–6 days (groups 3 and 4; Figures S2F and S2G). Conversely, the piggyBac transposed fluorescent proteins were perdurant, arguing against the possibility that non-specific toxicity caused loss of fluorescent protein expression rather than plasmid dilution (groups 3 and 4; Figures S2F and S2G).

TTA systems have shown promise in cell culture, but efficacy in the CNS has been inconsistent. To test the robust, non-leaky, inducible, and reversible characteristics of our pB-Tet-GOI system in vivo, pB-TRE-Bi-EGFP was introduced into ventricular zone (VZ) stem and progenitor cells using postnatal EP, a technique that allows for rapid expression of plasmid-based transgenes in these cells (Boutin et al., 2008; Breunig et al., 2012; Chesler et al., 2008). In the postnatal VZ, radial glia stem and progenitor cells differentiate into the mature cells of the striatum and provide neurons to the olfactory bulb (OB) (Alvarez-Buylla and Garcia-Verdugo, 2002; Carleton et al., 2003; Fernández et al., 2011). Five days after EP of pB-TRE-Bi-EGFP, half of the mice received a single dose of dox followed by brain removal at 1 hr, 1 day, 4 days, 6 days, 10 days, 2 weeks, 3 weeks, and 4 weeks (Figure 3A; Figure S3). The naive (un-stained) fluorescence of TagBFP2 and GFP was imaged to avoid artificial amplification of these signals (Figure 3B). Coronal brain sections showed nuclear TagBFP2 expression in the subventricular zone, striatum, and OB of all animals (Figures 3C–3L). Six days after treatment, robust GFP expression was observed in groups that received a single dox dose, but not in groups without dox (Figures 3C–3F). This robust expression persisted up to 14 days, decreasing at 3 weeks (Figures 3J and 3J1; Figure S3). Cells at 4 weeks after dox were indistinguishable from cells in animals without dox (Figures 3C–3E, 3K, and 3L), demonstrating the reversibility of transgene induction. Cell counts at the OB core of dox-treated animals revealed a decrease in the GFP+ cell number commencing after 14 days, while TagBFP2+ cell counts remained the same, indicating that the cells were still present but that their GFP expression turned off (Figure 3M). Analysis of the mean fluorescence intensity after the single dox dose revealed the most robust GFP expression at around 6 days, before declining to baseline at approximately 4 weeks (Figure 3N). In total, these data indicate that the pB-Tet-GOI system is a robust, inducible, and reversible method for transgene manipulation in vivo.

Non-invasively validating transgene expression could confirm appropriate delivery/expression of genetic elements
Figure 3. pB-Tet Is Non-leaky, Inducible, and Reversible in the Postnatal Mouse Brain

(A) Experimental timeline of dox administration after electroporation. Control mice received no dox administration.

(B) Schematic illustrating coronal planes of SVZ and OB imaged for panels (C)–(L). (C–L) Native unstained GFP and TagBF2 (pseudocolored red) fluorescence was analyzed in the VZ and OB. Significant GFP is observed only with dox.

Scale bar, 200 μm.

(M) TagBF2+ (all electroporated cells) and GFP+ (dox-responsive) cell counts in the OB core.

(N) Analysis of the mean fluorescence intensity of TagBF2+ and GFP+ cells in the OB.

SVZ, subventricular zone; Str, striatum; OB, olfactory bulb. Data points represent readings from a single litter used to minimize experimental variation that might result with comparing data across litters. Results are consistent with more than three independent experiments assessing dox activation kinetics in independent litters.

See also Figure S3.
and dox administration. We used a non-invasive imaging method with our well-characterized inducible Luc2P element (Figure 2B) in order to monitor dox induction of the pB-Tet-GOI system. Mice electroporated with a pB-TRE-Bi-mpClover-P2A-Luc2P-containing plasmid, along with pBase and rtTA-V10, were analyzed using in vivo luminescence imaging, which showed a significant induction of luminescence 1 week after dox administration (Figures 4A and 4B; see Figure S4A for timeline). After the initial imaging, several animals were used for histological examination, which demonstrated GFP+ cells in the dox-treated animals and no GFP+ expression in untreated animals (data not shown). One dox-treated animal not receiving a further dose subsequently displayed the basal luminescence reading of untreated littermates, while another littermate receiving another dose exhibited continued luminescence (Figure S4B). Finally, we electroporated littermates with a pB-TRE-Bi-mpClover-P2A-Luc2P-containing plasmid along with rTA-V10 (two mice) or tTA2-CA (three mice). As expected, luminescence was observed at similar levels in all animals, with the exception of the littermate that did not receive dox (Figures 4C and 4D). In summary, these data demonstrate the ability to monitor expression of the elements of our genetic system in a non-invasive manner.

**DISCUSSION**

Here, we characterize pB-Tet-GOI, a customizable, flexible genetic system that achieves stable transgene integration, along with inducible and reversible dox-mediated transgene expression in transduced cells and their resulting lineages without the need for viral packaging and generation of high viral titers. Specifically, we demonstrate the robust dox-mediated regulation of the latest-generation rTA and tTA2 variants and promoter elements, allowing for transgene expression in vitro and in vivo. Additionally, we provide tTA2-CA and i-tTA2-CA as a constitutive and inducible constitutive activator, respectively, of TRE-Bi for dox-insensitive expression of transgenes.
Through pBase-mediated transposition, stable cell lines can be rapidly generated. This technology not only facilitates the investigation of cell biology in vitro but also permits the in vivo induction of requisite transgenes from transplanted cell populations. Therapeutic applications include induced growth factor delivery from transplanted cells (Behrstock et al., 2006). Also, induced, direct differentiation in vivo permits the transplantation of precursor cells, which expand easier in culture and survive better post-transplantation, followed by dox-mediated maturation.

Our pB-Tet-GOI system allows for more precise investigations of gene function and cell lineages using in vivo EP with the ability to stably genetically trace and reversibly manipulate transgene expression. Notably, the combination of rtTA-V10 and i-tTA2 permits permanent cell lineage tracing after a single pulse. Thus, dox can be used as an orthogonal ligand for gene manipulation with other chemically induced genetic systems such as tamoxifen and RU486-induced transgenes. Finally, there is great potential to combine this system with INTRSECT technology, including alternate recombinases to achieve greater control over transgene manipulation (Fenno et al., 2014).

This genetic system is not limited to use in the CNS and is broadly applicable to many fields of biology. Any context requiring temporal control of transgenes may benefit from the use of the pB-Tet-GOI system. Further, we have recently confirmed that mir30 and miR-E-based small hairpin RNAs (Fellmann et al., 2013) can be similarly induced by our system, providing temporally controlled knockdown experiments (data not shown). Taken together, the pB-Tet-GOI tool kit will be useful for the investigation and elucidation of molecular mechanisms of biological systems in a wide variety of contexts. Further, it holds great promise for the fields of reprogramming and gene therapy.

**EXPERIMENTAL PROCEDURES**

**Mice and Electroporation**

CD1 mice were used according to the Cedars-Sinai Medical Center institutional animal care and use committee. EPs were performed as described previously (Breunig et al., 2012). Briefly, postnatal day 2 pups were placed on ice for ~8 min until unresponsive to tail pressure. A total of 1.2 μl of a plasmid cocktail of pBase (0.5 μg/μl [final]), transactivator (1 μg/μl [final]), and response plasmid (1 μg/μl [final]) diluted in Tri-EDTA buffer was injected into the left lateral ventricle. Fast green dye was added (10%v/v) to visualize injections. Platinum Tweezertrodes delivered five pulses of 120 V (50 ms; separated by 950 ms) from the ECM 830 System (Harvard Apparatus). SignaGel was applied to increase conductance. Mice were then warmed with a heat lamp for recovery.

**Cell Culture and Nucleofection**

HuNPCs (CNS010 cell line) were expanded as described previously (Svendsen et al., 1997, 1998). Briefly, HuNPCs were expanded as neurospheres in Stemline Media (Sigma S3194) with epidermal growth factor (Sigma E9644), leukemia inhibitory factor (Millipore LIF1010), and antibiotic-antimycotic (Life Technologies 15240-062). Spheres were dissociated for nucleofection with pB-Tet plasmids using the Lonza Nucleofector 2b device per the manufacturer’s recommendations. After nucleofection, media contained B-27 (Life Technologies 12587-010) without growth factors.

**Doxycycline**

Culture media contained dox (final concentration 100 ng/ml, Clontech 631311). Mice were gavaged with dox at 33 μg/g weight, using a 5-mg/ml stock solution.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, four figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2015.01.013.

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A Transposon-Mediated System for Flexible Control of Transgene Expression in Stem and Progenitor-Derived Lineages

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Figure S1. pB-Tet-mediated directed differentiation and utility for sorting of undifferentiated populations.

(A-C7) Vital imaging of HuNPCs reveals morphological changes in pB-Tet differentiating cells. HuNPCs were nucleofected with pB-TRE-Bi-EGFP (ctrl), pB-TRE-Bi-EGFP/HA-Dlx2, or pB-TRE-Bi-EGFP/Ngn2 and allowed to differentiate without growth factors for 6 weeks. EGFP autofluorescence was captured at denoted time points in live cells.

Scale Bar = 50μm

(D-F2) Immunocytochemical staining reveals altered differentiation patterns across pB-TRE-Bi-EGFP (ctrl), pB-TRE-Bi-EGFP/HA-Dlx2, or pB-TRE-Bi-EGFP/Ngn2. Yellow arrow indicates GFAP⁺ cell with polygonal astrocyte morphology (D-D2). Red arrows (E-E2) indicate TUJ1⁺ cells with small, interneuron-like or migratory neuron morphology. White arrows indicate TUJ1⁺ cells with long processes containing numerous varicosities (white arrowheads) (F-F2).

Scale Bar (D, E, F) = 50μm
Scale Bar (D1,2, E1, 2, F1, 2) = 25μm

(G, H, I) Camera-lucida tracing of GFP⁺ cells at two weeks indicating the presence of mainly morphological astrocyte-like cells in the pB-TRE-Bi-EGFP (ctrl) group with few morphological neuronal-like cells. Traced GFP⁺ cells revealed morphologically small, potentially migratory interneuron-like cells in the pB-TRE-Bi-EGFP/HA-Dlx2 group and large projection neuron-like cells in the pB-TRE-Bi-EGFP/Ngn2 group.

Scale Bar = 100μm

(J, K, L) FACS graphs indicating BFP⁺ populations in cells nucleofected with pB-TRE-Bi-EGFP (ctrl), pB-TRE-Bi-EGFP/HA-Dlx2, or pB-TRE-Bi-EGFP/Ngn2
(M-M$_2$) Sorted cells expanding as relatively pure populations of nucleospheres that are GFP$^+$, BFP$^+$. (pB-TRE-Bi-EGFP/Ngn2 shown)

Scale Bar = 500$\mu$m

(N-O$_2$) Nucleofected cells remain SOX2$^+$ in the absence of dox, and lose SOX2 expression after dox-induced differentiation. (pB-TRE-Bi-EGFP/Ngn2 shown)

Scale Bar = 50$\mu$m

**Figure S1 is linked to main Figure 1**

**Figure S2. Transactivator variants allow for flexible transgene control**

(A-E) Live images of N2a cells transfected with transactivator variants before luciferase cell lysis. Note: no dox was added to cells in columns A-E, and A$_2$-E$_2$. Cells in A$_3$-E$_3$ received a single pulse of dox immediately after imaging A$_3$ and before B$_3$ to activate rtTA-V10 and induce subsequent i-tTA2-CA constitutive activity.

Scale bar = 50$\mu$m

(F) Vital confocal imaging of nucleofected neural stem cells at day 6.

(G) Graph of fluorescent protein co-expression at 3 and 6 days post-nucleofection. Percentages are presented as the ratio of cells expressing each fluorescent protein as delineated in the graph legend (e.g. Group 1 is the percentage of mCHERRY cells that are EGFP positive).

Error bars represent mean ± SEM. **p<0.01, ***p<0.001. n = 3 biological replicates per condition per time point.

Scale bar = 100$\mu$m
Figure S3. pB-Tet-GOI is non-leaky, inducible and reversible in the mouse brain (full panel set related to Figure 3)

(A) Experimental timeline for postnatal electroporation and dox activation. (B) Brain regions analyzed in C-H1. (C-H1) Full set of fluorescence images for main Figure 3. Minimal EGFP expression is observed in littermates without dox. Examination of the brains of mice electroporated with pBase, pB-rtTA-V10, and pB-TRE-EGFP reveals detectable EGFP expression at 1 day. Littermates used to decrease experimental variation across litters. Results are consistent with >3 independent experiments investigating dox activation kinetics in separate litters.
Figure S4. Reversibility of non-invasive bioluminescence imaging of pB-Tet-GOI variants

(A) Timeline of experiments for littermates depicted in main Figure 4A and Supp. Figure 4C.

(B) Dox administration was continued (far right) or discontinued (far left) in littermates from Figure 4a.

Figure S4 is linked to main Figure 4
### Supplemental Table 1 - Antibodies used.

| Manufacturer | Catalog Number | Host Species | Antigen   | Concentration Used |
|--------------|----------------|--------------|-----------|--------------------|
| Immunostaining                                      |                |              |            |                    |
| Cell Signaling | C29F4          | Rabbit       | HA        | 1:1000            |
| Covance      | MMS-101R       | Mouse        | HA        | 1:1000            |
| Santa Cruz   | SC-19233       | Goat         | NGN-2     | 1:100             |
| Sigma        | T8660          | Mouse        | β-tubulin III | 1:2000    |
| Dako         | Z0334          | Rabbit       | GFAP      | 1:500             |
| Abcam        | 13970          | Chicken      | EGFP      | 1:5000            |
| Abcam        | 95038          | Goat         | V5        | 1:1000            |
| Invitrogen   | 46-0705        | Mouse        | V5        | 1:1000            |
| Western Blot                                        |                |              |            |                    |
| Covance      | MMS-101R       | Mouse        | HA        | 1:1000            |
| Santa Cruz   | SC-19233       | Goat         | NGN-2     | 1:1000            |
| Abcam        | 13970          | Chicken      | EGFP      | 1:10000           |
| Abcam        | 95038          | Goat         | V5        | 1:5000            |
| Invitrogen   | 46-0705        | Mouse        | V5        | 1:5000            |

### SUPPLEMENTARY EXPERIMENTAL PROCEDURES

**Cloning**

Plasmids were generated using In-fusion cloning (Clontech) along with standard molecular biology techniques to incorporate relevant cDNAs into pCAGGS (episomal) or pZGs (piggyBac)-based vectors. Details are available upon request.

**Tissue Preparation**

After anesthesia, mouse brains were isolated and immersion fixed in 4% ice-cold paraformaldehyde (PFA) overnight. Brains were then embedded in low melting point 4% agarose and sectioned at 70 μm on a vibratome.

**Subventricular Zone and Olfactory Bulb Imaging of Native TagBFP2 and EGFP Autofluorescence**

Unstained 70 μm SVZ and OB sections were mounted on glass slides and coverslipped. Confocal images were collected using a Nikon A1R inverted laser scanning confocal microscope. Immunohistochemical amplification of BFP and EGFP signals were omitted to preclude the artifactual, non-linear amplification and subsequent normalization of signal intensity observed following antibody staining. The number of cells in the olfactory bulb core was counted using a fixed-size grid centered on the terminus of the RMS. ImageJ was employed to count fluorescent cells and signaling intensities in an automated, unbiased fashion by employing the “Analyze Particles” function after calibration to ensure each cell was counted as a single particle.

**Imaging**

All confocal images were collected on a Nikon A1R inverted laser scanning confocal microscope with appropriate settings to sequentially image colors and avoid signal crosstalk. The exposure and saturation measures were utilized to capture the maximum dynamic range. Typically, after the exposure was set, the identical setting was reused for the subsequent samples in the group. Live images of HuNPCs were obtained on an EVOS digital fluorescence inverted microscope.

**Image Processing**

ND2 image files were initially imported into ImageJ for manipulation of confocal Z-stacks or for isolation of individual channels from single z-slices for subsequent editing in Adobe Photoshop CS6. Image curves were adjusted for
consistency of dynamic range and exposure in Photoshop CS6, cropped, and then imported into Adobe Illustrator CS6 for the preparation of final images.

**In vitro Phenotypic Quantification**

Cell phenotype percentages (TUJ1* and GFAP* of all GFP* and BFP* cells) were quantified in an unbiased manner by an observer blind to the treatment groups. TUJ1* and GFAP* quantification was done in triplicates (n= 3 coverslips per condition per timepoint); 100 GFP* cells per coverslip, performed in triplicate.

**Nucleofection**

Human Neural progenitor cell nucleofection was performed using the Amaxa Nucleofector 2b device according to manufacturer’s recommendations (Lonza AG). Unless otherwise noted, plasmids were added to the Lonza Nucleofection Solution in the following amounts (per reaction): Response plasmid(s): 7µg, pBase: 1µg, rtTA-V10: 3µg. For transposition experiment (Figure S2, F-G), plasmids were added to their respective groups in the following amounts (per reaction): pBase: 1µg, rtTA-V10: 3µg, all other plasmids: 7µg.

**Immunostaining of HuNPCs**

Immunocytochemistry was performed as previously described (Breunig et al., 2007b). Briefly, a primary antibody mixture was made in PBS-Triton (PBS, 0.3 % triton) with 3% normal donkey serum (NDS) and the desired primary antibodies at the ratios indicated in Supp. Table 1. Coverslips were incubated with the primary antibody mixture for at least 12 hours at 4°C, followed by three 5 min washes with PBS at room temperature. Secondary antibody mixtures were made with PBS-T and the appropriate secondary antibodies at a 1:1000 dilution (Jackson ImmunoResearch; conjugated with Alexa 405, Fitc, Alexa488, Dylight488, Alexa555, Dylight549, Alexa647, or Dylight649). The secondary antibody mixture was added to the coverslips and incubated at room temperature for 1 hour on a shaker. Coverslips were then washed in PBS and mounted on slides with anti-fade mounting gel medium (Invitrogen ProLong).

**Western Blot**

Mouse N2a cells were transfected with Lipofectamine 2000 (Invitrogen 11668019), following the manufacturer’s protocol. After transfection, the cells were grown for three days at 37°C. Cells were harvested by accutase incubation for 3 min at 37°C, followed by re-suspension in equal amount media, and centrifugation for 3 min at 3000 rpm. The resulting pellet was re-suspended in laemmli buffer and boiled for 15 min at 95°C. Protein concentrations were measured using a ThermoScientific Nano Drop. Protein separation was performed using SDS-PAGE separation and transferred onto nitrocellulose membranes, which were incubated overnight at 4°C using primary antibodies listed in Supp. Table 1 diluted in 5% milk in 0.1% PBS-Tween. All secondary antibodies (Li-cor IRDye®) were used at a 1:15000 dilution. Infrared detection was accomplished by the Li-Cor Odyssey® CLX Imaging System.

**FACS**

Once harvested cells reached approximately 80% confluency, media was removed and T75 cm² flasks were passaged with 2 mL of accutase at 37°C in 5% CO2 for 3 minutes. Accutase was neutralized with 2mL of media and cells were centrifuged at 1350 rpm for 3 minutes. Supernatant was removed and cells were resuspended in 2 mL of fresh media, using a P1000 to disrupt the cells in a careful yet vigorous manner. In increments of approximately 200 µL, cells were filtered through a 70 µm filter (BD Falcon). This filtrate was then placed on ice. Cells were then sorted at the Cedars-Sinai Flow Cytometry Core using a Beckman Coulter MoFlo sorter for blue fluorescence with tight gates making sure only the cells highly expressing BFP were collected, indicative of TagBFP-V5 expression from the response plasmid. Cells were collected into fresh media and kept cold on ice. Once all cells were sorted, the collected cells were pelleted, washed and placed into new T25 cm² flasks.

**IVIS Live Imaging**

Bioluminescence imaging of animals was performed at the Cedars-Sinai Medical Center Imaging Core Facility using the Xenogen Spectrum In Vivo Imaging System (IVIS). Animals were subcutaneously injected with 150mg/kg luciferin (VivoGlo®, Promega 1043). Following a 10 minute waiting period to allow for circulation of substrate, animals were imaged in the Xenogen IVIS while under isoflurane anesthesia. Luciferase expression was analyzed and quantified by subtracting the background noise (e.g. signal hindlimb regions distal from CNS) from each respective animal.
**Dual-Luciferase Reporter Assay**

Mouse N2a cells were transfected with Lipofectamine 2000 (Invitrogen 11668019), following the manufacturers protocol. Cells were grown in a 24-well plate as a monolayer in quadruplets (n=4 per condition) and imaged at respective time points followed by lysis and dual-luciferase activity assessment using the Promega Dual-Luciferase® Assay E910 according to the manufacturer’s protocol. Briefly, cells were passively lysed by adding 100μl of Promega® PLB buffer and shaking at room temperature for 15 minutes. 20μl of lysate and 50μl of Promega® LARII substrate was added to a 96-well assay plate. Firefly luciferase activity was measured using the Wallac Envision Manager Software with a 10 second delay per well. After firefly luciferase activity was measured, 50μl of Promega Stop & Glo® was added and Renilla activity was measured.

**Statistical Analysis**

Statistical analyses were carried out using Excel (Microsoft). We compared groups using two sample t tests. For cell counts, at least 100 cells per biological replicate were counted in all cases with the exception of Group 2 in Supplemental Figure S2, where 99 cells were counted.