One-Step *piggyBac* Transposon-Based CRISPR/Cas9 Activation of Multiple Genes

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Neural cell fate is determined by a tightly controlled transcription regulatory network during development. The ability to manipulate the expression of multiple transcription factors simultaneously is required to delineate the complex picture of neural cell development. Because of the limited carrying capacity of the commonly used viral vectors, such as lentiviral or retroviral vectors, it is often challenging to perform perturbation experiments on multiple transcription factors. Here we have developed a *piggyBac* (PB) transposon-based CRISPR activation (CRISPRa) all-in-one system, which allows for simultaneous and stable endogenous transactivation of multiple transcription factors and long non-coding RNAs. As a proof of principle, we showed that the PB-CRISPRa system could accelerate the differentiation of human induced pluripotent stem cells into neurons and astrocytes by triggering endogenous expression of different sets of transcription factors. The PB-CRISPRa system has the potential to become a convenient and robust tool in neuroscience, which can meet the needs of a variety of in vitro and in vivo gain-of-function applications.

INTRODUCTION

Neural cell fate determination is orchestrated by a tightly controlled transcription regulatory network. In the network, master regulatory genes or hub genes act in a predetermined sequence to govern proliferation and differentiation toward different lineages. The ability to manipulate expression of these genes simultaneously is a prerequisite for identifying important gene function and further delineating the complex picture of neural development.

Gene activation and suppression have been achieved by a variety of methods. Single gene perturbation is usually straightforward and relatively easy to carry out, while switching on or off the expression of several genes concurrently often requires multiple rounds of manipulation, which is frequently time consuming, labor intensive, and not always as efficient as one would anticipate. The CRISPR/Cas9 system has been developed and widely used as a tool for gene manipulation recently.1–7 In the system, Crispr RNA (crRNA) and trans-activating crRNA (tracrRNA) work together to bind to Cas9, a CRISPR-associated RNA-guided type II endonuclease. crRNA and tracrRNA (collectively called single-guide RNA, or sgRNA) then bring Cas9 endonuclease to the targeting site in the genome.

According to crystallography, Cas9 contains two lobes, a target recognition lobe and a nuclease lobe. The target recognition lobe is responsible for binding sgRNA and target DNA by recognizing the protospacer adjacent motif (PAM) sequence, while the nuclease lobe executes cleavage function on the target DNA.8,9 Taking advantage of the properties of recognition and binding of Cas9, a nuclease-deficient Cas9 (dCas9) was developed to mute its catalytic activity and maintain only the binding activity. dCas9 has been subsequently fused with effector domains such as transcription activators, allowing manipulation of gene expression at the transcription level.10–15

The initially designed system with dCas9-VP64-transactivating domain often requires multiple sgRNAs to activate just one gene. To enhance activation efficiency, several versions of CRISPRa platforms have been developed by recruiting various copies of VP16 or by hybridizing multiple transcription activators to dCas9. Examples include the repeating peptide array SunTag system;15 the three-component synergistic activation mediator (SAM) system;16 and the tripartite activator VP64, p65, and Rta (VPR) fusion protein system.17 Although these systems showed higher activation efficiency, they often would require the concurrent delivery of several transactivation components and sgRNAs to the same cells to achieve multiplex gene activation. A single all-in-one vector will expand potential applications of the transactivation platform, which will facilitate the evaluation of collaboration and interaction among multiple targeted genes.

One technical hurdle that hampers the construction of an all-in-one vector for stable activation is the limited payload capacity of the commonly used lentiviral or retroviral vectors, which hold a maximum cargo size of 7–8 kb, making it difficult to package multiple guide RNA-expressing fragments in one vector. To overcome this limitation, we decided to test whether the *piggyBac* (PB) transposon system, which could carry up to 100–200 kb of transgenes,18 would be able to fulfill such requirement. The PB system features transposons that can cut and paste transgenes into the genome where a TTAA sequence is

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found. In addition, the PB system has been demonstrated as an efficient tool in various applications, including induced pluripotent stem cell (iPSC) reprogramming and genome-wide mutagenesis.18–22

In the current work, we constructed PB-based vectors that contained highly efficient CRISPRa SAM components, as well as modular cassettes that allow for simultaneous expression of multiple sgRNAs. This platform allowed us to test a total of 72 sgRNAs for 24 genes of interest and identify the optimal sgRNAs for gain-of-function experiments. In addition, we have provided proof-of-principle applications by plugging in different sets of sgRNAs to the all-in-one vectors, and we were able to accelerate neural cell differentiation and cell type conversion in human iPSCs.

RESULTS
Identification of the Optimal sgRNAs for Gene Activation in the PB-CRISPRa System
To identify the best sgRNAs in the PB-CRISPRa system for each gene, we first created a human HEK293FT stable cell line that overexpressed MS2-p65-HSF1 (MSPH) and dCas9-VP64. MSPH and dCas9-VP64 were cloned into the PB backbone under the control of EF1α and CAG promoter, respectively (Figure 1A, renamed as PB-SAM). The HEK293FT cell line was transfected with PB-SAM and PBase, followed by hygromycin and blasticidin selection. The stable cell line obtained after selection was then renamed as 293FT-SAM. In 293FT-SAM, compared to untransfected cells (background control), the expression of MSPH and dCas9-VP64 was elevated by more than 200-fold for at least 3 months (the longest time point tested) without silencing (Figure 1B; Figure S1), indicating that 293FT-SAM cells could be used to test activation efficiency of sgRNAs in our PB-CRISPRa system.

Next we sought to evaluate the efficiency of the PB-CRISPRa system for activating endogenous gene transcription using the 293FT-SAM cell line. We chose 21 transcription factors (TFs) and three long non-coding RNAs (lncRNAs) that have been reported to be involved in the process of neural development (Table S1). We synthesized three sgRNAs per gene and tested 72 sgRNAs in total. For each gene, all three sgRNAs were designed to target ~200–1 bp upstream of its transcription start site (TSS).16” To quickly test the activation efficiency, we first co-transfected the 293FT-SAM cell line with all three sgRNAs. For 19 of 24 genes (79.2%), the mRNA expression level was increased by at least 2-fold (2- to 12,333-fold, mean fold increase was 763-fold; Figures 1C–1E and 1G), indicating that these sgRNAs could efficiently activate most genes of our interest respectively, although five genes (20.8%) failed to be activated in 293FT-SAM.

As the SAM system was reported to efficiently stimulate endogenous gene expression with only one sgRNA,16 we next evaluated the activation efficiency of individual sgRNAs for each gene, aiming to identify the best sgRNA for endogenous activation for each gene on our list. Compared to basal transcript level, an average of 130-fold increase was observed upon transfection of single sgRNAs, as shown by qPCR results. Specifically, mRNA expression level was increased by at least 2-fold (2- to 3,846-fold) for 43 of the total 72 sgRNAs (60%) tested, of which two sgRNAs (ASCL1-gRNA1 and ASCL1-gRNA2) caused mRNA expression level to increase by more than 1,000-fold and nine sgRNAs caused the corresponding mRNA expression to increase by 100- to 1,000-fold. Additionally, 17 sgRNAs were able to activate the gene expression by 10- to 100-fold and 15 sgRNAs by 2- to 10-fold (Figures 1C–1F). The sgRNA that was able to maximally activate endogenous gene expression in 293FT-SAM was identified to be the best sgRNA for subsequent experiments using our PB-CRISPRa system.

The PB-CRISPRa System Activates TFs and IncRNAs in Human iPSCs
We asked whether the optimal sgRNAs obtained from our screening in 293FT-SAM were able to activate the endogenous expression of TFs or IncRNAs in human cell types other than 293FT. We decided to test human iPSCs, cells that would be able to give rise to neural lineage cells among a variety of cell types of all three germ layers. Following the success in 293FT-SAM cells, we created an iPSC-SAM line by transfecting Cy2 iPSCs (NIH) with PB-SAM and PBase plasmids, followed by hygromycin and blasticidin selection. Similar to 293FT-SAM, the expression of PB-SAM mRNAs in iPSC-SAM was about 260-fold and 150-fold higher than in untransfected Cy2 iPSCs (Figure 2A; Figure S2). Hence, the PB-SAM-transfected Cy2 iPSC line was renamed as iPSC-SAM. Next, the optimal sgRNA that elicited the most robust gene activation in 293FT-SAM was introduced to iPSC-SAM cells. Consistent with the results obtained from 293FT-SAM, all genes were activated by the optimal sgRNAs, as endogenous gene expression level was elevated by at least 2-fold (2- to 2,470-fold) (Figures 2B–2D). Moreover, for 17 of the 24 genes (70.8%), the expression level after sgRNA transfection was increased by at least 10-fold (Figure 2B). These data suggest that, using the PB-CRISPRa system, the sgRNAs identified from the 293FT-SAM screening allowed for robust endogenous gene activation in human iPSCs as well.

As we briefly mentioned already, four TFs (NFIA, NFIB, NFIX, and SOX9) and an IncRNA (HAR1A) failed to be activated in 293FT-SAM cells. These genes, however, were successfully activated in iPSC-SAM. As shown in Figures 2C and 2D, these genes achieved an increase in expression of 8- to 110-fold, as determined by qPCR. As basal transcription level has been reported to contribute to endogenous gene activation by CRISPRa,16 we compared the basal expression level of all five genes, and we found that NFIA, NFIB, NFIX, and SOX9 had relatively higher basal transcription levels in 293FT than in iPSCs (Figure S3). To examine the potential correlation between the magnitude (fold increase in gene expression) of activation and basal transcription level in human iPSCs and 293FT cells, we performed a collective comparison of all 24 genes. As shown in Figures 2G, 2H, S3, and S4, iPSCs had a much higher fold increase in gene activation with relatively lower basal transcript level compared to 293FT cells. Linear regression analysis of the 24 genes showed a significant negative correlation between fold activation and basal transcript level in 293FT-SAM cells and human iPSC-SAM cells (Figures 2E and 2F; $r = 0.66, p = 0.0004$ in 293FT; $r = 0.76, p < 0.0001$ in iPSCs). A similar
Figure 1. Identification of the Best sgRNAs in Gene Activation by 293FT-SAM

(A) PB vectors used for creating 293FT-SAM. (B) Generation of 293FT-SAM cell line by co-transfecting 293FT with PB-SAM and PBase expression vectors. qPCR results showed that SAM components MS2-p65-HSF1 (MSPH) and dCas9-VP64 were stably expressed for at least 90 days. (C–E) qPCR results of gene expression activation of individual TFs (ASCL1, NEUROG1, NEUROG2, OLIG2, SOX10, LHX3, NIX1-2, MNX1, MYT1, FOXG1, LHX2, SOX8, OLIG1, SOX2, ISL1, SOX11, SOX9, POU3F2, NFIA, NFIB, and NFIX) and IncRNAs (RMST, HAR1B, and HAR1A) after transfecting 293FT-SAM cell line with corresponding sgRNA vectors. A total of three sgRNAs per gene was tested separately to evaluate their activation efficiency. The rightmost bars for each gene represent the extent of gene activation when all three sgRNA vectors for that gene were co-transfected. (F) A pie chart showing the range of fold change of gene expression after each sgRNA vector was transfected into the 293FT-SAM line. For each gene, three sgRNAs were tested separately. A total of 72 sgRNAs was tested. Basically, 43 sgRNAs were able to augment target gene expression by >2-fold. (G) For the 24 genes targeted, 19 genes were activated to be expressed at >2-fold. The qPCR results were normalized to GAPDH mRNA level. P2A, self-cleaving peptide P2A sequence; Hygro, hygromycin resistance cassette; Blast, blasticidin resistance cassette; NLS, nuclear localization signal. In (B)–(E), data are presented as mean ± SEM (n = 3).
Figure 1: iPSC-SAM and iPSC-SCM Analysis

(A) Expression of iPSC-SAM over time. The expression levels of dCas9-VP64 and MSPH are shown as fold activation over time.

(B) Circular genome viewer for iPSC-SAM. The expression levels of 24 genes are shown with different colors representing different abundance levels.

(C) Bar graph showing the fold activation of 9 genes over time. The genes are LHX2, LHX3, MNX1, OLG2, NFX1, SOX9, HART1, NGN1, and ASCL1.

(D) Another bar graph showing the fold activation of 9 genes over time. The genes are SOX8, OLG1, SOX1, NKCC1, MYT1, NFIB, SOX2, SOX10, RMB3, and HART1.

(E) Scatter plot showing the correlation between basal transcript level and fold activation in 293FT cells. The correlation coefficient is r = 0.66, P = 0.0004.

(F) Scatter plot showing the correlation between basal transcript level and fold activation in iPSCs. The correlation coefficient is r = 0.76, P < 0.0001.

(G) Box plots comparing fold activation between 293FT and iPSCs.

(H) Box plots comparing relative expression level (normalized to GAPDH) between 293FT and iPSCs.

(I) Box plots comparing ultimate gene expression level between 293FT and iPSCs.
correlation was also observed when data were pooled for all 24 genes in both cell types (Figure S5; \( r = 0.73, p < 0.0001 \)). Collectively, these data indicate that relatively lower basal transcript levels of the genes tested in iPSCs is one of the major factors that might contribute to higher activation magnitude induced by optimal sgRNAs. Interestingly, although iPSCs achieved a greater fold increase (i.e., ratio of gene expression level of post- versus pre-transactivation), there was no significant difference in the ultimate gene expression level between 293FT cells and iPSCs (Figure 2I), indicating that endogenous gene expression probably has an upper limit.

**All-in-One Vectors for Stable Activation of Multiple Genes in Single Cells**

To strengthen the application of the PB-CRISPRa system, we attempted to incorporate all components into one PB vector to enable simultaneous activation of multiple genes in single cells. As shown in Figure 3, the all-in-one PB vector contains EF1α promoter-driven MSPH, connected by CAG promoter-driven dCas9-VP64 and followed by multiple U6 promoter-driven sgRNAs. Additional U6-sgRNA expression cassettes could be incorporated by Multisite Gateway cloning as desired.

We first tested whether the all-in-one PB-CRISPRa vector was able to express all SAM components and stably activate endogenous expression of a single gene. To facilitate the test, we used an OLIG2-EGFP human iPSC knockin reporter, in which an EGFP cassette was knocked into one of the two alleles of OLIG2 by gene targeting and has been shown to faithfully reflect the endogenous expression of OLIG2 (S.L. and Y.L., unpublished data). The all-in-one vector containing OLIG2-sgRNA1 was transfected to the OLIG2-EGFP

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**Figure 3. A Schematic Representation of Constructing PB-CRISPRa All-in-One Vectors with Multisite Gateway Cloning Strategy**

The PB-SAM vector was amplified to become PB-SAM R1-R2 DEST vector containing attR1 and attR2 sites, a ccdB cassette, and a chloramphenicol resistance cassette (CmR). PB-sgRNA (MS2) vectors containing different sgRNA inserts were amplified to be attached to appropriate attB sites. All-in-one vectors were assembled by four-way LR reactions. P2A, self-cleaving peptide P2A sequence; Hygro, hygromycin resistance cassette; Blast, blasticidin resistance cassette; AmpR, ampicillin resistance cassette; KanR, kanamycin resistance cassette; NLS, nuclear localization signal; ITR, inverted terminal repeat sequence.
Figure 4. Stable Activation of Multiple Genes with All-in-One PB-CRISPRa Vectors

(A and B) An all-in-one PB-CRISPRa vector containing OLIG2-sgRNA1 was able to continuously activate EGFP (surrogate marker for endogenous OLIG2) expression in OLIG2-GFP human iPSC knockin reporter cell line, as indicated by the robust, native GFP signal from day 5 and day 28 post-transfection (A). The elevation of endogenous OLIG2 expression was validated by qPCR, which showed a 70- to 80-fold increase in OLIG2 gene expression starting from day 7 post-transfection that was maintained for at least 28 days (B). (C–F) Comparison of gene activation efficiency between all-in-one vectors and multiple single sgRNA vectors (one-by-one). The qPCR results showed...
reporter iPSC line, and EGFP expression was observed as early as day 1 after transfection. Starting from day 7 post-transfection, OLIG2 (EGFP) was activated in 35%–40% cells, and the expression level was elevated by about 80-fold as measured by qPCR. The activation was maintained for at least 4 weeks after transfection, indicating that the PB-CRISPRa system was able to stably activate expression of single genes in human iPSCs (Figures 4A and 4B; Figure S6).

To evaluate the efficiency of the all-in-one PB-CRISPRa vectors for concurrently activating multiple neural lineage-specific TFs, we generated a two-in-one NEUROG1 and NEUROG2 vector; a four-in-one NEUROG1, NEUROG2, LHx3, and ISL1 vector; a four-in-one SOX10, OLIG2, SOX8, and OLIG1 vector; and a four-in-one SOX9, NFIA, NFIB, and NFIX vector. To compare activation efficiency, 293FT cells were transfected with all-in-one PB CRISPRa vectors or co-transfected with two or four sgRNA vectors designed to activate single TFs (designated as one by one). Two days after transfection, expression of all genes was elevated. In addition, cells transfected with the all-in-one plasmids, especially the four-in-one vectors, had significantly higher activation efficiency than the one-by-one groups, with an upregulation of 5- to 1,050-fold increase in gene expression (Figures 4C–4F). Thus, with the all-in-one strategy, we were able to activate multiple TFs in the same cells more efficiently than with the separate one-by-one system.

**Rapid Generation of Neurons and Astrocytes from Human iPSCs via Endogenous Activation with All-in-One PB-CRISPRa Vectors**

NEUROG1 and NEUROG2 play important roles in neuronal development and cell fate determination.6–10 Exogenous overexpression of NEUROG2 alone or in combination with other TFs was able to direct or skew the fate of pluripotent stem cells, even mature somatic cells, toward neuronal lineages.11–12 To test whether the all-in-one system was able to convert human iPSCs to neurons by endogenous activation of NEUROG1 and NEUROG2, we transfected two human iPSC knockin reporter lines NEUROG2-mCherry and Doublecortin (DCX)-ZsGreen with an all-in-one PB-SAM-NEUROG1/2 gRNA vector and PBase. At 24 hr post-transfection, cells were treated with blasticidin and hygromycin B. Then 3 days after transfection, 30%–40% of cells started to show a bipolar morphology and expressed OLIG2 (mCherry, red) or DCX (ZsGreen, green), as observed under the fluorescence microscope (Figures 5A–5F). Within an additional 4 days, the morphology of the transfected cells highly resembled that of more mature neurons with longer neurites and extensive arborization. More than 80% of the transfected cells were confirmed by positive immunocytochemistry staining of NeuN, NF160, and MAP2, markers of mature neurons (Figures 5G–5O).

To expand the potential applications of the PB-CRISPRa all-in-one vectors, we also attempted the induction of human iPSCs toward the astrocyte lineage. iPSCs were transfected with an all-in-one PB-SAM-SOX9-NFIA-NFIB-NFIX gRNA vector and PBase. Treatment with blasticidin and hygromycin B was carried out 24 hr post-transfection to select cells that were successfully transfected with the all-in-one vector. At 14 days post-transfection, the majority of cells expressed multiple well-accepted astrocyte markers, CD44 (80%), S100B (95%), and GFAP (65%), indicating that the transfected cells were committed to the astrocyte lineage. Consistently, these cells did not express molecules of other lineages, such as A2B5, a glial progenitor marker, or β3 tubulin, a pan neuronal marker (Figure 6). In addition, concurrent activation of the four TFs seemed to be necessary for rapid induction toward the astrocyte lineage, as activation of only one TF (SOX9, NFIA, NFIB, or NFIX) failed to induce GFAP immunoreactivity from human iPSCs during the same induction period (data not shown).

**DISCUSSION**

In this work, we developed a PB-based CRISPRa system that allows for rapid sgRNA screening and stable activation of multiple TFs and lncRNAs. We tested the system in human iPSC lines and 293FT cells, and we compared the upregulation magnitude of gene expression level and basal transcript level, across genes and cell types. In addition, we showed that the PB-CRISPRa system could accelerate human iPSC differentiation into neurons and astrocytes by simultaneously augmenting endogenous expression of specific sets of TFs.

Two excellent examples of CRISPR-mediated gene activation systems, including the SAM46 and the VPR fusion protein systems,37 were reported to be able to regulate endogenous gene expression robustly. The SAM system was predicted to activate the endogenous expression of any given gene with only one sgRNA, as long as it targeted a PAM within 200 bp upstream of a TSS.16 To simplify vector design and reduce experimental variability, we decided to adapt the SAM platform to our multiplex transactivation system. We packaged all SAM system activation components into a PB vector, and we generated SAM overexpression stable cell lines for sgRNA screening and verification. Of the sgRNAs tested here, 60% (43 of 72) were able to increase mRNA expression by at least 2-fold in 293FT-SAM cells. When tested in human iPSCs, all of the optimal sgRNAs identified from 293FT-SAM achieved at least 2-fold activation as well (Figure 2). Thus, our strategy offers a convenient strategy for quick optimization of best sgRNAs for the activation of multiple endogenous genes, which provides additional feasibility and flexibility for gain-of-function experiments.

Several elegant tools have been developed to achieve multiplex sgRNA expression based on RNA processing mechanisms, including Csy4-cleavable cassettes,33,34 tRNA-sgRNA cassettes,45 and sgRNA-small hairpin RNA (shRNA) structure.46 Endonuclease Csy4 from
Pseudomonas aeruginosa recognizes a 28-nt RNA sequence and cleaves the target RNA. For multiplex sgRNA expression, sgRNAs are fused with the 28-nt Csy4-cleavable RNA sequence, and the transcript is processed into multiple sgRNAs with the addition of Csy4 protein. One limitation of this strategy is that the endonuclease Csy4 might be toxic to cells. The tRNA-sgRNA cassette approach takes advantage of the endogenous tRNA-processing system. Multiple tRNA-sgRNA units are assembled in a synthetic tRNA-sgRNA cassette, with each sgRNA containing a target-specific spacer, which can be cleaved by endogenous RNase to release mature sgRNAs and tRNA. The sgRNA-shRNA structure hijacks the endogenous shRNA processing system, and multiple sgRNAs and shRNAs are assembled in an sgRNA-shRNA structure format with interval sequences of the Drosha cutting site. Thus, the sgRNA-shRNA transcript is processed into functional sgRNAs and shRNAs by endogenous Drosha. These approaches, together with the present work that uses multiple Pol III U6 promoters in a single vector, provide a wide range of selections for simultaneous sgRNA expression for the purpose of CRISPR-mediated endogenous gene activation.

During development, master TFs dictate the gene expression regulatory network to drive cell fate determination, which provides a basis for direct conversion experiments across cell types, even different germ layers. Recently, several groups reported successful conversion of fibroblasts to neurons, myocytes, or iPSCs by endogenous activation of TFs using the CRISPRa systems. The CRISPRa system also allows for fast differentiation of pluripotent stem cells to endodermal lineages, trophoblast stem cells, extraembryonic endoderm cells, and neurons. However, there are at least two drawbacks in these systems: (1) most experiments required multiple sgRNAs to activate a single TF, and (2) dCas9-activator fragments were delivered separately. These limitations might decrease conversion efficiency. An all-in-one vector could help overcome the technical barriers and boost reprogramming efficiency. Our PB-CRISPRa system is able to stably activate multiple TFs by expressing corresponding optimal sgRNAs for multiple genes at the same time. Our data indicated that the all-in-one vectors achieved higher activation magnitude than the one-by-one vectors that relied on co-transfection of activators and multiple sgRNA plasmids. Using this system, we were able to accelerate directed neural lineage differentiation from human iPSCs (Figures 5 and 6). Thus, our PB-CRISPRa system is a necessary addition to the collection of gene activation tools.
Simultaneous perturbation of multiple TFs is a desired approach to unfold the collateral functions of TFs during development. Genetic manipulation of multiple genes in human cells has been more challenging to achieve than in animal models as it often requires multiplex targeting, which is difficult to perform even with multiple rounds of transfection. Indeed, technical limitations often restrict researchers going from single- or double-gene manipulation to multiple-gene perturbation. Our PB multiplex system has a large packaging capacity and is able to carry multiple components in a single plasmid. This is important as it brings extra flexibility in controlling the gene expression of groups of genes of interest.

One of the most appealing applications for the CRISPR system is the potential for gene therapy. Critical challenges for in vivo delivery of transgenes for gene therapy include limited cargo size and low efficiency. The PB transposon is a non-viral system with large cargo capacity. Previous reports showed that PB vectors were successfully delivered in vivo to the liver and the lung via mouse tail vein injections, and they maintained long-term gene expression. A recent work on rapid in vivo CRISPR library screening of tumor suppressor genes using the PB system further confirmed the high efficiency and broadened potential applications of the cooperation of PB and CRISPR systems. As a logical extension, our PB-CRISPRa system has the potential to be developed into a highly efficient gene delivery tool for in vivo gene activation in the future. When combined with previously validated reagents, such as nanoparticle, liposome, or electroporation, PB-CRISPRa also could activate genes in different target organs.

One limitation for PB vectors is that they are prone to integrate into transcription units of the genome. Integrated transgenes could function as oncogenes or may elicit immune responses in the target cells. Although re-expression of PBase could mediate the excision of integrated PB vectors, efficiency could be a challenge. Several procedures, such as inclusion of an inducible herpes simplex virus-1 thymidine kinase (hsv-TK) cassette in the PB vectors, allow for the selection of cells free of PB vector integration. In this case, cells with successful removal of PB vectors do not express TK-encoded thymidine kinase, and they will not convert Fialuridine (FIAU), the drug used for selection, to the toxic Fluorouracil (5-FU). To further enhance the excision efficiency, a hyperactive PBase was developed that had an increased excising efficiency in the human iPSC reprogramming process. The same PBase has also been effectively applied to gene therapy in vivo.

As 293FT cells have served as a workhorse for screening sgRNAs for multiple genes previously, we started out to test all of our 72 sgRNAs in 293FT cells first. However, the ability of the SAM system to activate a gene is dependent upon the epigenetic properties of that locus, which vary in different cell types. Therefore, it is possible that results from 293FT cells would not be predictive of human iPSCs. To address this question, we compared the activation efficiency of individual sgRNAs across the two cell types for selected genes (Figure S7). The overall trends were that the sgRNAs that were able to activate a target in 293FT cells also could efficiently activate it in human iPSCs. It is also interesting to note that the rank-order activity of the three sgRNAs for a given locus was similar, but not entirely the same, for the two cell types. For instance (Figure S7), the best sgRNAs that had the highest gene activation in MTY1, FOXG1, or POU3F1 were the same in the two cell lines, while the second or third ranked sgRNAs switched positions. Future experiments comparing additional loci across different cell types are warranted to fully reveal the effects of epigenetic properties of different loci.

Our data showed that the transactivation efficiency was higher with four-in-one constructs compared to the corresponding four one-by-one
constructs. This is probably due to the following reasons. First, the transfection efficiency for individual PB vectors varies. Second, the four-in-one strategy guaranteed the delivery of sgRNAs for activating all four TFs as well as SAM components (Figure 4), while the one-by-one strategy could not ensure concurrent delivery of SAM components and sgRNAs into the same cells. It would be interesting to determine whether there was a potential for cross-regulation among those genes. In that case, a multiplex-inducible system that could control the activation of individual TFs is needed. Assays that measure gene expression in single cells will facilitate the determination of such potential regulation.

In summary, we have built PB-CRISPRa, a platform that could quickly and stably activate multiple genes in human cells simultaneously. Our system has the potential to be readily scaled up for dissection of genome-wide complex transcription regulatory networks.

MATERIALS AND METHODS
Vector Construction
A vector containing EF1α promoter-driven MSPH SAM components and a vector containing CAG promoter-driven dCas9-VP64 components were obtained from Addgene (61426 and 61425). The PB backbone and PBase expression vectors were gifts from Dr. Sen Wu. To make the PB SAM vector (Figure 1A), MSPH fragments with a nuclear localization signal (NLS) and hygromycin resistance cassette and the dCas9-VP64 components with NLS and blasticidin resistance cassette were digested and ligated into PB backbone vector via Gibson assembly (New England Biolabs)59 (Figure 1A; Figure S8). The sgRNA (tracrRNA and crRNA) with MS2 stem loop fragment (Figure 1A) (Addgene, 61427) was digested and cloned into the PB vector via Gibson assembly. Sequence information of sgRNAs for 21 neural TFs and three lncRNAs is listed in Table S1. Sense and antisense strands of sgRNAs with appropriate overhangs for each gene were synthesized, incubated with T4 polynucleotide kinase (PNK) and T4 ligase buffer at 37°C for 30 min and 95°C for 5 min, and cooled down to room temperature within 2 hr. The resultant sgRNA insert of each gene was cloned into the PB-sgRNA (MS2) vector after six cycles of incubation at 37°C, for 5 min and 21°C for 5 min, respectively, with FastDigest BbsI and C14/C2 for 5 min, respectively, with FastDigest BbsI and C14/C2 for 30 min and 95°C for 5 min, and cooled down to room temperature within 2 hr. The resultant sgRNA insert of each gene was cloned into the PB-sgRNA (MS2) vector after six cycles of incubation at 37°C, for 5 min and 21°C for 5 min, respectively, with FastDigest BbsI and T7 ligase in the presence of Tango buffer, DTT, and ATP.50 Expression of sgRNA was driven by U6 promoter. All inserts were verified by sequencing.

All-in-one vectors containing expression cassettes of MSPH, dCas9-VP64, and sgRNAs for two to four genes desired to test were constructed using the Multisite Gateway strategy (Figure 3). Briefly, the PB-SAM vector described above was amplified to become PB-SAM R1-R2 DEST vector with attR1 and attR2 sites, ccdB cassette, and chloramphenicol resistance cassette (Cmr). PB-sgRNA (MS2) vectors containing different sgRNA inserts were amplified to be attached with appropriate attB sites (Figure 3). All-in-one vectors were assembled by four-way LR reactions as described previously.51 Primers used in Multisite Gateway cloning are listed in Table S2.

Cell Culture
HEK293FT cells were obtained from ATCC and cultured in high-glucose DMEM supplemented with 10% heat-inactivated fetal bovine serum, 1× GlutaMAX, and 1× non-essential amino acid (NEAA). A collection of human iPSC fluorescence protein reporters was created or obtained to facilitate the testing of the activation of neural lineage TFs or lncRNAs. Human NEUROG2-mCherry knockin reporter iPSC line was previously generated in our lab,23 and OLGJ2-EGFP iPSC knockin reporter line was created from human iPSC ND2-0 using a similar targeting strategy as previously described.23 Human DCX-ZaGreen iPSC reporter was obtained from the NIH. A non-engineered human iPSC line Cy2 was also obtained from the NIH. All human iPSCs described herein (NEUROG2-mCherry knockin reporter, OLGJ2-EGFP iPSC knockin reporter, DCX-ZaGreen reporter, and Cy2) were cultured in chemically defined mTeSR medium (STEMCELL Technologies) in feeder-free conditions and passaged every 4–5 days with Accutase (Innovative Cell Technologies) onto Matrigel- (Becton Dickinson) coated culture plates at a ratio of 1:4–1:8 with ROCK inhibitor Y-27632 (10 μM).

Generation of 293FT-SAM and Human iPSC-SAM Stable Cell Lines
To generate 293FT-SAM line, HEK293FT cells (1 × 10⁶ cells in one well of a six-well plate) were co-transfected with PB-SAM (1.5 μg) and PBase (0.5 μg) vectors using Lipofectamine 3000 (Life Technologies), following the manufacturer’s instructions. Then 48 hr after transfection, blasticidin (15 μg/mL) and hygromycin (200 μg/mL) were added to the culture medium, and a 293FT-SAM stable cell line was obtained after 2 weeks of selection. Similarly, to generate a human iPSC-SAM stable cell line, 1 × 10⁶ human Cy2 iPSCs (NIH) were co-transfected with PB-SAM (3.75 μg) and PBase (1.25 μg) vectors using Nucleofector 2B with human stem cell nucleofector kit (Lonza). Then 48 hr after electroporation, blasticidin (10 μg/mL) and hygromycin (37.5 μg/mL) were added to the culture medium, and the Cy2-iPSC-SAM stable cell line was obtained 2 weeks after selection. For both 293FT-SAM and Cy2-iPSC-SAM cells, the expression of the two SAM components was examined by qRT-PCR as described below.

Transactivation in Stable Cell Lines
Approximately 5 × 10⁵ 293FT-SAM cells (in one well of a 12-well plate) were transfeicted with 1 μg PB-MS2-sgRNA or sgRNA mass using Lipofectamine 3000. For experiments in human iPSC-SAM stable cells, 1 × 10⁶ cells were electroporated with 5 μg PB-sgRNA vector as described above. Then 48 hr after transfection, expression of the genes intended to activate was evaluated by qPCR. The sgRNAs that elicited the highest activation were selected for further experiments using normal human iPSCs (Cy2) or iPSC reporter lines.

Transactivation in Human iPSCs and iPSC Reporter Cell Lines with All-in-One Vectors
One million plain iPSCs or iPSC reporter cells were co-electroporated with all-in-one vectors (3.75 μg) and PBase vector (1.25 μg) as described above. OLGJ2-EGFP human iPSC reporter line was used.
to estimate OLIG2 activation. Similarly, NEUROG2-mCherry and DCX-ZsGreen human iPS cell lines were used for testing neuronal conversion, where NEUROG2 or DCX expression was estimated by mCherry or ZsGreen expression under a fluorescence microscope. One day after electroporation, iPSC mTeSR culture medium was changed to N2B27 medium containing DME/F12 and Neurobasal medium (1:1), with 1× N2 supplement, 1× B27 supplement, 1× NEAA, and 1× Glutamax, supplied with 20 ng/mL brain-derived neurotrophic factor (BDNF), 20 ng/mL glial cell-derived neurotrophic factor (GDNF), and 20 ng/mL neurotrophin-3 (NT3). Hygromycin (37.5 μg/mL) and blasticidin (8 μg/mL) were used for selection (Life Technologies). The Cy2 iPSC line was used for astrocyte induction, and astrocyte differentiation efficiency was estimated by GFAP immunocytochemistry. One day after electroporation, culture medium was switched to N2B27 medium supplied with 20 ng/mL recombinant bone morphogenetic protein 4 (BMP4), 8 ng/mL neurotrophic factor (CNTF, all from Life Technologies). Hygromycin (8 μg/mL) and blasticidin (8 μg/mL) were used for selection.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes eight figures and three tables and can be found with this article online at [http://dx.doi.org/10.1016/j.omtn.2017.06.007](http://dx.doi.org/10.1016/j.omtn.2017.06.007).

**AUTHOR CONTRIBUTIONS**

Conceptualization, S.L. and Y.L.; Methodology, S.L., H.X., and A.Z.; Investigation, S.L., H.X., A.Z., and D.L.; Writing – Original Draft, S.L. and Y.L.; Writing – Review & Editing, S.L. and Y.L.; Funding Acquisition, Y.L.; Resources, Y.L.; Supervision, Y.L.

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