Direct conversion of fibroblasts to functional neurons by defined factors

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Cellular differentiation and lineage commitment are considered to be robust and irreversible processes during development. Recent work has shown that mouse and human fibroblasts can be reprogrammed to a pluripotent state with a combination of four transcription factors. This raised the question of whether transcription factors could directly induce other defined somatic cell fates, and not only an undifferentiated state. We hypothesized that combinatorial expression of neural-lineage-specific transcription factors could directly convert fibroblasts into neurons. Starting from a pool of nineteen candidate genes, we identified a combination of only three factors, Ascl1, Brn2 (also called Pou3f2) and Myt1l, that suffice to rapidly and efficiently convert mouse embryonic and postnatal fibroblasts into functional neurons in vitro. These induced neuronal (iN) cells express multiple neuron-specific proteins, generate action potentials and form functional synapses. Generation of iN cells from non-neuronal lineages could have important implications for studies of neural development, neurological disease modelling and regenerative medicine.

The diverse cell types present in the adult organism are produced during development by lineage-specific transcription factors that define and reinforce cell-type-specific gene expression patterns. Cellular phenotypes are further stabilized by epigenetic modifications that allow faithful transmission of cell-type-specific gene expression patterns over the lifetime of an organism1–7. Recent work showing that four transcription factors are sufficient to induce pluripotency in primary fibroblasts demonstrated that fully differentiated cells can be induced to undergo dramatic cell fate changes5. Similarly, the transfer of somatic cell nuclei into oocytes, as well as cell fusion of pluripotent cells with differentiated cells, have proven to be capable of inducing pluripotency8–10. This remarkable transformation has been interpreted as a reversion of mature into more primitive developmental states, with a concomitant erasure of developmentally relevant epigenetic information10. Therefore, direct reprogramming between divergent lineages could be unique to reprogramming into an embryonic state, and might not be possible between different somatic cell states. However, cell fusion or forced expression of lineage-specific genes in somatic cells can induce traits of other cell types11,12. For example, the basic helix–loop–helix (bHLH) transcription factor MyoD (also called Myod1) can induce muscle-specific properties in fibroblasts but not hepatocytes13,14; ectopic expression of interleukin (IL)-2 and granulocyte–macrophage colony-stimulating factor receptors can lead to myeloid conversion in committed lymphoid progenitor cells15,16; expression of Cebpa in B cells or PU.1 (also called Sfpi1) and Cebpα in fibroblasts induces characteristics of macrophages17–19; deletion of Pux3 can induce B cells to de-differentiate towards a common lymphoid progenitor20; and the bHLH transcription factor neurogenin 3, in combination with Pdx1 and Mafa, can efficiently convert pancreatic exocrine cells into functional β-cells in vivo21. Here, we set out to determine whether specific transcription factors could directly convert fibroblasts into functional neurons.

A screen for neuronal-fate-inducing factors

Reasoning that multiple transcription factors would probably be required to reprogram fibroblasts to a neuronal fate, we cloned a total of 19 genes that are specifically expressed in neural tissues, have important roles in neural development, or have been implicated in epigenetic reprogramming (Supplementary Table 1). A pool of lentiviruses containing all 19 genes (19F pool) was prepared to infect mouse embryonic fibroblasts (MEFs) from TauEGFP knock-in mice, which express EGFP specifically in neurons22,23 (see Fig. 1a for experimental outline). Great care was taken to exclude neural tissue from the MEF preparation, and we were unable to detect evidence for the presence of neurons or neural progenitor cells in these cultures using immunofluorescence, fluorescence activated cell sorting (FACS) and polymerase chain reaction with reverse transcription (RT–PCR) analyses (Supplementary Fig. 1). However, uninfected MEFs did contain rare TuJ1-positive, TauEGFP-negative cells with fibroblast-like morphology, indicative of weak TuJ1 (that is, β-III-tubulin) expression in non-neuronal cells (Fig. 1b, c and Supplementary Fig. 1a). In contrast, 32 days after infection with the 19F pool, we detected TuJ1-positive cells with typical neuronal morphologies and bright TauEGFP fluorescence (Fig. 1d,e). Thus, some combination(s) of the genes in the 19F pool was capable of converting MEFs into iN cells.

We next set out to narrow down the number of transcription factors required for generation of iN cells. Given their important roles in neuronal cell fate determination24–26 we first tested the bHLH transcription factors Ascl1 (also known as Mash1) and Neuronl1 individually. Notably, we observed occasional TuJ1-positive, TauEGFP-negative cells with fibroblast-like morphology, indicative of weak TuJ1 (that is, β-III-tubulin) expression in non-neuronal cells (Fig. 1b, c and Supplementary Fig. 1a). In contrast, 32 days after infection with the 19F pool, we detected TuJ1-positive cells with typical neuronal morphologies and bright TauEGFP fluorescence (Fig. 1d,e). Thus, some combination(s) of the genes in the 19F pool was capable of converting MEFs into iN cells.

We therefore tested the neuron-inducing activity of Ascl1 in combination with each of the remaining 18 candidate genes (Supplementary Fig. 2a). Five genes (Brn2, Brn4 (Pou3f4), Myt1l, Zic1 and Olig2) substantially potentiated the neuron-inducing activity of Ascl1 (Supplementary Fig. 2a, b). Importantly, none of these five genes generated iN cells when tested individually (data not shown). Next, we tested whether combinatorial expression of these factors with Ascl1 could further increase the induction of neuron-like cells by infecting...
TauEGFP MEFs with a pool of Brn2, Myt1l, Zic1, Olig2 and Ascl1 viruses (5F pool). Given its close similarity to Brn2, we did not include Brn4 in the 5F pool. Twelve days after infection, we detected frequent Tuj1-positive cells (red) with highly complex morphologies (Fig. 1f). These 5F iN cells also expressed the pan-neuronal markers TauEGFP (h), NeuN (red, i) and MAP2 (red, j). k, Representative traces of membrane potential responding to step depolarization by current injection (lower panel). Membrane potential was current-clamped at around −65 mV. l, Representative traces of whole-cell currents in voltage-clamp mode; cells were held at −70 mV; step depolarization from −90 mV to +50 mV at 10-mV intervals was delivered (lower panel). The inset shows Na⁺ currents. m, Spontaneous action potentials recorded from a 5F MEF iN cell 8 days after infection. No current injection was applied. n–p, Twenty-two days after infection 5F MEF iN cells expressed synapsin (red, n) and vesicular glutamate transporter 1 (vGLUT1; red, o) or GABA (p). Scale bars: 5 μm (o), 10 μm (e, j, n, p) 20 μm (c, h, i) and 200 μm (f).

TauEGFP-positive cells on days 8, 12 and 20 after infection. For the majority of the iN cells analysed (85.1%, n = 47), action potentials could be elicited by depolarizing the membrane in current-clamp mode (Fig. 1k). Six cells (14.2%, n = 42) showed spontaneous action potentials, some as early as 8 days after transduction (Fig. 1m). These action potentials could be blocked by tetrodotoxin (TTX), a specific inhibitor of Na⁺ ion channels (Supplementary Fig. 3e). Moreover, in voltage-clamp mode we observed both fast, inactivating inward and outward currents, which probably correspond to opening of voltage-dependent K⁺- and Na⁺-channels, respectively, with a possible contribution of Ca²⁺ channels to the whole-cell currents (Fig. 1l and traces of membrane potential responding to step depolarization by current injection (lower panel)). Membrane potential was current-clamped at around −65 mV. l, Representative traces of whole-cell currents in voltage-clamp mode; cells were held at −70 mV; step depolarization from −90 mV to +50 mV at 10-mV intervals was delivered (lower panel). The inset shows Na⁺ currents. m, Spontaneous action potentials recorded from a 5F MEF iN cell 8 days after infection. No current injection was applied. n–p, Twenty-two days after infection 5F MEF iN cells expressed synapsin (red, n) and vesicular glutamate transporter 1 (vGLUT1; red, o) or GABA (p). Scale bars: 5 μm (o), 10 μm (e, j, n, p) 20 μm (c, h, i) and 200 μm (f).

Characterization of 5-factor iN cells
To explore whether iN cells have functional membrane properties similar to neurons, we performed patch-clamp recordings of

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Supplementary Fig. 3f). The resting membrane potentials ranged between −30 and −69 mV with an average of −55 mV on day 20 ($n = 12$, Supplementary Tables 2 and 3). Additionally, we asked whether these cells possessed functional ligand-gated ion channels. Induced neuronal cells responded to exogenous application of GABA ($\gamma$-aminobutyric acid), and this response could be blocked by the GABA receptor antagonist picrotoxin (Supplementary Fig. 3g). Thus, MEF-derived iN cells seem to exhibit the functional membrane properties of neurons and possess ligand-gated GABA receptors.

We then sought to characterize the neurotransmitter phenotype of iN cells. After 21 days of culture in minimal neuronal media, we detected vGLUT1-positive puncta outlining MAP2-positive neurites of some cells, indicating the presence of excitatory, glutamatergic neurons (Fig. 1o). In addition, we found iN cells labelled with antibodies against GABA, the major inhibitory neurotransmitter in brain (Fig. 1p). Some iN cells (9 out of ~500) contained the Ca$^{2+}$-binding protein calretinin, a marker for cortical interneurons and other neuronal subtypes (Supplementary Fig. 3c). No expression of tyrosine hydroxylase, choline acetyltransferase or serotonin was detected. The vast majority of iN cells were negative for peripherin, an intermediate filament characteristic of peripheral neurons (data not shown)\(^3\).

**Functional neurons from tail fibroblasts**

To evaluate whether iN cells could also be derived from postnatal cells, we isolated tail-tip fibroblasts (TTFs) from 3-day-old TauEGFP and Rosa26-rtTA mice\(^3\). Similar to our MEF cultures, we could not detect pre-existing neurons, glia, or neural progenitor cells (Supplementary Fig. 1a). Twelve days after infecting TTFs with the 5F pool, Tuji-positive iN cells with a complex, neuronal morphology could be readily detected (Fig. 2a). TTF iN cells expressed the pan-neuronal markers NeuN, MAP2 and synapsin (Fig. 2b, c, f). Electrophysiological recordings 12 days after infection demonstrated an average resting membrane potential of −57 mV (range: −35 to −70 mV, $n = 11$), firing of action potentials (81.8%, $n = 11$) (Fig. 2d),

![Figure 2](image1.png)

**Figure 2 | Efficient induction of neurons from perinatal tail-tip fibroblasts**

a, Tuji-stained tail-tip fibroblast (TTF) 13 days after infection with the 5F pool. b, c, TTF iN cells express the pan-neuronal markers MAP2 (b) and NeuN (c). d, Representative traces showing action potentials elicited at day 13 after infection. Nine of eleven cells recorded exhibited action potentials. e, Whole-cell currents recorded in voltage-clamp mode. Inward fast inactivating Na$^+$ currents (arrow) and outward currents can be observed. f–h, At 21 days after infection TTF iN cells express synapsin (red, f), vGLUT1 (red, g) and GABA (h). c, f and g are overlay images with the indicated marker (red) and Tuji (green). Scale bars: 20 μm (b, f, g), 100 μm (h), 200 μm (a).

![Figure 3](image2.png)

**Figure 3 | The 5F-pool-induced conversion is rapid and efficient**

a, Tuji-positive iN cells (red) show morphological maturation over time after viral infection. At day 13, TauEGFP expression outlines neuronal processes. b, FACS analysis of TauEGFP expression 8 and 13 days after infection. Control indicates uninfected TauEGFP MEFs. APC, allophycocyanin. c, Representative traces showing action potentials elicited from MEF iN cells at days 8, 12 and 20 after infection. Cells were maintained at a potential of approximately −65 to −70 mV. Step current injection protocols were used from −50 to +70 pA. Scale bar applies to all traces. d–g, Quantification of membrane properties in MEF iN cells at 8, 12 and 20 days after infection. Numbers in the bars represent the numbers of recorded cells. Data are presented as mean ± s.e.m. *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$ (Student’s $t$-test). AP, action potential; $C_m$, membrane capacitance; $R_m$, membrane input resistances; RMP, resting membrane potential. Action potential heights were measured from the baseline. h, BrdU-positive iN cells after BrdU treatment from day 0–13 or day 1–13 after transgene induction. i, Example of a Tuji (green) positive cell not labelled with BrdU (red) when added at day 0 after addition of doxycycline. Data are presented as mean ± s.d. j, Efficiency estimates for iN cell generation 13 days after infection (see Methods). Every bar represents an independent experiment. Doxycycline was added 48 h after plating in MEF experiment 1 and after 24 h in MEF experiments 2 and 3. Error bars indicate ± s.d. of cell counts. Scale bars: 10 μm (i), 100 μm (a).
Neuronal induction is fast and efficient

Next, we assessed the kinetics and efficiency of 5F iN conversion. In MEFs, Tuj1-positive cells with immature neuron-like morphology were found as early as 3 days after infection (Fig. 3a). After 5 days, neuronal cells with long, branching processes were readily detected, and over time increasingly complex morphologies were evident, suggesting an active process of maturation in newly formed iN cells (Fig. 3a). Similarly, we detected TauEGFP expression as early as day five (Supplementary Fig. 3h). The fraction of TauEGFP-positive cells remained similar at 8 and 13 days after infection, as determined by FACS analysis, indicating no de novo generation of iN cells after day 8 (Fig. 3b). Electrophysiological parameters such as action potential height, resting membrane potential, membrane input resistance and membrane capacitance also showed signs of maturation over time (Fig. 3c–g and Supplementary Tables 2 and 3).

To estimate the conversion efficiency, we first determined how many of the MEF-derived iN cells divided after induction of the viral transgenes by treating the cells with 5-bromodeoxyuridine (BrdU) after gene induction. The results showed that the vast majority of iN cells received the necessary complement of viral transgenes.

Neuronal induction is fast and efficient

Because iN cells exhibited the membrane properties of neurons, we next wanted to assess whether iN cells have the capacity to form functional synapses. To accomplish this we used two independent methods. First, we determined whether iN cells were capable of synaptically integrating into pre-existing neural networks. We used FACS to purify TauEGFP-positive iN cells and observed spontaneous and rhythmic network activity typical of cortical neurons in culture (Fig. 4a, b). Both excitatory and inhibitory postsynaptic currents (EPSCs and IPSCs) could be evoked after electrical stimulation delivered from a concentric electrode placed 100–150 μm away from the patched iN cells (Fig. 4b–d). In the presence of the N-methyl-D-aspartate (NMDA) and non-NMDA receptor channel blockers 6-cyano-7-nitroquinolinic acid-2,3-dione (CNQX) and d(-)-2-amino-5-phosphonovaleric acid (d-AP5), spontaneous IPSCs were reliably detected (Fig. 4c, upper panel). Evoked IPSCs could be blocked by further addition of picrotoxin (Fig. 4c, middle panel). Similarly, at a holding potential of −70 mV and in the presence of picrotoxin, fast-decaying EPSCs mediated by AMPA receptors could be evoked (Fig. 4d, middle panel). Conversely, at a holding potential of +60 mV (which relieves the voltage-dependent blockade of NMDA receptor-mediated currents) spontaneous EPSCs could be observed. At a holding potential of −70 mV, AMPA receptor (AMPA)-mediated EPSCs were monitored. When the holding potentials were set at +60 mV, both AMPA- and NMDA-mediated EPSCs could be recorded. The lower panel shows the short-term synaptic plasticity of both AMPAR- and NMDAR-mediated synaptic activities. Arrows indicate time of stimulation. A. Example of a TauEGFP-positive iN cell expressing MAP2 among cortical neurons. B. Higher magnification (×7) of area marked in a. C. Representative spontaneous post synaptic currents (PSCs) recorded from MEFF iN cells co-cultured with glia. D. Representative traces of evoked EPSCs. NMDAR-mediated EPSCs in the presence of 10 μM NBQX (2,3-dihydroxy-6-nitro-7-sulphamoylbenzo[f]quinolinic acid-2,3-dione), an AMPA-receptor antagonist, were recorded at a holding potential (V_h) of +60 mV. The AMPAR-evoked response was blocked by NBQX and AMPA. E. Current–voltage (I–V) relationship of NMDAR-mediated EPSCs (left panel); representative traces of evoked EPSCs at different V_h are indicated. The right panel shows the summarized I–V relationship. NMDAR EPSC amplitudes (I_{NMDA}) are normalized to EPSCs at a V_h of +60 mV (indicated by an asterisk, n = 5).

NMDAR EPSCs show ratifications at negative holding potentials, presumably because of the blockade of NMDAR by Mg^2+. Scale bars: 10 μm (a, e).
Mg²⁺ to NMDA receptors), slow-decaying NMDA-receptor-mediated EPSCs could be recorded (Fig. 4d, middle panel).

Moreover, synaptic responses recorded from iN cells showed signs of short-term synaptic plasticity, such as depression of IPSCs and facilitation of EPSCs during a high-frequency stimulus train (Fig. 4c, d, lower panels). The presence of synaptic contacts between iN cells and cortical neurons was independently corroborated by the immunocytochemical detection of synapsin-positive puncta surrounding MAP2-positive dendrites originating from EGFP-positive cells (Fig. 4e, f). We were also able to observe synaptic responses in similar experiments performed with iN cells derived from TTFs (Supplementary Fig. 4). These data demonstrate that iN cells can form functional postsynaptic compartments and receive synaptic inputs from cortical neurons.

Next we asked whether iN cells were capable of forming synapses with each other. To address this question we plated FACS-sorted TauEGFP-positive, MEF-derived 5F iN cells 8 days after infection onto a monolayer culture of primary astrocytes, which are thought to have an essential role in synaptogenesis. Importantly, we confirmed that these cultures were free of pre-existing Tuj1- or MAP2-positive neurons (data not shown). Patch-clamp recordings at 12–17 days after sorting indicated the presence of spontaneous postsynaptic currents in 5 out of 11 cells (Fig. 4g). Upon extracellular stimulation, evoked EPSCs could be elicited in a majority of the cells (9 out of 11 cells, Fig. 4h). Similar to iN cells cultured with primary cortical neurons, we were able to record both NMDA-receptor-mediated (9 out of 11 cells) and AMPA-receptor-mediated EPSCs (8 out of 11 cells; Fig. 4h, i). Interestingly, we were unable to detect obvious IPSCs in a total of 15 recorded 5F iN cells. These data indicate that iN cells are capable of forming functional synapses with each other, and that the majority of iN cells exhibit an excitatory phenotype.

Genes sufficient for neuronal conversion

As stated earlier, Ascl1 was the only gene from the 5F pool that was sufficient to induce neuron-like cells in MEFs. We next attempted to determine the relative contribution of each of the five genes by removing each gene from the pool and assessing the efficiency of iN cell generation. Surprisingly, only the omission of Ascl1 had a marked effect on induction efficiency (Supplementary Fig. 6a). Thus, we tested the effects of removing two genes at a time by evaluating all possible three-gene combinations. Our results indicated that either Ascl1 or both Brn2 and Myt1l must be present to generate iN cells (Fig. 5a). The most efficient conversions were achieved when Ascl1 and Brn2 were combined with either Myt1l (BAM pool) or Zic1 (BAZ pool). The efficiencies in these conditions were two- to three-fold higher than the 5F pool when the total amount of virus was kept constant (Fig. 5a–d). In this experiment the BAM iN cells appeared to have a more complex morphology than the BAZ cells (Fig. 5c, d and Supplementary Fig. 2b). MEFs infected with the 5F (Fig. 5e) and BAM (Fig. 5f) pools, e, iN staining of perinatal TTF iN cells 13 days after infection with the BAM pool. f, BAM-induced MEF iN cells express MAP2 (green) and synapsin (red) 22 days after infection.

Figure 5 | Defining a minimal pool for efficient induction of functional iN cells a, Quantification of Tuj1-positive iN cells from TauEGFP MEFs infected with different three-factor combinations of the five genes. Each gene is represented by the first letter in its name. A, Ascl1; B, Brn2; M, Myt1l; O, Olig2; Z, Zic1. Averages from 30 randomly selected visual fields are shown (error bars indicate ± s.d.). b–d, Representative images of Tuj1 staining of MEFs infected with the 5F (b), Ascl1 + Brn2 + Zic1 (BAZ; c) and Ascl1 + Brn2 + Myt1l (BAM; d) pools. e, Tuj1 staining of perinatal TTF iN cells 13 days after infection with the BAM pool. f, BAM-induced MEF iN cells express MAP2 (green) and synapsin (red) 22 days after infection. g, Representative traces of synaptic responses recorded from MEF-derived BAM (3F) iN cells co-cultured with glia after isolation by FACS. Vh, holding potential. At a Vh of −70 mV, AMPAR-mediated EPSCs were recorded; at a Vh of +60 mV, NMDAR-mediated EPSCs were revealed. h, Synaptic responses recorded from TTF-derived 3F iN cells. i, Representative traces of action potentials elicited from MEF-derived iN cells transduced with the indicated gene combinations, recorded 12 days after infection. Cells were maintained at a resting membrane potential of approximately −65 to −70 mV. Step current injection protocols were used from −50 to +70 pA. Traces in each subgroup (left or right panels) represent subpopulations of neurons with similar responses. Numbers indicate the fraction of cells from each group that were qualitatively similar to the traces shown. Right panels: representative images of Tuj1 staining after recordings from each condition. Scale bars: 20 μm (f) and 100 μm (b, i).

smaller subsets of the BAM pool to determine their functionality. In many Ascl1-induced cells, current injection elicited action potentials, but their properties appeared to be immature, consistent with their simple neurite morphology (Fig. 5i and Supplementary Fig. 2b). MEFs infected with Ascl1 and Brn2 or Myt1l generated more mature action potentials and displayed more complex neuronal morphologies. In contrast, the majority of BAM iN cells exhibited repetitive action potentials with more mature characteristics, and displayed the most
complex neuronal morphologies. Thus, it seems likely that Ascl1 alone is sufficient to induce some neuronal traits, such as expression of functional voltage-dependent channel proteins that are necessary for the generation of action potentials; however, co-infection of additional factors is necessary to facilitate neuronal conversion and maturation.

**Discussion**

Here we show that expression of three transcription factors can rapidly and efficiently convert mouse fibroblasts into functional neurons (iN cells). Although the single factor Ascl1 was sufficient to induce immature neuronal features, the additional expression of Brn2 and Myt1l generated mature iN cells with efficiencies of up to 19.5% (Supplementary Fig. 6b). Three-factor iN cells displayed functional neuronal properties such as the generation of trains of action potentials and synaptic formation. These transcription factors were identified from a total of 19 candidates that we selected because of their specific expression in neural cell types or their roles in reprogramming to pluripotency (see Methods).

Despite the heterogeneity of embryonic and TTF cultures, the highly efficient nature of this process effectively rules out the possibility that directed differentiation of rare stem or precursor cells with neurogenic potential can explain our observations. Future studies will have to be performed to unequivocally demonstrate that terminally differentiated cells such as mature B or T lymphocytes can be directly converted into neurons using this approach. It will now be of great interest to decipher the molecular mechanism of this fibroblast to neuron conversion. We assume that high expression levels of strong neural cell-fate-determining transcription factors can activate salient features of the neuronal transcriptional program. Auto-regulatory feedback and feed forward activation of downstream transcriptional regulators could then reinforce the expression of important cell-fate-determining genes and help to further stabilize the induced transcriptional program. Robust changes in transcriptional activity could also lead to genome-wide adjustments of repressive and active epigenetic features such as DNA methylation, histone modifications and changes of chromatin remodelling complexes that further stabilize the new transcriptional network. It is possible that certain subpopulations of cells are 'primed' to respond to these factors, depending on their pre-existing transcriptional or epigenetic states.

The majority of iN cells described in this report are excitatory and express markers of cortical identity. A low proportion of iN cells expressed markers of GABAergic neurons, but no other neurotransmitter phenotypes were detected. Our data indicate the intriguing possibility that additional combinations of neural transcription factors might also be able to generate functional neurons whose phenotypes remain to be explored. One of the next important steps will be to generate iN cells of other specific neuronal subtypes and from human cells.

Future studies will be necessary to determine whether iN cells could represent an alternative method to generate patient-specific neurons. The generation of iN cells is fast, efficient and devoid of tumorigenic pluripotent stem cell approaches, a key complication of induced pluripotent stem cell approaches in regenerative medicine. Therefore, iN cells could provide a novel and powerful system for studying cellular identity and plasticity, neurological disease modelling, drug discovery and regenerative medicine.

**METHODS SUMMARY**

Fibroblast isolation, cell culture and molecular cloning. Homozygous TauEGFP knock-in mice were purchased from the Jackson Laboratories and bred with C57BL/6 mice (Tacnic) to generate TauEGFP heterozygous embryos. MEFS were isolated from E14.5 embryos using a dissection microscope (Leica). Tail tips were sliced into small pieces, trypsinized and plated to derive fibroblast cultures. All fibroblasts were expanded for three passages before being used for experiments. Complementary DNAs for candidate genes were cloned into doxycycline-inducible lentiviral vectors, as described previously. MEFS were infected overnight and cultured in MEF media with doxycycline for 48 h before being transferred into N3 media with doxycycline.

**FACS analysis, cortical cultures, glia cultures.** For glia and cortical co-cultures cells were trypsinized 8 days after infection and EGFP-positive cells were isolated using a FACS Aria II (Becton Dickinson). EGFP-positive cells were replated on 7 days in vitro cortical cultures from wild-type p0 mice, or alternatively, passage three primary astrocytes isolated from p5 pups (as described previously). Electrophysiology and immunofluorescence analysis. Cells were analysed at indicated times after infection. Action potentials were recorded with current-clamp whole-cell configuration. The pipette solution for current-clamp experiments contained (in mM): 123 K-gluconate, 10 KCl, 1 MgCl₂, 10 HEPES, 1 EGTA, 0.1 CaCl₂, 1 KATP, 0.2 NaGTP and 4 glucose, pH adjusted to 7.2 with KOH. Membrane potentials were kept around −65 to −70 mV, and step currents were injected to elicit action potentials. Whole-cell currents including sodium currents, potassium currents were recorded at a holding potential of −70 mV, voltage steps ranging from −80 mV to +90 mV were delivered at 10 mV increments. Synaptic responses were measured as described previously. For immunofluorescence experiments, cells were fixed in 4% paraformaldehyde in PBS for 10 min. Antibodies were diluted to indicated concentrations (see Methods).

**Full Methods** and any associated references are available in the online version of the paper at www.nature.com/nature.

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Author Contributions T.V., A.O. and M.W. designed and conceived the experiments. T.V., Y.K. and M.W. produced the lentiviral vectors. T.V. and A.O. performed the lentiviral infections, isolated the fibroblasts and completed the molecular characterization of the iN cells. Z.P.P. and T.C.S. designed, performed and analysed the electrophysiological assays. T.V., A.O., Z.P.P., T.C.S. and M.W. wrote and edited the manuscript and produced the figures.

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METHODS

Embyronic fibroblast culture. Homozygous TauEGFP knock-in mice were purchased from the Jackson Laboratories and bred with C57BL/6J mice (Taconic) to generate BAC-TauEGFP heterozygous embryos. BALB/c mice were purchased from Taconic, and RA26-TTA mice were obtained from R. Jaensch.26 MEFs were prepared from E14.5 embryos under a dissection microscope (Leica). The head, vertebral column (containing the spinal cord), dorsal root ganglia and all internal organs were removed and discarded to ensure the removal of all cells with neurogenic potential from the cultures. The remaining tissue was manually dissociated and incubated in 0.25% trypsin (Sigma) for 10–15 min to create a single cell suspension. The cells from each embryo were plated onto a 15-cm tissue culture dish in MEF media (Dulbecco’s Modified Eagle Medium; Invitrogen) containing 10% fetal bovine serum (FBS; HyClone), β-mercaptoethanol (Sigma-Aldrich), non-essential amino acids, sodium pyruvate and penicillin/streptomycin (all from Invitrogen). Cells were grown at 37°C for 4–7 days until confluent, and then split once before being frozen. After thawing, cells were cultured on 15-cm plates and allowed to become confluent before being split onto plates for infections using 0.25% trypsin. Postnatal TTFs were prepared by removing the bottom third of tail from 3-day-old pups using surgical scissors. Cells were rinsed in ethanol, washed with HBSS (Sigma), and then dissociated using trypsin and 0.25% trypsin. TTFs were cultured in MEF media until confluent and passaged once before being pooled together and frozen down for further use.

Cell culture, molecular cloning and viral infections. We had three criteria for identifying candidates with neuron-inducing activity: (1) we reasoned that cell-fate inducing factors should be enriched in the gene category of transcriptional regulators. (2) We included factors previously involved in reprogramming to pluripotency (Klf4, c-Myc and Sox2). (3) We searched for genes specifically expressed in neural tissues. Those were selected based on published expression arrays of MEFs, embryonic stem cells and neuronal progenitor cells retrieved from the Gene Expression Omnibus database (GSE8024, http://www.ncbi.nlm.nih.gov/geo) and the EST Profile function of NCBI’s Unigene database (http://www.ncbi.nlm.nih.gov/unigene). cDNAs for the factors included in the 19 factor pool were cloned into lentiviral constructs under the control of the tetracycline operator27. Replication-incompetent, VSVg-coated lentiviral particles were packaged in 293T cells as described28. Passage three TauEGFP and BALB/c MEFs were infected in MEF media containing polybrene (8 μg ml⁻¹). After 16–20 h in media containing lentivirus, the cells were switched into fresh MEF media containing doxycycline (2 μg ml⁻¹) to activate expression of the transduced genes. After 48 h in MEF media with doxycycline (Sigma), the media was changed every 2–3 days for the duration of the culture period. For BrdU experiments, 10 μM BrdU was added to the culture media and was maintained throughout media changes until the cells were fixed.

Immunofluorescence, RT–PCR and flow cytometry. Neuronal cells were defined as cells that stained positive for TuJ1 and had a process at least three times longer than their cell body. For immunofluorescence staining, cells were washed with PBS and then fixed with 4% paraformaldehyde for 10 min at room temperature. Cells were then incubated in 0.2% Triton X-100 (Sigma) in PBS for 30 min at room temperature. After washing twice with PBS, cells were blocked in a solution of PBS containing 4% BSA and 1% FBS. Fields of cells for staining were outlined with a PAP pen (Dako). Primary and secondary antibodies were typically applied for 1 h and 30 min, respectively. Cells were washed three times with PBS between primary and secondary staining. For anti-BrdU staining, cells were treated with 2 N HCl in PBS for 10 min and washed twice with PBS before permeabilization with Triton X-100 (Sigma). The following antibodies were used for our analysis: goat anti-choline acetyltransferase (Millipore, 1:100), rabbit anti-GABA (Sigma, 1:4,000), rabbit-GFAP (DAKO, 1:4,000), mouse anti-BrdU (Abcam, 1:1,000), rabbit anti-ChAT (Millipore, 1:1,000), mouse anti-Sox2 (R&D Systems, 1:50), rabbit anti-serotonin (Biogenesis, 1:100), rabbit anti-Tuj1 (Covance, 1:1,000), mouse anti-Tuj1 (Chemicon, 1:100), goat anti-Bn2 (Santa Cruz Biotechnology, 1:100), anti-mouse-BrdU (Becton Dickinson, 1:3,5), mouse anti-calretinin (Dako, 1:100), sheep anti-tyrosine hydroxylase (Pel-Freez, 1:100), E028 rabbit anti-synapsin (gift from T. Sudhof, 1:500), guinea-pig anti-vGLUT1 (Millipore, 1:2,000), mouse anti-GAD6 (Developmental Studies Hybridoma Bank (DSHB), 1:500), mouse anti-Pax3 (DSHB, 1:250), mouse anti-Pax6 (DSHB, 1:30), mouse anti-Pax7 (DSHB, 1:250), mouse anti-Nkx2.2 (DSHB, 1:100), mouse anti-Olig1 (NeuroMab, 1:100), FITC- and Cy3-conjugated secondary antibodies were obtained from Jackson Immunoresearch. Alexa-488-, Alexa-546- and Alexa-633-conjugated secondary antibodies were obtained from Invitrogen. TauEGFP-expressing cells were analysed and sorted on a FACS Aria II (Becton Dickinson). Flow cytometry data were analysed using Flowjo (Tree Star). After sorting, cells were plated on cortical cultures or glia cultures derived from neonatal brains. Cells were kept in 50% N3 media and 50% growth media (see media composition below) and 2 μg ml⁻¹ doxycycline for 1 week before being switched to growth media without doxycycline until electrophysiological analysis was completed. For RT–PCR analysis, RNA was isolated using Trizol (Invitrogen) following the manufacturer’s instructions, treated with DNase (NEB) and 1.5 μg was reverse-transcribed with Superscript II (Invitrogen). PCR was performed using the following primers: Sox2, forward 5’-TGGATCTTGTTGCGGAT-3’; Sox10, forward 5’-GACGAGGTGCGGAGG-3’; reverse 5’-CGCTGTCTACCTTGTCATC-3’; β-actin, forward 5’-CGTTGGCGCGCTACAGGA-3’; reverse 5’-CTTAAGGGTCAGGGGGGC-3’. PCR products were analysed on a 1% gel.

Efficiency calculation. The following method was used to calculate the efficiency of neuronal induction. The total number of TuJ1+ cells with a neuronal morphology, defined as cells having a circular, three-dimensional appearance that extend a thin process at least three times longer than their cell body, were quantified 12 days after infection. This estimate was based on the average number of IN cells present in 30 randomly selected ×20 visual fields. The area of a ×20 visual field was then measured, and we used this estimated density of IN cells to determine the total number of neurons present in the entire dish. We then divided this number by the number of cells plated before infection to get the percentage of the starting population of cells that adopted neuron-like characteristics.

Cortical cultures. Primary cortical neurons were isolated from newborn wild-type mice as described29,30 with modifications. Briefly, cortices were dissociated by papain (10 U ml⁻¹, with 1 μM Ca²⁺ and 0.5 μM EGTA) digestions and plated on Matrigel-coated circle glass coverslips (11 mm). The neurons were cultured in vitro in growth media consisting of MEM (Invitrogen) supplemented with B27 (Invitrogen), glucose, transferrin, FBS and Ara-C (Sigma).

Glia cell isolation. Forebrains were dissected from postnatal day five wild-type mice and were manually dissociated into ~50 μm² pieces in a total of 2 ml of HBSS. A total of 500 μl of 2.5% trypsin and 1% DNase were added and dissociated tissue was incubated at 37°C for 15 min. Solution was mixed every 5 min. The supernatant was then transferred into 1.5 ml FBS. A total of 4 ml HBSS, 500 μl 2.5% trypsin, and 500 μl DNase were again added to the remaining dissociated tissue and incubated at 37°C for 15 min, mixing every 5 min. The supernatant was again removed and added to the FBS-containing solution. Using a pipette, the remaining tissue was further dissociated and passed through a 70 μm nylon mesh filter (BD Biosciences) into the FBS-containing solution. The cell mixture was then spun at 1,000 r.p.m. for 5 min and re-suspended in HBSS media. Glia cells were passaged three times before culturing with MEF or TTF-derived IN cells. Contaminating neurons in p3 glia cell cultures could not be detected when stained for either TuJ1 or MAP2.

Electrophysiology. Recordings were performed from MEF- and tail-cell-derived IN cultures at 8, 12 and 20 days after viral infection, or ~7 days after co-culturing with cortical neurons. Spontaneous or evoked synaptic responses were recorded in the whole-cell voltage-clamp mode. Evoked synaptic responses were triggered by 1-ms current injection through a local extracellular electrode (FHC concentric bipolar electrode, Catalogue number CBAEC75) with a Model 2100 Isolated Pulse Stimulator (A-M Systems), and recorded in whole-cell mode using a Multiclamp 700B amplifier (Molecular Devices)31. Data were digitized at 10 kHz with a 2 kHz low-pass filter. The whole-cell pipette solution for synaptic current recordings contained (in mM): CaCl₂ 135, HEPES 10, EGTA 1, Mg-ATP 4, Na₂GTP 0.4, and KCl 340–314, pH 7.4. The bath solution contained (in mM): NaCl 140, KCl 5, CaCl₂ 1, MgCl₂ 2, HEPES 10, and glucose 10, pH 7.4. IPSCs were pharmacologically isolated by addition of 50 μM m-AP5 and 20 μM CNXQ to the bath solution. EPSCs were pharmacologically isolated by addition of 30 μM picrotoxin. Data were analysed using Clampfit 10.02 (Axon Instruments). Action potentials were recorded with the current-clamp whole-cell configuration. The pipette solution for current-clamp experiments contained (in mM): 123 K-glucurate, 10 KCl, 1 MgCl₂, 10 HEPES, 1 EGTA, 0.1 CaCl₂, 1 K₂ATP, 0.2 Na₂GTP, and 4 glucose, pH adjusted to 7.2 with KOH. Membrane potentials were kept around ~60 to ~70 mV, and step currents were injected to elicit action potential. Whole-cell currents including sodium currents and potassium currents were recorded at a holding potential of ~70 mV; voltage steps ranging from ~80 mV to ~90 mV were delivered at 10-mV increments.

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