Generation of highly enriched V2a interneurons from mouse embryonic stem cells

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Challenges in parsing specific contributions to spinal microcircuit architecture have limited our ability to model and manipulate those networks for improved functional regeneration after injury or disease. While spinal interneurons (INs) have been implicated in driving coordinated locomotor behaviors, they constitute only a small percentage of the spinal cord and are difficult to isolate from primary tissue. In this study, we employed a genetic strategy to obtain large quantities of highly enriched mouse embryonic stem cell (ESC)-derived V2a INs, an excitatory glutamatergic IN population that is defined by expression of the homeodomain protein Chx10 during development. Puromycin N-acetyltransferase expression was driven by the native gene regulatory elements of Chx10 in the transgenic ESC line, resulting in positive selection of V2a INs after induction and treatment with puromycin. Directly after selection, approximately 80% of cells are Chx10+, with 94% Lhx3+; after several weeks, cultures remain free of proliferative cell types and mature into normal glutamatergic neurons as assessed by molecular markers and electrophysiological methods. Functional synapses were observed between selected ESC-derived V2a INs and motor neurons when co-cultured, demonstrating the potential of these cells to form neural networks. While ESC-derived neurons obtained in vitro are not identical to those that develop in the spinal cord, the transgenic ESCs here provide a unique tool to begin studying V2a INs in isolation or for use in vitro models of spinal microcircuits.

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1. Introduction

Neural networks in the form of central pattern generators (CPGs) are capable of generating rhythmic motor outputs that are essential to a range of sophisticated locomotor behaviors, but the microcircuit architecture involved has been much harder to characterize due to the diversity of cell types and lack of spatial organization in the spinal cord (Arber, 2012; Rybak et al., 2015). This is not only detrimental to our understanding of CPG circuitry but also hinders manipulations of those networks for improved functional outcomes after disease or trauma (Courtine et al., 2008; Harkema, 2008). Recent work using genetic ablation and ex vivo electrophysiological characterization of isolated spinal cord preparations has helped identify unique transcriptional markers to define the spinal interneuron (INs) populations that comprise these local spinal circuits (Arber, 2012; Azim et al., 2014; Crone et al., 2008; Gosgnach et al., 2006; Jessell, 2000; Kiehn, 2006; Lanuza et al., 2004; Zhang et al., 2008). However, the dependence on animal models precludes high-throughput pharmacological testing or in vitro modeling of spinal circuitry which may aid in the development of targeted therapeutics that promote neural regeneration and plasticity. Here we describe a method to generate large quantities of highly enriched INs from embryonic stem cells (ESCs), focusing on the acquisition of V2a INs. V2a INs are defined by expression of the homeodomain protein Chx10 and are involved in CPG and propriospinal networks in the spinal cord and respiratory centers of the hindbrain (Al-Mosawie et al., 2007; Azim et al., 2014; Crone et al., 2008, 2012; Dougherty and Kiehn, 2010a, 2010b; Lundfald et al., 2007; Peng et al., 2007). They are an ipsilaterally projecting glutamatergic premotor population with conserved locomotor functions in zebrafish and mice (Crone et al., 2008; Dougherty and Kiehn, 2010b; Kimura et al., 2006). V2a INs are distributed homogeneously along the rostrocaudal axis of the spinal cord in early mouse embryos but are localized to the ventral horn in the adult (Dougherty and Kiehn, 2010a; Francis et al., 2013). Genetic ablation studies have demonstrated their role in coordinating left–right alternation and
skilled-reaching, as well as modulation of locomotor variability and rhythmic breathing (Azim et al., 2014; Crone et al., 2008, 2009, 2012; Dougherty and Kiehn, 2010a; Zhong et al., 2010, 2011).

Robust "highly enriched" neuronal cultures are desirable because they can provide mechanistic insights otherwise confounded by mixed culture conditions. V2a INs, among others, are difficult to isolate from primary tissue in part because they make up a relatively small fraction of the total cells in the spinal cord (Crone et al., 2008). Self-renewing pluripotent cells, such as ESCs, are an attractive alternative to sorting primary tissue because they can be differentiated into a variety of cell types in large quantities for in vitro study or transplantation. By adapting established motor neuron (MN) differentiation protocols (Wichterle et al., 2002), we have previously shown that directed differentiation of ESCs into V2a INs is possible by exposing embryoid bodies (EBs) to retinoic acid (RA); a weak sonic hedgehog (Shh) agonist, purmorphamide; and a Notch-inhibitor, DAPT (Brown et al., 2014). However, despite our ability to derive V2a INs from ESCs, post-mitotic Chx10− cells constitute only ~15% of the total cell population post-induction, which is further diluted as glial cells proliferate with time (Brown et al., 2014). Methods including fluorescence-activated cell sorting (FACS) and magnetic-activated cell sorting have been used to isolate single cell populations, but they are limited by the availability of antibodies to lineage-specific surface antigens, which have not been identified for many ventral IN populations, and require dissociation processes that can be harmful to mature neurons. While FACS can be used with transgenic reporter or lineage tracing cells, it can significantly compromise the viability of mature neurons and retains the potential for contamination.

Transgenic selection of desired ESC-derived populations has proven to be an effective method to generate isolated populations of a variety of cell types, including progenitor motor neurons (pMNs) and MNs (Anderson et al., 2007; Li et al., 1998; Marchetti et al., 2002; McCreedy et al., 2012, 2014a; Soria et al., 2000). Using lineage-specific promoters to drive antibiotic resistance, differentiation of the transgenic ESC line and subsequent antibiotic treatment results in highly enriched cultures that persist through maturation in vitro and in vivo after transplantation (McCreedy et al., 2012, 2014a, 2014b). In this study, we generated a selectable "Chx10-Puro" ESC line and investigated whether it could be used to obtain V2a INs that were comparable to endogenous V2a IN populations.

2. Methods

2.1. ESC culture

Transgenic and RW4 mouse ESCs were cultured on gelatin-coated T-25 flasks in complete media, consisting of Dulbecco's Modified Eagle Medium (DMEM; Life Technologies #11965–092, Carlsbad, CA) containing 10% newborn calf serum (Life Technologies #16,010–159), 10% fetal bovine serum (Life Technologies #26140–079), and 1 × Embryonic Nucleosides (Millipore #ES-008-D, San Francisco, CA). ESCs were passaged every two days at a 1:5 ratio in fresh complete media containing 1000 U/mL leukemia inhibitory factor (LIF; Millipore #ESG1106) and 100 μM β-mercaptoethanol (BME; Life Technologies #21985–023).

2.2. Chx10-Puro selection vector

The targeting vector was constructed from a Gateway-compatible plasmid (pStart-K; Addgene #20346, Cambridge, MA) using a 750 bp SalI-Ascl fragment containing 5′ untranslated sequences and a 750 bp Ascl-Not1 fragment containing 3′ genomic sequences of Chx10 exon 1. A PAC/pGKneo dual resistance cassette was inserted between the two arms as previously described (McCreedy et al., 2012). The dual resistance cassette contains from 5′ to 3′: Asc1 site, Kozak sequence, puromycin cassette with bgh polyA signal (PKO-Select Puro; Agilent Genomics, Santa Clara, CA), floxed phosphoglycerate kinase I promoter driving the neomycin phosphotransferase gene (PGK-neo) with bgh polyA signal, and Ascl site (Kozak, 1986; Thomas and Capecci, 1987; Wu et al., 2008). Gateway recombination with the LR Clonase II Kit (Life Technologies #11791) was used to transfer aTTL-flanked regions into the pWS-TK3 vector, which contains the thymidine kinase gene for negative selection (Supplementary Fig. 1A) (Wu et al., 2008).

2.3. Electroporation and clonal analysis

A CRISPR/Cas9 system was used to insert our resistance cassette into the Chx10 locus with high efficiency. The vector containing the Chx10 guide RNA (gChx10) was generated by the Genome Engineering Core at Washington University in St. Louis. The Chx10 gRNA sequences target the ATG start site of Chx10 exon 1 and were placed into a derivative of Addgene plasmid #43,860; no common SNPs were found and off target profiles were excellent with at least 3 bp of mismatch between the target and any other site in the genome, which dramatically reduces the probability of an off target cut (Veres et al., 2014). The p3s-Cas9HC vector contains the Cas9 open reading frame (Addgene plasmid #43945) (Cho et al., 2013), 1 × 104 RW4 ESCs were resuspended in electroporation buffer (20 mM HEPES pH 7.5, 137 mM NaCl, 5 mM KCl, 0.7 mM Na2HPO4, and 6 mM dextrose) with 8 μg Chx10-Puro selection vector, 1 μg Cas9 vector, and 1 μg gChx10 vector. Cells were electroporated at 0.23 kV and 960 μF in a 0.4 cm cuvette (Bio-Rad #165–2081, Hercules, CA) then plated on a 100 mm gelatin-coated petri dishes in complete media with LIF and BME for the cells to recover overnight. For the remainder of the expansion, cells were exposed to media containing LIF, BME, 150 nM flurididine (FIU, Moravek #M251, Brea, CA) and 40 μg/ml geneticin (G418, Life Technologies #10131) which was replaced every 2 days. After 10 days, single colonies were picked and plated into individual wells of a gelatin-coated 96-well plate. Clones were screened for correct insertion by junction PCR (Chx10Forward: GCCA, Chx10Reverse: GCCCAGAGGCCTTCCACTCTGTT-GCT) and expanded accordingly. The amplification region spanned from the endogenous 5′ UTR (outside of the Chx10-Puro selection vector homology arm) into the PAC cassette to confirm targeted insertion (Supplemental Fig. 1B). Fifty-six transgenic clones were screened for puromycin resistance in Chx1010 interneurons following V2a induction (see V2a induction and V2a selection).

Transgenic clones with puromycin-resistant Chx10− INs were analyzed for PAC copy number by quantitative real-time polymerase chain reaction (qRT-PCR) using a customized TaqMan Copy Number Assay; mouse Tert was used as the endogenous control and the assay was prepared according to the manufacturer's instructions. Calculations were completed using CopyCaller Software (v2.0, Applied Biosystems). The calculated copy number of each clone was normalized to the calculated copy number for the Hb9-Puro ESC line, which was previously determined to contain a single PAC insertion (McCreedy et al., 2014a).

2.4. Cre excision

The PGK promoter and neomycin phosphotransferase genes were removed from an individual Chx10-Puro clone using the pTurboCre plasmid (gift from Timothy Ley). The pTurboCre plasmid (10 μg) and an mRFP-expressing plasmid (1 μg) were prepared for transfection using Lipofectamine 3000 (Life Technologies #L3000-001) according to the manufacturer's instructions; cells were transfected for 24 h. ESC cultures were then dissociated and plated sparsely (3 × 104 cells) in a 100 mm gelatin-coated petri dish with complete media, LIF, and BME. Single colonies were picked after 10 days and plated in 96-well gelatin coated plates. When confluent, the ESCs were split and tested for sensitivity to neomycin by exposure to 40 μg/ml geneticin for 5 days; those not sensitive were discarded. A single transgenic clone was used as the Chx10-Puro ESC line used for the remaining studies.
2.5. V2a induction

V2a INs were obtained from ESCs using a “2−/4+” induction protocol. EBs were formed by transferring 1 x 10⁶ ESCs into 10 mL of DFK5 media on an agar-coated 100 mm petri dish for two days (2−). DFK5 is a DMEM/F12 base media containing 5% Knockout Serum Replacement (Life Technologies #10828–028), 50 μM Nonessential Amino Acids (Life Technologies #11140–050), 100 μM BME, 1:100 100 × Insulin–Transfer- rin–Selenium (Life Technologies #41400–045), 100 μM BME, 1:200 100 × EmbryoMax Nucleosides. EB media was then replaced with 10 mL DFK5 containing 10 nM retinoic acid (RA; Sigma #R2625) and 1 μM purmorphamine (EMD Millipore #540223) for another two days (2−). For the final 2 days (4+), the media was replaced with DFK5 containing 10 nM RA, 1 μM purmorphamine and 5 μM N-[N-(3,5-difluorophenacetyl-l-alanyl)]-(S)-phenylglycine-t-butyl-ester (DAPT; Sigma #D95942). Induced V2a control cultures, the Shh antagonist cyclopamine (Cyc, 1 mM; Sigma #C4116) was used instead of purmorphamine.

2.6. V2a selection and culture

To determine cell viability after selection, transgenic ESCs induced using the 2−/4+ protocols were dissociated with 0.25% Trypsin–EDTA (Life Technologies #25200–056) for 10 min with agitation and quenched with complete media containing 0.001% DNase (Sigma #DN25). Dissociated cells were counted and centrifuged at 1200 rpm for 5 min then re-suspended in a selection media of DFK5NB containing B-27 supplement (Life Technologies #35050), 100 μM EDTA. Cells were induced using the 2−/4+ protocol were rinsed twice with Tyrode’s solution (150 mM NaCl, 4 mM KCl, 2 mM CaCl₂, 1 mM MgCl₂, 10 mM glucose, and 10 mM HEPES, at pH 7.4). Whole-cell electrodes were filled with internal solution (140 mM K-glucuronate, 10 mM NaCl, 5 mM MgCl₂, 0.2 mM EGTA, and 10 mM HEPES, at pH 7.4). Whole-cell electrodes were filled with internal solution (140 mM K-glucuronate, 10 mM NaCl, 5 mM MgCl₂, 0.2 mM EGTA, and 10 mM HEPES, at pH 7.4).

2.7. Post-selection cell viability

To determine the cell viability after 24 h of selection with puromycin, cells induced using the 2−/4+ protocol were rinsed twice with DFK4 media to remove debris then incubated for 30 min in fresh DFK5 media containing calcein-AM (Life Technologies # C481). Fluorescent images were captured using a MICROfire camera attached to an Olympus IX70 inverted microscope. Quantification was completed by flow cytometry (n = 4 biological replicates, see Flow cytometry).

2.8. qRT-PCR

qRT-PCR was performed and analyzed as previously described (Brown et al., 2014). Briefly, cDNA was synthesized using a high capacity RNA-to-cDNA Kit (Invitrogen). The cDNA was combined with TaqMan Gene Expression Assays (Applied Biosystems, Carlsbad, CA) and TaqMan Fast Advanced Master Mix (Applied Biosystems). qRT-PCR was performed using a Step One Plus Applied Biosystems thermocycler with the following protocol: 95 °C for 20 s; 40 cycles of 95 °C for 1 s and 60 °C for 20 s. There were n = 3 technical replicates completed per run and n = 4 biological replicates for each condition. For PAC expression, samples were treated with DNase (Qiagen #7924, Valencia, CA) prior to analysis.

2.9. Immunocytochemistry

Immunocytochemistry staining was performed as previously described (McCreedy et al., 2014a). Antibodies used include mouse anti-Chx10 (1:1000, Santa Cruz sc-374151, Santa Cruz, CA), mouse anti-Lhx3 (1:1000, Lmi3, Developmental Studies Hybridoma Bank (DSHB) #67.4E12, Iowa City IA), mouse anti-Hb9 (1:20, DSHB #81.5C10), mouse anti-Lim1/2 (1:50, DSHB #4F2), mouse anti-Evx1 (1:50, DSHB #99.1–324), rabbit anti-beta-tubulin III (1:1000, βtubill, Covance #181100, Princeton, NJ), rabbit anti-Ki67 (1:250, Abcam # ab15580), guinea pig anti-Vglut2 (1:2500, EMD Millipore #AB2251), mouse anti-MAP2 (1:250, EMD Millipore #AB5622), rabbit anti-mouse-SV2 (1:100, DSHB #SV2), and secondary Alexa Fluor conjugated goat antibodies (1:200, Life Technologies). Nuclei were counterstained with Hoechst (1:1000, Life Technologies #H3569). For cell counting experiments, live cells were identified by phase contrast and Hoechst; only cells with apparent processes were included. Cell counts were performed using Imagej (NIH); there were n = 4 biological replicates for each condition with >450 cells counted per replicate.

2.10. Flow cytometry

A modified 2−/4+ protocol was used to obtain data on early expressing transcription factors in selected ESC-derived neuronal cultures. Briefly, EBs were dissociated at 2−/3− and 5 x 10⁶ cells were plated in individual wells of a laminin-coated 24-well plate in DFK5 media containing 10 nM RA, 1 μM purmorphamine, 5 μM DAPT, and 2 or 4 μg/mL puromycin for 24 h. Surviving cells were dissociated with 0.25% Trypsin–EDTA for 5 min and quenched with complete media. Staining was conducted using the Transcription Buffer Set (Becton Dickinson #562.725, Franklin Lakes, NJ) according to the manufacturer's protocols with antibodies as described in “Immunocytochemistry.” Data was collected using a BD Canto II Flow Cytometer (Becton Dickinson). Between 10,000 and 100,000 events were recorded per condition (n ≥ 4 biological replicates per condition) and analyzed using FlowJo software (Flowjo, Ashland, OR); debris was removed from analysis using forward scatter versus side scatter and Hoechst versus forward scatter plots. Gating parameters were set using control groups stained only with secondary antibodies.

2.11. Single-cell electrophysiology

Cells were induced using the 2−/4+ protocol, dissociated onto laminin coated 35 mm dishes at 2 x 10⁵ and selected for 24 h with puromycin. Cultures were grown for up to 10 days post-selection in DFK5NB media with growth factors prior to recording. Cultures were perfused with Tyrode’s solution (150 mM NaCl, 4 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 10 mM glucose, and 10 mM HEPES, at pH 7.4). Whole-cell electrodes were filled with internal solution (140 mM K-glucuronate, 10 mM NaCl, 5 mM MgCl₂, 0.2 mM EGTA, and 10 mM HEPES, at pH 7.4, supplemented with 5 mM Na-ATP and 1 mM Na-GTP). The open tip resistance was 2–5 MΩm. External solutions were delivered by local perfusion from an 8-barrelled pipette. Voltage-gated channel antagonists tetrodotoxin (TTX, 500 nM), tetraethylammonium chloride (TEA, 30 mM) and 4-aminopyridine (4-AP, 5 mM) were dissolved in Tyrode’s solution. Ligand-gated channel agonists were dissolved at 100 μM in 160 mM NaCl, 2 mM CaCl₂, 10 mM HEPES. Currents were recorded under voltage clamp with an Axopatch 200 A amplifier, filtered at 1 kHz and digitized at 10 kHz using pClamp software (Molecular Devices, Sunnyvale, CA). Evoked synaptic currents were detected by simultaneous whole-cell recordings from adjacent pairs of cells in Chx10-Puro/Olig2-Puro mixed cultures. Brief steps to 0 mV from a holding potential of −80 mV elicited fast inward sodium currents that escaped voltage control in the axon of presynaptic cells and evoked excitatory postsynaptic currents in the adjacent cell, which was held at −80 mV.
2.12. V2a and pMN co-culture

Mixed Chx10-Puro and Olig2-Puro cultures were established to assess synapse formation between V2a INs and MNs. A constitutively active fluorescent reporter was knocked into the Rosa26 domain of the Chx10-Puro line as described above using the Ai9 plasmid (gifted by Hongkui Zeng, Addgene plasmid #22799) (Madisen et al., 2010), pTurboCre, the CaSk9 plasmid, and Rosa26 guide RNAs (gR26) generated by the Genome Engineering Core at Washington University in St. Louis. Clones were screened for correct insertion by junction PCR (R26Forward: TCCCAAATGCCTGCTGACGT; CAGReverse: CCATCGCTGACAAAA TAAT) and expanded accordingly. The previously established Olig2-Puro ESC line contains the PAC gene under the expression of Olig2, which marks the pMN domain (McCreedy et al., 2012). The Olig2-Puro and Chx10-Puro ESCs were induced according to their respective 2−/4+ protocols (McCreedy et al., 2012), dissociated onto laminin (Life Technologies #23017–015) coated 35 mm dishes at 2 × 10⁶ at a 1:1 ratio and selected for 24 h in DFK5NB with 4 μg/mL puromycin. The media was replaced with DFK5NB containing B27, GlutaMAX, NT-3, GDNF, and BDNF and was replenished as needed every two days for up to 7 days. Neurobasal media containing the supplements and growth factors was used for the remaining duration of culture up to 4 weeks.

2.13. Statistical analysis

Statistical analyses were performed using Statistica software (v5.5, StatSoft, Tulsa, OK). Significance was determined using Scheffe’s post hoc test for analysis of variance (ANOVA) with confidence as indicated. Average values reported with error bars are the standard deviation or standard error of the mean (SEM) as indicated.

3. Results

3.1. Puromycin resistance coincides with Chx10 expression in transgenic ESC line

In order to isolate V2a INs from a heterogeneous population of differentiated ESCs, we developed a genetic strategy to positively select for cells expressing the defining transcription factor Chx10 (Fig. 1).

CRISPR-assisted homologous recombination was used to target the first exon of the mouse Chx10 locus for puromycin N-acetyltransferase (PAC) insertion, thereby placing antibiotic resistance under the control of the native Chx10 gene regulatory elements. A single clone with targeted genomic insertion (Supplementary Fig. 1B) and one copy of the PAC gene (Supplementary Fig. 1C) was treated with Cre recombinase to remove the neomycin selection cassette. The resulting Chx10-Puro ESC line was used for all subsequent analyses in this study.

To determine whether Chx10 expression resulted in the expression of PAC in the Chx10-Puro line, Chx10-Puro ESCs were differentiated using a 2−/4+ V2a induction protocol—cells were aggregated for two days to form EBs, then exposed to RA, purmorphamine, and DAPT for four days. Unmodified RW4 ESCs and cultures induced using the Shh antagonist Cyc were used as controls. qRT-PCR demonstrated that expression is activated with Chx10 expression in the Chx10-Puro line. When Shh signaling was suppressed with Cyc. This suggests that PAC expression is activated with Chx10 expression in the Chx10-Puro line.

To confirm cell viability post-selection, cultures were dissociated following the induction and plated on laminin-coated wells in DFK5NB media containing puromycin for 24 h (Fig. 2B). In the absence of puromycin, cells were viable in all conditions as evidenced by calcein AM staining (Fig. 2C). Following puromycin selection, viable cells were only apparent in induced Chx10-Puro cultures (Fig. 2C,H). The addition of puromycin killed all RW4 cultures as well as Chx10-Puro cultures induced with Cyc. Analysis by flow cytometry indicated 12.0 ± 3.6% and 11.0 ± 3.6% of cells were viable after selection with 2 μg/mL and 4 μg/mL puromycin, respectively, compared to 85.81 ± 3.79% viable cells in unselected cultures (p < 0.0001) (Fig. 2D). Surviving cells demonstrated neuronal morphology, including phase bright cell bodies and neurite extensions that are consistent with V2a induction (Fig. 2E).

3.2. V2a IN markers are enriched in selected Chx10-Puro cells

Immunocytochemistry and flow cytometry were used to confirm V2a IN identity in selected cultures using antibodies for the pan-neuronal marker (beta tubulin III (β-tubIII)) and defining transcription

Fig. 1. Schematic showing PAC insertion into the Chx10 gene locus. (1) gChx10 guide RNAs mediate double stranded break at the ATG start codon. (2) Homologous recombination incorporated targeting vector cassette into the first exon. PAC expression was driven by the native Chx10 upstream promoters. The ubiquitously expressed PGK promoter drove neomycin phosphotransferase to screen cells with the incorporated dual cassette. (3) Cre recombinase was used to remove the PGK-neo cassette. (4) The final Chx10-Puro construct expresses PAC when Chx10 is expressed.
Factors for V2a INs (Chx10, Lhx3), motor neurons (Hb9, Lhx3), V1 INs (Lim1/2), and V0 INs (Evx1, Lim1/2). The p2 progenitor marker Lhx3 is expressed in both V2 INs and immature MNS (Sharma et al., 1998), but expression of Lhx3 in post-mitotic cells has been used as a marker for V2a INs as it drives expression of Chx10 in certain contexts (Crone et al., 2008; Tanabe et al., 1998; Thaler et al., 1999). To capture early, transient expression of these transcription factors, a shorter, modified 2−/3+ induction protocol was used to ensure high protein expression post-selection for detection and quantification (Fig. 3A).

Neuronal enrichment was quantified by counting the number of cell nuclei that co-localize with β-tubIII. Approximately 86.81 ± 5.40% of cells were β-tubIII+ in unselected control cultures; the percentage increased significantly to 98.27 ± 0.25% or 100% (p < 0.05) following selection with 2 μg/mL or 4 μg/mL puromycin respectively. (Fig. 3B). Analysis of the population distribution by flow cytometry confirmed that selection constitutes an enrichment of V2a cell identity (Fig. 3C, Supplementary Fig. 2). While unselected cultures contained a significant majority of cells were Chx10+ (Fig. 3D) and/or Lhx3+ (Fig. 3E) after treatment with puromycin (p = 0.0001). The combination of staining and quantification suggest that there are negligible quantities of V0, V1 (p < 0.0001) or MNS (p < 0.05) that survive selection (Fig. 3C, Supplementary Fig. 3).

While the intent of this study was to generate spinal V2a INs using Chx10 as the definitive transcription factor for identification, Chx10 is also expressed in the developing retina (Dhomen et al., 2006; Rowan and Cepko, 2004). The expression of two retinal developmental markers, Rax and Six3 (Loosi et al., 1999; Muranishi et al., 2012), was assessed and they were not significantly upregulated in selected cultures compared to ESC controls, but Chx10 expression was significantly upregulated, as expected (Supplementary Fig. 4A−B). Cultures were stained for Isl1, which is a developmental marker for MNS in the spinal cord, but in combination with Chx10 and Lhx3 is also a marker for retinal bipolar cells (Elshatory et al., 2007). In selected cultures, 3.09 ± 0.93% and 0.47 ± 0.55% of cells were Isl1+ in cultures treated with 2 μg/mL and 4 μg/mL puromycin respectively, which were significantly lower (p < 0.05) compared to 8.10 ± 1.78% of cells in unselected induced cultures (Supplementary Fig. 4C–D). Interestingly, using a higher concentration of puromycin resulted in a lower percentage of cells expressing V2a markers (Fig. 3C), but also significantly reduces the number of cells that express Isl1 compared to using 2 μg/mL puromycin (p < 0.05). These more highly enriched (4 μg/mL puromycin) cultures were used for all maturation and electrophysiology experiments.

3.3. Selected Chx10-Puro neurons are post-mitotic and achieve functional maturity

Because neuronal differentiation of ESCs can give rise to proliferative cell types that reduce the purity of the culture, selected Chx10-Puro cultures were evaluated for mitotic activity. An antibody for the proliferation marker Ki67 was used to detect and quantify mitotic cells via immunocytochemistry and flow cytometry (Fig. 4A, B Supplementary Fig. 2). Ki67+ cells comprised 15.13 ± 2.58% of cells observed in unselected cultures, which was significantly decreased when treated with 2 μg/mL (1.80 ± 0.10%) or 4 μg/mL (0.83 ± 0.29%) puromycin (p < 0.0001). Unselected cultures also visibly showed a rapid increase in the number of cell nuclei over 2 weeks (Fig. 4D).

Most V2a INs in vivo express excitatory vesicular glutamate transport-2 (Vglut2) (Lundfeld et al., 2007). To confirm functional maturation of selected Chx10-Puro cultures, cells were induced using the 2−/4+ protocol, dissociated onto laminin-coated plates and selected for 24 h with puromycin, then grown for 14 days post-selection in DFK5NB media with growth factors (Fig. 4C). Selected cells abundantly express Vglut2 and the mature dendritic marker, microtubule associated protein 2 (MAP2) (Fig. 4D). Vglut2+ neurons appear to be the majority of cells in selected cultures compared to only a small fraction in unselected cultures. Taken together, these data suggest that we have generated highly enriched, post-mitotic V2a INs.

3.4. Selected V2a INs demonstrate mature electrophysiological profiles

Whole cell current and voltage clamp recordings were performed to evaluate the functional maturity of cells in puromycin-selected cultures up to 12 days post-selection (d12). Cell capacitance, which is proportional to surface area, was low on d2 (17.1 ± 1.95 pF, n = 5) but increased to approximately 30 pF by d3 and remained relatively constant through d12 (Table 2). Input resistance was highest on d2 (2.22 ± 0.66 GΩ, n = 5) and declined as cells matured (415 ± 80 MΩ, d12, n = 7). Cell resting potentials were relatively depolarized at early time points after selection, but by d9−d12 the mean Vrest was approximately −50 mV. When membrane potential was maintained near −60 mV by steady current injection under current clamp nearly all
cells with neuronal morphology were capable of firing action potentials upon stimulation with a depolarizing current pulse (Fig. 5A). Of these, 35.7% fired single action potentials, 35.7% fired several action potentials near the beginning of an 800 msec pulse but then adapted, while the remaining 28.6% exhibited tonic firing throughout the depolarizing pulse. Under whole-cell voltage clamp, depolarizing steps from $-80$ mV elicited fast inward currents that were sensitive to blocking by the voltage-gated sodium channel antagonist tetrodotoxin (TTX, 0.5 μM). Traces in Fig. 5B show TTX-sensitive current obtained by subtracting current recorded in TTX at each test potential from current recorded in control solution. In addition, all cells displayed transient and sustained outward currents mediated by voltage-gated potassium channels (Fig. 5C, D). Consistent with previous work in other cell types (Bean, 2007), the organic cation tetraethylammonium (TEA) blocked sustained currents, whereas transient currents were selectively inactivated at a holding potential ($V_h$) of $-40$ mV (Fig. 5C) or by application of 4-aminopyridine.

**Table 1**

Quantification of IN subtypes by flow cytometry. Flow cytometry analysis of ventral IN subpopulations in control and puromycin-selected cultures. Values presented as mean ± standard deviation. * denotes $p < 0.0001$ compared to control group. # denotes $p < 0.05$ compared to control group.

<table>
<thead>
<tr>
<th></th>
<th>Chx10</th>
<th>Lhx3</th>
<th>Hb9</th>
<th>Evx1</th>
<th>Lim1/2</th>
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<tbody>
<tr>
<td>Control</td>
<td>27.50 ± 9.21</td>
<td>53.93 ± 1.30</td>
<td>8.60 ± 4.52</td>
<td>23.80 ± 4.17</td>
<td>15.16 ± 1.53</td>
</tr>
<tr>
<td>2 μg/ml</td>
<td>80.37 ± 3.27*</td>
<td>93.96 ± 0.62*</td>
<td>1.17 ± 1.06*</td>
<td>0.61 ± 0.66*</td>
<td>3.21 ± 0.40*</td>
</tr>
<tr>
<td>4 μg/ml</td>
<td>70.25 ± 12.90*</td>
<td>86.97 ± 5.83*</td>
<td>1.97 ± 1.42*</td>
<td>2.26 ± 1.16*</td>
<td>0.76 ± 0.64*</td>
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</table>
The transient outward currents required less depolarization for activation as shown in the plots of peak and steady-state (SS) outward currents as a function of test potential (Fig. 5D).

To test for functional neurotransmitter receptors, cells were exposed to 100 μM selective agonists for AMPA/kainite, NMDA, glycine and GABA. Agonist activated inward currents were observed while holding cells at a fixed potential of −80 mV (Fig. 6A). As the cells matured from d3 to d20 or more, the amplitude of these agonist-gated currents increased (Fig. 6B). Agonist-evoked currents were also recorded as the membrane potential was slowly ramped between −110 and +110 mV (Fig. 6C). Currents elicited by the inhibitory transmitters GABA and glycine reversed polarity near the estimated equilibrium potential for chloride ions (−54 mV) given the composition of our internal and external solutions. In contrast, the currents evoked by NMDA and kainate reversed much closer to 0 mV, consistent with monovalent cation permeability.

Overall, the membrane properties of our selected Chx10-Puro cells are broadly consistent with previous electrophysiological characterization of V2a interneurons in acute tissue slice and isolated spinal cord preparations (Dougherty and Kiehn, 2010a; Zhong et al., 2010). Recordings from native neurons in these preparations have revealed a variety of action potential patterns including tonic, adapting and single spike firing, as are also seen in our cultures.

### 3.5. Synapses observed in mixed V2a and pMN culture

The ability of ESC-derived neuronal populations to form synapses is critical for their use in cell therapy or for in vitro modeling of neural

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**Table 2**

Electrophysiological properties of selected Chx10-Puro cells. Values presented as mean ± SEM (number of cells) (* denotes significantly different from d2–4 by t-test, Mann–Whitney rank sum test, or Z-test). Cell capacitance and input resistance were determined from 10 mV voltage clamp steps from a holding potential of −80 mV. First spike latency, amplitude, absolute amplitude, threshold and half-width were determined for the first spike recorded at threshold depolarization. In addition, 800 msec depolarizations that elicited spiking with an average frequency of 10 Hz were used to measure 1st latency, instantaneous frequency from the first inter-spike interval (ISI), and frequency adaptation (ratio of last to 1st ISI). Sag in voltage responses was determined for 800 msec hyperpolarizing current injections from −60 mV.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>d2–4</th>
<th>d9–12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Capacitance (pF)</td>
<td>33.1 ± 3.2 (19)</td>
<td>29.8 ± 2.1 (38)</td>
</tr>
<tr>
<td>Input resistance (GOhm)</td>
<td>1.7 ± 0.24 (19)</td>
<td>0.57 ± 0.12 (38)*</td>
</tr>
<tr>
<td>Tau (msec)</td>
<td>52.1 ± 7.1 (19)</td>
<td>22.8 ± 9.1 (38)*</td>
</tr>
<tr>
<td>V rest (mV)</td>
<td>−37.6 ± 1.5 (18)</td>
<td>−51.2 ± 1.8 (28)*</td>
</tr>
<tr>
<td>% V rest &lt; −50 mV</td>
<td>5.60%</td>
<td>53.5%</td>
</tr>
<tr>
<td>Rheobase (pA)</td>
<td>13.1 ± 6.3 (7)</td>
<td>49.7 ± 12.0 (21)*</td>
</tr>
<tr>
<td>1st spike latency (msec)</td>
<td>91.1 ± 22.4 (7)</td>
<td>76.1 ± 15.0 (21)</td>
</tr>
<tr>
<td>1st spike amplitude (mV)</td>
<td>70.4 ± 7.1 (7)</td>
<td>92.5 ± 4.1 (21)*</td>
</tr>
<tr>
<td>1st spike overshoot (mV)</td>
<td>11.9 ± 6.8 (7)</td>
<td>31.4 ± 4.1 (21)*</td>
</tr>
<tr>
<td>1st spike threshold (mV)</td>
<td>−38.9 ± 1.9 (7)</td>
<td>−37.4 ± 1.4 (21)</td>
</tr>
<tr>
<td>1st spike width (msec)</td>
<td>6.5 ± 0.9 (7)</td>
<td>2.7 ± 0.3 (21)*</td>
</tr>
<tr>
<td>10 Hz 1st latency (msec)</td>
<td>41.9 ± 5.9 (4)</td>
<td>31.4 ± 3.1 (14)</td>
</tr>
<tr>
<td>10 Hz 1st frequency (Hz)</td>
<td>11.3 ± 0.5 (4)</td>
<td>13.9 ± 1.3 (14)</td>
</tr>
<tr>
<td>10 Hz frequency adaptation</td>
<td>1.3 ± 0.03 (4)</td>
<td>1.4 ± 0.14 (14)</td>
</tr>
<tr>
<td>10 Hz after potential (mV)</td>
<td>−2.4 ± 2.7 (4)</td>
<td>−0.15 ± 0.62 (14)</td>
</tr>
<tr>
<td>Sag at −90 mV (mV)</td>
<td>2.4 ± 1.0 (4)</td>
<td>2.4 ± 0.72 (14)</td>
</tr>
<tr>
<td>Peak I Na (nA)</td>
<td>−1.87 ± 0.27 (16)</td>
<td>−2.29 ± 0.22 (29)*</td>
</tr>
<tr>
<td>Vm for peak I Na (mV)</td>
<td>−26.6 ± 2.0 (16)</td>
<td>−32.6 ± 1.6 (29)*</td>
</tr>
</tbody>
</table>

(data not shown). To test for functional neurotransmitter receptors, cells were exposed to 100 μM selective agonists for AMPA/kainite, NMDA, glycine and GABA. Agonist activated inward currents were observed while holding cells at a fixed potential of −80 mV (Fig. 6A). As the cells matured from d3 to d20 or more, the amplitude of these agonist-gated currents increased (Fig. 6B). Agonist-evoked currents were also recorded as the membrane potential was slowly ramped between −110 and +110 mV (Fig. 6C). Currents elicited by the inhibitory transmitters GABA and glycine reversed polarity near the estimated equilibrium potential for chloride ions (−54 mV) given the composition of our internal and external solutions. In contrast, the currents evoked by NMDA and kainate reversed much closer to 0 mV, consistent with monovalent cation permeability.

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### 3.5. Synapses observed in mixed V2a and pMN culture

The ability of ESC-derived neuronal populations to form synapses is critical for their use in cell therapy or for in vitro modeling of neural
circuitry. Endogenous V2a INs primarily synapse onto MNs, but have also been observed to synapse onto other V2a INs. To model these interactions, selected ESC-derived V2a INs and pMNs were co-cultured for up to 4 weeks and examined for functional connectivity (Fig. 7A–D). A previously established Olig2-Puro ESC line was used to obtain a highly enriched pMN culture, which gives rise to a variety of cell types including MNs, oligodendrocytes, and astrocytes. The presence of glia has been shown to be beneficial to spontaneous electrical activity and for improved synapse formation in culture (Boehler et al., 2007; Pfrieger and Barres, 1997). A constitutively-active red fluorescent reporter was knocked into the Chx10-Puro line to identify V2a INs in co-culture conditions (Fig. 7E–G); MNs were identified by their distinct morphology (Fig. 7B). Robust staining for synaptic vesicles (SV2) was observed in both V2a INs and MNs after 3 weeks; 98.67 ± 1.03% of Vglut2+ cells in these cultures co-stained with SV2 (Fig. 7H–J).

To test for functional synaptic connections, we performed simultaneous whole-cell recordings from adjacent pairs of cells on d21 to d31 (Finley et al., 1996). Both cells were held at −80 mV and brief (5–10 msec) voltage steps to 0 mV were delivered at 0.2 Hz, first to one cell and then the other. As shown in Fig. 8, the voltage steps in a labeled Chx10-Puro cell elicited inward sodium currents that escaped from voltage control in the presynaptic axon and evoked excitatory postsynaptic currents (EPSCs) in the adjacent Chx10-Puro cell. In all, we obtained successful simultaneous recordings from 9 cell pairs with 18 potential presynaptic cells, including 13 labeled Chx10-Puro cells and 5 unlabeled cells. Upon stimulation, most of the cells evoked EPSCs (11 of 13 labeled Chx10-Puro cells and 3 of 5 unlabeled cells) with an average amplitude of 520 ± 170 pA and a mean synaptic delay of 2.0 ± 0.2 msec from the peak of presynaptic inward current to the initial rise of the EPSC. The remaining 2 labeled Chx10-Puro cells and 2 unlabeled Olig2-Puro cells failed to evoke any postsynaptic response, suggesting either they did not make functional contact with the adjacent cell or they did not release a transmitter capable of eliciting fast postsynaptic currents. Importantly, the AMPA/kainate receptor antagonist NBQX (30 μM) blocked synaptic transmission mediated by all of the presynaptic Chx10-Puro cells, as well as the 3 unlabeled presynaptic cells, confirming that they exhibit an excitatory glutamatergic phenotype. Superfusion with the GABAA receptor antagonist bicuculline methiodide (200 μM) did not affect evoked synaptic transmission in any of our paired recordings, but did block spontaneous IPSCs that were observed in several of the recorded cells in mature Chx10-Puro/Olig2-Puro mixed cultures. Similarly, NBQX also blocked spontaneous EPSCs observed while recording from individual cells in mixed cultures.

4. Discussion

4.1. Chx10-Puro mouse ESC line yields highly enriched V2a INs

While optimized induction protocols have been developed to obtain several ventral spinal populations from ESCs (Brown et al., 2014; McCreedy et al., 2012, 2014a; Wichterle et al., 2002; Xu and Sakiyama-Elbert, 2015), the heterogeneity of resulting cultures make
the study or transplantation of single cell types unfeasible. By using defining transcription factor promoters to drive puromycin resistance in ESC-derived cell populations, our lab and others have been able to positively select for enriched populations of beta cells, cardiomyocytes, endothelial cells, neurepithelial progenitors, pMNs, and MNs (Anderson et al., 2007; Li et al., 1998; Marchetti et al., 2002; McCreedy et al., 2012, 2014a; Soria et al., 2000). Here we generated a Chx10-Puro ESC line and confirmed that, after induction and puromycin treatment, the vast majority of viable cells were Chx10+ and Lhx3+. By modulating the concentration of puromycin, contamination due to other ventral cell types and Chx10+/Lhx3+/Isl1+ retinal bipolar cells could be reduced. Within two weeks, selected Chx10-Puro cells mature into Vglut2+ glutamatergic neurons; Vglut2 presents as punctate staining along the neurite surface, however strong cytosolic staining was also observed at early time points. This may be caused by an upregulation of protein expression during early stages of maturation, or, in selected cultures, due to a lack of supportive cells that aid in maturation. Within four weeks of culture (Fig. 7E–G), the cytosolic component is reduced. We also report a higher yield of Chx10+ cells in unselected cultures than previously published (Brown et al., 2014), likely because the cells are maintained as EBs for the duration of the induction.

Our data suggests 80–90% purity in selected Chx10-Puro cultures, but it is probable that quantification by flow cytometry is an underestimation of true population counts and the purity of the selected cultures is likely to be higher than we observed due to transcription factor inactivation. While increasing the concentration of puromycin was expected to improve the yield of cells positive for V2a markers, instead we saw a downward trend despite the elimination of potential retinal cells marked by Isl1 expression and other ventral markers. A rapid decrease in the number of Chx10 and Lhx3 expressing cells was also observed with time (data not shown). Given that selected cultures are not overtaken by proliferative cell types and remain primarily glutamatergic neuronal cells when mature, it is conceivable that when selected with 4 μg/mL puromycin, early V2a INs and retinal cells die before PAC expression is sufficient to counter puromycin activity, and that the surviving cells analyzed begin to lose V2a marker expression but represent a more mature V2a population. This is at odds with endogenous cells in animal models, where immunohistochemistry has been used to identify Chx10+ and Lhx3− cells in spinal cord slices into maturity (Al-Mosawie et al., 2007; Crone et al., 2008, 2009; Dougherty and Kiehn, 2010a, 2010b; Lundfald et al., 2007). Lineage tracing using cre-recombinase would enable a clearer delineation of total subtype yields in ESC-derived cultures. Selected Chx10-Puro V2a INs also exhibited morphological and electrophysiological heterogeneity consistent with observations ex vivo, which seems to imply at least some degree of diversity (Al-Mosawie et al., 2007; Dougherty and Kiehn, 2010a, 2010b; Kimura et al., 2006). Several dozen transcription factors have been detected to identify discrete ventral IN subpopulations in the last few years alone (Francius et al., 2013), including at least one functionally distinct subpopulation of Chx10+ cells, a Shox2+/Chx10+ population, termed V2d (Dougherty et al., 2013). The availability of markers to classify V2 IN subpopulations is of major interest and would help to further characterize the purified population.

4.2. Functional activity in purified ESC-derived V2a populations mimics ex vivo data

Most native V2a interneurons in acute spinal cord preparations appear capable of spiking repeatedly throughout the duration of a prolonged depolarizing pulse, although a substantial minority exhibit other firing patterns including burst firing that adapts and single spiking (Dougherty and Kiehn, 2010a; Zhong et al., 2010). Our selected Chx10-Puro cells also exhibit a range of action potential firing, as well as other electrophysiological properties that are generally consistent with work on native V2a cells. Maturing selected Chx10-Puro cells expressed receptors for the major fast excitatory and inhibitory spinal neurotransmitters, suggesting that they should be capable of functional integration into spinal circuits.

V2a INs in vivo play a critical role in CPGs, networking with other cell types in order to achieve coordinated locomotion (Azim et al., 2014; Crone et al., 2008, 2009, 2012; Zhong et al., 2010, 2011). In the present study, we demonstrate the ability of selected Chx10-Puro V2a INs to make functional excitatory synapses onto each other, as well as onto presumptive MNs. In the instances where there was no apparent connection between Olig2-Puro cells and adjacent labeled Chx10-Puro cells, it is possible that the unlabeled Olig2-Puro cells were MNs releasing acetylcholine, which did not acutely activate channels and thus did not produce a fast post synaptic current. The connections observed during paired recordings were likely monosynaptic because of their large
amplitudes and short synaptic delays. In all cases tested, the AMPA/kainate receptor antagonist, NBQX, produced complete block of evoked transmission, demonstrating that the transmitter released by presynaptic Chx10-Puro cells activates postsynaptic glutamate receptors. Thus, our paired recordings provide conclusive evidence that selected Chx10-Puro cells are glutamatergic.

Previous work by Zhong et al. (2010) provided evidence for selective electrical coupling between V2a INs with similar action potential firing characteristics, a feature that might help to coordinate rhythmic activity. We saw no examples of electrical coupling in our paired recordings either from 9 cell pairs in mature Chx10-Puro/Olig2-Puro co-cultures (d20 to d31), or in preliminary recordings from 6 pairs in d10 Chx10-Puro-Olig2-Puro co-cultures.

Fig. 7. Chx10-Puro and Olig2-Puro co-culture. (A) Schematic of 2−/4+ induction protocol for Chx10-Puro and Olig2-Puro cells, followed by selection and up to four weeks of co-culture. Media is switched from DFK5NB to Neurobasal after one week. (B) Phase contrast and (C, D) Hoechst staining of mixed culture demonstrates variety of cell morphologies and cell debris. (E) TdTomato allows for identification of Chx10-Puro cells. (F, G, H) Vglut2 staining appears almost exclusively in V2a INs. (I, J) SV2 staining is prevalent in most neurons and indicative of synaptic activity. Scale bars = 200 μm.

Fig. 8. Evoked excitatory synaptic transmission between selected Chx10-Puro cells. Simultaneous whole-cell recordings of pre- and postsynaptic currents from two adjacent Chx10-Puro cells 31 days after puromycin selection. Voltage steps to 0 mV from a holding potential of −80 mV elicited inward sodium current, and uncompensated capacity transients, in the presynaptic cell. Excitatory synaptic currents (EPSCs) were evoked in the postsynaptic cell with a delay of 3.2 msec, measured from the peak of presynaptic sodium current to the initial rise of the EPSC. Exposure to the AMPA/kainate receptor antagonist NBQX (30 μM) completely blocked the EPSC, while the GABA<sub>A</sub> receptor antagonist bicuculline methiodide (0.2 mM) had no effect. Each panel shows 3 superimposed traces in each condition. Presynaptic stimuli were delivered at 0.2 Hz; postsynaptic holding potential, −80 mV.
Puro cultures; however, we were not able to characterize the spiking properties of these cells to determine whether or not both cells in a pair displayed similar firing patterns.

4.3. Chx10-Puro cells for in vitro modeling

While most in vitro studies that use neuronal cell cultures have investigated functional properties, advances in culture methods, geometric and spatial patterning, instrumentation, and signal processing make possible “lab on a chip” technologies for brain and spinal cord (Wheeler and Brewer, 2010). V2a INs occupy a significant role in the CPG and, compared to other IN types, have well characterized network properties that make them an attractive population with which to design an experimental platform (Dougherty and Kiehn, 2010a). A minimalist approach could be taken, whereby V2a INs are observed in isolation, but transcription-factor driven selectable ESC lines could also be used modularly to generate complex cultures that remain well defined. We demonstrate the feasibility of such an approach here using both the Chx10-Puro and Olig2-Puro ESC lines to create a simple model of V2a-MN interactions. Because ESC induction protocols can be manipulated to enrich for distinct anatomical regions (Lippmann et al., 2015; Okada et al., 2004), there is a significant degree of flexibility possible using selectable lines.

A general caveat in the use of pluripotent stem cells for modeling or therapy is the gap between neurons obtained in the dish and those that develop normally in vivo. As evidenced by the selected Chx10-Puro cultures generated in this study, there remain some phenotypic and functional differences between ESC-derived V2a INs and endogenous cells. Some variation is expected, especially given the abnormal isolation of these neurons during a critical period of maturation, and it is possible that a longer duration of maturation or optimization of culture conditions might improve comparability. However, it is similarly possible that the significant differences in development cannot be overcome and the cells here merely mimic V2a IN properties while incapable of the complex functional activity expected of endogenous V2a INs. Compounding the issue of characterization is the absence of comprehensive genetic profiles to identify functionally distinct IN subtypes. Creating tools to investigate these potential differences is beneficial to our understanding of spinal cord development and for improving stem cell therapies.

5. Conclusions

By knocking PAC into the Chx10 locus of mouse ESCs, the addition of puromycin to differentiated ESC cultures killed Chx10 negative cells. V2a IN enrichment was evident directly after selection of differentiated “Chx10-Puro” cells and persisted through maturation into functional glutamatergic neurons. Electrophysiology demonstrated that selected V2a INs are capable of spontaneously responding to a range of agonists. Finally, our selected V2a INs form synapses that selected Chx10-Puro V2a INs, while not identical to native V2a INs, are broadly comparable and can be a powerful resource for in vitro modeling of neural networks, investigation of neuronal development and diversification, and targeted cell therapies.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.expneurol.2016.01.011.

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