Rapid and efficient generation of oligodendrocytes from human induced pluripotent stem cells using transcription factors

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Edited by Brigid L. M. Hogan, Duke University Medical Center, Durham, NC, and approved February 1, 2017 (received for review August 30, 2016)

Rapid and efficient protocols to generate oligodendrocytes (OL) from human induced pluripotent stem cells (iPSC) are currently lacking, but may be a key technology to understand the biology of myelin diseases and to develop treatments for such disorders. Here, we demonstrate that the induction of three transcription factors (SOX10, OLIG2, NKX6.2) in iPSC-derived neural progenitor cells is sufficient to rapidly generate O4⁺ OL with an efficiency of up to 70% in 28 d and a global gene-expression profile comparable to primary human OL. We further demonstrate that iPSC-derived OL disperse and myelinate the CNS of $Mbp^{sh/shi} Rag^{-/-}$ mice during development and after demyelination, are suitable for in vitro myelination assays, disease modeling, and screening of pharmacological compounds potentially promoting oligodendroglial differentiation. Thus, the strategy presented here to generate OL from iPSC may facilitate the studying of human myelin diseases and the development of high-throughput screening platforms for drug discovery.

human iPSC | oligodendrocytes | forward patterning | myelination | disease modeling

ligodendrocytes (OL) play a key role in myelin-related dis-O asses, including multiple sclerosis (MS), leukodystrophies, as well as periventricular leukomalacia, and there is an increasing awareness of their potential role in neurodegenerative diseases (e.g., multiple system atrophy and amyotrophic lateral sclerosis) or traumatic spinal cord injury (1-6). OL form and maintain the myelin sheaths that insulate axons and organize the distribution of axonal voltage-gated ion channels, a prerequisite for conduction of action potentials and trophic support of axons. Demyelination in MS contributes to axonal damage and disease progression (7). Immunosuppressive or immunomodulatory therapies, including complete ablation of the immune system by radiation and chemotherapy, prevent new inflammatory lesions that underlie clinical relapses but do not arrest disease (8). Therapies promoting remyelination represent a promising new treatment strategy to protect and restore axonal integrity and neurologic function (4). The development of such therapeutics is hampered, at least in part, by the limited availability of human OL. Stem cell technologies are a promising tool to circumvent this problem. Availability of OL derived from human induced pluripotent stem cells (iPSC) would permit studies to delineate mechanisms regulating repair by endogenous myelin lineage cells and provide a source of autologous cells for replacement therapy. Such cells would also provide new opportunities to identify pathological mechanisms underlying de- or dysmyelinating diseases. However, to date only a few protocols have resulted in the successful generation of human stem cell-derived OL. Furthermore, these protocols require long culture periods (70–150 d) to obtain $O4^+$ OL and show limited efficiency (9–12).

Here, we describe an efficient strategy that facilitates and optimizes the generation of human $O4^+$ OL from human iPSCderived neural progenitor cells (NPC) (13). This approach yields up to 70% O4⁺ OL within 28 d of differentiation, using a combination of three transcription factors (TF), namely SOX10, OLIG2, and NKX6.2. Furthermore, 30% of the O4⁺ iPSC-derived OL (iOL) differentiate into mature myelin basic protein-positive (MBP⁺) OL within 7 additional days. The global gene-expression pattern of O4⁺ OL is comparable to that of human primary OL (pOL). The induced human iOL produce myelin-like structures around nanofibers or iPSC-derived neurons. After transplantation into MBP-deficient shiverer mice (*Shi/Shi Rag2^{-/-}*) iOL disperse widely and myelinate host axons in the developing central nervous system (CNS), as well as the adult demyelinated spinal cord.

Significance

Understanding of myelin diseases and development of new treatment options are at least partly hampered by the limited availability of human oligodendrocytes. Induced pluripotent stem cells (iPSC) may be an ideal tool to circumvent this problem; however, rapid and efficient protocols to generate oligodendrocytes from human iPSC are currently lacking. The induction of the transcription factors SOX10, OLIG2, and NKX6.2 in iPSC-derived neural progenitors accelerates oligodendroglial differentiation significantly resulting in up to 70% of O4⁺ oligodendrocytes within 28 d. These oligodendrocytes myelinate the CNS during development and after demyelination, and are suitable for pharmacological screens and disease modeling. The strategy presented herein will markedly facilitate the studying of human myelin diseases and the development of screening platforms for drug discovery.

This article is a PNAS Direct Submission.

Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, www.ncbi.nlm.nih.gov/geo (accession no. GSE79914).

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1614412114/-/DCSupplemental.

PNAS PLUS

Author contributions: M.E. and T.K. designed research; M.E., S.M., and L.S. performed research; S.V., A.-L.H., Q.-L.C., A.S., K.-P.K., C.B., A.M., G.H., R.M.J., J.A., J.S., H.Z., H.R.S., and A.B.-V.E. contributed new reagents/analytic tools; A.B.-V.E. designed animal experiments; M.E., S.M., and M.G. analyzed data; and M.E., A.B.-V.E., and T.K. wrote the paper.

Conflict of interest statement: M.E. and T.K. have a pending patent application for the oligodendroglial differentiation protocol.

Furthermore, iOL can be exploited for disease modeling and to test the potential of pharmacological compounds in promoting oligodendroglial differentiation.

Results

Identification of OL Lineage Inducing TF in Human NPC. Human PSC present a valuable source for the generation of myelinogenic OL for research and autologous cell-replacement therapies (9-12). Currently, available protocols are technically challenging and time-consuming, limiting the application of these cells for research and regenerative medicine. In these protocols, NPC are rapidly and efficiently derived from human PSC, which is followed by the rate-limiting step of oligodendroglial specification and differentiation. Therefore, we first aimed to identify TF accelerating the oligodendroglial specification and differentiation from human iPSC-derived NPC. We performed literature data mining and selected a set of seven TF, which are enriched in OL compared with other neural lineages (2, 14), and are required for oligodendroglial specification (15, 16): ASCL1, MYT1, NKX2.2, NKX6.1, NKX6.2, OLIG2, and SOX10. Coding sequences for these proteins, as well as red fluorescent protein (RFP), were individually cloned into a tetracycline (tet)-inducible lentiviral vector. Human iPSC-derived NPC, which can be frozen and cost-efficiently expanded, as previously described (17), were transduced with a combination of lentiviruses expressing one of the TF candidates and the reverse tet-controlled transactivator (rtTA). Among all TF candidates, only SOX10 was capable of inducing O4 (9.99 \pm 0.81%), a highly specific marker of late-stage oligodendroglial progenitor cells (OPC) and OL, after 14 d of exposure (Fig. 1*B*). Controls, including NPC either uninfected or infected with RFP only, did not yield any O4⁺ cells (Fig. 1*A*). We subsequently determined the oligodendroglial induction capacity of SOX10 in combination with any of the remaining six TF. We identified OLIG2 as a factor that substantially increased the SOX10-mediated oligodendroglial lineage commitment (Fig. 1 *C* and *G*), whereas ASCL1 and MYT1 significantly decreased the number of O4⁺ cells (Fig. 1 *D* and *G*). The combination of SOX10 and OLIG2 with NKX6.2, a TF associated with oligodendroglial maturation, significantly increased further the portion of O4⁺ cells (Fig. 1 *E*, *F*, and *H*). Thus, we concluded that the ectopic expression of SOX10, OLIG2, and NKX6.2 (subsequently referred to as SON) was the most efficient combination of TF tested here to induce OL from iPSC-derived NPC.

Ectopic Expression of TF Accelerates Oligodendrogenesis from iPSC-NPC.

To further enhance the generation of human iOL, we generated a polycistronic lentiviral expression vector containing SON and RFP as a reporter gene under control of the retroviral spleen focusforming virus (SFFV) promoter (Fig. 2*A*). iPSC-derived NPC (13) were infected with SON/RFP-expressing lentivirus (Fig. 2*B*). After induction of SON, a two-step differentiation protocol was sufficient to derive increasing numbers of iOL over 28 d (Fig. 2*B*). One day after viral transduction, NPC medium was replaced by glial induction medium (GIM) containing different supplements, including PDGF, SAG (smoothened agonist), and thyroid hormone (Triiodo-L-Thyronine, T3; 10 ng/µL). After 4 d, this medium was replaced by



Fig. 1. Screening for oligodendroglial lineage inducing TF in human NPC. Human iPSC-derived NPC were infected with individual OL-specific TFs or RFP control virus. (A-F) OL-lineage commitment of infected iPSC-derived NPC was analyzed 14 d after transgene induction by immunostaining using the OL-specific antibody O4 (green). Nuclei were counterstained with Hoechst (blue). (A) Control cultures did not express the O4 epitope. (B) SOX10 was the only tested TF inducing O4⁺ OL. (C) Addition of OLIG2 enhanced the OL-lineage commitment, whereas (D) ASCL1 led to a decreased number of O4⁺ iOL. (E) Coexpression of SOX10, OLIG2, and NKX6.2 increased the number of O4⁺ cells, (F) accompanied by the appearance of iOL with a more mature oligodendroglial morphology. (Scale bars, 50 µm in A-E and 25 µm in F.) (G and H) Quantification of O4⁺ iOL over all cells with indicated TF combinations 2 wk after transgene induction. Data are presented as mean of replicates from three independent experiments + SD. One-way ANOVA with Bonferroni's multiple comparisons test was used as statistical test (*P < 0.05, **P < 0.01, ***P < 0.001).

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Fig. 2. SON induce a rapid and efficient oligodendroglial lineage commitment. (A) Schematic presentation of the lentiviral expression vector used for the polycistronic expression of SON. (B) Schematic summary of the differentiation protocol using NPCM, GIM, and DM. (C-F) Representative immunofluorescence images of different NPC and OL markers during differentiation. Nuclei were counterstained with Hoechst (blue). (C) iPSC-derived NPC homogenously expressed the neural progenitor marker NESTIN (green) and SOX1 (red). (D) Seven days after transgene induction, NG2⁺ and O4⁺ oligodendroglial lineage cells were detected. (E) By day 28, iOL expressed the O4-epitope, the more mature OL marker GALC, and presented with a branched morphology. (F) Further maturation led to the emergence of MBP⁺ mature iOL forming myelin sheaths. (Scale bars, 50 µm.)

differentiation medium (DM) containing higher concentrations of T3 (60 ng/µL) and lacking PDGF and SAG. To ensure the reproducibility of our protocol, all experiments were performed with four independent NPC lines derived from three different iPSC lines and one embryonic stem cell (ESC) line. All NPC lines homogeneously expressed the neural stem cell marker SOX1 and NESTIN (Fig. 2*C*). Seven days after SON induction, OL cultures comprised mainly NG2[±] glial progenitor cells (82.26 ± 2.36%) together with O4[±] cells (8.7 ± 3.0%) (Fig. 2*D*). At day 28 after SON induction, the majority of cells expressed the OL marker O4 (65.5 ± 11.1%) and presented with a more complex morphology identified by an increased number of primary and secondary processes (Fig. S1). Additionally, cells started to express more mature oligodendroglial markers, like GALC (19.19 ± 1.46%) and MBP (30.37 ± 7.87% of O4[±] cells) by days 28 and 35, respectively (Fig. 2 *E* and *F*; see also Fig. 5 *D* and *E*).

To assess the kinetics, efficiency, and yield of SON-mediated oligodendroglial lineage specification, we conducted weekly flow cytometry analyses of the O4 epitope expression during differentiation (Fig. 34). As a control, NPC were infected with RFP expressing lentivirus and cultured under the same differentiation conditions. All NPC lines tested were found to perform similarly with respect to iOL generation, starting from $8.7 \pm 3.0\%$ O4⁺ cells at day 7 to $65.5 \pm 11.1\%$ O4⁺ cells by day 28 (Fig. 3B). In contrast, only $1.4 \pm 0.5\%$ O4⁺ cells were identified in RFP-transduced cell cultures (Fig. 3B). The protocol was highly efficient and reproducible among all cell lines, illustrated by the quantification of O4⁺ cells at day 28, ranging from $62.1 \pm 9.5\%$ (ESC–NPC) to $79.0 \pm 14.8\%$ (iPSC–NPC-3) (Fig. 3C).

(ESC-NPC) to 79.0 \pm 14.8% (iPSC-NPC-3) (Fig. 3*C*). Yields of O4⁺ iOL (total O4⁺ cell number/starting NPC cell number) ranged from 133.70 \pm 24.83% at day 14 to 241.20 \pm 19.07% at day 28 (Fig. 3D). These data suggest an expansion of SON-expressing cells during our differentiation protocol. Identification of proliferative cells using Ki-67 revealed a proliferation rate of 35% among RFP⁺ cells at day 14, which declined to 10% by day 28, confirming that the transgene-expressing cell population further expanded during differentiation (Fig. 3E). Interestingly, the proliferation capability was retained in 20% of O4⁺ iOL at day 14 and diminished to 5% by day 28 (Fig. 3 F and G). Next, we tested whether iOL cultures can be expanded by a prolonged exposure to a glial expansion medium (GEM) postlentiviral transduction. Compared with GIM, GEM additionally contained FGF2 (5 ng/mL) and completely lacked IGF-1. SON-transduced NPC were cultured in GEM for either 4, 8, or 12 d and subsequently transferred to DM for an additional 28 d. FACS analyses of the iOL cultures after 0, 14, and 28 d in DM revealed that although exposure to GEM for 12 d reduced the differentiation efficiency (Fig. S24), yields of $O4^+$ iOL were significantly increased (Fig. S2B).

Furthermore, flow cytometry analyses exhibited the presence of an $O4^+/RFP^-$ cell population in SON-transduced cultures (Fig. 3A), which comprised up to 50% of the $O4^+$ cell population and which could be confirmed by immunocytochemical (ICC) analysis (Fig. 3H), suggesting transgene silencing in a subset of iOL. To analyze whether iOL become independent from trans-

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Next, we determined the influence of SON overexpression on the lineage commitment of NPC. ICC analysis of SON infected NPC cultures compared with RFP-infected control cultures at day 28 revealed a decreased number of SOX1⁺ NPC (Fig. 3*I* and Fig. S4*A*) and a significant switch from neuronal to oligodendroglial cell fate (Fig. 3*J* and Fig. S4*B*). In contrast, the astroglial lineage commitment was not affected (Fig. S4 *C* and *D*).

Global Gene-Expression Profiling Demonstrates That iOL Resemble Primary Human Adult OL. To further characterize the cellular identity of iOL, we compared the global gene-expression profiles of O4⁺ iOL purified at day 28 of differentiation with human MBP⁺ pOLderived from surgically resected brain samples from adult patients (Fig. S5), as well as with iPSC-derived NPC before induction of SON. As a negative control, we used gene-expression values of undifferentiated iPSC. The unbiased hierarchical clustering clearly demonstrated that iOL and pOL exhibit highly comparable geneexpression signatures and form a distinct cluster significantly segregating from NPC and iPSC (Fig. 4A). When we compared neural lineage-specific gene sets, we identified a strong up-regulation of OLspecific genes, such as OLIG1, MOG, and MBP in iOL compared with NPC, whereas NPC-related genes, including SOX1, PAX6, and PAX7, were down-regulated in iOL (Fig. 4 B and C). Because we compared O4-sorted iOL with MBP⁺ pOL, it is not surprising that iOL expressed some OPC-specific genes, such as PDGFRA and ST8SIA1, to a higher extent than pOL, indicating a more immature cell identity of iOL (Fig. S64). Coherently, pOL exhibit a stronger expression of some mature ÓL-specific genes, such as MAG and MOBP, compared with O4⁺ iOL. To further analyze the influence of ectopic SON expression on the oligodendroglial lineage commitment of NPC, we determined differentially expressed genes in iOL compared with the original NPC population. This analysis revealed 755 commonly up- and 955 commonly down-regulated genes among all iOL cell lines (Fig. S6 B and C). Gene Ontology (GO) terms associated with up-regulated genes in iOL include categories such as "cell adhesion," "myelin sheath," "axon ensheathment," "myelin," and "regulation of action potential." Conversely, GO terms associated with down-regulated genes include categories such as "cell cycle," "DNA replication," "mitosis," and "nucleoplasm" (Table S1).

These results indicate that ectopic expression of SON induces an oligodendroglial gene-expression profile comparable to native human adult OL.





iOL Differentiate into Mature MBP-Expressing OL in Vitro and Produce Myelin-Like Sheaths. Next, we assessed the terminal differentiation potential of iOL in vitro. At day 35, iOL cultures contained many highly branched O4⁺ cells (Fig. 5A), as well as mature OLexpressing CNP (2',3'-cyclic nucleotide 3'-phosphodiesterase) (32.82 \pm 2.80%) (Fig. 5B). Additionally, $30.37 \pm 7.87\%$ of O4⁺ cells differentiated into mature MBP⁺ iOL with myelin-like sheaths (Fig. 5 D and E) of which $36.62 \pm 8.93\%$ coexpressed MAG (Fig. 5C). To evaluate the myelinogenic capability of iOL in vitro, we purified O4⁺ iOL using magnetic cell separation (MACS) at day 21 and cultured them for 14 d on 3D cell culture surfaces with aligned nanofibers. ICC analysis of mature MBP+ iOL in these cultures revealed the extension of multiple processes along the nanofibers, with some of these extensions wrapping around the nanofibers (Fig. 5F). Evidence for ensheathment of axons in vitro was evaluated in cocultures of O4+ iOL with iPSCderived neurons. After 3 wk, the cultures exhibited myelin-like sheaths surrounding the axons, identified by confocal analysis of MBP and TUJ1 expression (Fig. S7A). Three-dimensional reconstruction of confocal optical sections in high magnification showed colabeling of neuronal processes (TUJ1) with MBP (Fig. 5 G and H), which was further evaluated by orthogonal projections clearly displaying the formation of MBP⁺ structures around neuronal processes (Fig. S7B). Control cultures completely lacked these MBP⁺ structures. These data clearly illustrate the capability of iOL to mature into MBP+ OL and to ensheath neuronal processes in vitro.

iOL Myelinate the Developing Brain and Remyelinate the Demyelinated Spinal Cord of Dysmyelinating Mice. The differentiation of iOL into myelin-forming OL was further validated by grafting day 14 Fig. 3. Quantification of oligodendroglial lineage cells after SON induction. (A) Representative flow cytometry analyses for the expression of O4 and RFP in control and SON cultures 7 and 28 d after transgene induction. (B) Quantification of O4⁺ cells in control and SON cultures 1 to 4r wk after transgene induction. Data are presented as mean of replicates from four independent experiments, each using NPC derived from an independent human PSC line + SD. (C) Quantification of O4⁺ iOL at day 28 derived from one human ESC and three independent iPSC lines. Data are presented as mean of replicates from three to five independent differentiation experiments per cell line + SD. (D) Quantification of O4⁺ iOL yields at days 14 and 28 after transduction. Data are presented as mean of replicates from three independent differentiation experiments, each using an independent iPSC-derived NPC line + SD. (E) Quantification of Ki-67⁺ transgene-expressing cells (RFP) and of (F) Ki-67⁺ /O4⁺ iOL at days 14 and 28 after induction. Data are presented as mean of replicates from three independent differentiation experiments + SD (**P < 0.01, ***P < 0.001). (G) Immunostaining of iOL for O4 (purple) and the proliferation marker Ki-67 (green) at day 14 after transduction. (H) Representative immunofluorescence image of O4⁺ iOL (green) 28 d after transduction either expressing (filled arrowhead) or silencing (empty arrowhead) the transgenes. (Scale bars, 40 μ m.) (/) Quantification of SOX1⁺ iPSC-derived NPC and (J) TUJ1⁺ neurons in control and SON cultures at day 28. Data are presented as mean of replicates from three independent differentiation experiments + SD. Student's t test was performed for statistical analysis (***P < 0.001).

MACS-purified O4+ iOL in the immune- and MBP-deficient Shi/ Shi Rag2^{-/-} mouse CNS. To facilitate the identification of transplanted cells, iOL coexpressed RFP as a reporter. These mice carry a deletion of seven exons of the MBP gene and completely lack MBP protein expression. Because of the lack of endogenous MBP expression in Shi/Shi Rag2^{-/-} mice, myelin generated by transplanted cells can be easily identified by MBP immunohistochemistry and by detection of myelin compaction, as well as the presence of the major dense line by electron microscopy (EM) (18, 19). To address developmental myelination, cells were grafted bilaterally and rostrally to the corpus callosum of newborn mice. Analysis of sagittal sections 16 wk postgrafting (wpg) indicated the presence of numerous MBP⁺ myelin profiles associated with RFP⁺ and human nuclei-positive (STEM101⁺) cells (Fig. 6 A and B). Higher magnification using confocal microscopy showed that iOL, identified by combined human nuclei (STEM101⁺) and cytoplasmic (STEM121⁺) antigens to highlight their cellular contours, extended processes frequently connected to MBP+ myelin, thus further validating the donor origin of the myelin (Fig. 6 C and G). MBP^+ myelin generated by grafted iOL wrapped around SM312⁺ host axons (Fig. 6D). The presence of human-derived myelin surrounding SM312⁺ axons was further validated combining MBP detection with human NOGO-A, which identifies late stages of mature OL (Fig. 6 E and F). The human-derived myelin internodes were often associated with the paranodal marker CASPR flanking Ankyrin-G nodal structures, demonstrating functionality of the human cell-derived myelin (Fig. 6H). Some of the animals were also used for ultrastructural analysis of myelin compaction. In control Shi/Shi Rag2-/mice, myelin sheaths were thin and noncompacted (Fig. S8). In

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Fig. 4. Global transcriptional profiling of iOL. (A) Hierarchical clustering of whole-genome expression profiles of iPSC (black), iPSC-derived NPC (green), iOL (28 d after transduction) (red), and human adult OL pOL (blue) revealed a strong correlation between iOL and pOL. (*B* and *C*) Pairwise scatterplot analysis of log_{2} -adjusted global gene-expression values of iPSC-derived NPC and their corresponding iOL (n = 10). Genes presenting with a <twofold difference in gene expression are illustrated in gray. (*B*) Characteristic OL-enriched genes were up-regulated in iOL, whereas (*C*) characteristic NPC-enriched genes were down-regulated.

mice that received iOL, numerous normal compacted myelin sheaths with alternating major dense lines and intermediate lines were observed (Fig. 6 I-K), validating unambiguously that iOL have the capacity to differentiate into functional myelin-forming cells in vivo. We quantified the percentage of myelinated axons over axons with a diameter >1 μ m in grafted (n = 3) vs. nongrafted mice (n = 3) by EM analyses. We did not detect a difference in the percentage of myelinated axons (grafted mice: $69.7 \pm 3.28\%$ vs. nongrafted mice $73.9 \pm$ 5.1%; P value: 0.2892). This result is most likely because of the competition between endogenous and grafted cells and the relatively early time point after transplantation of human iOL, which are known to differentiate more slowly than rodent cells. However, in the corpus callosum of transplanted mice, 12% of the axons were myelinated by grafted cells, as determined by the percentage of axons with compacted myelin over the total numbers of myelinated axons. Moreover, the g-ratio in grafted mice was significantly smaller compared with nongrafted mice $(76.8 \pm 1.8 \text{ vs. } 89.5 \pm 1.36; P \text{ value: } 0.0003)$

To address the ability of iOL to remyelinate demyelinated axons, RFP⁺ iOL were grafted into the dorsal funiculus of adult *Shi/Shi* Rag2^{-/-} mice spinal cord. Although *Shi/Shi* Rag2^{-/-} mice are hypomyelinated, numerous axons are surrounded by MBP⁻ non-compacted myelin (20) (Fig. S8). To optimize the number of demyelinated axons accessible for remyelination, mice were injected with lysophosphatidyl-choline (LPC) 48 h prior grafting to induce demyelination (21, 22). Twelve weeks postgrafting, immunolabeling of serial cross-sections for nucleic STEM101 and cytoplasmic RFP, together with MBP, revealed MBP⁺ donut-like myelin structures, indicating that grafted iOL not only colonized and remyelinated the lesion site, but also myelinated the entire neuraxis, including ventral and dorsal white and gray matter (Fig. S9 *A*, *C*, and *D*).

The differentiation of human grafted cells into various neural cell types was assessed by colabeling of human nucleic STEM101 and RFP with cell-specific markers of oligodendroglial cells (OLIG2, CC1), neurons (NeuN), and astrocytes (GFAP) (Figs. S9B and S10). Quantification of the various populations at the lesion site indicated that about 81% of the grafted human cells generated OLIG2⁺ oligodendroglial cells with 25.64% \pm 4.65% OPCs (OLIG2⁺/CC1⁻) and 56.06% \pm 3.11% mature OL (OLIG2⁺/CC1⁺) (Fig. S10G). The remaining population (19%) differentiated into NeuN⁺ neurons (2.66% \pm 0.66%) and GFAP⁺ astrocytes which could not be quantified because of highly interdigitated processes, but which by deduction represent most likely 17% of the grafted cells. That the majority of the

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Human iOL Respond to Compounds Promoting Oligodendroglial Differentiation. Identification of drugs inducing remyelination via promotion of oligodendroglial differentiation presents a promising approach for the treatment of demyelinating disorders like MS. Thus, we assessed whether iOL can be used to identify compounds promoting oligodendroglial differentiation. We selected six drug candidates (miconazole, clobetasol, benztropine, indometacin, clemastine, and oxybutynin), which have been previously described to promote differentiation or myelination of rodent OL (1, 23-25). iOL were cultured in a minimum growth medium lacking differentiation-promoting factors and were treated with either vehicle [0.01% (vol/vol) DMSO] as a negative control, T3 as a positive control, or the drug candidate dissolved in DMSO at three different concentrations (0.5, 1, 5 µM) (Fig. 7 and Fig. S13). In DMSOtreated control cultures, $14.01 \pm 2.89\%$ O4⁺ iOL were observed in minimum DM after 21 d of culture, whereas addition of T3 resulted in the doubling of O4⁺ cells (28.25 \pm 3.47%). Several drug candidates performed as well as T3 and demonstrated a dose-dependent



increase of O4⁺ cells (Fig. 7*A*). Interestingly, clemastine and oxybutynin failed to promote the formation of O4⁺ iOL. Furthermore, we observed a toxic influence of miconazole, benztropine, and clemastine on iOL at higher concentrations. Quantification of mature MBP⁺ iOL revealed a fourfold increase in the presence of T3 compared with DMSO control (Fig. 7*B*). The effect was most striking with 1 μ M miconazole, inducing an almost 10-fold increase of MBP⁺ cells compared with DMSO control cultures. Clobetasol and benztropine enhanced the formation of MBP⁺ mature iOL comparable to T3-treated cultures. These data demonstrate that iOL can be used to identify compounds that promote differentiation into O4⁺ as well as maturation into MBP⁺ OL.

iOL from Patients Carrying the N279K Microtubule-Associated Protein Tau Mutation Have a Higher Susceptibility to Oxidative Stress-Induced Cell Death. Next, we wanted to determine whether iOL can be used for in vitro disease modeling. The microtubuleassociated protein tau (MAPT) is developmentally expressed in OL (26, 27) and mutations in *MAPT* have been associated with frontotemporal dementia with Parkinsonism linked to chromosome 17 (FTDP-17), a disease also characterized by pathological changes in white matter (28–30). Therefore, we generated iOL from two iPSC clones from one patient carrying the N279K *MAPT* mutation associated with FTDP-17 (17) and compared these to their isogenic controls (31). Additionally, we included another independent control iPSC line.

After 28 d of differentiation, O4+ iOL harboring the N279K mutation (MAPT-OL) were morphologically indistinguishable from their gene-corrected control cell lines (MAPT-GC-OL) (Fig. 8A) and featured similar differentiation efficiencies among all cell lines included (Fig. 8B). We next set out to investigate whether the N279K MAPT mutation induces an altered expression of tau isoforms in iOL. We purified O4⁺ iOL using FACS before RNA sample preparation. Analysis of MAPT expression revealed mutation-specific significantly higher levels of 4R tau compared with MAPT-GC-OL (Fig. 8C), which is in line with observations in iPSC-derived neurons and brains of FTDP-17 patients harboring this mutation (17, 32). FTDP-17 patients display widespread neurodegeneration because of increased cellular vulnerability. Therefore, we investigated whether MAPT-OL are more susceptible to oxidative stress induced by rotenone, an inhibitor of the mitochondrial complex I. Exposure of MAPT and MAPT-GC-OL for 48 h to rotenone revealed an enhanced vulnerability of MAPT-OL to oxidative stress identified by an increased number of cleaved CASPASE-3⁺ iOL in MAPT-cultures (Fig. 8D). This effect was obvious in all tested concentrations of rotenone (100, 250, and 500 nM), leading to an average increase of cell death of $48.9 \pm$ 18.7% in MAPT-OL (Fig. 8E).

Discussion

Previously established protocols using in vitro patterning to derive OL from murine iPSC require fewer than 30 d (33). Additionally, it has been shown that mouse fibroblasts can be directly converted into OPC by the forced expression of Sox10, Olig2, and Zfp536 or Nkx6.2, resulting in ~15% O4⁺cells after 20 d (2, 14). However, the generation of human OL from iPSC or ESC is more challenging and characterized by much longer culture periods (70-150 d), limited efficiencies, and variable reproducibility. Although the protocols to generate human OL have been further optimized to reduce culture times and increase efficiency, they still require at least between 60 and 130 d of culture to generate OPC from iPSC/ ESC-derived NPC and only a small percentage of cells become MBP⁺ mature OL (9-11). One of the first studies demonstrating the successful derivation of human OL from iPSC was published in 2013 (11) using a modified in vitro-patterning strategy from ESC-based protocols (34, 35). However, the transition from pre-OPC to OPC was hampered; by day 150, 20-40% of the cells expressed the OPC marker CD140a, whereas the late OPC marker O4 was only present in 5-10%. Interestingly, the application of low levels of oxygen (3%) during the differentiation of PSC to OL could greatly enhance oligodendroglial lineage commitment, resulting in around 40% O4⁺ cells at day 135 (10). The use of retinoic acid from the beginning of differentiation, followed by the application of sonic hedgehog agonist SAG could further increase the oligodendroglial lineage commitment, resulting in up to 70% of O4⁺ OPCs at day 75 (9). To accelerate oligodendroglial differentiation that is orchestrated by TF, we tested individual and combinations of TF previously shown to be involved in oligodendroglial differentiation in rodents (15, 36-40). Our results have defined a combination of three different TF that efficiently induces iOL and indeed overcomes the rate-limiting steps of the transition from a neurogenic to a glial progenitor and accelerates oligodendroglial differentiation. In line with an earlier study, in which overexpression of SOX10 alone was sufficient to induce oligodendroglial commitment (~22% O4⁺ cells 14 d after transduction) in neural progenitors derived from the human fetal brain (16), in our experiments SOX10 was the only TF that induced expression of O4 in iPSC-derived NPC, demonstrating that SOX10 is one key TF to induce oligodendroglial lineage commitment. However, combination of SOX10 with OLIG2 and NKX6.2 further enhanced the commitment into the oligodendroglial lineage, resulting in a significantly higher percentage of O4⁺ cells 14 d after induction. Our protocol was highly efficient and reproducible, resulting in 50-70% O4⁺ cells and yields of 220-260% after 28 d using three different iPSC-derived NPC lines and a single ESC-derived NPC line.

The myelinating capacity of iOL was tested in vitro and in vivo. In vitro iOL ensheathed the neuronal processes of iPSC-derived

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neurons, as well as nanofibers confirming that physical properties of axons are sufficient to induce wrapping of axons, as it has been described for rodent OL (41, 42). The coculture of iOL with nanofibers facilitates the identification of compounds that exclusively promotes axon ensheathment without potentially modulating molecular axonal signaling. We did not detect CASPR accumulations, indicative for paranode formation in iPSC-derived neuron/ iOL cocultures, suggesting that distinct axonal signaling cascades required for the formation of paranodes and nodes were not activated in iPSC-derived neurons. However, CASPR⁺ paranodes were easily detectable in vivo after transplantation into *Shi/Shi Rag2^{-/-}* mice. The transplanted cells not only efficiently myelinated the forebrain in newborn *Shi/Shi Rag2^{-/-}* mice, but also remyelinated the adult demyelinated spinal cord. Proof of the donor origin of the myelin was provided by immunohistochemistry using humanspecific nuclei and cytoplasmic markers, allowing us to highlight the connection of donor cell processes to myelin internodes surrounding host axons, and the integration of these myelin internodes into nodal and paranodal elements. We provided additional proof of the donor origin of myelin by the detection at the ultrastructural level of thick, compact myelin with the major dense line, a hallmark of wild-type myelin compared with thin, noncompacted, and dense line-null shiverer myelin (18, 19). It is known that despite the lack of MBP, the number of axons ensheathed by noncompacted myelin increases progressively in shiverer mice over time. In the gracile fasciculus in the dorsal spinal cord, the number of myelinated axons increases from 56% 2 wk after birth to up to 77% at 20 wk (20). Injection of lysolecithin in the dorsal funiculus results in complete demyelination of these axons (22). (Re)myelination, although more efficient in the lesion, was not limited to the dorsal funiculus because MBP+ exogenous myelin was found throughout the spinal cord (over 22 mm) with numerous myelin-

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PNAS | Published online February 28, 2017 | E2249

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Fig. 7. iOL are suitable to test the differentiation promoting effects of selected compounds. Quantification of (A) O4⁺ and (B) MBP⁺ iOL after treatment with either vehicle, T3, or the drug candidate dissolved in DMSO at three different concentrations (0.5, 1, 5 μ M) for 21 d in minimum DM. Data are presented as mean of replicates from three independent experiments + SD. One-way ANOVA with Dunnett's multiple-comparisons test was performed for statistical analysis comparing the mean of each sample with DMSO control (**P* < 0.05, ***P* < 0.01, ****P* < 0.001). 0* = Toxic culture condition.

ated axons in gray matter and ventral white matter, suggesting that grafted cells dispersed widely in their adult host environment. Similar observations were reported for transplanted human fetal neural progenitors (21), indicating that O4⁺ iOL are as migratory as human fetal NPC in adult *Shi/Shi Rag2^{-/-}* mice. Compared with the developing CNS, the adult CNS represents an environment with impaired tissue plasticity and trophic support. Whereas the colonization and remyelination potential of murine iPSC-derived NPC has already been described in this environment (22), this study demonstrates that human iPSC-derived oligodendroglial lineage cells have similar capabilities. However, one has to keep in mind that the endogenous cells are compromised because of the lack of MBP expression in contrast to the grafted cells. Therefore, the grafted cells may have a competitive advantage and not only remyelinate demyelinated axons but also replace the endogenous pool of myelinating cells. In the adult spinal cord, the majority of iOL differentiated into OL (81%), rarely into neurons (2%), and few into astrocytes (estimated to be 17%), confirming previous data obtained with iPS-derived O4⁺ cells (9) or O4⁺ human fetal cells (11) grafted in the newborn shiverer mouse brain. Interestingly iOL-derived neurons were observed only in gray matter surrounding the lesion, and iOL-derived astrocytes were detected only in the demyelinated area but not in the ventral or lateral white matter tracks, suggesting that their differentiation is highly regulated by cell-specific environmental cues.

The iPSC technology is an emerging tool for drug development. Promotion of remyelination represents until now an unmet treatment strategy for patients with MS. Without SON, less than 2% of NPC differentiated into $O4^+$ OL. Importantly, SON transduced NPC cultured in minimal medium lacking T3 and growth factors (except dbcAMP), resulted in ~14% of O4⁺ OL. Addition of T3 increased the efficiency to 28% and culturing in DM to 40% after 21 d (as shown in Figs. 3B and 7, respectively). These data demonstrate that SON is sufficient to induce an oligodendroglial cell fate; however, growth factors are necessary to further increase maturation of the cells. The increase in differentiation by exogenous factors is prerequisite to use iOL for

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pharmacological screens aimed at the identification of compounds promoting remyelination. In large compound screens using primary or iPSC-derived OL from rodents, a number of Food and Drug Administration-approved drugs has been identified that were able to promote oligodendroglial differentiation in vitro and remyelination in vivo (1, 24, 25). To determine whether iOL may be suitable for pharmacological screens, we cultured iOL in the presence of compounds identified in earlier rodent studies (1, 23-25). In contrast to these earlier studies, some but not all of these compounds increased the number of O4+ iOL in a dose-dependent manner and were at least as effective as T3, a known promoter of oligodendroglial differentiation. Furthermore, only a subset of these drugs enhanced the maturation of O4⁺ iOL into MBP⁺ mature OL, suggesting that the compounds affect different stages of oligodendroglial differentiation. Miconazole demonstrated the strongest effect on iOL; this is in line with an earlier publication by Najm et al., in which they reported a strong differentiation promoting effect of miconazole on OL (1). However, we observed a toxic effect with a fivefold-higher concentration, suggesting that miconazole might have a narrow range of efficacy. Our observations thus suggest that there are species-specific differences between rodent and human OL that could be relevant for drug screens aiming at identifying compounds that promote oligodendroglial differentiation.

To determine whether iOL are suitable for disease modeling, we characterized in a proof-of-concept study the phenotypes of iOL derived from a patient diagnosed with an inherited form of FTD. FTD is characterized by cortical degeneration of the frontal and temporal lobe that in 15-20% of patients with an inherited form of FTD is a result of mutations in the MAPT gene that encodes the MAPT located on chromosome 17q21. The neuropathology of FTDP-17 patients with mutations in the MAPT gene is characterized by tau⁺ inclusions in neurons and glia, including OL (for review, see ref. 43). Furthermore, extensive myelin pathology can be observed in patients with FTD (28–30). In OL tau regulates and stabilizes the microtubule network that is also involved in the transport of RNA granules, for example, those containing MBP mRNA. Knockdown of tau or mutated tau in rodent OL impairs process outgrowth and the differentiation into MBP⁺ myelinating mature OL (44, 45). Therefore, we assessed whether changes in OL may directly contribute to the white matter pathology observed in FTDP-17 patients. In iOL from patients with a N279K mutation in the MAPT gene, we observed as expected, significantly increased expression levels of the 4R tau isoform. Furthermore, we observed an increased susceptibility to cell death induced by respiratory stress compared with gene-corrected control cell lines, similar to that reported in iPSC-derived neurons from the same patient (17). These data suggest that MAPT mutations in OL may directly contribute to myelin pathology and thus to disease progression in patients with FTDP-17.

In summary, our studies demonstrate that a combination of three TFs, namely SOX10, OLIG2, and NKX6.2, greatly accelerates the generation of OL from iPSC-derived NPC and that these cells are suitable for disease modeling and pharmacological screens. Thus, our protocol should significantly facilitate the development of high-throughput screening platforms and the study of human myelin diseases and repair strategies using patient-derived iPSC.

Methods

Culturing of Human PSC. The iPSC included in this study have previously been generated and characterized (13, 17). iPSC colonies were maintained on a layer of mitotically inactivated mouse embryonic fibroblasts in human ESC medium consisting of knockout DMEM (Invitrogen) with 20% (vol/vol) Knockout Serum Replacement (Invitrogen), 1 mM β -mercaptoethanol (Invitrogen), 1% nonessential amino acids (NEAA; Invitrogen), 1% penicillin/ streptomycin/glutamine (PAA), freshly supplemented with 5 ng/mL FGF2 (Peprotech). PSC were split at ratios of 1:6–1:8 every 7 d by mechanic disaggregation with 1 mg/mL collagenase IV (Invitrogen).

Generation and Culturing of Human NPC. NPC were derived from human PSC by treatment with small molecules as previously described (13, 17). Methodological details are provided in *SI Methods*.

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Fig. 8. MAPT-OL exhibit mutation related phenotypes. (A) Immunostaining for O4 (green) demonstrating differentiation of iPSC carrying the N279K *MAPT* mutation (MAPT1, MAPT2) and genetic corrected controls (MAPT1 GC, MAPT2 GC) into iOL. Nuclei were counterstained with Hoechst (blue). (Scale bar, 25 μ m.) (*B*) Flow cytometry-based quantification of O4⁺ iOL after 28 d of differentiation in *MAPT* mutation cultures, genetic corrected cultures, and an independent healthy control culture. Data are presented as mean of replicates from three independent experiments + SD. (C) Quantitative RT-PCR analysis on control, MAPT gene-corrected, and MAPT mutated iOL cultures for 4R tau isoforms containing exon 10. Expression levels were normalized to total tau expression and control lines. Data are presented as mean of replicates from three independent experiments + SD. One-way ANOVA with post hoc Tukey test was performed for statistical analyses (***P* < 0.01). (*D*) Quantification of cleaved CASPASE 3⁺ iOL in control and MAPT cultures after 48 h of either vehicle [0.01% (vol/vol) DMSO] or rotenone treatment. Data are presented as mean of replicates from three independent experiments + SD. One-way ANOVA with post hoc Tukey test was performed for statistical analysis (**P* < 0.05, ***P* < 0.01). (*E*) All results combined after normalization by setting all control cultures to 100%, show that *MAPT* N279K causes a higher sensitivity to oxidative stress. Error bars present SD. Student's *t* test was performed for statistical analysis (***P* < 0.001).

Lentiviral Vector Construction and Production of Lentiviral Particles. Details including the cloning strategies are provided in *SI Methods* and Fig. S14.

Transduction of NPC for TF Screening. Human NPC were seeded with a density of 1×10^5 cells per well in 12-well plates, allowed to attach overnight, and transduced with equal volumes of concentrated Lenti-rtTA and 1-TF virus particle supplemented with 5 µg/mL protamine sulfate (Sigma) in fresh NPC expansion medium (NPCM). The 2-TF infections were done by mixing equivalent volumes of Lenti-rtTA, pLV-TetO-SOX10 and 1-TF virus particle for infection. For 3-TF infections, the volume of each virus was reduced by one quarter and equivalent volumes of Lenti-rtTA, pLV-TetO-SOX10, pLV-TetO-SOLIG2, and 1-TF were mixed for NPC transduction. Further details are provided in *SI Methods*.

Oligodendroglial Differentiation. For oligodendroglial differentiation, human NPC were seeded with a density of 1×10^5 cells per well in 12-well plates, allowed to attach overnight, and transduced with concentrated SON lentiviral particle and 5 µg/mL protamine sulfate in fresh NPCM. Viral medium was removed after 24 h and replaced with GIM. After 4 d, GIM was replaced by DM. Details are provided in *SI Methods*.

ICC. Details of ICC are provided in SI Methods and Table S2.

Flow Cytometry Analysis. Cells were enzymatically detached and singularized by accutase treatment for 10 min at 37 °C. Following washing with PBS, singularized cells were resuspended in PBS/0.5% BSA buffer and filtered through a 70- μ m cell strainer (BD Falcon). After determination of cell number, cells were incubated with mouse IgM anti–O4-APC antibody (Miltenyi Biotec) following the manufacturer's protocol. Stained cells were washed, resuspended in PBS/0.5% BSA buffer (5 × 10⁶ cells/mL), and immediately sorted using a FACSAria cell sorter (BD Biosciences). Debris, doublets, and aggregates were excluded by appropriate gating using forward and side scatter. Additionally, DAPI was used for dead cell exclusion. Unstained cells were used to set background fluorescence and undifferentiated human NPC were used as negative controls.

Neuronal Differentiation, in Vitro Myelination Assay, and 3D Culture. Human iPSC-derived NPC were differentiated into neurons as previously described (17). To assess the in vitro myelination capacity of iOLs, O4⁺ cells were purified at differentiation day 21 by magnetic cell sorting using anti-O4 MicroBeads (Miltenyi Biotec) following the manufacturer's protocol. Purified iOLs were added to 21-d-old neuronal cultures derived from iPSC-derived NPC populations or nanofibers. Details are provided in *SI Methods*.

Isolation of Adult Human pOL. Brain tissue was obtained from adults undergoing surgical resections as treatment for nontumor-related intractable epilepsy in accordance with the guidelines set by the Biomedical Ethics Unit of McGill University. As described previously (46), tissue specimens were enzymatically digested and placed on a linear 30% Percoll density gradient (Pharmacia Biotech). Further details are provided in *SI Methods*.

Whole-Genome Expression Analysis. Details including the microarray data processing are provided in *SI Methods*.

Cell Transplantation. Details of cell transplantation into *Shi/Shi Rag2^{-/-}* dysmyelinating immunodeficient mice are provided in *SI Methods*. Animal experiments were performed according to European Community regulations and INSERM ethical committee (authorization 75-348; 20/04/2005) and were approved by the local Darwin ethical committee.

Immunohistochemistry and EM. The methodology for immunohistochemistry and EM is described in detail in *SI Methods*.

Compound Screen. To assess the sensitivity of iOL toward differentiation promoting drugs, iPSC-derived NPC were transduced with concentrated SON virus particle, as described above. Viral medium was removed after 24 h and replaced with GIM lacking T3. The end of the virus infection period was termed day 0. Further details including the differentiation with drug candidates are provided in *SI Methods*.

Generation and Characterization of N297K MAPT NPC and Isogenic Controls. The N279K MAPT iPSC-derived NPC included in this study have previously been generated and characterized (17). Frozen NPC, termed FTDP-17-1-I and

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FDTP-17-1-II in the aforementioned publication were thawed at passages 10–14 and designated as MAPT-1 and MAPT-2 in this study.

Quantitative RT-PCR. Details of RT-PCR including a list of primers are provided in *SI Methods* and Tables S3.

Stress-Induced Cell Death. Details are described in SI Methods.

Quantifications. Quantifications were performed as described in detail in SI Methods.

Statistics. Data of at least three independent differentiation experiments are presented as mean + SD. Statistical significance was determined by Student's *t* test and with one-way ANOVA, respectively.

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ACKNOWLEDGMENTS. We thank Jacqueline Trotter (Molecular Cell Biology, Department of Biology, Johannes Gutenberg University Mainz) for critical reading and helpful discussion of the manuscript, and Martina Sinn and Ingrid Gelker for excellent technical support. This study was supported by German Research Foundation Grant SFB-TR128-B7; Ku1477/6-1 (to T.K.); Interdisciplinary Clinical Research Center, Münster Grant IZKF; KuT3/012/15 (to T.K.); an operating grant from the Multiple Sclerosis Society of Canada (to J.A.); Cluster of Excellence REBIRTH (A.S.); Multiple Sclerosis Society of the United States Grant RGPA-1501-02553; and INSERM, the program "Investissements d'Avenir" ANR-10-IAIHU-06 and ANR-11-INBS-0011–NeurATRIS. S.M. is supported by a Du Pré grant from the Multiple Sclerosis International Foundation, Fondation des Treilles, Ecole de Neurosciences de Paris, and Electricité de France Foundation. A.M. is supported by the European Leukodystrophy Foundation.

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