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Patient iPSC-Derived Neurons for Disease Modeling of Frontotemporal Dementia with Mutation in CHMP2B

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SUMMARY

The truncated mutant form of the charged multivesicular body protein 2B (CHMP2B) is causative for frontotemporal dementia linked to chromosome 3 (FTD3). CHMP2B is a constituent of the endosomal sorting complex required for transport (ESCRT) and, when mutated, disrupts endosome-to-lysosome trafficking and substrate degradation. To understand the underlying molecular pathology, FTD3 patient induced pluripotent stem cells (iPSCs) were differentiated into forebrain-type cortical neurons. FTD3 neurons exhibited abnormal endosomes as well as impaired mitochondrial function. Moreover, mitochondria of FTD3 neurons displayed defective cristae formation, accompanied by deficiencies in mitochondrial respiration and increased levels of reactive oxygen. In addition, we provide evidence for perturbed iron homeostasis, presenting an in vitro patient-specific model to study the effects of iron accumulation in neurodegenerative diseases. All phenotypes observed in FTD3 neurons were rescued in CRISPR/Cas9-edited isogenic controls. These findings illustrate the relevance of our patient-specific in vitro models and open up possibilities for drug target development.

INTRODUCTION

Frontotemporal dementia (FTD) is the second most common form of early-onset (<65 years) dementia, accounting for 5%–8% of total dementia cases. FTD is a neurodegenerative disorder with cognitive impairment affecting the frontal and/or temporal lobes of the brain associated with progressive brain atrophy (Rossor et al., 2010). FTD is clinically, neuropathologically, and genetically heterogeneous. One gene affected in familial cases is the charged multivesicular body protein 2B (CHMP2B) located on chromosome 3 (FTD3). Patients display global cortical and central brain atrophies, with no apparent amyloid plaque formation or conclusive hyperphosphorylated tau aggregates (Isaacs et al., 2011). Early behavioral changes include apathy, restlessness, disinhibition, and hyperorality. Late-stage behavioral changes include stereotype behavior, mutism, and dystonia (Isaacs et al., 2011). CHMP2B is a component of the endosomal sorting complex required for transport III (ESCRT-III) complex, which facilitates recycling or degradation of cell surface receptors (Chassefeyre et al., 2015). As such, the FTD3-causing mutation of CHMP2B affects functionality of the endosome. Mouse and Drosophila FTD3 animal models have yielded valuable in vivo insights into the dysfunction of the endosomal lysosomal system and pathologic progression (Ahmad et al., 2009; Ghazi-Noori et al., 2012). However, transgene integration and species-specific differences may contribute to observed phenotypes in such models. Hence, there is an emerging need for human FTD3 models. In addition, studying how CHMP2B, as a rare mutation, contributes to neurodegenerative disorders has not yet attracted broader attention. Consequently, a CHMP2B mutant cellular model could provide further insights into common underlying dysfunction of biological pathways, disrupted or disturbed in neurodegenerative diseases and thereby linking different forms of neurodegeneration.

The availability of viable neurons from patient brains, at least in part, limits the investigation of the mechanism of
**A**

- **ssODN 90 bp**
  - Silent Mutation
  - CHMP2B DNA
  - Exon 6
  - 31489G>C Mutation
  - Gene-corrected hiPSCs
  - FTD3 hiPSCs

**B**

- hiPSCs Seeding:
  - Undifferentiated hiPSCs
  - Neuroepithelial sheet
  - Neural rosette formation
  - Neuronal maturation

- Medium:
  - DMEM/F-12:Neurobasal (1:1) + N2 + B27

- Growth factor/inhibitor:
  - LDN193189 + SB431542
  - bFGF
  - B/GDNF + db-cAMP + Ascorbic Acid

- Coating:
  - Matrigel
  - Poly-L-Ornithine/Laminin

**C**

- **TUJ1**
  - Relative mRNA levels

- **VGLUT1**
  - Relative mRNA levels

- **FOXG1**
  - Relative mRNA levels

**D**

- **Average neurite length (μm)**

- **Soma size (μm²)**

(legend on next page)
neurodegenerative pathogenesis. In this context, human-induced pluripotent stem cells (iPSCs) provide invaluable access to study the disease progression in neurons derived from patient iPSCs and facilitate the development of new therapies (Ehrlich et al., 2015; Rasmussen et al., 2014). Meanwhile, recent advances of state-of-the-art genome engineering technique CRISPR/Cas9 (Ran et al., 2013) have had a tremendous impact allowing for gene correction in patients who are carriers of disease-causing single-point mutations. Such genetically edited iPSCs are ideal isogenic controls for the patient-derived iPSCs, allowing to precisely dissect the significance of the disease-causing mutation while maintaining the patient’s own genetic background.

To study the cellular and molecular events of FTD3, we derived a disease model using human iPSCs from patients carrying the 31449G > C mutation in CHMP2B and isogenic controls generated via the CRISPR/Cas9 system with subsequent in vitro neuronal differentiation. Our model not only confirmed partly prior-observed pathologies in animal models such as endosome dysregulation (Ghazi-Noori et al., 2012), but also revealed additional disease-relevant phenotypes in FTD3 by way of abnormal mitochondrial ultrastructure and function. Moreover, RNA sequencing (RNA-seq) uncovered that the CHMP2B mutation manifests in dysregulated expression of key genes associated with Parkinson’s disease (PD) and Alzheimer’s disease (AD) and indicated that perturbed iron homeostasis is one of the underlying disease mechanisms.

RESULTS

Generation of FTD3 iPSCs and Genetically Corrected Isogenic Controls

Skin fibroblasts from two symptomatic FTD3 patients (H150 and H151) and one presymptomatic FTD3 patient (H242) were reprogrammed using non-integrative episomal plasmids. All of these individuals are from the same family, and one independent healthy control from an unrelated family was also included. Individual FTD3 iPSC lines were assessed for pluripotency features via qPCR (Figure S1A) and immunocytochemistry (ICC) (Figure S1B). Assessment of pluripotency potential was confirmed via embryoid body formation and subsequent spontaneous differentiation (Figure S1C). The FTD3-causing mutation 31449G > C (located at the splice acceptor site in CHMP2B, at the border of intron 5 and exon 6) was repaired in the FTD3 iPSC lines using the CRISPR/Cas9 system (Figure 1A). The presence of altered CHMP2B mRNA, as formerly shown in Urwin et al. (2010) was confirmed via qPCR and showed expression of the intron 5 and delta-10 isoforms only in the patient iPSC. These abnormal transcripts, generated via the CHMP2B mutation affecting the splice acceptor site, are completely absent in the isogenic controls (Figure 1A). All iPSC and genetically corrected isogenic lines were determined to be karyotypically normal (Figure S1D).

Differentiation of FTD3 iPSCs to Forebrain Cortical Neurons

We employed a modified dual SMAD protocol (Figure 1B) to generate forebrain-specific cortical neurons. Flow cytometry analyses demonstrated the neural progenitor cell (NPC) cultures to be comparable with respect to NESTIN and SOX1 expression (Figure S2A). All NPCs expressed the forebrain markers Pax6 and OTX2 (Figure S2B). ICC revealed the majority of the matured neurons to be glutamatergic (immunopositive for VGlut1 [Figure S2C] and co-positive for MAP2, TAU, and TuJ1 [Figure S2C]). Approximately 10% of neurons were GABAergic (Figure S2C), while approximately 5% were GFAP-positive astrocytes, with a minor population (2%–3%) comprising tyrosine hydroxylase-positive dopaminergic neurons (Figure S2C). Since quantification via ICC is challenging we performed qPCR for TuJ1, VGlut1, and FOXG1 in order to estimate the comparability in regard to the neural cell composition of the investigated FTD3 iPSC- and isogenic iPSC-derived neurons. All lines showed comparable amounts of expression with no significant differences, which is also indicative that the mutation in CHMP2B does not affect the differentiation potential toward neurons (Figure 1C). Neurons exhibited spiking activity and action potentials, which occurred both spontaneously
Mito Norm

- **A**
  - Area of individual endosome
  - FTD3 neurons
  - Isogenic controls
  - Independent control

- **B**
  - Relative endosome : cytoplasma area ratio
  - FTD3 neurons
  - Isogenic controls
  - Independent control

- **C**
  - Top 10 endosome-associated genes differentially expressed
  - Log2 fold change FTD3/ISO
  - Genome wide corrected FDR 5%

- **D**
  - Top 10 mitochondria-associated genes differentially expressed
  - Log2 fold change FTD3/ISO
  - Genome wide corrected FDR 5%

- **E**
  - Relative mitochondria : cytoplasma area ratio
  - FTD3 neurons
  - Isogenic controls
  - Independent control

- **F**
  - Mito Cristaeless

- **G**
  - MitoTracker
  - FTD3 neurons
  - Isogenic controls

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FTD3 Neurons Display Abnormal Mitochondria

Given that mitochondrial defects are commonly observed in neurodegenerative diseases, we assessed whether such a phenotype is evident in FTD3 neurons. While we noted no apparent difference in size of mitochondria between FTD3 neurons and their isogenic controls, we did observe a significant increase in abnormal mitochondria within neurons compared with only 7% in isogenic controls (Figure 3A). Thus, FTD3 neurons are exposed to higher levels of oxidative stress, and targeted gene editing can reverse the disease phenotype. Subsequently, we investigated whether the increased oxidative stress impacts cell viability, but we detected no evidence for increased apoptosis in FTD3 neurons (Figure 3B).

FTD3 Neurons Display Impaired Mitochondria Function and Increased Oxidative Stress

Prompted by our observations of aberrant mitochondrial morphology, we sought to test the functionality of such FTD3 mitochondria by assessing their basal respiration rate, ATP-linked respiration, proton leak, and reserve capacity. Our analyses uncovered a reduced basal respiration rate in FTD3 neurons as well as a reduced maximal capacity (Figure 3E). Consistent with mitochondrial dysfunction manifesting in oxidative stress, we observed up to 20% occurrence of oxidative stress in FTD3 neurons compared with only 7% in isogenic controls (Figure 3A). Thus, FTD3 neurons are exposed to higher levels of oxidative stress, and targeted gene editing can reverse the disease phenotype. Subsequently, we investigated whether the increased oxidative stress impacts cell viability, but we detected no evidence for increased apoptosis in FTD3 neurons (Figure 3B).

Global Gene Expression Analyses Confirm Defects in the Endosome, Mitochondrial Dysfunction, and Increased Oxidative Stress

To obtain an overview of gene expression changes in CHMP2B-dependent neurodegeneration, we performed
RNA-seq analyses, which revealed 3,205 differentially expressed genes in FTD3 neurons compared with their isogenic controls. Heatmap analyses showed that neurons derived from iPSCs of all three FTD3 individuals clustered together, while isogenic controls clustered together with the independent healthy individual (Figure 4A), suggesting that the isogenic control lines reverted to a healthy control gene expression profile after repair of the CHMP2B mutation. To identify which cellular components are primarily affected in FTD3 neurons, we performed gene ontology (GO) term analysis. We found enrichment of the GO term for postsynaptic membrane, synaptic membrane, neuron projection, and synaptic vesicle membrane (Figure 4C), and depletion of early endosome membrane, lysosomal lumen, and endocytic vesicle membrane (Figure 4D).

With the aim to identify additional genes associated with the main phenotypes observed in FTD3 neurons, we extracted the top ten differentially regulated genes functionally linked to neurodegeneration and associated with endosomes (Figure 2D), mitochondria (Figure 2H), and oxidative stress (Figure 3C). Strikingly, APOE ranked among the ten most downregulated endosome-associated genes (Figure 2D). APOE4, an isoform of APOE, is a well-known risk factor for sporadic AD (sAD) (Corder et al., 1993) and, consistent with a putative role in FTD3 pathophysiology, some sAD patients present with enlarged early endosomes (Cataldo et al., 2000).

Most interestingly, we observed an upregulation of leucine-rich repeat kinase 2 (LRRK2) among the mitochondria-related genes (Figure 2H). LRRK2 is one of the five genes mutated in familial forms of PD. Previous studies reported mitochondrial fragmentation due to increased expression of LRRK2 (Wang et al., 2012), pointing to an association between upregulated LRRK2 expression and cristae dysgenesys in FTD3 neurons. LRRK2 was also detected in the oxidative stress group (Figure 3C) and is upregulated in some cases of sporadic PD (Cho et al., 2013) accompanied by elevated levels of oxidative stress (Nguyen et al., 2011). MPO and APOD, both implicated in AD (Figure 3C) (Maki et al., 2009; Martinez et al., 2012), were strongly downregulated in FTD3 neurons.

Imbalance of Iron Homeostasis Represents a Unifying Factor Linking Endosome, Mitochondrial Dysfunction, and Oxidative Stress

Via our in silico analyses, we identified mis-expression of genes encoding proteins or receptors known to be involved in iron homeostasis. Among these were TRPC6 (12-fold upregulated), HFE (8-fold downregulated), and ABCG2, HP, and CP (all 2-fold downregulated) (Figure 4B). A potential imbalance of iron homeostasis might plausibly induce intracellular ferrous iron (Fe2+) accumulation, damaging mitochondria and inducing stress, to be further exacerbated by the defects in the endosome compartment contributing to increased amounts of trapped Fe3+ within endosomes (Figure 4E). Supporting this hypothesis, cytoplasmic Fe2+ levels are significantly increased in FTD3 neurons versus isogenic control neurons (Figure 4F). This increase in intracellular iron was consistent in several independent neural differentiation experiments, and the increased expression in TRCP6 was also confirmed via western blot analysis (Figure 4B), further substantiating our hypothesis for impaired iron homeostasis triggered by altered expression and protein amounts of key regulators.

Taken together we have shown that FTD3 neurons have enlarged early endosomes, display mitochondria dysfunction and perturbed iron homeostasis, with subsequent increase in oxidative stress.

DISCUSSION

The biological function of CHMP2B has primarily been investigated in animal models, with the attendant caveat that observed phenotypes are potentially species specific or contributed to by transgene integration. Here, to preclude such issues, we developed an FTD3 patient-derived disease model by way of iPSCs generation, genome editing, and in vitro neuronal differentiation techniques.

We identified enlargements of particularly the early endosome population in FTD3 neurons, indicating that perturbed endo-lysosomal fusion and degradation of substrates is key to the underlying FTD3 pathophysiology. Cellular

Figure 3. Impaired Mitochondrial Function and Oxidative Stress in FTD3 Neurons

(A) CellRox green and SYTOX red flow cytometry assay of live oxidative stressed neurons. n = 3 with three replicates in each experiment, significant differences are indicated by \( *p < 0.01 \) and \( **p < 0.001 \).

(B) CellEvent Caspase-3/7 green and SYTOX red flow cytometry assay of apoptotic neurons.

(C) Total number of oxidative stress-associated genes and the top ten differentially expressed genes (on the basis of fold-change) within an FDR of 5%, n.s., not significant.

(D and E) Mitochondrial function profile of FTD3 neurons (H150, H151 and H242) and isogenic controls (H150 ISO, H151 ISO, H242 ISO).

(D) oxygen consumption rate (OCR) was measured before (basal) and after the addition of oligomycin (oligom), carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone (FCCP), and a mixture of rotenone (rot) and antimycin A (antimA), respectively; (E) mitochondrial respiration parameters were calculated based on the OCR measurements shown in (D) as described in the Supplemental Experimental Procedures. n = 3 with three replicates in each experiment, significant differences are indicated by \( *p < 0.05 \), \( **p < 0.01 \), and \( ***p < 0.001 \).
Down-regulated GO terms (cellular component) enrichment

Up-regulated GO terms (cellular component) enrichment

TRPC6 Western Blot

Relative intensity to β-Actin

FTD3 neurons
Isogenic controls

Fe⁺⁺ Assay

P = 0.0426

(legend on next page)
components, including mitochondria, are degraded in the endosome; in amyotrophic lateral sclerosis, a progressive lysosomal deficit is causative for impaired degradation of damaged mitochondria in motor neurons (Xie et al., 2015). Defective recycling of large organelles, such as mitochondria, is the earliest detectable impairment in a chain of events initiated by defective recycling (Cannizzo et al., 2012), which clearly links endosome and mitochondrial dysfunction.

FTD3 neurons displayed abnormal mitochondrial localization and cristae formation, manifesting in impaired functionality typical for neurodegenerative diseases. Defective mitochondria show impaired ATP generation and release of deleterious chemicals such as reactive oxygen species (ROS), potentially inducing cell death. We contend that increased ROS release contributes to increased organelle damage, which cannot be appropriately ameliorated through degradation owing to the observed endosome defects, resulting in a self-perpetuating cycle of cellular damage.

Strikingly, in FTD3 neurons we identified a robust, significant upregulation of LRRK2 encoding the dardarin protein. One function of dardarin is the modulation of intracellular trafficking via recruitment of RAB7 (Greggio et al., 2006). Mutant CHMP2B impairs the recruitment of RAB7, crucial for endosome-lysosome fusion (Urwin et al., 2010). This relationship between CHMP2B, RAB7, and dardarin is supported by studies showing that dardarin negatively regulates RAB7-dependent localization of the lysosome (Dodson et al., 2012). Therefore, upregulation of LRRK2 as a consequence of CHMP2B mutation might contribute to deficient endo-lysosomal trafficking and fusion. Dardarin also plays a role in mitochondrial functionality and positioning, since induced expression of wild-type LRRK2 induces mitochondrial fragmentation via increased fission and upregulation of DLPL, which is critical for correct mitochondrial distribution (Wang et al., 2012).

Furthermore, our analysis revealed APOE to be significantly downregulated in FTD3 neural cells. The cells analyzed via RNA-seq are a mixed population of neurons and glial cells, with an up to 90% enrichment of VGlut1+ neurons. Due to this mixed population of cells we were still able to detect APOE via RNA-seq, even though it is mainly expressed in astrocytes. APOE is a major cholesterol carrier in neurons that supports lipid transport and brain injury repair, and there is a key requirement for APOE in facilitating the clearance of soluble amyloid beta (Abeta) (Jiang et al., 2008). Furthermore, APOE suppresses microglia activation (Laskowitz et al., 2001), and reduced APOE levels might potentially trigger inflammatory events. Together, this suggests that downregulation of APOE in FTD3 neurons might be a contributing factor in the microglia activation seen in postmortem FTD3 brains. Interestingly, we observed another phenotype restricted to glial cells. It has previously been reported for FTD3 mouse models and patient brains that these display p62 inclusions, which are indicative for impaired autophagy, oxidative stress, and neurotoxicity (Ghazi-Noori et al., 2012; Holm et al., 2007). We identified these p62 inclusions only in the S100β+ glial cell population and not in our TUJ1+ neurons (Figure S4B). Both the APOE and the p62 phenotypes clearly show that, even though several disease features can be replicated in our FTD3 neurons, other neural cell types need to be investigated in the future to understand the full complexity and pathophysiology of the disease.

Based on the differentially expressed key components and increase of intracellular iron in FTD3 neurons, we present an additional pathological feature of FTD3, which is associated with an imbalance in iron homeostasis (Figure 4E). Iron uptake principally proceeds through iron bound to transferrin mediated via the transferrin receptor or via direct uptake of Fe2+ facilitated by a variety of receptors including TRPC6 (Mwanjewe and Grover, 2004). We propose that upregulation of TRPC6 manifests in increased uptake of Fe2+, while downregulation of HFE (which competes with Tf), results in uptake of more ferric iron which can further be reduced into Fe2+. In contrast, export of heme-bound iron via ABCG2 or unbound Fe2+ through FPN can be reduced. Concordant with this model, we confirmed in several individual differentiation experiments a small but significant increase of intracellular iron within 5 weeks of neuronal differentiation in FTD3 neurons. Excessive amounts of intracellular Fe2+ are toxic to

Figure 4. Transcriptional Analysis of FTD3 Neurons Versus Isogenic Controls and Quantification of Intracellular Fe2+ Content
(A) Heatmap and hierarchical clustering analysis of gene expression profile of FTD3 neurons (H150, H151, H242) and isogenic controls (H150 ISO, H151 ISO, H242 ISO) as well as one independent healthy control (H256).
(B) qPCR validation of genes of interest from RNA-seq results and western blot relative quantification of TRPC6 protein expression.
(C and D) GO term enrichment analysis of differentially expressed genes via ClueGO v1.4 software. Functionally grouped network with terms as nodes linked based on their kappa score level (≥0.5), where only the label of the most significant term per group is shown. The node size represents the term enrichment significance. Functionally related groups partially overlap.
(E) Diagram of key dysregulated genes involved in intracellular iron homeostasis.
(F) Fe2+ assay and concentration normalized to total cellular protein. n = 3 with three replicates in each experiment, significant differences are indicated by *p < 0.05.
cells and trigger the production of ROS. Moreover, Fe^{2+} and hydrogen peroxide engage in the so-called Fenton reaction, producing ferric iron and highly reactive hydroxide, which damages DNA, proteins, and lipids in the cell (Altamura and Muckenthaler, 2009).

FTD3 neurons present with enlarged early endosomes, increased cristaless mitochondria, intracellular iron accumulation, and oxidative stress. All of these phenotypes were rescued via genetic correction of the CHMP2B mutation.

We propose that in FTD3 neurons two events occur in parallel: (1) mutant CHMP2B causes an imbalance in iron homeostasis triggering mitochondrial impairment, which consequently triggers ROS and neuronal damage, and (2) mutant CHMP2B-dependent enlargement of endosomes, which causes defects of the endo-lysosomal degradation and recycling pathway, including recycling of receptors for iron uptake, results in accumulation of defective mitochondria, imbalanced iron homeostasis, ROS, and neuronal damage. Future studies will determine the exact role of mutant CHMP2B on a molecular level within these cellular events.

In conclusion, our findings indicate a possibility of targeting iron imbalances therapeutically in efforts to intervene in this cycle of damage to attempt treatment of FTD3. Furthermore, the CHMP2B-dependent human iPSC-derived neurons can be added to the list of in vitro cellular model systems that serve as a tool to study not only common mechanisms of pathogenesis, such as mitochondrial defects and oxidative stress, but also the effects of iron accumulation in neurodegenerative diseases.

**EXPERIMENTAL PROCEDURES**

**Cell Culture, iPSC Generation, and Neuronal Differentiation**

iPSCs were generated via episomal reprogramming (Addgene, 27077, 27078, 27080) from human dermal fibroblasts obtained from skin biopsies. Neuronal differentiation followed a modified dual SMAD protocol.

**Genome Editing**

Generation of isogenic controls from the three FTD3 iPSC lines was achieved via the CRISPR/Cas9 system (Addgene, 62988) using single-strand donor oligonucleotides as a template.

**Transmission Electron Microscopy**

Cells were seeded on 13 mm Thermorax plastic coverslips (Nunc, 174950) coated with poly-O-Lysine/laminin. Ultra-thin (50–70 nm) sections were prepared, contrasted with 2% uranyl acetate (PolyScience, 21447) and lead citrate, and examined using a Philips CM100 transmission electron microscope equipped with a Morada camera.

**RNA-Seq Analysis**

For deep RNA-seq, libraries of FTD3 neurons, isogenic controls, and an independent healthy control were prepared and sequenced on a HiSeq 2000 Sequencing System (Illumina) following the manufacturer’s protocol.

**Statistical Analysis**

Data are presented as mean ± SE. n is the number of independent differentiations of NPCs (from the cryobank) to neurons. Significance of data was evaluated by Student’s t test or two-way ANOVA with Bonferroni post hoc test. Unless noted otherwise, p < 0.05 was considered statistically significant.

For further details on Experimental Procedures, see the Supplemental Information.

**ACCESSION NUMBERS**

The RNA-seq data described in this study have been deposited in the NCBI GEO database under accession number GEO: GSE92340.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, four figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2017.01.012.

**AUTHOR CONTRIBUTIONS**


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**REFERENCES**

Supplemental Information

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A

Isogenic Control

FTD3 Neuron

Axon
Axon

Nucleus

Nucleus

* * x

x x

B

FTD3 Neurons

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Isogenic Controls

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SUPPLEMENTAL FIGURE LEGENDS

Figure S1. Basic Characterisation of iPSCs, Related to Figure 1. (A) qPCR analysis of pluripotency genes in FTD3 iPSCs relative to H1 human ESCs. (B) ICC of FTD3 iPSCs for pluripotency marker proteins. Scale bars, 200 μm for all. (C) ICC of FTD3 iPSCs with βIII-TUBULIN (ectoderm), α-SMA (mesoderm), and AFP (endoderm) after EB differentiation. Scale bar, 200 μm for all. (D) Karyotype analysis of FTD3 iPSCs and isogenic controls showing no abnormalities.

Figure S2. Forebrain Identity of NPCs and Differentiation into Cortical Neurons, Related to Figure 1. (A) Flow cytometry assay for NESTIN and SOX1 in the FTD3 NPCs and isogenic controls. (B) Immunocytochemistry (ICC) assay of NPCs for NESTIN and forebrain progenitor protein markers OTX2 and PAX6. Scale bar, 50 μm. (C) ICC confocal images of FTD3 neurons and isogenic controls: dendritic neuronal protein marker MAP2 coupled with axonal protein marker TAU, glial marker GFAP, cortical markers TBR1 and CTIP2, axonal neuronal protein marker TUJ1 coupled with glutamatergic neuron marker VGLUT1, GABAergic neuron marker GABA and dopaminergic marker TH. Scale bars, 50 μm for all and 15 μm for TUJ1 coupled with VGLUT1.

Figure S3. Functional Characterization of Cortical Neurons, Related to Figure 1. (A) Intracellular calcium kinetics in FTD3 neurons (H150, H151, H242) and isogenic controls (H150 ISO, H151 ISO, H242 ISO). (B), (C), (D) and (E) Electrophysiological properties of FTD3 neurons and isogenic controls: (B) action potentials activated by current steps; (C) spontaneous elicited action potentials; (D) whole-cell currents activated by a series of voltage steps: in some recordings, excitatory postsynaptic potentials (EPSP) were observed at negative potentials as demonstrated in the enlarged portion of the recording; (E) representative image of patched cells. Scale bar, 300 μm
Figure S4. TEM Overview Image of Neurons and p62 Aggregates Characterization in the Neural Culture, Related to Figure 2-4. (A) Images represent initial segment of an axon and a portion of the nucleus. In the FTD3 neurons, endosome-like structures (*) and mitochondria (x) accumulate, and the arrangement of microtubule structures (indicated by triangle arrow) is perturbed compared with isogenic controls. Scale bar, 2 μm for all. (B) Immunostaining for p62 and the neuronal marker protein TUJ1, glial maker protein S100β in FTD3 neural cultures and isogenic control. Red box indicates area with cells having p62 positive aggregates. DAPI was used a nuclear marker. Scale bar, 20 μm for all.
**Table 1A**

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**Table 1B**

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SUPPLEMENTAL EXPERIMENTAL PROCEDURES

Ethical Statement
The study was approved by the Ethics Committee of the Capital Region of Denmark (H-4-2011-157), and written informed consent was obtained from each participant before enrollment.

Cell Culture, iPSC Generation and Neuronal Differentiation
Human dermal fibroblasts cultures were obtained from skin biopsies from two symptomatic (H150, H151) and one presymptomatic (H242) FTD3 individuals (all from the same family), as well as one healthy unrelated individual (H256). The FTD3 individuals were carriers of the G-to-C transition at the 5’ acceptor splice site of CHMP2B exon 6 on chromosome 3 (Skibinski et al., 2005). Fibroblasts were cultured in DMEM supplemented with 1% GlutaMAX™, 10% FBS and 1% Penicillin-Streptomycin. For the generation of iPSCs, one microgram of reprogramming plasmid mixtures was electroporated into $1 \times 10^5$ fibroblasts using the Neon® transfection system (Thermo Fisher Scientific, USA) with a 10μL kit according to the manufacturer’s instructions. The plasmid mixtures used in this experiment were pCXLE-hOCT3/4, pCXLE-hSK and pCXLE-hUL (Addgene, 27077, 27078, 27080) (Okita et al., 2011). Electroporation conditions were as followed: 1,200 V, 20 ms, 2 time pulses. After transfection, $5 \times 10^4$ cells were distributed onto 35-mm dishes coated with Matrigel (BD Biosciences, USA) and cultured in DMEM, 1% GlutaMAX™, 10% FBS without bFGF and Penicillin-Streptomycin. The culture medium was replaced the next day and supplemented with bFGF, Penicillin-Streptomycin and cultured at 5% O₂, 5% CO₂ and 90% N₂. On day 3 to 4 depending on confluence, the culture medium was replaced with ESC medium consisting of Essential 8™ basal medium with supplement (Thermo Fisher Scientific, A1517001), 100 μM Sodium Butyrate and 0.1% Penicillin-Streptomycin. The colonies were manually picked 21–28 d
after plating, and those colonies morphologically similar to human embryonic stem cells (ESCs) were selected for further cultivation and evaluation. When confluent, iPSC were passaged 1:3-6 with 0.5 mM EDTA on to Vitronectin™ (Thermo Fisher Scientific, A14700)-coated 6-well dishes. The hiPSC clones used in this study are summarized in Table S1A. Mutation status was confirmed by Sanger sequencing.

iPSC lines were differentiated toward neural-specific progenies in 6-well plate wells (NUNC, USA) pre-coated with Matrigel® (BD Biosciences, 354277) in a 21% O₂ 5.0% CO₂ 37°C incubator employing a modified dual SMAD protocol (Shi et al., 2012). Cells were expanded in E8 medium until around 100% confluence was reached. Neural induction was initiated by changing the medium to neural basic medium consisted of 50% DMEM/F-12 medium, 50% Neural basal medium (Thermo Fisher Scientific, 10888-022), N2 (Thermo Fisher Scientific, 17502-048), B27 without Vitamin A (Thermo Fisher Scientific, 12587-010), supplemented with SB431542 a small molecule inhibitor of the TGFβ pathway (Selleck, S1067) and LDN193189 an inhibitor of the BMP pathway (Selleck, S2618). The duration of this initial neural induction was 10 to 12 days. Subsequently, cells were split onto poly-O-Lysine/laminin (Sigma, L2020) pre-coated 6-well plate wells and remained in the neural induction medium overnight. Expansion of the neural progenitor cells, for cryopreservation and subsequent terminal differentiation was achieved by proliferating these cells in neural basic medium supplemented with 10 ng/ml bFGF. Neuronal maturation was achieved by addition of the following media: Neural basic medium supplemented with 20 ng/ml BDNF (Cell Guidance Systems, GFH1-10), 10 ng/ml GDNF (Cell Guidance Systems, GFH2-10), 50 μM db-cAMP (Sigma, D0627) and 200 μM L-Ascorbic acid 2-phosphate (Sigma, A8960). One week after terminal differentiation cells were replated again for further maturation.

**Genome Editing**
Generation of isogenic controls was achieved via the CRISPR/Cas9 system using single-strand donor oligonucleotides (ssODNs) as templates for single nucleotide repairs.

Isogenic gene-corrected controls were obtained using the CRISPR/Cas9 system in combination with ssODNs as homologous templates covering the mutation site. The ssODNs were designed with silent mutations that both prevent recutting of the CRISPR/Cas9 and create a restriction enzyme recognition site for SacII to enable screening for recombinant clones. The CRISPR/Cas9 vectors were generated following the protocol from Ran et al. (Ran et al., 2013). At a confluency of 80%, the iPS cells growing in a 10-cm plate were incubated with Accutase for 5 – 10 minutes at 37 °C. iPS cells were harvested and centrifuged. 2 million cells were co-transfected with 10 µg of the CRISPR/Cas9 plasmid (Addgene, 62988) and 1 µL of 100 ng/µL of the ssODNs. We used a 4D nucleofector (program CA167) from Amaxa in combination with the P3 Primary Cell Kit for transfection. Transfected cells were plated on a 10 cm dish coated with Matrigel. After one day, Puromycin (Thermo Fisher Scientific, A1113803) was added for 2 days in a 1:50000 dilution. After selection, Puromycin was withdrawn. Around 4 - 8 days after selection, resistant iPS colonies were picked and transferred to a cell culture 96 well plate. When the wells became confluent, iPS cells were split into a DNA isolation plate and a 96 well cell culture dish for further cultivation. DNA was isolated using the prepGEM™ Kit (zyGEM, PTI0500). Subsequently, a PCR was performed with primer pair P1-FW and P1-RV. PCR products were digested with SacII for 30 – 60 minutes at 37 °C. Positive clones were sequenced to confirm gene-editing of CHMP2B.

Quantitative qPCR

RNA was extracted using RNeasy® Plus Mini Kit (Qiagen, 74134) according to the manufacturer’s protocol. cDNA was synthesized from 1 µg of total RNA in 20 µL reaction using iScript™ cDNA synthesis Kit (BIO-RAD, 1708890). After synthesis, the cDNA was diluted five
times with double distilled water and stored at −20 °C. Quantitative RT-PCR (qPCR) reactions were done in triplicates using the FastStart Lightcycler 480 SYBR Green I Master (Roche, 04707516001) on LightCycler® 480 real-time PCR system (Roche, Switzerland). cDNA samples were subjected to PCR amplification with primers for OCT4, SOX2, NANOG, GDF3 and GAPDH (Table S1B). A commercial ESCs line H1 (Wicell, USA) was used as a positive control.

Quantification of CHMP2B RNA Transcripts

Half a microgram of RNA was subjected to first strand synthesis using random hexamer primers and SuperScript III Reverse Transcriptase (Invitrogen) in accordance with the manufacturer’s instructions. PCR reactions were assembled using 5 µl of Brilliant III ultra-fast SYBR Green qPCR master mix (Agilent Technologies), 0.5 µl of forward primer (10 µM), 0.5 µl of reverse primer (10 mM), 2 µl of water, and 2 µl of cDNA template. The PCR program was as follows: 2 min at 50°C, 10 min at 95°C and then 40 cycles of 15 s at 95°C, 30 s at 58-60°C (dependent on primers) and 1 min at 72°C. To distinguish the three different isoforms of CHMP2B, a common forward primer was used with either one of the three isoform specific reverse primers. The sequences of the primers were listed in Table S1B. For normalization, primers detecting the housekeeping gene EIF6 were used. CHMP2B specific primers were used at an annealing temperature of 58°C, whereas the EIF6A primer pair was used at 60°C.

The relative abundance of the CHMP2B WT transcript between the different cell lines was calculated by normalization to EIF6 employing the relative standard curve method. The relative abundance of the three different isoforms of CHMP2B within each cell line was calculated by constructing standard curves from serial dilutions of a single plasmid holding all three amplicons in a stoichiometry of 1:1:1. This way we were able to calculate the exact amount of each isoform in relation to one another within in each cell line using the relative standard curve method.
**Immunocytochemistry and Confocal Microscopy**

For the immunofluorescence staining, cells were fixed with 4% formaldehyde in Dulbecco's Phosphate-Buffered Saline (DPBS) for 15 min, permeabilized with 1% Triton X-100 in DPBS for 15 min, and blocked with 2% bovine serum albumin in DPBS for 1 h. Thereafter, cells were incubated with the following primary antibodies (Table S1C) for 1 h. Primary antibodies were detected using secondary antibodies conjugated to Alexa Fluor 488 (1:500; Molecular Probes), Alexa Fluor 594 (1:500; Molecular Probes) and Alexa Fluor 647 (1:500; Molecular Probes). Cells were washed in DPBS and stained with DAPI (Sigma-Aldrich, D9542). Fresh DPBS was replaced and cells were imaged immediately using sequential line scanning with a Leica TCS SP5 II inverted CLSM or Z-stack recording with a Zeiss LSM 710 CLSM equipped with UV laser, Argon laser and AOBS beam splitter.

**Flow Cytometry**

To verify purity and similar proportions of marker positive hNPCs from each differentiated cell line, intracellular markers of hNPCs were analyzed by flow cytometry. When the hNPCs reached 80%–90% confluence, cells were dissociated using TrypLE™ Express (Thermo Fisher Scientific, 12605010) and incubated with Fixation/Permeabilization Working Solution (eBioscience, 88-8824) at room temperature for 60 mins. After centrifuging, the supernatant was discarded and cell pellets were resuspended in 1× Permeabilization Buffer containing monoclonal antibodies against NESTIN (BD Pharmingen, 560393) and SOX1 (BD Pharmingen, 561592). After 45 min of incubation in the dark at room temperature, cells were washed with fresh 1× Permeabilization Buffer, and resuspended in Flow Cytometry Staining Buffer (eBioscience, 00-4222) at a density of 1×10^6 cells/ml. Subsequently, they were analyzed by the CSampler™ Flow Cytometer (BD
Unlabeled samples were used as controls for gating. Data was analyzed using the FlowJo software (version 10.0).

**Whole-cell Patch Clamp Recording**

Whole-cell current-clamp and voltage-clamp measurements were performed on FTD3 neurons and isogenic controls after terminal differentiation from hNPCs for 5 weeks by using a MultiClamp 700B amplifier and MultiClamp Commander (Axon Instruments, Molecular Devices, USA). Patch pipettes were fabricated from borosilicate glass capillaries (1.5mm O.D., Fisher Scientific) on a gravity puller (Model PP-830, Narishige, Japan) and had a final tip resistance of 2–3 MΩ. Recordings were performed at 36±1°C in extracellular solution containing (in mM): NaCl, 125; HEPES, 10; NaH2PO4, 1.25; MgCl2, 1; CaCl2•2H2O, 2; glucose, 25; pyruvic acid, 3 (pH 7.4, 290 mOsm) and patch pipettes were filled with intracellular solution containing (in mM): K-gluconate, 135; NaCl, 7; MgCl2, 2; HEPES, 10; Na2-ATP, 2; Na2-GTP, 0.3 (pH 7.2, 270 mOsm). The recording conditions were modified from Shi et al., 2012 (Shi et al., 2012). Spontaneous mini excitatory/inhibitory postsynaptic or action potential activity was monitored for 1 min. The membrane potential was adjusted to approximately -60 mV followed by stepwise current injections from 0 pA to +50 pA. Voltage-clamp recordings were performed from a holding potential of -110 mV. Currents were activated by a series of depolarizing steps ranging from -90 mV to +30 mV. All analog signals were acquired at 50 kHz, filtered at 6 kHz, digitized with a Digidata 1440 converter (Axon Instruments, Molecular Devices) and stored using pClamp10 software (Axon Instruments, Molecular Devices). Recordings were corrected for a -14 mV junction potential and data were analyzed using Clampfit 10 software (Axon Instruments, Molecular Devices).
Recordings of Intracellular Calcium Kinetics

Recordings of intracellular calcium kinetics were performed on FTD3 neurons and isogenic controls after terminal differentiation from hNPCs for 5 weeks. Neurons cultured in 96-well plates were loaded for 1 h at 37°C with 100 μL of the fluorescent calcium indicator, Calcium 5 (Molecular Devices, USA) dissolved in Hanks Balanced Salt Solution (HBSS; Thermo Fisher Scientific) with 20 mM HEPES and 1.26 mM CaCl₂ to 1/3 of the concentration suggested by provider. Recordings were performed at 37°C in a FDSS 7000 fluorescence kinetics plate reader (Hamamatsu Photonics, Japan). The excitation wavelength was 480 nm; the emission wavelength was 540 nm and the sampling frequency 1 Hz. Compound dilutions were prepared in assay buffer and 20 μl applied with 10 min intervals. Baseline fluorescence was recorded for 10 min before application of 300 μM glutamate/10 μM glycine, 100 μM Serotonin (5-hydroxytryptamine, 5-HT), 25 mM K⁺, 100 μM GABA, 100 μM Dopamine (DA), and 300 μM acetylcholine (Ach). The fluorescence was normalized to the first data point of each of the traces.

Morphometric Analyses

Images of beta tubulin III-immunoreactive cells were recorded (12 images/cell line) using a bright field microscope (×200) connected to a digital camera (Leica). Average neurite length and soma size were assessed on coded images (experimenters ‘blind’ to sample identity) using the software ImageJ (version 1.50e) and the plugin NeuronJ (version 1.4.3). Only cells displaying an intense beta tubulin III immunostaining and a neuronal morphology (well-preserved cell soma and ≥ 1 primary neurite) were included in the analyses. Results are expressed as mean values ± SEM from 129-515 neurons per cell line.

Transmission Electron Microscopy
Cells were seeded on 13 mm Thermanox™ plastic coverslips (NUNC, 174950) coated with poly-O-Lysine/laminin. Cells were fixed with 3% glutaraldehyde (Merck, 1042390250) in 0.1 M Na-phosphate buffer, pH 7.4, post-fixed in 1% osmium tetroxide in 0.1 M Na phosphate buffer, dehydrated stepwise in a graded ethanol series, and embedded in Epon (TAAB, T031). Semi-thin (2 μm) sections were cut with a glass knife (KnifeMaster II, LKB Bromma 7800) on an ultramicrotome (Leica Ultracut, Leica Microsystems, Wetzlar, Germany), stained firstly with 1% toluidine blue O (Millipore, 1159300025) and 1% Borax (LabChem, LC117101). Ultra-thin (50 nm to 70 nm) sections were sectioned with a diamond knife (Jumdi, 2 mm) on an ultramicrotome (Leica Ultracut), contrasted with 2% uranyl acetate (Polyscience, 21447) and lead citrate, and examined using a Philips CM100 transmission electron microscope equipped with a Morada camera.

**Oxidative Stress Assay**

Reactive oxygen species (ROS) production was quantified in FTD3 neurons and isogenic controls using CellROX® Green Flow Cytometry Assay kit (Thermo Fisher Scientific, C10492). Staining was carried out according to the manufacturer’s instructions, and stained cells were dissociated from the plate using Accutase and immediately analyzed by flow cytometry using CSampler™ Flow Cytometer (BD Accuri™, USA).

**Apoptosis Assay**

Activated Caspases 3 and 7 in apoptotic cells was quantified in FTD3 neurons and isogenic controls using CellEvent® Caspase-3/7 Green Flow Cytometry Assay kit (Thermo Fisher Scientific, C10427). Staining was carried out according to the manufacturer’s instructions, and labeled cells were dissociated from the plate using Accutase and immediately analyzed by flow cytometry using
CSampler™ Flow Cytometer (BD Accuri™, USA).

Seahorse XF®96 Mitochondria Respiration Assay

Oxygen consumption rate (OCR) was measured using a Seahorse XF®96 Extracellular Flux Analyzer (Seahorse Biosciences, USA). FTD3 hNPCs and isogenic controls were seeded in a Seahorse 96-well cell culture microplate at a density of 6.5×10³ cells/well for neuronal maturation. On the day of the assay, the culture media was changed to unbuffered DMEM (pH 7.4) supplemented with 2.5 mM glucose and the cells were equilibrated for 10 min at 37 °C in a CO₂-free incubator. The pH of the reagents used to test mitochondrial function was adjusted to 7.4 on the day of assay. The OCR measurement cycle consisted of 3-min mix and 3-min measurement of the oxygen level. Test of mitochondrial function was initiated by three baseline OCR measurement cycles. These were followed by the sequential injection of the inhibitor of ATP synthase, Oligomycin (2 µM final concentration); the uncoupling agent, Carbonyl cyanide 4-(trifluoromethoxy)phenylhydrazone (FCCP, 1 µM); a mixture of the inhibitors of mitochondrial complexes I and III, Rotenone (0.5 µM) and Antimycin A (0.5 µM) with one OCR measurement cycle in between each injection and two final measurement cycles. Oxygen consumption rates were automatically recorded and calculated by the Seahorse XF® 96 software, Wave. Subsequent to the Seahorse analysis, the protein content was measured for each well using the Pierce assay with bovine serum albumin as standard. Results are expressed as mean values ± SEM from 12 wells/plate obtained in triplicate for each condition. Mitochondrial respiration parameters were calculated as follows: basal respiration, non-mitochondrial respiration (minimum measurement after rotenone/antimycin injection) was subtracted from the last measurement obtained before oligomycin injection; proton leak, non-mitochondrial respiration was subtracted from the measurement after oligomycin injection; maximal respiration, measurement obtained after FCCP
injection; *spare respiratory capacity*, basal respiration subtracted from maximal respiration; *ATP production*, measurement after oligomycin injection was subtracted from the last measurement before oligomycin injection.

**MitoTracker® Assay**

Cells for MitoTracker® Red CMXRos (Molecular Probes, M7512) analysis were cultured on 0.17 μm glass coverslips coated with poly-O-Lysine/laminin. When ready for staining, cells were incubated with 50 nM MitoTracker® in DMEM/F-12 medium for 15 min in 37 °C. Following, fixation in 4% paraformaldehyde in DPBS, permeabilization in 0.5% Triton X-100 in DPBS for 15 min, incubation with DAPI for 10 min in room temperature. In between each step the samples were washed 3 times in DPBS. After the last wash, the glass coverslips were mounted with mounting media and sealed onto slides. Images were acquired by laser scanning confocal microscopy (Zeiss LSM 710).

**RNA Sequencing Analysis**

For deep RNA sequencing, libraries of FTD3 neurons, isogenic controls and one independent healthy control were prepared and sequenced independently on HiSeq 2000 Sequencing System (Illumina, USA). For library construction, after extracting the total RNA from samples, mRNA and non-coding RNAs are enriched by removing rRNA from the total RNA with kit. By using the fragmentation buffer, the mRNAs and non-coding RNAs are fragmented into short fragments (about 200-500 nt), then the first-strand cDNA is synthesized by random hexamer-primer using the fragments as templates, and dTTP is substituted by dUTP during the synthesis of the second strand. Short fragments are purified and resolved with EB buffer for end reparation and single nucleotide A (adenine) addition. After that, the short fragments are connected with adapters, then
the second strand is degraded using UNG(Uracil-N-Glycosylase) finally. For data processing, Illumina Casava1.8 software was used for base calling. Clean reads were obtained by filtering reads with adaptor sequences and low quality reads. Clean reads that aligned to rRNA were removed. After removing reads aligned to rRNA, we used SOAP2 to align clean reads to the hg19 UCSC RefSeq (RNA sequences, GRCh37) (para: -m 0 -x 1000 -s 40 -l 32 -v 5 -r 2 -p 4). Gene level expression was performed by transforming uniquely mapped transcript reads to TPM. Normalized differentially expressed genes were detected. Genes whose expression changes are more than 1.5 folds with FDR < 5% identified in FTD3 neurons compared to isogenic controls were submitted to Gene ontology (GO) enrichment analysis. GO enrichment analysis of assembled transcripts was performed with ClueGO v1.4 (Bindea et al., 2009) using the following parameters: use GO term fusion; enrichment/depletion two-sided hypergeometric statistical test; correction method: Benjamini-Hochberg; GO term range levels: 3-8; minimal number of genes for term selection: 10; minimal percentage of genes for term selection: 10%; \( \kappa \)-score threshold: 0.5; general term selection method: smallest \( p \) value; group method: \( \kappa \); minimal number of subgroups included in a group: 3; minimal percentage of shared genes between subgroups: 50%.

For qPCR validation of the RNA sequencing results, we performed independent neuronal differentiation experiments and examined key gene expression, and all primer sequences for these genes are listed in Table S1B.

**Western Blotting**

Cell pellets from FTD3 neurons and isogenic controls were lysed in mPER™ mammalian protein extraction reagent (Thermo Fisher Scientific, 78501) containing protease inhibitor (Complete tablets, Roche 11873580001) and phosphatase inhibitors (PhosSTOP tablets, Roche 04906845001). 10 \( \mu \)g of protein were separated by NuPAGE™ Novex™ 4-12 % Bis-Tris mini gel
(Thermo Fisher Scientific, NP0335BOX) followed by immunoblotting with rabbit anti-TRPC6 (Alomone Labs, ACC-017) and mouse anti-β-Actin (Sigma, A5441) primary antibodies, donkey-anti-rabbit IRDye® 680LT (LI-COR, 926-68023) and donkey-anti-mouse IRDye® 800CW (LI-COR, 32212) secondary antibodies. Immunoblots were developed by Odyssey® Fc Imaging System (LI-COR) and data was processed using the software ImageStudio version 5.2.5. Expression levels of TRPC6 were normalized to β-Actin.

**Ferrous Iron Assay**

Cell pellets were obtained from FTD3 neurons and isogenic controls. An iron assay kit (Abcam, ab83366) was subsequently used to quantify ferrous iron (Fe^{2+}) in the neurons according to manufacturer's instructions. Briefly, cell pellets were homogenized with 120 µL iron assay buffer and supernatants were collected. 20 µL cell supernatant was used for the assay and volume was adjusted to 100 µL with iron assay buffer. Then, 100 µL of the iron probe solution was added producing a Fe^{2+}-Ferene S complex that absorbs light at 593 nm. Infinite® 200 PRO Multimode Reader (TECAN, Switzerland) was used to detect absorbance at this wavelength.

**Statistical Analysis**

Data are presented as mean ± standard errors (S.E.). “n” is the number of independent differentiations of NPCs (from the cryobank) to neurons. Significance of data was evaluated by Student’s t test or two-way ANOVA with Bonferroni post hoc test. Unless noted otherwise, P<0.05 was considered statistically significant.
Supplemental References


