Niemann-Pick type C1 patient-specific induced pluripotent stem cells display disease specific hallmarks

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Abstract

Background: Niemann-Pick type C1 disease (NPC1) is a rare progressive neurodegenerative disorder caused by mutations in the NPC1 gene. In this lysosomal storage disorder the intracellular transport and sequestration of several lipids like cholesterol is severely impaired, resulting in an accumulation of lipids in late endosomes and lysosomes. The neurological manifestation of the disease is caused by dysfunction and cell death in the central nervous system. Several animal models were used to analyze the impaired pathways. However, the underlying pathogenic mechanisms are still not completely understood and the genetic variability in humans cannot be reflected in these models. Therefore, a human model using patient-specific induced pluripotent stem cells provides a promising approach.

Methods: We reprogrammed human fibroblasts from a NPC1 patient and a healthy control by retroviral transduction with Oct4, Klf4, Sox2 and c-Myc. The obtained human induced pluripotent stem cells (hiPSCs) were characterized by immunocytochemical analyses. Neural progenitor cells were generated and patch clamp recordings were performed for a functional analysis of derived neuronal cells. Filipin stainings and the Amplex Red assay were used to demonstrate and quantify cholesterol accumulation.

Results: The hiPSCs expressed different stem cell markers, e.g. Nanog, Tra-1-81 and SSEA4. Using the embryoid body assay, the cells were differentiated in cells of all three germ layers and induced teratoma in immunodeficient mice, demonstrating their pluripotency. In addition, neural progenitor cells were derived and differentiated into functional neuronal cells. Patch clamp recordings revealed voltage dependent channels, spontaneous action potentials and postsynaptic currents. The accumulation of cholesterol in different tissues is the main hallmark of NPC1. In this study we found an accumulation of cholesterol in fibroblasts of a NPC1 patient, derived hiPSCs, and neural progenitor cells, but not in cells derived from fibroblasts of a healthy individual. These findings were quantified by the Amplex Red assay, demonstrating a significantly elevated cholesterol level in cells derived from fibroblasts of a NPC1 patient.

Conclusions: We generated a neuronal model based on induced pluripotent stem cells derived from patient fibroblasts, providing a human in vitro model to study the pathogenic mechanisms of NPC1 disease.

Keywords: Niemann-Pick Type C1, Induced pluripotent stem cells, Lysosomal storage disorder, Neural progenitor cells, Neural differentiation
Background

Human induced pluripotent stem cells (hiPSCs) feature three major advantages in the field of stem cell research. First, cells can be obtained by reprogramming different somatic cells [1-4] without raising ethical concerns, as it is the case with embryonic stem cells. Second, the pluripotent potential of the cells offers the opportunity to differentiate them into each cell of the body, e.g. motor neurons [5], cardiomyocytes [6], pancreatic insulin-producing cells [7], or male germ cells [8]. Third, iPSCs and subsequently differentiated cells have the same genetic information as the donor cells. Different diseases have already been modeled by using human iPSCs, e.g. Parkinson disease [9], metabolic liver disorders [10], retinal degeneration [11], Huntington disease [12], and mucopolysaccharidosis type IIIB, a fatal lysosomal storage disorder [13], and have been successfully utilized e.g. in drug screening [14].

Taken together, these characteristics of the cells are excellent prerequisites to model diseases in vitro. However, no in vitro model for Niemann-Pick disease Type C1 (NPC1) based on hiPSC cells is currently available. NPC1 is a rare progressive neurodegenerative disease caused by mutations in the NPC1 gene located on chromosome 18q11 encoding for a 1278-amino acid intracellular membrane glycoprotein [15-17]. It is inherited in an autosomal recessive manner and shows a prevalence of 1:120,000 live births [18]. A mutation in the NPC1 gene leads to an impaired lipid transport and sequestration resulting in e.g. a cholesterol accumulation in the late endosome and lysosome [19]. The clinical manifestation varies from neonatal icterus and hepatosplenomegaly in early childhood, cerebellar ataxia, seizures, gelastic cataplexy, and vertical supranuclear palsy in adolescence, to progressive neurological degradation, psychosis, and dementia in adulthood [18]. The symptoms are diverse and show intrafamilial variability [18,20].

The pathogenic mechanisms ultimately leading to a massive degeneration and loss of neurons in the CNS, especially Purkinje cells in the cerebellum, are not exactly understood. Most of our knowledge regarding NPC1 is based on cell models like human fibroblasts [21-23] and animal models like mouse [24], cat [25], and fruit fly [26]. Studies using these models have neither revealed the mechanisms leading to the selective massive degeneration of neurons nor found drugs, which can efficiently halt disease progression. Although the function of NPC1 in lipid trafficking is evolutionary highly conserved [27], the widely used murine BALB/c NPC1 model [28] cannot exactly reproduce human pathology. For example, neurofibrillary tangles composed of tau protein, which are seen in human NPC1 neurons, are absent in this model reflecting obvious biochemical and physiological differences [29]. Thus, studies utilizing disease-specific human neurons hold great promise to significantly increase our knowledge in understanding the pathological mechanism leading to massive neuronal degeneration. Recently, a human neuronal NPC1 model was reported based on multipotent adult stem cells [30].

In our study, we generated patient-specific induced pluripotent stem cells from a NPC1 patient and a healthy individual. The hiPS cell lines were differentiated into neural progenitor cells and subsequently differentiated into functional neurons to gain a human neuronal model of NPC1 disease.

Methods

Cell culture

Human dermal fibroblast cell lines GM18436 and GM05659 (Coriell Institute for Medical Research, Camden, USA) were obtained by skin biopsies from one-year old male Caucasian donors. GM18436 exhibits compound heterozygous mutations in the NPC1 gene (c.1628delC and GLU612ASP), representing a frameshift mutation and a missense mutation, respectively. The mutations lead to a non-functional protein as demonstrated by cholesterol esterification assay [31]. Fibroblast cell line GM05659 is obtained from a healthy donor. In the following cells of the cell line GM18436 will be referred to as mutNPC1 and cells of the cell line GM05659 will be referred to as wtNPC1. Cells were cultivated in fibroblast medium containing DMEM high glucose, 10% FBS and 1% Penicillin/ Streptomycin. Mitotically inactivated mouse embryonic fibroblasts (GlobalStem, Rockville, USA) for increased plating efficiency. HiPS cells were cultured on a feeder cell layer in iPSC medium containing DMEM/ F12, 20% knockout serum replacement, 1% Penicillin/ Streptomycin, 1% GlutaMAX, 1% MEM non essential amino acids, 0.2% 2-mercaptoethanol, and 10 to 15 ng/ ml hFGF-2 (Globalstem, Rockville, USA). hiPS cells on matrigel (BD Biosciences, Heidelberg, Germany) were cultured in mTESR1 medium (Stemcell Technologies, Grenoble, France). Medium was changed daily and cells were passaged weekly using 10 μM ROCK inhibitor Y-27632 (Stemgent, Cambridge, USA) for increased plating efficiency.

HiPS cells growing on a feeder cell layer were split mechanically weekly using pulled glass hooks by performing the cut and paste technique. Cells growing on matrigel were harvested enzymatically using 1 mg/ml dispase (Stemcell Technologies, Grenoble, France) for 7 min and large bore tips to break down large clumps according to manufacturer’s recommendations. HEK293FT cells (Invitrogen, Darmstadt, Gernany) used to obtain the viral vectors were cultivated in fibroblast
medium without Penicillin/ Streptomycin. All cells were cultivated at 37°C in a saturated humidity atmosphere containing 5% CO₂.

**Generation of retroviruses**

Retroviral pMIG vectors containing the cDNA of the human genes Oct4, Sox2, Klf4, and c-Myc were used as described recently [32]. Briefly, 3×10⁶ HEK293FT cells per 10 cm-dish were seeded onto ten dishes and incubated overnight. A solution containing 2.5 μg retroviral vector encoding for GFP and one of the transcription factors (Sox2, Klf4, Oct4, or c-Myc) was incubated with 0.25 μg VSV-G and 2.25 μg Gag-Pol in X-tremeGENE9 (Roche, Mannheim, Germany)/ DMEM High Glucose mixture (1:4) which was added to each of the dishes. Medium was renewed after 18 h and cells were incubated further for 48 h. Subsequently, the virus-containing medium was collected and passed through a 0.45 μm filter. To concentrate the virus, the medium was centrifuged at 70.000 × g at 4°C for 90 min, resuspended in 0.1 to 1 ml DMEM medium, and stored at −80°C. All four vectors contained a GFP sequence thus enabling titering by determining the percentage of GFP positive HEK293FT cells using FACS. Therefore, 1×10⁵ HEK293FT cells were seeded per 12-well in Penicillin/ Streptomycin free fibroblast medium containing 5 μg/ml protamine sulfate and concentrated virus in the following volumes: 6.25 μl, 12.5 μl, 25 μl, and 50 μl. After 48 h cells were washed with PBS containing Ca²⁺/ Mg²⁺, trypsinized and centrifuged for 5 min at 500 × g. Pellet was resuspended in 100 μl PBS without Ca²⁺/ Mg²⁺ and fixed by adding 100 μl of 4% paraformaldehyde for 15 min. Afterwards, the percentage of GFP positive cells was determined via FACS analysis.

**Transduction of human fibroblasts**

For transduction, 1×10⁶ fibroblasts were seeded per cavity of a 6-well plate and cultured for 18 h in fibroblast medium without Penicillin/ Streptomycin. Afterwards, fibroblast medium without Penicillin/ Streptomycin was supplemented with a volume of retrovirus of Sox2, Oct4, Klf4 (corresponding to 70 – 80% infection efficiency), and c-Myc (corresponding to 40–50% infection efficiency) in the presence of 5 μg/ml protamine sulfate. Cells were incubated for 48 h. Subsequently, medium was aspirated and cells were washed twice with PBS containing Ca²⁺/ Mg²⁺. Transduced cells were trypsinized and reseeded onto a gelatin coated 6 cm-dish. The next day, medium was replaced with iPS medium supplemented with 0.5 mM valproic acid to further increase the efficiency of reprogramming. Medium was changed daily and valproic acid was omitted after seven days.

**Generation of hiPS cell lines**

Initial hiPS colonies were routinely observed after three to four weeks. For further cultivation, they were picked using a 100 μl pipette tip and a pulled glass hook. Single colonies were transferred to the cavities of a 24-well plate, coated with 0.1% gelatin and 45.000 feeder cells/ cm². After 4 to 7 days of proliferation hiPS colonies were mechanically divided into two to four pieces and further expanded. Within six weeks each single hiPS colony was expanded to obtain different clones.

**Karyotyping**

Karyotyping was performed by Giemsa Trypsin banding. In short, colonies were incubated with a colcemid solution (10 μg/ ml in HBSS) for three hours to arrest cells in metaphase. Cells were treated with trypsin (0.25%) and the enzymatic reaction was stopped with Ammoniam solution (Invitrogen, Darmstadt, Germany). Cells were centrifuged at 300 × g for 10 min and the pellet was resuspended in 4 ml hypotonic potassium chloride solution (5.62%). Cells were incubated for 5 min at 37°C and centrifuged at 300 × g for 10 min. The cells were resuspended and fixed in 5 ml glacial acetic acid and methanol (1:3) and subsequently centrifuged for 7 min at 350 × g. This step was repeated once. Finally, most of the supernatant was removed and cells were resuspended. Cell suspension was dropped onto cold slides and dried at 100°C for 1 h. Giemsa solution (5%) was added and incubated for 5 min. Slides were washed in distilled water two times, dried at room temperature and sealed with cover slips.

**Sequencing**

Genomic DNA of fibroblasts, iPS cells grown on matrigel or neural progenitor cells were isolated using AllPrep Kit (Qiagen, Hilden, Germany) according to the manufacturer’s recommendations. Exon regions were amplified using HotStart Taq (Qiagen, Hilden, Germany) as follows: 95°C for 15 min followed by 13 cycles of 94°C for 30 s, 66°C for 30 s with 1.5°C decrease/ cycle and 72°C for 20 s; followed by 8 cycles of 94°C for 30 s, 46.5°C for 30 s with 1°C increase/ cycle and 72°C for 20 s; followed by 13 cycles of 94°C for 30 s, 66°C for 30 s with 1.5°C decrease/ cycle and 72°C for 20 s; followed by 11 cycles of 94°C for 30 s, 54.5°C for 30 s with 1°C increase/ cycle and 72°C for 20 s. Products were purified using ExoSAP Kit (USB Europe GmbH, Staufen, Germany) according to manufacturer’s recommendations. Sequence analysis was performed on a 3130XL Genetic Analyzer (Applied Biosystems, Carlsbad, USA).

**Alkaline phosphatase staining**

HiPSCs were cultivated on a feeder cell layer for five days. Medium was removed, cells were washed with PBS.
and fixed with ice-cold methanol (100%) for 10 min at −20°C. Methanol was removed and cells were washed with PBS. Subsequently, cells were incubated at room temperature for 15 min with the staining solution: 75% distilled water, 10% sodium chloride solution (1 M), 10% Tris solution (1 M, pH 9.8), 5% magnesium chloride solution (1 M), and NBT/BCIP solution (1:50, Roche, Mannheim, Germany). Staining solution was removed and cells were washed with distilled water. Microphotographs were taken using a Nikon Eclipse TS100 (Nikon, Düsseldorf, Germany).

Immunocytochemistry
Cells were fixed at room temperature for 15 minutes in 4% paraformaldehyde, washed with PBS and stored in 0.02% NaN₃ at 4°C. Immunocytochemistry was performed for Nanog (1:100, rabbit IgG polyclonal), Oct4 (1:100, rabbit IgG polyclonal), SSEA3 (1:100, rat IgM), SSEA4 (1:100, mouse IgG₂), Tra-1-60 (1:100, mouse IgG), Tra-1-81 (1:100, mouse IgM, all Stemgent, Cambridge, USA), Smooth muscle actin (SMA, 1:50, mouse monoclonal, Dako, Glostrup, Denmark), alpha-fetoprotein (alpha FP, 1:500, mouse monoclonal IgG, Sigma-Aldrich, Hamburg, Germany), Nestin (1:100, mouse monoclonal, R&D, Wiesbaden, Germany), MAP2ab (1:100, mouse monoclonal, Chemicon, Schwalbach, Germany), TuJ1 (1:100, mouse monoclonal Tu-20, Santa Cruz biotechnology, Heidelberg, Germany) and Sox-2 (1:200, rabbit monoclonal, Abcam, Cambridge, UK). Blocking and permeabilization was carried out using 0.3% Triton X-100 and 5% normal goat serum on ice. Directly before injection, cell suspension was resuspended in 0.25% trypsin/EDTA. After 1 min, the reaction was stopped by adding 2 ml of fibroblast medium and centrifuged again for 2 min at 200 × g. The pellet was resuspended in 1 ml of 0.25% trypsin/EDTA. After 1 min, the reaction was stopped by adding 2 ml of fibroblast medium and centrifuged again for 2 min at 200 × g. Cells were resuspended in 140 μl of cold DMEM/F12 and stored on ice. Directly before injection, cells were injected subcutaneously into the flank of the hind limb. After 8–12 weeks, when tumors were clearly visible, the animals were sacrificed and tumors were removed. Tumor tissue was fixed in 4% formalin for 12 to 18 hours and embedded in paraffin for subsequent staining.

H&E staining of tumor sections
4 μm thick tumor tissue sections were deparaffinized in xylol for 10 min and a descending ethanol concentration for 5 min each. Afterwards, the sections were washed in distilled water and stained with Mayer’s hematoxylin (Merck, Darmstadt, Germany) for 1 min. The slides were washed again twice and then dehydrated using an ascending ethanol concentration and xylol. Slides were mounted in Mowiol-DABCO. Microphotographs were taken with a Biozero 8000 microscope system (Keyence, Hamburg, Germany).

Neural differentiation
To differentiate hiPS colonies into neural direction, the colonies were cut, transferred to Poly-L-ornithine...
(15 μg/ml)/ laminin (10 μg/ml) coated dishes, and cultivated for 10 days in medium consisting of Neurobasal, DMEM/F12, 1xN2 supplement, 1xB27 supplement, GlutaMAX (2 mM) complemented with mouse recombinant noggin Fc-chimera (500 ng/ml, R&D, Wiesbaden, Germany), SB431542 (20 μM, Sigma-Aldrich, Hamburg, Germany) and hFGF-2 (5 ng/ml, GlobalStem, Rockville, USA). Neural rosettes were manually isolated using pulled glass hooks, gently trypsinized, and seeded as single cells on Poly-L-ornithine/ laminin coated dishes in medium consisting of Neurobasal, DMEM/F12, 1×N2, 1×B27, and GlutaMAX (2 mM) supplemented with hFGF-2 (10 ng/ml) and hEGF (10 ng/ml, Peprotech, Hamburg, Germany). Neural progenitor cells were seeded at high densities (100–150,000 cells/cm²) and passed one day after reaching confluence using Trypsin/ Benzonase. Differentiation was induced by seeding the cells at a density of 50,000 cells/cm² and withdrawal of growth factors in the presence of BDNF (20 ng/ml, Peprotech, Hamburg, Germany).

Patch clamp recordings
Patch clamp recordings were performed using an EPC-10 amplifier (Heka, Lambrecht, Germany). Patch pipettes were pulled from borosilicate glass tubing (Harvard Apparatus, Holliston, USA). The internal solution contained (mM): KCl 130, NaCl 10, HEPES 10, EGTA 11, MgCl₂×6H₂O 1, CaCl₂×H₂O 2, Mg-ATP 2. pH was adjusted to 7.2. When filled, electrodes had a resistance of 6–8 MΩ. Cell cultures were continuously superfused with an extracellular solution, consisting of (mM): NaCl 125, KCl 2.5, CaCl₂×H₂O 2, MgCl₂×6H₂O 1, NaHCO₃ 26, NaH₂PO₄×H₂O 1.25, glucose×H₂O 25. Solution was continuously bubbled with carbogen to maintain a pH of 7.4. Recordings were made in the whole cell configuration with holding potentials (VH) of −60 or −80 mV. Current voltage responses were evoked by applying 100 ms voltage steps from −60 mV to +50 mV in 10 mV increments. Data were filtered at 3 kHz, digitized and stored on-line using Pulse (Heka, Lambrecht, Germany). Na⁺ and K⁺ currents were identified via their I-V relationship. Na⁺ currents were antagonized in some experiments with TTX (1 μM). Current clamp mode was used to apply current steps to induce action potentials or to measure spontaneous action potentials. Postsynaptic currents were measured in the voltage clamp mode at a VH of −60 mV. Mini Analysis 6 (Synaptosoft, USA) was used to analyse recordings of post-synaptic currents. Data are given as mean ± SEM.

Filipin staining
Filipin is a polyene antibiotic which binds to free cholesterol and is widely used to analyze the sequestration of unesterified cholesterol in NPC1-deficient cells. Therefore, cells were fixed with 4% paraformaldehyde (in PBS) for 15 min, washed with PBS and incubated at room temperature for 45 min in the dark with a staining solution containing 100 μg/ml Filipin (Polsciences, Eppelheim, Germany) in PBS. Cells were washed twice with PBS for 5 min. Slides were mounted using Mowiol-DABCO and sealed with cover slips. After 12 h of drying in the dark at room temperature, fluorescence pictures were taken with a Biozero 8000 microscope (Keyence, Hamburg, Germany).

Amplex red assay
To quantify the amount of cholesterol we used the Amplex Red cholesterol assay [33]. Fibroblasts, iPS cells grown on matrigel and neural progenitor cells were harvested in PBS/ SDS (0.1%) at room temperature, sheared through a 27 G needle and cholesterol levels were determined using the Amplex Red cholesterol assay kit (Molecular Probes, Darmstadt, Germany) according to the manufacturer’s instructions. Protein concentrations in lysates were measured using the bicinchoninic acid assay (BCA, Pierce, USA).

Statistical analysis
Analysis of the data was carried out with GraphPad Prism 5 (GraphPad Software Inc., USA). Data are given as mean ± SEM. Unless otherwise stated, unpaired t-tests were used to test for significance, with * = p<0.05 and ** = p<0.01, *** = p<0.001.

Results
Reprogramming of mutNPC1 and wtNPC1 fibroblasts
We reprogrammed fibroblasts originating from a male patient with an early-infantile form of NPC1 characterized by massive accumulation of unesterified cholesterol in lysosomal and late endosomal structures. Cells derived from this donor will be referred to as mutNPC1 and cells derived from age- and sex-matched fibroblasts of a healthy individual will be referred to as wtNPC1.

After three to four weeks of cultivation, the first hiPSC colonies appeared characterized by their embryonic stem (ES) cell-like morphology, e.g. round to oval shape with a sharp border and a high nuclear to cytoplasm ratio (Figure 1A-D). Mechanically isolated colonies were expanded to hiPSC lines on irradiated mouse embryonic fibroblasts and later also on matrigel (Figure 1C,D). The morphology of mutNPC1 and wtNPC1 hiPSCs was similar in both culture systems. The karyotype of the cells was analyzed to rule out any chromosomal abnormalities, which may have arisen during reprogramming, where our hiPSCs displayed a normal karyotype (Figure 1E,F). Sequencing of the hiPSCs revealed that the mutations in the NPC1 gene were maintained (data not shown).
Pluripotency of mutNPC1 and wtNPC1 hiPSCs

HiPSCs derived from of mutNPC1 and wtNPC1 human fibroblasts were characterized regarding their pluripotency. First, we analyzed the alkaline phosphatase (AP) expression. All hiPSCs colonies demonstrated strong AP expression (Figure 2A,B). The expression of several transcription factors and surface markers was determined by immunocytochemistry. HiPSCs displayed a high expression of the transcription factors Nanog (Figure 2C,D) and Oct4 (Figure 2E,F). The glycosphingolipids SSEA3 and SSEA4 (Figure 2G-J), were strongly expressed as well as the keratan sulfate antigens Tra-1-60 and Tra-1-81 (Figure 2K-N). No obvious differences between mutNPC1 and wtNPC1 cells in marker expression could be observed. The spontaneous differentiation by embryoid body (EB) formation into cells of all three germ layers was also used to verify the pluripotency (Figure 3A-H). Herein, cells from all three germ layers were identified, thus proving the pluripotency of the hiPSCs in vitro. The induction of teratoma was used as an in vivo pluripotency assay. The hiPSCs induced teratomas in immunodeficient mice, and the analysis of the tumors revealed tissues of all three germ layers (Figure 3I-N).

Neuronal differentiation of mutNPC1 and wtNPC1 hiPSCs

In a last step, we generated neural progenitor cells, which were positive for Nestin and Sox2 (Figure 4A,B,E,F). Differentiated neural progenitor cells expressed neuronal markers like MAP2 (Figure 4C,G), and Tuj1 (Figure 4D,H) demonstrating the neuronal phenotype of the cells. Furthermore, we proved the differentiation into functional neuronal cells by means of patch clamp recordings. In these experiments we observed voltage dependent Na+ and K+ channels (NaV, and Kv) (Figure 4I) after three to four weeks of differentiation, where inward currents could be blocked by TTX (Figure 4K). Although the cells exhibited NaV, they did not demonstrate any
Figure 2 Detection of pluripotency markers in hiPSCs.

hiPSCs-derived from fibroblasts of NPC1 patient (mutNPC1) and unaffected control (wtNPC1) showed a strong alkaline phosphatase (AP) staining (A,B) and expressed a set of pluripotency markers (C-N) demonstrating the pluripotent state of the cells (scale bars = 100 μm).

Figure 3 Embryoid body formation and teratoma induction.

Free floating embryoid bodies (EBs) of mutNPC1 and wtNPC1 hiPSCs were generated (A,B), which subsequently spontaneously differentiated (C-H). EBs of both cell lines were positive for markers of the three germ layers, namely Nestin (C,D) for ectoderm, smooth muscle actin (SMA) (E,F) for mesoderm, and alpha fetoprotein (alpha FP) (G,H) for endoderm. Nuclei were stained by DAPI, shown in blue, staining of Nestin, SMA and alpha FP is shown in green (scale bars A-H = 100 μm). HiPSCs of the mutNPC1 and wtNPC1 cell lines were injected in immunodeficient mice and tumors were extracted after 8–12 weeks. MutNPC1 and wtNPC1 hiPSCs induced teratomas contained structures typical for the ectoderm (I,J), mesoderm (K,L), and endoderm (M,N; scale bars I-N = 50 μm).
spontaneous action potentials in the current clamp mode. But, we observed spontaneous action potentials after 7–8 weeks of differentiation (Figure 4L). In addition, we recorded spontaneous postsynaptic currents. An example of a mutNPC1 cell is shown in Figure 4M. The analysis of the decay kinetics (Figure 4N) revealed a rise time of 2.1 ± 1.1 ms. Analysing the decay kinetics we found a group of post synaptic currents best fitted by a mono-exponential function (τ: 3.5 ± 0.4 ms, Figure 4N, black trace) with a mean amplitude of 29.5 ± 1.1 pA, and a group best fitted
by a bi-exponential function (τ1: 9.9 ± 1.9 ms, τ2: 124.7 ± 23.8 ms, Figure 4N, red trace) with a mean amplitude of 68.1 ± 7.1 pA.

**mutNPC1 cells accumulated cholesterol**
The hallmark of NPC1 is abnormal cholesterol trafficking resulting in an accumulation of cholesterol. Free cholesterol can be visualized by Filipin. An analysis of the cholesterol distribution in mutNPC1 fibroblasts, iPSCs, and derived neural progenitor cells (Figure 5A,C,E) revealed an accumulation of cholesterol. In contrast, an accumulation was not detectable in the fibroblasts, iPSCs, or neural progenitor cells of the wtNPC1 counterpart (Figure 5B,D,F). As a next step, we used the Amplex Red assay [33] to confirm and quantify the observed cholesterol accumulations. The experiments revealed a significantly increased cholesterol content in mutNPC1 cells in comparison to wtNPC1 cells (Figure 5G) (fibroblasts: mutNPC1: 13.7 ± 0.5 μg vs. wtNPC1: 7.3 ± 0.3 μg; iPSCs: mutNPC1: 11.6 ± 0.6 μg vs. wtNPC1: 8.9 ± 0.7 μg; neural progenitor cells: mutNPC1: 23.6 ± 0.9 μg vs. wtNPC1: 14.2 ± 1.6 μg).

**Discussion**
In this study we aimed to reprogram fibroblasts originating from a NPC1 patient with an early-infantile form of

![Figure 5](image-url)
the disease. Therefore, we used retroviruses expressing Oct4, Klf4, Sox2, and c-Myc in combination with GFP. These factors have been described previously to be efficient in generating hiPSCs [34]. The retroviral particles used in this study were successfully used to reprogram skin fibroblasts of Parkinson’s disease into hiPSCs [32]. HiPSC colonies were chosen based on an absent GFP-signal indicating a silenced expression of transcription factors [35] and were subcultured to stable hiPSC lines. The obtained mutNPC1 and wtNPC1 hiPSC lines were characterized by their ES-cell like morphology, the expression of alkaline phosphatase, and the pluripotency markers Nanog, Oct4, SSEA3, SSEA4, Tra-1-60, Tra-1-81. A comparable expression of Tra-1-81 was reported for NPC1 knock-down and control ES cells [36]. However, comparative analyses of additional pluripotency markers were not performed in this study. We found no obvious differences between mutNPC1 and wtNPC1 cells in pluripotency marker expression. These results are in accordance with other studies dealing with patient-specific induced pluripotent stem cells in a variety of other diseases [9]. However, to our knowledge this is the first study describing the expression of pluripotency markers SSEA3, SSEA4, and Tra-1-60 in pluripotent cells harboring disease-causing mutations in the NPC1 gene. In addition, the widely known risk of chromosomal abnormalities, potentially occurring during iPS generation and expansion [37], did not arise in our cell lines as proved by karyotyping. The spontaneous differentiation by embryoid body (EB) formation [38] into cells of all three germ layers and the induction of teratomas in immunodeficient mice [39] further demonstrated the pluripotent state of the mutNPC1 and wtNPC1 hiPSCs.

We further differentiated the hiPSCs into neural progenitor cells to generate a suitable in vitro disease model. So far, two human cellular neural models based on NPC1 knockdown have been reported. These include SH-SY5Y neuroblastoma cells [40] and human embryonic stem cells [36], which resemble the phenotype only in some aspects of the NPC1 disease. Therefore, they are not an appropriate model to analyze the influence of specific mutations in a patient-specific (epi)genetic background. Here, we generated homogenous neural progenitor cells based on mutNPC1 and wtNPC1 hiPSCs, which were positive for neural markers Nestin and Sox2 [41]. In contrast, Ordonez et al. [36] obtained a homogenous population of neural stem cells from control hESCs but not from NPC1 knock-down hESCs. The authors speculate that these findings might be based on the genetic background of the cells [36].

Our differentiated neural progenitor cells expressed the neuronal markers MAP2 and TuJ1. We did not observe obvious morphological differences between mutNPC1 and wtNPC1 neuronal cells. In contrast, distortion of neuronal shape and extensive growth of ectopic neurites have been reported for multipotent adult stem cells (MASCs) derived cells [30]. However, a further detailed analysis of our neural progenitor cells and derived neuronal cells will be performed to analyze changed morphology of neuronal cells and perturbances of proliferation, described for murine neural stem cells [42]. In a first set of experiments, we demonstrated the differentiation of neural progenitor cells into functional neuronal cells by means of patch clamp recordings. We observed voltage dependent Na+ and K+ channels, where inward currents were blocked by TTX. Spontaneous action potentials and postsynaptic currents could only be observed in cells differentiated for 7–8 weeks, indicating the maturation time of human neural progenitor cells to functional neurons, as described for interneurons derived from hiPSCs [43,44]. The here recorded spontaneous postsynaptic currents displayed different amplitudes and the time constants of the current decay could be fitted with mono-exponential or bi-exponential functions. The differences between the time constants may indicate different types of synaptic input, where currents with small amplitudes and fast mono-exponential decay suggest excitatory and events with larger amplitudes and slow bi-exponential decay suggest inhibitory input [45,46]. These preliminary results indicate that the differentiated cells are able to build up chemical synapses. This is of special interest as recent studies described disturbed transmitter release in NPC1 deficient mice, where a higher rate of glutamate release was observed leading to higher frequency of excitatory postsynaptic currents [47]. Thus, our cells provide a platform to study such alterations in synaptic transmission in human neuronal cells gained from different individuals. Ultimately, these results demonstrate a maturation into functional neuronal cells, where future studies will focus on the nature of the expressed voltage and ligand gated ion channels in mutNPC1 and wtNPC1 neuronal cells.

Our neural progenitor cells were analyzed regarding their impaired cholesterol trafficking by Filipin. It visualizes free cholesterol and is routinely used for human dermal fibroblasts in the diagnostics of the NPC1 disease [20]. We found clear cholesterol accumulation in mutNPC1 fibroblasts, hiPSCs, and derived neural progenitor cells. In contrast, such an accumulation was not observed in the fibroblasts, hiPSCs, or neural progenitor cells of the wtNPC1 counterpart. The accumulation pattern of cholesterol in the herein described cells was comparable to accumulations described in a NPC1 knock-down mouse model [48], and SH-SY5Y neuroblastoma cells [40]. Recently, a neural model based on multipotent adult stem cells was described [30]. The neural differentiated progeny of these cells, demonstrated a comparable accumulation of cholesterol, where
this derivation method is only applicable to early passages (<3 passages) of fibroblasts, potentially limiting its use with characterized cell lines from cell repositories.

Finally, we used the Amplex Red assay to confirm and quantify the observed cholesterol accumulations in our cells. These experiments revealed significantly increased cholesterol content in mutNPC1 cells in comparison to wtNPC1 cells, which was conserved in fibroblasts, hiPSCs, and derived neural progenitor cells. To our knowledge this is the first analysis of cholesterol distribution using Filipin staining and cholesterol quantification in hiPSCs and neural progenitor cells derived from human NPC1 deficient fibroblasts.

Conclusion

In this study we generated, for the first time, induced pluripotent stem cells derived from fibroblasts of a NPC1 patient. The cells demonstrated an accumulation of cholesterol, resembling the phenotype of NPC1 deficient cells, and can provide an in vitro model of NPC1. We are convinced that the here reported hiPSCs and the derived neural progenitor cells are an excellent model to study the influence of the specific mutation on the phenotype, e.g. consequences of a misfolded NPC1 protein. Moreover, the cells provide the opportunity to analyze the consequences of a NPC1 mutation on the patient-specific (epi)genetic background, and will thus serve to elucidate further the pathogenic mechanisms of this fatal lysosomal storage disorder.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

MT: conception and design, collection and / or assembly of data, data analysis and interpretation, manuscript writing. RH: conception and design, collection and / or assembly of data, data analysis and interpretation, manuscript writing. PS: collection and / or assembly of data, manuscript writing. AR: conception and design, manuscript writing, final approval of manuscript. MJF: conception and design, collection and / or assembly of data, data analysis and interpretation, manuscript writing. CK: collection and / or assembly of data, manuscript writing. AR: conception and design, collection and / or assembly of data, data analysis and interpretation, manuscript writing. PS: collection and / or assembly of data, manuscript writing. MT: conception and design, collection and / or assembly of data, data analysis and interpretation, manuscript writing. AR: conception and design, manuscript writing, final approval of manuscript. MJF: conception and design, collection and / or assembly of data, data analysis and interpretation, manuscript writing, final approval of manuscript. All authors read and approved the final manuscript.

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