Induced pluripotent stem cell-derived motor neurons from amyotrophic lateral sclerosis (ALS) patients carrying different superoxide dismutase 1 mutations recapitulate pathological features of ALS

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Abstract

Background: Investigations of the pathogenic mechanisms in motor neurons (MNs) derived from amyotrophic lateral sclerosis (ALS) disease-specific induced pluripotent stem (iPS) cell lines could improve understanding of the issues affecting MNs. Therefore, in this study we explored mutant superoxide dismutase 1 (SOD1) protein expression in MNs derived from the iPS cell lines of ALS patients carrying different SOD1 mutations.

Methods: We generated induced pluripotent stem cell (iPSC) lines from two familial ALS (FALS) patients with SOD1-V14M and SOD1-C111Y mutations, and then differentiated them into MNs. We investigated levels of the SOD1 protein in iPSCs and MNs, the intracellular Ca²⁺ levels in MNs, and the lactate dehydrogenase (LDH) activity in the process of differentiation into the MNs derived from the controls and ALS patients’ iPSCs.

Results: The iPSCs from the two FALS patients were capable of differentiation into MNs carrying different SOD1 mutations and differentially expressed MN markers. We detected high SOD1 protein expression and high intracellular calcium levels in both the MN and iPSCs that were derived from the two SOD1 mutant patients. However, at no time did we observe stronger LDH activity in the patient lines compared with the control lines.

Conclusions: MNs derived from patient-specific iPSC lines can recapitulate key aspects of ALS pathogenesis, providing a cell-based disease model to further elucidate disease pathogenesis and explore gene repair coupled with cell-replacement therapy. Incremental mutant expressions of SOD1 in MNs may have disrupted MN function, either causing or contributing to the intracellular calcium disturbances, which could lead to the occurrence and development of the disease.

Keywords: Amyotrophic lateral sclerosis; Induced pluripotent stem cell; SOD1 gene mutation; Motor neuron; SOD1 aggregation toxicity of misfolded SOD1 in familial ALS (FALS).

Introduction

Amyotrophic lateral sclerosis (ALS) is a rapidly progressing, fatal, neurodegenerative disease for which no effective treatment exists. The exact ALS pathogenesis remains unclear, but recent multiple molecular events and genetic discoveries have provided important insights that implicate common pathological processes across the spectrum of ALS. Superoxide dismutase 1 (SOD1) has been one of the most studied mutations owing to the early discovery of causative gene and the widest applications to murine models. Considerable evidence over recent years emphasizes that SOD1 mutations cause the disease via one or more toxic properties. Protein aggregation, mitochondrial dysfunction, and Ca²⁺ dysregulation were related to the progression of pathogenic discoveries, there were no therapies that succeeded in translation of experimental observations into the clinic [3]. In part, this reflects lack of appropriate human cell-based models to demonstrate the disease mechanisms and identify promising therapeutics before long and expensive clinical trials. Human induced pluripotent stem cell (iPSC) systems with naturally occurring human pathology could complement existing murine models, potentially facilitating clinical translation and bridging this gap. Our and others’ researches have previously reported that motor neurons (MNs) derived from ALS-iPSC partially elucidate the mechanisms of mutant protein-related ALS disease [6-9].

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It has recently been reported that the presence of misfolded proteins, specifically those causing protein aggregation, may disrupt several intracellular mechanisms and trigger neurotoxicity in ALS.[10] These protein aggregates exhibit intense immunoreactivity with antibodies against SOD1; they develop before the onset of the clinical disease, and in some cases, represent the earliest sign of the disease. Remarkably, as we learned using models of mutant SOD1-mediated ALS, mitochondria compromised in ALS patients is obvious from several studies that have been conducted using cellular or animal models.[11] ALS-SOD1 alters the mitochondrial protein composition and decreases the protein import into mitochondria, which then comprises part of the mitochondrial damage.[12] Moreover, mitochondrial abnormalities disturb the calcium homeostasis, as the higher intracellular calcium levels lead to possible mitochondrial Ca\(^{2+}\) overload.[13] It is important to assess whether these changes in animal models also can occur in iPSC, including the targeted types that are mainly implicated in the disease, namely MNs and astrocytes.

In the present study, we reprogrammed skin fibroblast cells into iPSCs from two FALS patients with SOD1-V14M and SOD1-C111Y mutations and from control subjects. Then, these iPSCs were differentiated into MNs. Next, we investigated the levels of the SOD1 protein in both iPSCs and MNs and measured intracellular Ca\(^{2+}\) levels in the MNs. Analysis of these assays in iPSC-derived neural cells from ALS patients carrying different mutations provided insight into the convergence of the cellular and molecular mechanisms in different familial types of ALS, highlighting the importance of protein aggregation and calcium dysregulation in ALS.

Methods

Ethical approval

The study was performed in accordance with the Helsinki Declaration of 1975, as revised in 2000 (5). It was approved by the Ethical Committee of Peking University Third Hospital (IRB00006761-L-2010055). Written informed consent was obtained from all participants.

Participants

The clinical data were collected and the SOD1 gene mutations were screened from our FALS database. Then, the selected probands with SOD1-V14M and SOD1-C111Y mutations were analyzed. All ALS patients met the diagnosis of the El Escorial revised criteria.[14] Additionally, skin fibroblasts were donated from four age-matched and sex-matched healthy controls of Han Chinese descent with no previous personal or family history of neurodegenerative disease, which were used to generate iPSCs. The p.V14M mutation was observed in a 21-year-old woman, who started with weakness of the lower limb and the course of the disease lasted about 109 months. The patient carrying the p.C111Y mutation was a 70-year-old female diagnosed with ALS. She reported a 20-month history of progressive upper limb muscle weakness and the course of the disease lasted about 48 months. She died of respiratory failure.

Derivation of patient-specific fibroblast cells and mutation detection

The SOD1-V14M and SOD1-C111Y mutations were detected using polymerase chain reaction (PCR) and direct sequencing of genomic DNA. We isolated 3 mm dermal explants by a skin punch biopsy of the two FALS patients with the mutations SOD1-V14M and SOD1-C111Y. The mutation fibroblast outgrowth from the explants were passaged with trypsin, and then frozen for 1 to 2 weeks. A TIANamp Genomic DNA Kit (Qiagen, Valencia, CA, USA) was used to extract the patients’ genomic DNA; genomic sequencing of the patients’ DNA was performed by the BGI Genomics Co., Ltd. (Shenzhen, Guangdong, China).

Cell culture

Human fibroblast cells were cultured in fibroblast medium (FM) (Dulbecco’s modified Eagle’s medium [DMEM]; Gibco, Carlsbad, CA, USA), supplemented with 10% fetal bovine serum (FBS) (HyClone, Logan, UT, USA), 1 mmol/L glutamine, and 1% penicillin/streptomycin (Chemicon InternationalInc, Temecula, CA, USA). Human embryonic stem (ES) cells and iPSCs were cultured in a standard human embryonic stem cell (hESC) medium as described previously.[8,15] The human iPSC derivation medium was the same as the human iPSC culture medium, except that the concentration of fibroblast growth factor beta (bFGF) (PeproTech Inc, Cranbury, NJ, USA) was 10 ng/mL.

Retroviral production and iPSC cell generation

We used a previously described protocol to generate iPSCs[16] Retroviruses (human octamer-binding transcription factor 4 (OCT4), SRY (sex determining region Y)-box 2 (SOX2), Kruppel-like factor 4 (KLF4), and myelocytomatosis cellular oncogene (c-MYC) were introduced into fibroblast cells. After 48 h of transfection, medium containing virus was collected and concentrated by centrifugation for 2 h at 22,000 rpm, 4 °C. The viral pellet was re-suspended in 1 mL FM and used to infect 5 × 10^4 fibroblast cells in a well of a 12-well plate. After 8 to 12 h of infection, we replaced the fresh FM with infected fibroblast cells. After 3–4 days, the cells were trypsinized, and the mitomycin C (MMC)–treated Mouse embryonic fibroblast (MEF) cells were seeded in 100 cm culture dishes. The medium we used was human iPSCs derivation medium. After 3 to 4 weeks, iPSC colonies appeared, and these were mechanically passed every 3 to 4 days.

Assessment of iPSCs pluripotency

Pluripotency of iPSCs was examined using previously protocols.[8] We confirmed pluripotency ALS-iPS and control-iPSCs by expression of pluripotency markers stage-specific embryonic antigen-4 (SSEA-4), tumor-related antigen-1-60 (TRA1-60), tumor-related antigen-1-81 (TRA1-81), Nanog, OCT3/4, SOX2 and reverse transcription PCR (RT–PCR) with three germ-layer differentiation verified by reduced expression 1 (REX1), developmental pluripotency associated protein 4 (DPPA4), Nanog, and OCT4 expression. We also performed bisulfite...
treatment of genomic DNA and karyotype analysis. Finally, all clones were subcutaneously injected into the groin of severe combined immunodeficiency mice to carry out an assessment based on their ability to form teratoma in vivo.

**Immunofluorescence staining**

The cells growing on slides were fixed in 4% paraformaldehyde (PFA) for 30 min, and then permeabilized with 0.5% TritonX-100 for 15 min. Slides were blocked in aldehyde (PFA) for 30 min, and then permeabilized with secondary antibodies (AlexaFluor®, Invitrogen, Carlsbad, California, USA) for 1 h and incubated in primary antibodies overnight at 4°C, and subsequently incubated with secondary antibodies (AlexaFluor®, Invitrogen, Carlsbad, California, USA) for 1 h at room temperature. Imaging was performed using a Leica confocal microscope (Leica TCS SP8 MP, Chicago, IL). The primary antibodies included: SSEA-4 (1:500; Millipore, Billerica, MA, USA), TRA1-60 (1:500; Millipore, Billerica, MA, USA), TRA1-81 (1:500; Millipore, Billerica, MA, USA), Nanog (1:500; Cosmobio, Tokyo, Japan), OCT3/4 (1:500; Santa Cruz Biotechnology, Santa Cruz, CA, USA), SOX2 (1:500; Santa Cruz Biotechnology, Santa Cruz, CA, USA), TUJ1 (1:1000; Covance, Princeton, NJ, USA), HB9 (1:100; Developmental Studies Hybridoma Bank (DSHB), Iowa City, IA, USA), and ISL1/2 (1:200; DSHB, Iowa City, IA, USA).

**MN differentiation from iPSCs**

We differentiated ALS-iPS and control-iPSCs into MNs based on methods that were slightly modified from previous ones.[6,7,18] Briefly, iPSCs were placed into ultra-low adherent culture dishes and treated with embryoid body (EB) medium (DMEM/F12, 2% B27 and 1% N2; Life Technologies, Carlsbad, California, USA), 10 μmol/L SB-431542 (Sigma-Aldrich, St. Louis, MO, USA) and, additionally, with 1 μmol/L retinoic acid (Sigma-Aldrich, St. Louis, MO, USA) and 50–100 ng/mL smoother agonist (SHH; PeproTech Inc, Cranbury, NJ, USA) at a particular time. On the tenth day of EB formation, these colonies were plated onto poly-DL-ornithine and laminin (PO/LAM)-coated (Sigma-Aldrich, St. Louis, MO, USA) plates. To allow for the maturation of MNs, after 20 days of differentiation, the cultures were dissociated into single cells with papain (Sigma-Aldrich, St. Louis, MO, USA) and seeded onto PO/LAM-coated slides in Neurobasal® medium, containing 2% B27 and 1% N2, with 10 ng/mL of brain-derived neurotrophic factor (BDNF; R&D Systems, Minneapolis, MN, USA) and glial cell line-derived neurotrophic factor (GDNF; R&D Systems, Minneapolis, MN, USA).

**Lactate dehydrogenase (LDH) assays for neurons**

ALS and control lines could be differentiated in parallel to enable direct comparison. Over the 13 days of differentiation, the cell culture medium was collected once a week for the 13 days of differentiation from the SOD1-V14M (L1 and L6) and SOD1-C111Y (Y3 and Y8), and control lines (C3 and C12; D1). LDH activity (mU/mL) was calculated for each cell type using LDH assay kits (Abcam, Cambridge, UK). LDH activity was designed against the differentiation date, and linear models were generated by comparisons between the cell types.

**Western blot for SOD1 expression in iPSCs and MNs**

MN cells in the 26 to 28 days of differentiation were lysed in the radio-immunoprecipitation assay (RIPA) buffer with protease inhibitors cocktail (Roche, Basel, Switzerland). Thirty micrograms of protein was separated by 12% Tris-glycine sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE), and then transferred to nitrocellulose membranes and probed with anti-SOD1 (1:2000; Abcam) and anti-α-Tubulin (1:1000; Sigma-Aldrich, St. Louis, MO, USA). The densitometric analysis was performed using a CDP-Star chemiluminescent detection system (Applied Biosystems, Foster City, CA, USA).

**Measurement of intracellular calcium (Ca2+) in MNs**

Intracellular Ca2+ generation was assessed using the Ca2+-specific fluorescent dye Fluo 3-AM (Beyotime, Beijing, China). The MNs, plated on confocal dishes, could reach optimal confluence in the 28th day of differentiation, at which time they were washed three times with PBS and loaded with 5 μmol/L Fluo 3-AM at 37°C for 45 min. Then they were washed with PBS, and observed under a Leica confocal microscope (Leica TCS SP8 MP, Chicago, IL). Fluo 3-AM was observed at a wavelength excitation of 488 nm.

**Statistical analysis**

All of the data was reported as mean ± standard deviation of the three independent experiments, and were analyzed by one-way analysis of variance (ANOVA) and Student’s t test. Significance level was set at P < 0.05. Semiquantitative immunofluorescence analysis of MMP and intracellular calcium was performed with Leica Application Suite X (LAS X) software.

**Results**

**Generation of ALS-iPSCs from fibroblast cells of FALS patients with SOD1-V14M and SOD1-C111Y mutations**

We obtained dermal explants by biopsy from the two FALS patients and established their fibroblast cell lines [Figure 1A]. We recruited a total of six individuals into this study: two patients diagnosed with FALS carrying the p. V14M mutation and p.C111Y mutations in the SOD1 gene; and four healthy individuals with no history of neurological disease (CONT). Patient-derived fibroblast cells were infected with the four Yamanaka factors—KLF4, SOX2, OCT3/4, and c-MYC.[19-21] Approximately 20 days after infection, several colonies appeared with morphology similar to that of hESCs [Figure 1B]. At least 4–6 iPSC differentiations were performed for each line, but 2–4 lines per patient among total lines were thoroughly characterized and shown to be fully reprogrammed to pluripotency. Based on this analysis, 1–2 lines from each
carrier were selected for further characterization: 4 lines (from four healthy individuals,) 2 from SOD1-V14M member, and 2 from SOD1-C111Y member.

**Pluripotency of iPS cell lines**

We identified whether the induced cells had the characteristics of typical hESCs. We focused only on the exhaustive characterization of four clones, though we originally isolated four iPSC clones from the SOD1 mutant cells and four iPSC clones from the controls. Pluripotency results of the iPSC lines show that they contain tissue of all three germ layers, such as intestinal epithelium, cartilage, and keratinized epithelium; scale bar = 100 μm. (f) The karyotyping results of V14M-iPS-6 and C111Y-iPS-3 indicate that these cells are normal 46, XX human karyotype. (g) Quantitative RT-PCR results show that the endogenous pluripotent genes’ expression levels of OCT4, NANOG, DPPA4, and REX1 were similar to human ES cells, which is much higher than their initial fibroblast cells. (h) Both iPSC cell lines express human ES cells’ nuclear markers, such as OCT3/4, SOX2, and NANOG, and are also positive for the cell-surface antigens SSEA4, TRA-1-60, and TRA-1-81; scale bar = 50 μm. (i) Bisulfite genomic sequencing of NANOG promoters of these two iPSC cell lines. The white and black circles represent the unmethylated and methylated CpGs, respectively. For each iPSC cell line, ten samples were sequenced, and each row indicates a repeat. SOD1: Superoxide dismutase 1; ALS: Amyotrophic lateral sclerosis; iPSC: Induced pluripotent stem; RT-PCR: Reverse Transcription-Polymerase Chain Reaction; ES: Embryonic stem.
this was not observed in their initial fibroblasts [Figure 1I]. In summary, this data demonstrated that fibroblasts were successfully reprogrammed into a pluripotent state and that the two SOD mutations in iPSCs did not disturb with pluripotency.

Neural differentiation of ALS patient-derived iPSCs

To verify whether there were obvious defects in the MNs generated from the two SOD1-iPSCs, we induced MN differentiation based on a previously published method with slight modifications.[8,17] Schematic protocol for the differentiation process is illustrated in Figure 2A. Floating cultivation of EBs was detected at day 6 [Figure 2B]. After 14 days of differentiation, we found neural, progenitor-like outgrowths from the plated EBs that interacted with each other [Figure 2B]. After 28 days of differentiation, MN-like cells were found in our cultures [Figure 2B]. Immunofluorescence co-staining was performed with neuron-specific class III b-tubulin (TUJ1), HB9, and ISL1 to confirm whether that these cells were MNs. As a result, we found that the HB9-positive cells or ISL1-positive cells co-expressed the MN-specific markers Tuj1 [Figure 2C]. Interestingly, we found no significant difference in the number of ISL1, Tuj1, and HB9 between patients and controls at day 28 (Figure 2D, Tuj1/HB9: \( F = 0.7390, P > 0.05 \); HB9/4′,6-diamidino-2-phenylindole[DAPI]: \( F = 0.2740, P > 0.05 \); Figure 2E, Tuj1/ISL1 \( F = 0.0847, P > 0.05 \); ISL1/DAPI: \( F = 0.4593, P > 0.05 \); Tukey’s Multiple Comparison Test, one-way ANOVA). Thus, our results confirmed the similar ability of control and ALS iPSC to give rise to MNs.

Comparable viability of neurons derived from all iPSC lines

Having established the equivalent MN cultures from both ALS-iPSC lines and the control-iPSC, we investigated if there were any differences in cell viability between the neurons derived from those from the ALS-iPSCs and the control group. We made quantitative analyses of cell viability using the cellular release of LDH. Analysis of LDH activity in SOD1-V14M and SOD1-C111Y showed no greater LDH activity in the patients at any time during the 4 weeks in culture, compared with the control lines [Figure 3A; \( P < 0.05 \)]; however, the level of LDH activity in the differentiation process decreased [Figure 3A].

SOD1 protein levels with specific mutation in iPSCs and MNs

A cytoplasmic SOD1 mutant inclusions are a likely key pathological feature of ALS.[12] Thus, we next detected SOD1 protein levels by Western blotting, and discovered that the MNs and iPSCs in SOD1-V14M and SOD1-C111Y exhibited higher SOD1 levels than the levels of the controls (Figure 3B; Figure 3C: \( F = 28.5100, P < 0.05 \), Tukey’s Multiple Comparison Test, one-way ANOVA; Figure 3D: \( F = 50.2700, P < 0.05 \), Tukey’s Multiple Comparison Test, one-way ANOVA).

Intracellular Ca\(^{2+}\) levels in MNs with specific mutation

It remains unclear whether the different SOD1 mutant that was linked to mitochondrial dysfunction occurs in ALS pathophysiology. Mitochondrial dysfunction is associated with calcium dysregulation.[13] In the present study, as expected, ALS-MNs had significantly higher intracellular Ca\(^{2+}\) than the controls’ (Figure 3E and F: \( F = 215.4000, P < 0.05 \), Tukey’s Multiple Comparison Test, one-way ANOVA). Similarly, there was a difference in the SOD1-V14M and SOD1-C111Y mutations (Figure 3E; \( P < 0.05 \), Tukey’s Multiple Comparison Test). Taken together, we found that mitochondrial dysfunction occurs in ALS patients with the different SOD1 mutant.

Discussion

Disease models are indispensable tools for elucidating disease mechanisms which can be used to identify novel therapeutics. As limited availability of primary patient cells in small quantities, transgenic animals or transformed cell lines were usually used. Unlike conventional models, iPSCs, derived from patients with genetic mutation naturally occurring in ALS patients, are likely to manifest disease phenotypes. Whereas, there are still some problems that hinder utilization of disease models using human ES cells, particularly owing to the genetic variation among patients and the heterogeneity of target neural types for epigenetic changes. To overcome genetic variation, we generated iPSCs from ALS patients with different mutations (SOD1-V14M and SOD1-C111Y), and then differentiated these iPSCs into MNs damaged in ALS. Further, we used a patient-specific iPSC model to recapitulate the pathology of SOD1-related disorders. Similar to other iPSC studies of carrying ALS-associated mutations, none of the two different SOD1 mutations affected the differentiation of ALS-iPSCs into MNs.

Various ALS-SOD1 mouse models and FALS patients have been analyzed previously.[24-26] In our study, MNs and iPSCs from patients carrying different mutations exhibited much higher SOD1 protein levels compared to the controls’. However, we did detect cytoplasmic mislocalization and the formation of FUS-immuno-positive inclusions aggregates in the MNs differentiated from ALS patient-specific iPSCs carrying the FUS-P525L mutation.[8] This result is contrary to previous reports, in which the SOD1 level is not increased in human ALS MNs, alike to the unchanged SOD1 level in spinal cord homogenates from ALS patients.[27] However, the exact mechanism of ALS patients with lower SOD1 level in human ALS MNs compared to previously reported in mouse models presents similar phenotypes that is still not known.

Next, we performed metabolic assays to monitor the intracellular Ca\(^{2+}\). As expected, the MNs had significant changes in intracellular Ca\(^{2+}\) assays than the controls’. Mitochondria plays a crucial role in Ca\(^{2+}\) signaling in MNs.[28] Additionally, SOD1 proteins can disrupt fundamental Ca\(^{2+}\) signaling pathways in MNs, and Ca\(^{2+}\) itself can directly or indirectly impact many ALS-related proteins and cellular processes.[4] Mitochondrial dysfunction and damage occur in both SALS and FALS. Yet, it is still unknown whether there is a causal relationship between mitochondrial alteration and mutant proteins for recently discovered ALS-linked mutations such as FUS,
Figure 2: Differentiation of human iPS cells into motor neurons. (A) Overview of human iPS cell differentiation into MNs. EB medium (DMEM/F12, containing 2% B27 and 1% N2, supplemented with 1% NEAA, 200 nmol/L dorsomorphin, 10 μmol/L SB-431542), RA retinoic acid, low and high SHH 50-100 ng/mL sonic hedgehog, PO/LAM, neurobasal medium (2% B27 and 1% N2, with 10 ng/mL BDNF and GDNF). (B) The representative image of morphology of EBs at day 6, neural progenitors at day 14, and motor neuron-like cells at day 28 of controls (control cell line: cell line C3) and ALS patients (SOD1-V14M: cell line: L6); scale bar 1.00 mm. (C) Immunostaining of control and patient-specific MN cultures; motor neurons co-express TUJ1/HB9 and TUJ1/ISL1, nuclei are counterstained with DAPI, shown in blue; scale bar = 50 μm. (D) Quantitative analyses of cells co-express positive for TUJ1/HB9. Bars represent average with SEM as error bars. Data for CONT is the average of 4 iPSC lines, for ALS from 4 iPSC lines. No significant differences were found in the ability of iPSC to generate neurons after 28 days of differentiation. (E) Quantitative analyses of cells co-express positive for TUJ1/ISL1, IPS: Induced pluripotent stem; MNs: Mechanisms in motor neurons; EB: Embryoid body; BDNF: Brain-derived neurotrophic factor; GDNF: Gliarial cell line-derived neurotrophic factor; SEM: Standard error of mean; CONT: Healthy individuals with no history of neurological disease; PO/LAM: Poly-DL-oromithine and laminin; NEAA: Non-essential amino acid; DAPI: 4', 6-diamidino-2-phenylindole.
TDP-43, and chromosome 9 open reading frame 72 (C9ORF72); however, this has been studied in great detail with mutations. IPSC-derived MNs exhibit disturbances in mitochondrial morphology and motility, which is dependent on the presence of the SOD1A4V mutation.\(^{29}\) Mutant SOD1 aggregates interacted with Bcl-2 in spinal cord mitochondria, affecting mitochondrial function, and ultimately led to cell death.\(^{30}\) Cytoplasmic aggregates of SOD1 might inhibit conductance of voltage-dependent anion channel (VDAC1), decreasing the supply of ADP to the mitochondria for ATP synthesis, which induces mitochondrial dysfunction.\(^{11,31}\) Moreover, the mitochondria in both mouse and human spinal cords with SOD1 mutations increased susceptibility to oxidative stress and structural damage, which ultimately led to the release of cytochrome c.\(^{11,32}\) The heterogeneity of genetic and phenotype, as well as the failure to identification of a candidate therapeutic reinforce the opinion that ALS is a multifactorial neurodegenerative disease. Also, the analysis needs to contain information as to whether the multiple manifestations of toxicity of mutant proteins in SOD1 iPSC-derived neurons are linked to a particular mutation, since in our study multiple lines and patients need to be analyzed.

In summary, we obtained iPSC lines from two FALS patients with SOD1-V14M and SOD1-C111Y mutations. These ALS-iPSC lines are pluripotent and capable of differentiation into MNs. Increased SOD1 protein, and calcium dysregulation, were observed in SOD1 iPSC-derived neurons. Multiple manifestations of the toxicity of mutant proteins in SOD1 iPSC-derived neurons could facilitate the identification of useful combination therapies for further testing.

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Conflicts of interest

None.

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