α-Synuclein oligomers induce early axonal dysfunction in human iPSC-based models of synucleinopathies

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α-Synuclein (α-Syn) aggregation, proceeding from oligomers to fibrils, is one central hallmark of neurodegeneration in synucleinopathies. α-Syn oligomers are toxic by triggering neurodegenerative processes in vitro and in vivo models. However, the precise contribution of α-Syn oligomers to neurite pathology in human neurons and the underlying mechanisms remain unclear. Here, we demonstrate the formation of oligomeric α-Syn intermediates and reduced axonal mitochondrial transport in human neurons derived from induced pluripotent stem cells (iPSC) from a Parkinson’s disease patient carrying an α-Syn gene duplication. We further show that increased levels of α-Syn oligomers disrupt axonal integrity in human neurons. We apply an α-Syn oligomerization model by expressing α-Syn oligomer-forming mutants (E46K and E57K) and wild-type α-Syn in human iPSC-derived neurons. Pronounced α-Syn oligomerization led to impaired retrograde axonal transport of mitochondria, which can be restored by the inhibition of α-Syn oligomer formation. Furthermore, α-Syn oligomers were associated with a subcellular relocation of transport-regulating proteins Miro1, KLC1, and Tau as well as reduced ATP levels, underlying axonal transport deficits. Consequently, reduced axonal density and structural synaptic degeneration were observed in human neurons in the presence of high levels of α-Syn oligomers. Together, increased dosage of α-Syn resulting in α-Syn oligomerization causes axonal transport disruption and energy deficits, leading to synapse loss in human neurons. This study identifies α-Syn oligomers as the critical species triggering early axonal dysfunction in synucleinopathies.

α-synuclein | oligomers | axonal transport | synucleinopathies | neurodegeneration

Physiologically, α-synuclein (α-Syn) is a presynaptic protein, which, when mutated or modified, is causative for a group of neurodegenerative disorders termed “synucleinopathies.” The pathological feature of synucleinopathies is α-Syn aggregates that evolve from small oligomers to fibrillar aggregates and accumulate in intracytoplasmic and intraneuritic deposits called Lewy bodies (LB) and Lewy neurites (LN), respectively (1). The presence of LB and LN is the major hallmark of synucleinopathies. For example, a widespread cortical Lewy pathology is evident in the majority of Parkinson’s disease (PD) cases carrying an α-Syn gene duplication (Dupl) (2). Although fibrillar α-Syn is the main component of LB and LN, α-Syn oligomers have been recently reported to characterize early stage lesions of synucleinopathies in PD patients’ postmortem brains (3), emphasizing their role in disease onset by yet largely unknown mechanisms.

In rodents, α-Syn overexpression and oligomerization resulted in axonopathy, neuritic and synaptic degeneration preceding neuronal death (4–8). Considered to be early pathogenic events of synucleinopathies, they might be attributed to cytoskeletal defects. Indeed, α-Syn oligomers specifically inhibit tubulin polymerization (9, 10), and α-Syn accumulation accompanies disruption of microtubules in cell lines (11, 12). We previously revealed a critical role of α-Syn oligomers for kinesin-microtubule motility in a cell-free system (13). Furthermore, impaired microtubule-dependent trafficking was associated with α-Syn aggregates in zebrafish and rodent neuronal models (11, 14, 15). Notably, alterations of axonal transport proteins were reported in neurons with α-Syn inclusions in postmortem PD brains (16) and in rat striatum and substantia nigra upon overexpression of the human A53T mutant α-Syn (8). Of note, oligomers of amyloid-β, a crucial protein for neurodegeneration in Alzheimer’s disease, selectively impaired α-Syn function.

Significance

α-Synuclein (α-Syn) aggregation underlies neurodegeneration in synucleinopathies. However, the nature of α-Syn aggregates and their toxic mechanisms in human pathology remains elusive. Here, we delineate a role of α-Syn oligomeric aggregates for axonal integrity in human neuronal models of synucleinopathies. α-Syn oligomers disrupt anterograde axonal transport of mitochondria by causing subcellular changes in transport-regulating proteins and energy deficits. An increase of α-Syn oligomers in human neurons finally results in synaptic degeneration. Together, our data provide mechanistic insights of α-Syn oligomeric toxicity in human neurons. Taking into account that α-Syn oligomers and axonal dysfunction are characteristic for early neurodegeneration in synucleinopathies, our data might deliver targets for therapeutic interference with early disease pathology.
fast transport of mitochondria (17) and induced neuritic dystrophy (18), suggesting a particular toxicity of oligomeric protein assemblies to neurite function. However, the specific role of α-Syn oligomers in the regulation of axonal transport and involvement of axonal transport aberrations in human synucleinopathies remains elusive.

In the present study, we analyze the impact of α-Syn oligomers generated by an α-Syn dosage increase on axonal transport in human neurons. First, we demonstrated an impaired axonal transport of mitochondria accompanied by increased α-Syn aggregation and formation of α-Syn intermediates in neurons derived from induced pluripotent stem cells (iPSC) of a PD patient with α-Syn Dupl. To better understand the impact of α-Syn aggregation, we established a model that allows the investigation of distinct α-Syn species in human iPSC-derived neurons by introducing wild-type human α-Syn (WTS) or its oligomer-forming mutants (E46K and E57K). Specific transport of anterograde mitochondrial transport was observed upon pronounced α-Syn oligomerization in E46K and E57K neurons and was partly rescued by inhibiting α-Syn oligomer formation. α-Syn oligomer-induced transport alterations were accompanied by reduced ATP levels, a pathologic subcellular relocation of proteins regulating axonal transport, and an apparent synaptic degeneration. In summary, we demonstrate a link between α-Syn oligomers and dysfunctional mitochondrial axonal transport with a consequent synaptic degeneration in human neurons as an early phenotype of synucleinopathies.

Results
Reduction of Mitochondrial Axonal Transport and Enhanced α-Syn Aggregation in Human Neurons Carrying an α-Syn Dupl. To investigate whether a cell-autonomously elevated intracellular level of α-Syn affects axonal transport in human neurons, we generated iPSC from fibroblasts of a PD patient carrying an α-Syn Dupl and from healthy individuals [controls (Ctrls); Fig. 1A]. Ctrls and Dupl iPSC differentiated equally well into neurons based on expression of the neuronal marker β3-tubulin (Tubb3, green). (Scale bar: 100 μm.) (C) Elevated α-Syn protein expression in Dupl neurons (three NPC lines: 1.1 and 1.2 from the Dupl1 iPSC clone and 2.1 from the Dupl2 iPSC clone) compared with Ctrls (three iPSC clones from two individuals: 1.1 and 1.2 from Ctrl1 and 2.1 from Ctrl2) by Western blot (WB). (D) Neurons were differentiated in microfluidic devices for 20 d, and transport of Mito-DSRed-positive mitochondria was measured (Experimental setup). Representative kymographs of Ctrl and four Dupl lines: three NPC lines as representative Ctrl and Dupl lines. Increase of insoluble α-Syn (P3 fraction) in Dupl neurons was observed (Fig. 1B). 

We next asked whether axonal transport alterations in Dupl neurons occur coincidently with α-Syn aggregation. We assessed α-Syn solubility in Dupl neurons by sequential extraction of proteins first with detergent-free buffer (soluble fraction S1) and subsequently with buffers containing detergents of increasing solubilization capacity, resulting in Triton X-100–soluble (S2), SDS-soluble (S3), and urea/SDS-soluble (P3) fractions (Fig. 1H). An increase of α-Syn in the P3 fraction and thus a decreased solubility of α-Syn in Dupl neurons was observed (P < 0.01 for the P3 fraction, Fig. 1H), indicating an ongoing α-Syn aggregation in these neurons. A further assessment of the composition of α-Syn aggregates by sucrose density gradient centrifugation using a calibration with preformed characterized α-Syn species (19) revealed an increased signal in several fractions enriched for α-Syn aggregation intermediates in Dupl neurons (red boxes, Fig. 1I). These results imply a tight link between axonal transport alterations of mitochondria and enhanced formation of early α-Syn intermediate species in human neurons.

increased α-Syn Oligomerization Coincides with Impaired Anterograde Axonal Transport. Aggregation of α-Syn is a multistep process involving different aggregation states with intermediate oligomeric species as initial steps. To understand the early stage pathology and to dissect the toxic nature of α-Syn intermediate species in human neurons, we aimed to decipher the effect of α-Syn oligomers on axonal transport. We thus modeled the process of α-Syn...
aggregation by introducing WTS, the familial α-Syn mutant E46K, and the nonnaturally occurring oligomer-prone E57K α-Syn mutant (6) into Ctrl iPSC-derived neurons by lentiviral (LV) infection (Fig. 2A). Mito-ΔsRed was encoded by each LV vector to visualize mitochondria (Fig. 2A). The efficiency of LV infection was similarly high under all conditions (97–100%, SI Appendix, Fig. S1A), resulting in comparable expression levels of WTS and α-Syn mutants in human iPSC-derived neurons (SI Appendix, Fig. S1B). Sequential protein extraction revealed a significant increase of α-Syn levels in the less- and insoluble fractions (S2, S3, and P3) in α-Syn mutants, in particular E57K, compared with WTS (P value of the interaction between α-Syn variants and relative fraction size ≤ 0.001, Fig. 2B), pointing toward a strong aggregation process of α-Syn in E46K- and E57K-expressing neurons. To further verify the presence of α-Syn oligomers, we performed size exclusion chromatography (SEC) calibrated with preformed oligomeric species of recombinant α-Syn to assign α-Syn monomers and oligomers of different sizes to SEC fractions. This analysis showed significantly enhanced levels of α-Syn in the fraction enriched for trimers in E46K and E57K neurons compared with WTS (Fig. 2C). Thus, the expression of E46K and E57K in human neurons increases the proportion of small α-Syn oligomers.

In conditions of increased α-Syn oligomerization, we observed clustering of thickened and short mitochondria in E46K and E57K neurites compared with thin and elongated mitochondria within control neurites (asterisk, Fig. 2D). A mixture of short and, partly clustered mitochondria was found in WTS neurons (Fig. 2D). The mitochondrial phenotype observed in E46K and E57K neurons coincided with significantly decreased amounts of ATP measured in these neurons, but not in WTS neurons (Fig. 2E).

Furthermore, α-Syn oligomerization propensities were translated into profound differences in mitochondrial axonal transport in human neurons (representative kymographs, Fig. 2F). Specifically, the movement incidence was significantly reduced in WTS, E46K, and E57K neurons (Fig. 2G), whereas the presence of α-Syn oligomers significantly changed the directionality of movement toward decreased frequency of anterograde transport in E46K and E57K neurons (Fig. 2I). Together, while increased dosage of α-Syn due to either genetic or LV overexpression results in an overall reduction of mitochondrial movement frequency, α-Syn oligomers specifically decrease the anterograde axonal transport accompanied by reduced ATP levels.

Interfering with α-Syn Oligomer Formation Improves Mitochondrial Axonal Transport. To better link α-Syn oligomers and dysfunctional mitochondrial anterograde transport, we used the de novo-developed compound NPT100-18A that reduces toxicity in PD models by interfering with α-Syn oligomer formation (20, 21). NPT100-18A restored mitochondrial movement incidence in WTS, E46K, and E57K neurons (Fig. 2H and SI Appendix, Fig. S2A) and improved movement directionality toward increased anterograde motility without influencing retrograde transport frequency in α-Syn mutant neurons (Fig. 2J and SI Appendix, Fig. S2B).

In line with impaired movement incidence, expression of WTS, E46K, and E57K led to significantly reduced numbers of mitochondria within axons compared with Mock, while the mitochondrial numbers within axons were increased by NPT100-18A in E46K and E57K, but not in WTS neurons (SI Appendix, Fig. S2C). Accordingly, impaired maximal anterograde velocities were increased by NPT100-18A in E46K and E57K, but not in WTS neurons (Fig. 2K, Left).

Furthermore, reduced mean velocities and decreased frequencies of fast moving (>0.3 μm/s) mitochondria in anterograde direction were detected in E46K and E57K neurons compared with Mock (Fig. 2K and SI Appendix, Fig. S2D, respectively). Whereas mean anterograde velocity was improved only in E57K neurons (SI Appendix, Fig. S2D), NPT100-18A significantly reduced the amount of slow-moving (0.1 μm/s), increased the amount of middle-speed (0.3 μm/s), and restored to a control level the amount of fast-moving (>0.3 μm/s) anterograde mitochondria in E46K and E57K neurons (Fig. 2K, Middle).

In WTS neurons, NPT100-18A inhibited the rate of slow-moving and increased the rate of middle-speed mitochondria, but did not influence the numbers of fast-moving anterograde mitochondria, which were not altered compared with Mock in

![Figure 2](https://example.com/figure2.png)

**Fig. 2.** Increased α-Syn oligomerization and impaired anterograde axonal transport in iPSC-derived neurons expressing α-Syn mutants. (A) Control LV- and -ΔsRed-infected neurons (20 d of differentiation) were infected with WTS or α-Syn mutant EViK (E46K or E57K) and Mito-ΔsRed lentivirus. (B) An increase of insoluble α-Syn (Triton X-100-soluble [S2], SDS-soluble [S3], and urea/SDS-soluble [P3] fractions) was observed in E57K and E46K neurons. S1, a detergent-free soluble fraction. (C) Soluble α-Syn multimers were analyzed by size exclusion chromatography. Soluble α-Syn trimers (fr) were significantly increased in E46K and E57K lysates compared with WTS. D, dimers; HMWD, higher-molecular-weight oligomers; M, monomers; T, tetramers. (D) Clustering (asterisks) and shortening of mitochondria in neurites with α-Syn oligomers (E46K, E57K) compared with elongated mitochondria in control neurons (Mock). Scale bar: 500 nm. (E) ATP levels were significantly reduced in neurons expressing the α-Syn mutants E46K and E57K. (F) Representative kymographs. (G) Less frequently moving mitochondria were found in all α-Syn overexpressing neurons and (H) can be restored by NPT100-18A. Positive delta indicates an increase in negative delta indicates a decrease of a value in NPT100-18A compared with DMSO (diluent control in all NPT100-18A experiments). (I) Neurons expressing α-Syn mutants E46K and E57K revealed a lower incidence of anterograde axonal transport, which was significantly increased by NPT100-18A. (J) Decreased maximal (max) anterograde velocities in WTS, E46K, and E57K neurons, which were improved by NPT100-18A in α-Syn mutant neurites (Left). NPT100-18A reduced slow-moving (0.1 μm/s) and increased middle-speed (0.3 μm/s) anterograde mitochondria frequencies in WTS, E46K, and E57K neurites (Middle). Significant increase of fast-moving (>0.3 μm/s) anterograde mitochondria by NPT100-18A was observed in E46K and E57K neurites (Middle). No changes of retrograde axonal transport were detected independently of NPT100-18A usage (Right). *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001, ****P ≤ 0.0001. Data are presented as mean ± SD in H, J, and K and as mean ± SEM in C. Data from two independent iPSC clones from Ctrl1 are presented collectively in H, J, and K.
the absence of the compound (Fig. 2K). Mean anterograde velocity revealed only a slight, not significant, reduction in WTS compared with Mock neurons and was not affected by NPT100-18A (SI Appendix, Fig. S2D). No changes of retrograde mitochondrial axonal transport and no influence of NPT100-18A on this transport were found (Fig. 2K and SI Appendix, Fig. S2D, Right). Of note, the highest efficiency of NPT100-18A in improving anterograde mitochondrial transport in α-Syn oligomer-prone mutant neurons was accompanied by its ability to reduce insoluble α-Syn levels in these neurons, indicative of reduced α-Syn aggregation (SI Appendix, Fig. S2E).

Thus, the highest effectiveness of NPT100-18A toward anterograde axonal transport in neurons with strong α-Syn aggregation and higher levels of small oligomers (as evident in E46K and E57K neurons) indicates that small α-Syn oligomers have a significant effect on mitochondrial axonal transport.

α-Syn Oligomerization Is Associated with Alterations in Adaptor Motor Proteins. Axonal transport of mitochondria involves adaptor proteins that either facilitate transport by attaching mitochondria to the motor proteins via Mitochondrial Rho GTPase 1 (Miro1) and Kinesin Light Chain 1 (KLC1) or impair motility of axonal mitochondria by docking them to microtubules via Syntaphilin (SNPH) (22). In line with impaired movement incidence of mitochondria, neuritic levels of Miro1 were significantly lower compared with levels in the soma in WTS, E46K, and E57K neurons (Fig. 3A). Interestingly, SNPH levels in neurites, especially proximal at the soma, were significantly increased in E46K neurons (Fig. 3B). Notably, significant reduction of KLC1 in neurites over 50 μm in length was determined in α-Syn oligomer-containing E46K and E57K neurons (Fig. 3C), corresponding to a severe reduction of anterograde transport velocities in these neurons.

The axonal microtubule-associated protein Tau is known to selectively reduce kinesin-dependent transport (23). Interestingly, Tau protein expression was significantly increased in E46K and E57K neurons compared with control and WTS neurons with the highest level in the perinuclear regions (asterisks, Fig. 3D). The perinuclear regions of E46K and E57K neurons were also enriched for phosphorylated Tau (pTau) clumps (asterisks, Fig. 3F), indicating Tau pathology.

Thus, KLC1 reduction in neurites in conjunction with increased Tau pathology is specifically associated with impaired anterograde axonal transport and elevated levels of α-Syn oligomers.

α-Syn Oligomers Affect the Axonal and Synaptic Integrity in Human Neurons. α-Syn oligomer-associated transport and energy deficits might affect the axonal compartment and synapses in human neurons. Axonal fiber densities, visualized by Tubb3 staining, were reduced in E46K and E57K neurons compared with controls (Mock) (Fig. 4A and B).

Before analyses of the synaptic compartment, the maturity of differentiated neurons was confirmed by whole-cell patch clamp analysis showing similar firing behavior of the neurons for evoked action potentials (SI Appendix, Fig. S3A). Ultrastructural characterization of the synaptic compartment identified intact synapses containing presynaptic vesicles and postsynaptic densities (arrows) in Mock and WTS neuronal cultures (Fig. 4C). In contrast, barely detectable postsynaptic densities, irregular presynaptic membrane inclusions, and dystrophic synaptic structures were characteristic for E46K and E57K neuronal cultures (Fig. 4C). Consistently, the analysis of isolated synaptosomes (SI Appendix, Fig. S3B) determined reduced amounts of the presynaptic proteins: synapsin I, synaptosomal-associated protein 25 (SNAP25), and synaptophysin (SYN38) in synaptosomal preparations of E46K and E57K neuronal cultures compared with control (SI Appendix, Fig. S3 D and E). The synaptic nature of isolated synaptosomes was confirmed by their double positivity for presynaptic and postsynaptic markers, synapsin I, and postsynaptic density 95, respectively (SI Appendix, Fig. S3C). Interestingly, reduced levels of kinesin 1 were found in synaptosomal compartments of E46K and E57K cultures (SI Appendix, Fig. S3E). These data together reveal profound structural abnormalities of axonal and synaptic compartments in α-Syn oligomer-containing neurons that are in line with impaired anterograde axonal transport and ATP deficit in these neurons.

In conclusion, increased dosage of α-Syn impairs axonal transport due to the critical impact of α-Syn oligomers on axonal biology. Thus, we propose a model of neurodegeneration in synucleinopathies, where critical levels of α-Syn oligomers reduce anterograde axonal transport of mitochondria due to Tau pathology and redistribution of kinesin adaptor proteins. Transport deficits co-occur with morphological and functional alterations of mitochondria upon pronounced α-Syn oligomerization, finally leading to synaptic degeneration in human neurons (Fig. 4D).

Discussion

Aberrations of axonal transport have been speculated to contribute to neurodegeneration in PD and synucleinopathies (8, 16, 24). This study directly measures axonal transport in human PD-specific neurons in conjunction with elevated α-Syn levels and α-Syn aggregation intermediates. In iPSC-derived neurons of a PD patient with an α-Syn Dupl, mitochondrial axonal transport deficits were accompanied by increased formation of α-Syn aggregation intermediates. To define which of the α-Syn species has the deleterious effect on axonal transport, we co-expressed WTS overexpression with specific α-Syn oligomer-forming mutants. We identified small α-Syn oligomers as the crucial species involved in functional impairment of axons in human neurons. Transport defects could be recovered by inhibiting α-Syn oligomer formation. α-Syn oligomers promoted a subcellular relocation of transport-regulating proteins and resulted in energy...
Materials and Methods

Extended experimental procedures are described in SI Appendix, SI Materials and Methods. All vectors are listed in SI Appendix, Table S1, and antibodies are listed in SI Appendix, Table S2.

Cells and Cell Culture. Human iPSC from a PD patient with α-Syn Dupl were provided by Douglas Galasko, University of California, San Diego (clones SD1-R-C3 (Dupl1) and SD1-R-C11 (Dupl2)). This sample was from a female PD patient with a disease onset at the age of 58 with a progressive disease

Impaired transport of mitochondria is associated with axonal degeneration in mouse dopamine neurons (34, 35) and synaptic dysfunction (22). Indeed, four independent PD models revealed axonal transport deficits, impaired levels of presynaptic proteins (synapsin I, SNAP25, and SYN38) in synaptosomes, and ultrastructurally degenerated synaptic structures in neurons with the strongest disruption of mitochondrial axonal transport and pronounced α-Syn oligomerization. α-Syn overexpression and aggregation are known to lead to “vacant synapses” (36), and synaptic degeneration was found in a mouse model overexpressing the E57K mutant of α-Syn (5). Accordingly, the overexpression of human WTS or mutant α-Syn led to alterations of axonal transport proteins, axonopathies, and synaptic dysfunction preceding neuronal loss in rat brain in vivo (7, 8). In line with this, α-Syn oligomers were recently shown to cause synaptic impairment in rat primary neuronal cultures without inducing a prominent toxicity (37). Thus, α-Syn oligomers are the critical species promoting axonopathy and synaptic loss probably due to an impaired axonal transport.

Our study shows that impaired anterograde axonal transport of mitochondria was accompanied by reduced Miro1 and KLC1 levels in axons and a soma-proximal increase of Tau and pTau levels in α-Syn oligomer-containing neurons. Dysregulation of Miro1 caused mitochondrial transport abnormalities and impaired anterograde axonal transport, attributed to reduced abundance of Miro1 and KLC1 within axons accompanied by increased levels of SNPH and Tau pTau. These changes result in a profound degeneration of synapses.

deficits. Finally, increased levels of α-Syn oligomers resulted in a profound degeneration of synaptic structures.

The analysis of α-Syn aggregation in α-Syn Dupl neurons confirms and expands previous studies, which showed increased α-Syn expression or α-Syn accumulation in iPSC-based PD neuronal models carrying a triplication of α-Syn gene (Trip1) (25–28) or a leucine-rich repeat kinase 2 (LRKK2) mutation (29, 30). α-Syn oligomers and amyloid aggregates in α-Syn Trip1 neurons caused lysosomal dysfunction due to disrupted hydrolysis trafficking (28). Our report shows an increased formation of α-Syn aggregation intermediates and defective axonal transport of mitochondria in iPSC-derived neurons of a PD patient carrying an α-Syn Dupl. Our data indicate that elevated amounts of α-Syn and accumulation of α-Syn aggregation intermediates are pathologically connected with alterations of mitochondrial axonal transport.

A recent gene expression study revealed that iPSC-derived neurons most resemble fetal brain tissue, thereby suggesting particular suitability of iPSC-based models to study disease predisposition (31). Interestingly, α-Syn oligomer-induced mitochondrial motility changes and the formation of oligomers in α-Syn Dupl neurons were observed at early neuronal differentiation stages in our study. This might reflect abnormalities in developing neurons characterized by increased α-Syn oligomer levels. The developmental abnormalities in turn may determine their lower resilience to stressors during aging. Earlier human iPSC-based studies found significant alterations in mitochondrial motility in iPSC-derived neurons from sporadic and LRRK2 mutant PD patients (32, 33). In line with this, in animal models, α-Syn aggregation, caused by either α-Syn mutant or WTS overexpression or by extracellular α-Syn preformed fibrils, affected the expression level of axonal transport proteins and disrupted axonal transport of mitochondria or autophagosomes (8, 14, 15). While extracellular spreading of α-Syn aggregates may be a prominent mechanism of pathology propagation, cell-autonomous α-Syn aggregation is an initiating event in synucleinopathies. Therefore, our data emphasize that a disruption of axonal mitochondrial transport by endogenously formed α-Syn aggregation intermediates represents an early cellular phenotype in synucleinopathies.
course who developed dementia. iPSC from two healthy Caucasian in-
dividuals with no history of neurologic disease served as controls
Ctrl1,1, clone UKER1E4-R1-016; Ctrl1.2, clone UKER133Q-R1-106; and
Ctrl1.2, clone UKER1E4-R1-016, previously reported in ref. 42).
All experiments with human iPSC-derived cells were reviewed and
approved by the Institutional Ethics Review Board (Nr. 4120:
Generation of human neuronal models of neurodegenerative
diseases). Written in-
formed consent was received from the participants before inclusion in the
study at the movement disorder clinics at the Department of Molecular Neu-
rology, Universitätssklinikum Erlangen, Friedrich-Alexander-Universität
Erlangen-Nürnberg and at the Department of Neurosciences, University of
California, San Diego.
Materials and Data Availability. All antibodies and materials are commercially available. The source of vectors is provided in SI Appendix, Table S1. All experimen-
tal procedures are described in detail in SI Appendix, Materials and
Methods.

2. Konno T, Ross OA, Puschmann A, Dickson DW, Wszolek ZK (2016) Autosomal domi-
α-synuclein oligomers reveals previously undetected pathology in Parkinson’s
4. Winner B, et al. (2012) Role of α-synuclein in adult neurogenesis and neuronal mat-
5. Rockenstein E, et al. (2014) Accumulation of oligomer-prone α-synuclein exacerbates synaptic
axonal transport proteins combined with striatal neuroinflammation preceede dopa-
mineergic neuronal loss in a rat model of αVA synucleinopathy. J Neurosci 29:
3635–3673.
depolymerization potentiates alpha-synuclein oligomerization. Front Aging Neurosci 1:5.
13. Prots I, et al. (2013) α-Synuclein oligomers impair neuronal microtubule-kinesin inter-
15. Volpicielli-Daley LA, et al. (2014) Formation of α-synuclein Lewy neurite-like aggre-
gates in axons impedes the transport of distant endosomes. Mol Biol Cell 25:
4010–4023.
17. Rui Y, Zheng QJ (2016) Amyloid β-oligomers elicit mitochondrial transport defects and
fragmentation in a time-dependent and pathway-specific manner. Mol Brain 9:79.
able amyloid β-oligomers in culture and in vivo: Prevention by scyllo-inositol.
Neurabol Dis 8:152–163.
a human iPSC-based model of familial Parkinson’s disease. Proc Natl Acad Sci USA
114:E3639–E3688.
24. Lamberts JT, Hildebrandt EN, Brundin P (2015) Spreading of α-synuclein in the face of
axonal transport deficits in Parkinson’s disease: A speculative synthesis. Neurobiol Dis
77:276–283.
accumulate α-synuclein and are susceptible to oxidative stress. PLoS One 6:e26159.
pairs neuronal differentiation and maturation in Parkinson’s patient-derived induced
omal dysfunction occurs through disruptions in protein trafficking in human mid-
29. Nguyen HN, et al. (2011) LRK22 mutant iPSC-derived DA neurons demonstrate in-
human iPSC models of genetic and sporadic Parkinson’s disease. EMBO Mol Med 4:
380–395.
with schizophrenia. Mol Psychiatry 20:361–368.
32. Hsieh CH, et al. (2016) Functional impairment in miro degradation and mitophagy is
33. Cooper O, et al. (2012) Pharmacological rescue of mitochondrial deficits in iPSC-
derived neural cells from patients with familial Parkinson’s disease. Sci Transl Med
4:141ra90.
specifically impairs mitochondrial transport in dopamine axons. J Neurosci 31:
7212–7221.
and Parkinson-independent degeneration of respiratory chain-deficient dopa-
36. Scott DA, et al. (2010) A pathologic cascade leading to synaptic dysfunction in alpha-
trafficking of vesicles, mitochondria, and endoplasmic reticulum: Implications for
tion in models of Parkinson’s disease: A speculative synthesis. J Neurosci 31:
953–968.
8:267–280.