Ubiquitin-independent function of optineurin in autophagic clearance of protein aggregates

Jelena Korac¹, Veronique Schaeffer², Igor Kovacevic², Albrecht M. Clement³, Benno Jungblut⁴, Christian Behl³, Janos Terzic¹ and Ivan Dikic^{1,2,*}

¹Department of Immunology and Medical Genetics, School of Medicine, University of Split, Soltanska 2, 21000 Split, Croatia ²Institute of Biochemistry II and Buchmann Institute for Molecular Life Sciences, School of Medicine, Goethe University, Theodor-Stern-Kai 7, 60590 Frankfurt (Main), Germany

³Institute for Pathobiochemistry, University Medical Center, Johannes Gutenberg University, Duesbergweg 6, 55099 Mainz, Germany ⁴Department of Cardiac Development and Remodeling, Max-Planck-Institute for Heart and Lung Research, Parkstrasse 1, 61231 Bad Nauheim, Germany

*Author for correspondence (ivan.dikic@biochem2.de)

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Summary

Aggregation of misfolded proteins and the associated loss of neurons are considered a hallmark of numerous neurodegenerative diseases. Optineurin is present in protein inclusions observed in various neurodegenerative diseases including amyotrophic lateral sclerosis (ALS), Huntington's disease, Alzheimer's disease, Parkinson's disease, Creutzfeld-Jacob disease and Pick's disease. Optineurin deletion mutations have also been described in ALS patients. However, the role of optineurin in mechanisms of protein aggregation remains unclear. In this report, we demonstrate that optineurin recognizes various protein aggregates via its C-terminal coiled-coil domain in a ubiquitin-independent manner. We also show that optineurin depletion significantly increases protein aggregation in HeLa cells and that morpholino-silencing of the optineurin ortholog in zebrafish causes the motor axonopathy phenotype similar to a zebrafish model of ALS. A more severe phenotype is observed when optineurin is depleted in zebrafish carrying ALS mutations. Furthermore, TANK1 binding kinase 1 (TBK1) is colocalized with optineurin on protein aggregates and is important in clearance of protein aggregates through the autophagy-lysosome pathway. TBK1 phosphorylates optineurin at serine 177 and regulates its ability to interact with autophagy modifiers. This study provides evidence for a ubiquitin-independent function of optineurin in autophagic clearance of protein aggregates as well as additional relevance for TBK1 as an upstream regulator of the autophagic pathway.

Introduction

Accumulation of misfolded proteins and the associated loss of neurons are considered as a hallmark of many neurodegenerative diseases such as Parkinson's disease, Alzheimer's disease, Huntington's disease (HD), amyotrophic lateral sclerosis (ALS), Creutzfeld-Jacob disease, Pick's disease as well as several types of spinocerebellar ataxias (Aguzzi and O'Connor, 2010). Two main proteolytic systems are responsible for the degradation of proteins in the cell; the ubiquitin-proteasome system (UPS) and the autophagy-lysosomal pathway. Misfolded and/or damaged proteins are usually successfully degraded by the UPS. If the amount of proteins targeted to the UPS surpasses the degradation efficiency rate of the proteasome, subsequent oligomerization and accumulation of proteins activate autophagy (Tyedmers et al., 2010). The autophagy-lysosomal degradation machinery can also, to some extent, clear protein aggregates (Korolchuk et al., 2010).

Macroautophagy (hereafter referred to as autophagy) is an intracellular process responsible for the degradation of protein aggregates, long-lived proteins, defective organelles and intracellular pathogens (Dikic et al., 2010; Chen and Klionsky, 2011; Mizushima et al., 2011). During autophagy, a double-membrane vesicle (the autophagosome) engulfs the cargo and

subsequently fuses with the lysosome which degrades the cargo from the autophagosome. Over the last few years it has been shown that autophagic membranes can selectively target cargo via ubiquitin-like microtubule-associated protein light chain 3 modifiers [MAP1(LC3)] and gamma-aminobutyrate receptor associated protein (GABARAP), which act as receptor molecules (Kirkin et al., 2009a; Kraft et al., 2010). For example, autophagy of protein aggregates is coordinated by specific autophagy adaptor proteins p62 and NBR1, which simultaneously bind to polyubiquitinated protein aggregates and LC3 (Pankiv et al., 2007; Kirkin et al., 2009b). Recently, other autophagy adaptor proteins were reported, such as Alfy, Nix, NDP52, optineurin and FYCO1, all of which have been demonstrated to play a crucial role in autophagy of protein aggregates (Filimonenko et al., 2010), defective mitochondria (Novak et al., 2010) and bacteria Salmonella enterica (Thurston et al., 2009; Wild et al., 2011) or in the general movement of autophagosomes by binding to microtubules (Pankiv et al., 2010), respectively.

Optineurin (OPTN) is a 577 amino acid protein of versatile functions which interacts with a variety of proteins involved in membrane and vesicle trafficking (Rab8, huntingtin, myosin VI), cellular morphogenesis, NF- κ B regulation, signal transduction,

transcription regulation and cellular division control (Hattula and Peränen, 2000; Anborgh et al., 2005; Sahlender et al., 2005; Zhu et al., 2007; del Toro et al., 2009; Kachaner et al., 2012). Mutations in OPTN gene have been associated with normaltension glaucoma, primary open-angle glaucoma (Sarfarazi and Rezaie, 2003; Chalasani et al., 2009), Paget's disease (Albagha et al., 2010; Chung et al., 2010) and ALS (Maruyama et al., 2010; Del Bo et al., 2011; van Blitterswijk et al., 2012). Recently, OPTN was characterized as an autophagy adaptor protein which regulates selective autophagy of ubiquitin-coated cytosolic Salmonella enterica. This OPTN function depends on the phosphorylation of its LC3-interacting motif by the protein kinase TANK-binding kinase 1 (TBK1) (Wild et al., 2011). OPTN has been shown to colocalize with protein inclusions in ALS (Maruyama et al., 2010), HD (Schwab et al., 2012) and many other neurodegenerative diseases (Osawa et al., 2011). However, little is known about the role of OPTN in neurodegenerative disorders, particularly in respect to the autophagy-mediated degradation of misfolded protein inclusions.

In this report, we use different aggregation-prone protein models: short fragment of the protein huntingtin carrying an extended polyglutamine (polyQ) mutation (htt ex1 Q103) and superoxide dismutase 1 (SOD1) protein carrying point mutations at position Gly-93 (SOD1 G93C or SOD1 G93A), commonly used as models of HD and ALS, respectively (Witan et al., 2008; Filimonenko et al., 2010). We show that OPTN recognizes htt ex1 Q103 and SOD1 G93C aggregates through its C-terminal coiled-coil domain in a ubiquitin-independent manner and actively participates in the degradation of both htt ex1 Q103 and SOD1 G93C inclusions. Furthermore, we provide evidence for the hypothesis that the degradation of htt ex1 Q103 and SOD1 G93C aggregates is regulated by OPTN via phosphorylation of OPTN Ser177 by TBK1. Finally, OPTN depletion in wild-type zebrafish causes a motor axonopathy phenotype similar to the phenotype reported in a zebrafish model of ALS generated by overexpression of mutant form of SOD1.

Results

OPTN interacts with aggregating proteins involved in ALS and HD

Recently, mutations in the gene encoding OPTN have been associated with some familial and sporadic forms of ALS. OPTN is frequently detected in protein inclusions in neurons of ALS and HD patients (Maruyama et al., 2010; Schwab et al., 2012). To address the possible interaction of OPTN with protein aggregates involved in neurodegenerative pathologies, we transiently transfected HeLa cells with GFP-SOD1 G93C mutant. This mutation has been identified in ALS patients and is responsible for familial cases of ALS (Gaudette et al., 2000). We also used HeLa cells stably transfected with CFP-htt ex1 Q103 as a wellknown model of HD (Filimonenko et al., 2010). The interaction between OPTN and SOD1 G93C or htt ex1 Q103 aggregates was studied using co-immunostaining, immunoprecipitation and binding experiments.

Confocal analyses indicated that OPTN readily colocalized with both SOD1 G93C and htt ex1 Q103 inclusions (Fig. 1A, upper panels). Similarly, we observed a colocalization of ubiquitin, p62 and LC3 with SOD1 G93C and htt ex1 Q103 aggregates (Fig. 1A). To further confirm OPTN interaction with SOD1 G93C inclusions, we co-expressed wild-type SOD1 (SOD1 wt) or mutant SOD1 G93C with a wild-type HA-OPTN

(HA-OPTN wt) in HeLa cells. Wild-type HA-OPTN coimmunoprecipitated only with the aggregation-prone SOD1 G93C mutant and not with wild-type SOD1 (Fig. 1B). Moreover, fractionation experiments, separating soluble and insoluble fractions of the cells, revealed that a significant amount of endogenous OPTN was found in the insoluble fraction of HeLa cells expressing GFP-SOD G93C while the portion of insoluble OPTN detected in cells transfected with the empty vector or with the wild-type GFP-SOD1 was found to be less prominent (Fig. 1C). Co-immunoprecipitation experiments of soluble GFP-SOD1 G93C with endogenous OPTN were performed to determine if OPTN interacted with the soluble monomeric and oligomeric forms of mutant GFP-SOD1 G93C present in the soluble fraction of HeLa cells. Mutant GFP-SOD1 G93C but not wild-type SOD1 co-immunoprecipitated with endogenous OPTN from the soluble fraction (Fig. 1D). To test if the interaction of OPTN with SOD1 G93C mutant was direct, we carried out a GST pull-down binding assay with either purified MBP-OPTN and GST-SOD1 G93C or GST-OPTN and His-SOD1 G93C. Neither purified MBP-OPTN nor GST-OPTN interacted with mutant GST-SOD1 G93C or His-SOD1 G93C, respectively (Fig. 1E,F). Finally we tested whether OPTN directly interacted with aggregated mutant SOD1 G93C using a dot blot assay. We induced aggregation of His-SOD1 G93C in vitro (Yerbury et al., 2013) and mature aggregates were incubated with purified GST-OPTN. OPTN did not bind to mature SOD1 G93C aggregates. The results obtained suggest that OPTN indirectly recognized soluble monomeric or oligomeric forms of mutant SOD1 G93C but also the aggregated form of SOD1 (Fig. 1B-G).

The C-terminal coiled-coil domain of OPTN is responsible for binding to protein aggregates in a ubiquitinindependent manner

Next, we mapped the domain of OPTN responsible for the interaction with aggregation-prone proteins. We co-expressed various fragments of OPTN with GFP-SOD1 G93C in HeLa cells. The N-terminal fragment of OPTN (1-315) did not colocalize with GFP-SOD1 G93C aggregates (Fig. 2A, right panel), while the C-terminal fragment (297-577) colocalized with the inclusions (data not shown). Further fragmentation of the Cterminal part of OPTN showed that the isolated coiled-coil domain (amino acids 454-520) interacted with GFP-SOD1 G93C protein aggregates whereas a deletion of the same domain abolished this interaction (Fig. 2A, right panel), suggesting that the C-terminal coiled-coil domain is crucial for OPTN binding to GFP-SOD1 G93C protein inclusions. Co-expression of the aforementioned fragments of OPTN with wild-type SOD1 demonstrated that OPTN did not form aggregates in the absence of mutant SOD1 G93C (Fig. 2A, left panel).

OPTN contains the UBAN domain, an ubiquitin binding domain present in various proteins such as NEMO and ABIN-3 (Wagner et al., 2008; Husnjak and Dikic, 2012). It was previously shown that the E478G mutation in UBAN domain of OPTN abolished OPTN binding to ubiquitin (Wild et al., 2011). We observed that OPTN E478G mutant was still able to colocalize with GFP-SOD1 G93C protein aggregates, suggesting that OPTN interacts with protein aggregates in a ubiquitin-independent manner. Moreover, the LC3-interacting motif (LIR) mutant, OPTN F178A, unable to bind LC3/GABARAP modifiers, had no influence on OPTN localization to GFP-SOD1 G93C protein

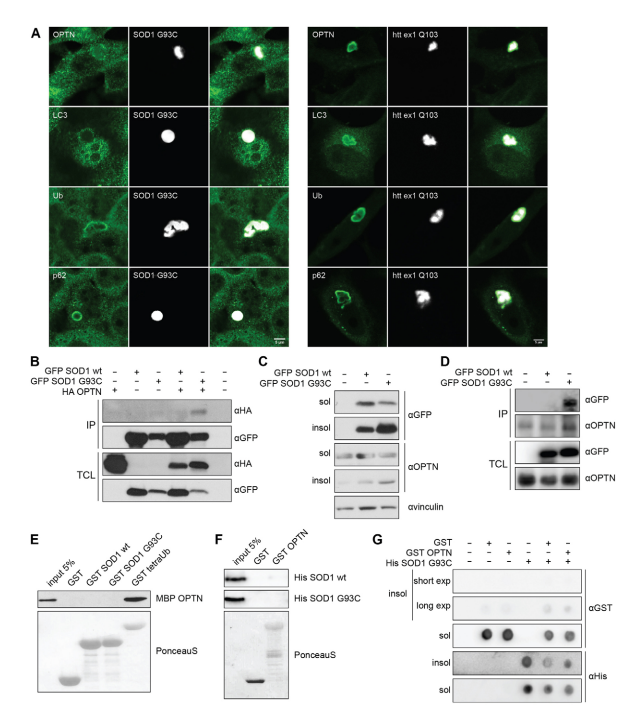


Fig. 1. OPTN interacts with aggregating proteins involved in ALS and HD. (A) Endogenous OPTN, LC3, ubiquitin (Ub) and p62 colocalized with GFP-SOD1 G93C aggregates in transiently transfected HeLa cells for 48 hours (left) and with CFP-htt ex1 Q103 aggregates in stable transfected HeLa cells (right). Scale bars: $5 \ \mu$ m. (**B**) Co-immunoprecipitation experiment using HeLa cells co-transfected with wild-type GFP-SOD1 or GFP-SOD1 G93C and HA-OPTN. Total cell lysates (TCL) were subjected to immunoprecipitation using anti-GFP antibody and then probed with anti-HA antibody. (**C**) Immunoblot analysis of cellular fractions using antibodies against vinculin, GFP and OPTN. HeLa cells transfected with empty vector, wild-type GFP-SOD1 and GFP-SOD1 G93C were lysed in NP40 lysis buffer at 48 hours after transfection and the lysates separated into soluble and insoluble fractions. OPTN showed enrichment in insoluble fractions together with SOD1 G93C aggregates. (**D**) Co-immunoprecipitation using anti-OPTN antibody and then probed with anti-GFP antibody. (**E**) GST pull-down assay of purified wild-type GST-SOD1 or G93C mutant. Wild-type MBP-OPTN did not interact with either wild-type GST-SOD1 or G93C mutant. (**F**) GST pull-down assay of purified wild-type His-SOD1 or G93C mutant using GST and wild-type GST-OPTN. Neither wild-type His-SOD1 nor G93C mutant interacted with wild-type GST-OPTN. (**G**) Aggregated His-SOD1 G93C mutant was incubated with either anti-His or anti-GST antibody.

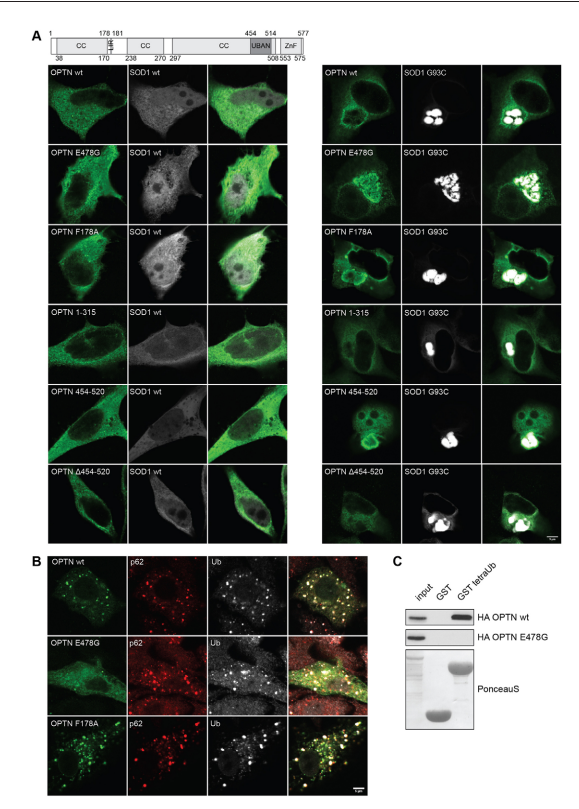


Fig. 2. C-terminal coiled-coil domain of OPTN is responsible for binding to proteins aggregates in a ubiquitin-independent manner. (A) Schematic representation of OPTN domain structure. Numbers indicate the amino acids. Transient co-transfection was performed with the indicated fragments of HA-OPTN and wild-type GFP-SOD1 (left) or GFP-SOD1 G93C (right) in HeLa cells. Wild-type HA-OPTN, HA-OPTN F178A, E478G and 454-520 localized to GFP-SOD1 G93C aggregates. Expression of neither HA-OPTN 1-315 nor Δ454-520 led to colocalization with aggregates. (B) HeLa cells transfected with wild-type GFP-OPTN, GFP-OPTN F178A and GFP-OPTN E478G were treated 24 hours after transfection with puromycin for 2 hours, fixed and stained with anti-ubiquitin and anti-p62 antibodies. Wild-type GFP-OPTN and GFP-OPTN F178A showed localization with puromycin-induced aggregates whereas GFP-OPTN E478G showed lack of colocalization. (C) GST pull-down assay of wild-type HA-OPTN and HA-OPTN E478G using GST and GST-tetraUb. Wild-type HA-OPTN interacted with tetraUb whereas HA-OPTN E478G showed lack of interaction with tetraUb. Scale bars: 5 μm.

aggregates. We obtained similar results when we expressed the aforementioned OPTN fragments in stably transfected CFP-htt ex1 Q103 HeLa cells (data not shown), confirming that OPTN recognizes protein aggregates through its C-terminal coiled-coil domain in a ubiquitin-independent manner.

We also investigated colocalization of OPTN with puromycininduced ubiquitin and p62 positive aggregates (Pankiv et al., 2007) by monitoring the localization of wild-type HA-OPTN, HA-OPTN E478G and HA-OPTN F178A in HeLa cells treated with puromycin for 2 hours (Fig. 2B). We observed that wild-type HA-OPTN and HA-OPTN F178A colocalized with protein inclusions, but HA-OPTN E478G colocalization could not be detected. The results obtained suggest that OPTN interacts with puromycin induced ubiquitin and p62 positive aggregates through the UBAN domain in a ubiquitin-dependent manner, similarly to its interaction with *Salmonella enterica* (Wild et al., 2011). The GST pull-down binding assay confirmed that wild-type HA-OPTN binds ubiquitin chains, while the OPTN E478G mutation abolished this binding (Fig. 2C), as previously described (Wild et al., 2011).

OPTN depletion increases protein aggregation

Following the confirmation of an indirect interaction between OPTN and protein aggregates, we investigated the functional relevance of OPTN in protein aggregation. For this purpose, we established HeLa cell lines stably expressing a control non-targeting shRNA (ctrl shRNA) or shRNAs directed against OPTN (shOPTN #3 and shOPTN #4). We transiently transfected OPTN knock-down and shRNA control cells with GFP-SOD1 G93C or GFP-htt ex1 Q103 for 48 hours and quantified the aggregates of GFP-SOD1 G93C and GFP-htt ex1 Q103 (representative pictures for GFP-SOD1 G93C shown in Fig. 3A). OPTN depletion induced a significant increase of GFP-SOD1 G93C or htt ex1 Q103 aggregates (Fig. 3A,B). Aggregation observed by immunofluorescence was further confirmed by investigation of the soluble and insoluble fractions. Immunoblot analysis showed an increase of insoluble GFP-SOD1 G93C in HeLa cells depleted of OPTN (Fig. 3C). Similarly, transient depletion of OPTN using two different siRNA targeting OPTN (siOPTN #1 and siOPTN #2) for 48 hours in HeLa cells stably expressing CFP-htt ex1 Q103 led to an increased number of cells containing aggregates as well as insoluble CFP-htt ex1 Q103 compared to control cells expressing OPTN (Fig. 3D,E). Inversely, overexpression of HA-OPTN had no significant effect on the number of GFP-SOD1 G93C (Fig. 3F) or GFP-htt ex1 Q103 inclusions (data not shown). The results obtained indicate that the lack of OPTN increases the accumulation of GFP-SOD1 G93C and GFP-htt ex1 Q103 protein aggregates.

OPTN modulates protein aggregation via autophagy

In light of the recently established role of OPTN in selective autophagy of ubiquitin-coated *Salmonella enterica* (Wild et al., 2011) we investigated the role of OPTN in the autophagy clearance of protein aggregates. Stable ctrl shRNA and shOPTN HeLa cells were transfected with GFP-SOD1 G93C or GFP-htt ex1 Q103 and treated with DMSO (control), rapamycin (autophagy activator) and bafilomycin A1 (autophagy inhibitor). A significant decrease in GFP-SOD1 G93C and GFP-htt ex1 Q103 aggregates in ctrl shRNA HeLa cells treated with rapamycin was observed (Fig. 4A,B). In contrast, treatment with bafilomycin A1 caused a twofold increase of the number of GFP-SOD1 G93C and GFP-htt ex1 Q103 inclusions (Fig. 4A,B), which was comparable with the augmentation of protein aggregates induced by OPTN depletion in shOPTN HeLa cells. In OPTN-depleted HeLa cells rapamycin and bafilomycin A1 had no effect on the quantity of cells with GFP-SOD1 G93C and GFP-htt ex1 Q103 protein inclusions. Autophagy activation or inhibition was confirmed by assessing the levels of LC3-I/LC3-II in total cell lysates (Fig. 4C). In order to demonstrate a specific role of OPTN in the aggregation phenotype observed in cellular models we performed a reconstitution of OPTN-depleted shOPTN HeLa cells with wild-type OPTN, OPTN F178A (LC3-binding-deficient OPTN) and OPTN E478G (ubiquitinbinding-deficient OPTN). Both wild-type and OPTN E478G resulted in a decrease in aggregation of GFP-SOD1 G93C to levels similar to ctrl shRNA HeLa cells (Fig. 4D). However, OPTN F178A was not able to rescue the levels of GFP-SOD1 G93C (Fig. 4D). The results observed indicate that OPTN regulates GFP-SOD1 G93C and GFP-htt ex1 Q103 aggregation through autophagy via binding to LC3.

Next, we treated ctrl shRNA and shOPTN HeLa cells transfected with GFP-SOD1 G93C and GFP-htt ex1 Q103 with the proteasomal inhibitor MG132 for 24 hours (Fig. 4E). Inhibition of proteasomal degradation significantly increased the number of cells with protein aggregates in ctrl shRNA HeLa cells but also in OPTN depleted HeLa cells, indicating that the effect of OPTN on protein aggregation is independent of ubiquitin-proteasome system.

TBK1 phosphorylates OPTN on protein aggregates

As the targeting of protein aggregates to the autophagy-lysosome pathway requires the LIR motif of OPTN, we investigated whether TBK1-mediated phosphorylation of Ser177, adjacent to the LIR motif, is involved in the regulation of the interaction of OPTN with protein inclusions (Wild et al., 2011). We showed that TBK1 colocalized with protein aggregates in HeLa cells transiently co-transfected with myc-TBK1 and GFP-SOD1 G93C or GFP-htt ex1 Q103 (Fig. 5A). OPTN phosphorylated at Ser177 (phospho S177 OPTN) colocalized with GFP-SOD1 G93C inclusions. As no phospho S177 OPTN was detected on GFP-SOD1 G93C aggregates in HeLa cells co-transfected with TBK1-targeted siRNA, we hypothesized that phosphorylation of OPTN on protein aggregates is mediated by TBK1 (Fig. 5B).

Additionally, we analyzed spinal cord sections of transgenic mice expressing human mutant SOD1 G93A (Gurney et al., 1994) at three different stages of the pathology. We examined 5-month-old (no phenotype), 7-month-old (early-stage of the disease) and 10-monthold (end-stage of the disease) human mutant SOD1 G93A transgenic mice. We confirmed the colocalization of phospho S177 OPTN and TBK1 with SOD1 G93A protein aggregates (Fig. 5C) at all phenotype stages investigated even in asymptomatic 5-month-old SOD1 G93A mice where only sparse SOD1 aggregates were observed (Gurney et al., 1994). Occasionally, ubiquitinated SOD1 aggregates were not co-stained with phospho S177 OPTN or TBK1 (Fig. 5C). These results demonstrate a colocalization of TBK1 and phospho S177 OPTN with SOD1 G93A aggregates in this mouse model of ALS, suggesting that OPTN targeting of SOD1 G93A aggregates to the autophagy-lysosome pathway is regulated by phosphorylation of OPTN by TBK1.

TBK1 plays a role in autophagic degradation of protein aggregates

We further investigated the role of TBK1 in autophagic degradation of protein aggregates. Depletion of TBK1 in ctrl

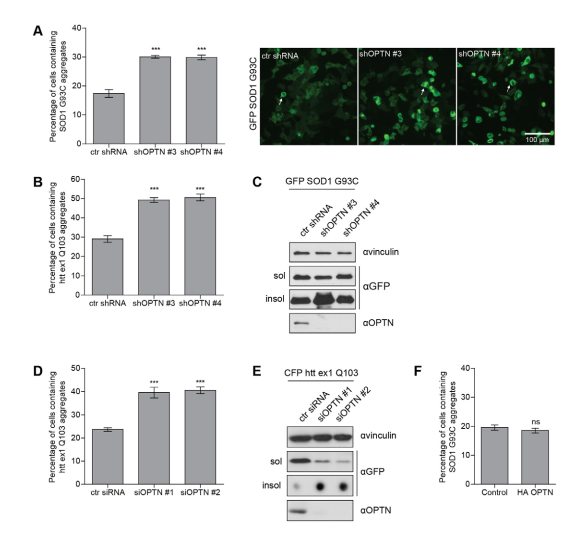


Fig. 3. OPTN depletion increases protein aggregation. (A) Left: quantification of transfected cells containing GFP-SOD1 G93A aggregates in ctrl shRNA and two different shRNA OPTN-depleted HeLa cell lines transiently transfected with GFP-SOD1 G93C for 48 hours. Right: representative images of each group. Arrows indicate typical transfected cells containing GFP-SOD1 G93C aggregates. Scale bar: 100 μ m. (B) Quantification of transfected cells containing GFP-tht ex1 Q103 aggregates in ctrl shRNA and two different shRNA OPTN-depleted HeLa cell lines. (C) Immunoblot analysis of cellular fractions using antibodies against vinculin, GFP and OPTN. HeLa cells transfected with GFP-SOD1 G93C in ctrl shRNA and two different shRNA OPTN-depleted HeLa cell lines were lysed in NP40 lysis buffer at 48 hours after transfection and lysates separated into soluble and insoluble fractions. The quantity of GFP-SOD1 G93C increased in the insoluble fraction in shRNA oPTN-depleted HeLa cells. (D) Quantification of aggregates in ctrl siRNAs targeting OPTN for 48 hours. (E) Immunoblot analysis of cellular fractions using antibodies against vinculin, GFP and OPTN. HeLa cells stably expressing CFP-htt ex1 Q103 and transfected with ctrl siRNAs targeting OPTN were lysed in NP40 lysis buffer at 48 hours after transfection and lysates separated into soluble and insoluble fractions using antibodies against vinculin, GFP and OPTN. HeLa cells stably expressing CFP-htt ex1 Q103 and transfected with ctrl siRNAs targeting OPTN were lysed in NP40 lysis buffer at 48 hours after transfection and lysates separated into soluble and insoluble fractions was analyzed by SDS-PAGE and the insoluble fraction was applied on a dot-blot apparatus. The amount of CFP-htt ex1 Q103 in the insoluble fraction increased in HeLa cells transfected with siRNA targeting OPTN. (F) Quantification of transfected cells containing GFP-SOD1 G93C aggregates in HeLa cells transiently co-transfected with empty vector (control) or HA-OPTN for 48 hours. All data are shown as mean valu

shRNA HeLa cells induced an increase in GFP-SOD1 G93C aggregation similar to the increase observed after silencing of OPTN. In contrast, TBK1 silencing had no additional effect in shOPTN HeLa cells (Fig. 6A). These results indicate a common pathway of modulation of protein aggregation shared between TBK1 and OPTN. The silencing of TBK1 induced decreased levels of the lipidated form of LC3 (LC3-II), representing a decreased incorporation of LC3 into autophagosomal membranes and a decreased autophagic activity (Fig. 6B).

Overexpression of myc-TBK1 in ctrl shRNA HeLa cells decreased GFP-SOD1 G93C aggregation (Fig. 6C) in a way

comparable to rapamycin-treated cells (Fig. 4A). Interestingly, the overexpression of myc-TBK1 in shOPTN HeLa cell partially decreased GFP-SOD1 G93C aggregation, indicating potential activation of autophagic degradation of GFP-SOD1 G93C by TBK1 in the absence of OPTN. In contrast, the kinase-inactive TBK1 mutant K38A had no impact on GFP-SOD1 G93C aggregation (Fig. 6C), indicating that the kinase activity of TBK1 could be responsible for its effect on protein aggregation. The results obtained suggest an important role of TBK1 phosphorylation of OPTN and potentially other proteins involved in degradation of protein aggregates via the autophagy-lysosome pathway.

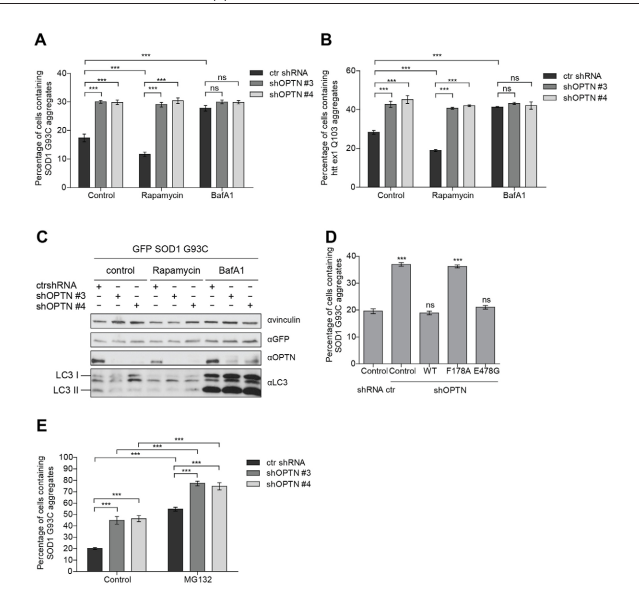
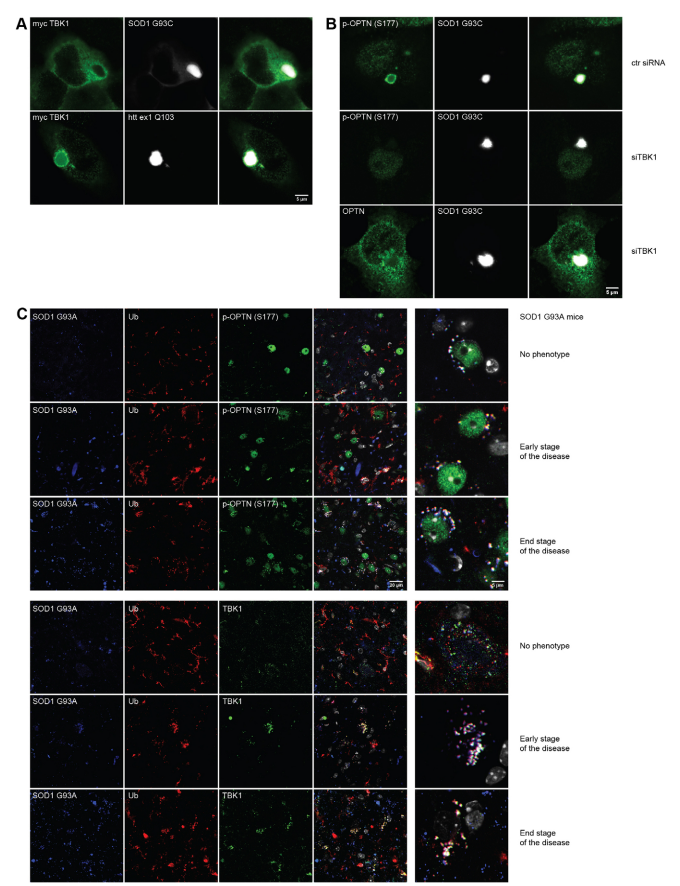


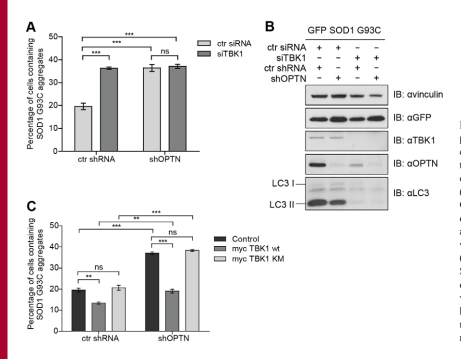
Fig. 4. OPTN modulates protein aggregation via autophagy. (A,B) Quantification of transfected cells containing GFP-SOD1 G93C (A) or GFP-htt ex1 Q103 (B) aggregates in HeLa cells transiently transfected with control shRNA or two different shRNA silencing OPTN (shOPTN) and treated with DMSO (control), 2 μ M rapamycin for 48 hours or 200 nM bafilomycin A1 (BafA1) for 24 hours. (C) Immunoblot analysis using antibodies against vinculin, GFP, OPTN and LC3. Cells were lysed in RIPA buffer and analyzed by SDS-PAGE. LC3 levels confirmed autophagy activation by rapamycin and autophagy inhibition by bafilomycin A1. (D) Quantification of transfected cells containing GFP-SOD1 G93C aggregates in control shRNA or two different shOPTN HeLa cell lines and treated with DMSO (control) for 48 hours or 10 μ M MG132 for 24 hours. (E) Quantification of transfected cells containing GFP-SOD1 G93C aggregates in OPTN-depleted HeLa cells co-transfected with GFP-SOD1 G93C and empty vector or various shRNA-resistant OPTN constructs: wild-type OPTN, OPTN F178A and OPTN E478G. Wild-type OPTN and OPTN E478G mutant rescued OPTN-depleted HeLa cells whereas OPTN F178A failed to rescue. All data are shown as mean values \pm s.e.m.; ***P<0.001; ns, not significant.

OPTN depletion in zebrafish (*Danio rerio*) causes motor axonopathy and increases motor axonopathy induced by mutant SOD1 overexpression

In order to investigate the effects of OPTN depletion on protein aggregation *in vivo*, we employed a well-validated zebrafish model of ALS overexpressing the SOD1 G93A mutant characterized by the development of a specific and dose-dependent axonopathy (Lemmens et al., 2007; Laird et al., 2010; Xi et al., 2011). In order to deplete OPTN levels in zebrafish, two different antisense morpholino oligonucleotides (AMO) targeting OPTN were used and the zebrafish were analyzed 48 hours after fertilization. Both AMOs induced an

efficient silencing of OPTN, and depletion of OPTN in wild-type zebrafish resulted in a specific phenotype characterized by embryos with curved tails (Fig. 7A) and delayed and/or no ability to swim away after touch (data not shown). This phenotype was also observed in zebrafish overexpressing SOD1 G93A without silencing of OPTN (Fig. 7A). We also observed shorter and abnormally branched motor axons measured by labeling embryos with the synaptic vesicle marker, SV2 (Lemmens et al., 2007). Knockdown of OPTN led to a decrease in motor axon length and an increase in abnormal motor axon branching (Fig. 7C). These data indicate that OPTN depletion causes motor axonopathy in zebrafish embryos. Similarly, co-injection of OPTN AMO and





mRNA coding for FLAG-SOD1 G93A produced a more pronounced decrease in motor axon length in zebrafish embryos as well as an increase in abnormal motor axon branching (Fig. 7D,E).

Discussion

OPTN has been reported to colocalize with protein inclusions in a wide range of neurodegenerative diseases such as ALS (Maruyama et al., 2010), HD (Schwab et al., 2012), Alzheimer's disease, Parkinson's disease, Creutzfeldt-Jakob disease and Pick disease (Osawa et al., 2011). Moreover, recent studies described OPTN deletion mutations in ALS patients (Maruyama et al., 2010; Del Bo et al., 2011; Sugihara et al., 2012; Tumer et al., 2012; van Blitterswijk et al., 2012). Despite the existing observations, little is known about the potential function of OPTN in neurodegenerative diseases. Our data suggest an important role for OPTN in protein aggregation, accumulation and degradation. We also provide insights into the regulation of OPTN function in autophagic degradation of protein aggregates involved in the pathogenesis of ALS and HD.

Fig. 5. TBK1 phosphorylates OPTN on protein aggregates. (A) Cotransfection of myc-TBK1 and GFP-SOD1 G93C or GFP-htt ex1 Q103 for 48 hours in HeLa cells showed localization of myc-TBK1 on GFP-SOD1 G93C and GFP-htt ex1 Q103 aggregates. (B) Phosphorylated Ser177 OPTN [p-OPTN (S177)] localized to GFP-SOD1 G93C aggregates in HeLa cells cotransfected with control siRNA whereas co-transfection with siTBK1 abolished phosphorylated Ser177 OPTN colocalization with GFP-SOD1 G93C inclusions. Endogenous OPTN localized to GFP-SOD1 G93C aggregates in HeLa cells co-transfected with siRNA targeting TBK1. (C) Immunohistochemical staining of the spinal cord of 5-month-old (no phenotype, asymptomatic), 7-month-old (early stage of the disease) and 10month-old (end stage of the disease) transgenic mice expressing human SOD1 G93A showed that SOD1 G93A proteins aggregates were immunopositive for TBK1 and phosphorylated Ser177 OPTN in motor neurons. Gray, DAPI staining. Fig. 6. TBK1 plays a role in autophagy degradation of protein aggregates. (A) Quantification of transfected cells containing GFP-SOD1 G93C aggregates transiently cotransfected with control siRNA or siRNA targeting TBK1 in control shRNA or OPTN-depleted HeLa cells. (B) Immunoblot analysis using antibodies against vinculin, GFP, TBK1, OPTN and LC3. Control shRNA or OPTNdepleted HeLa cells co-transfected with GFP-SOD1 G93C and control siRNA or siRNA targeting TBK1 for 48 hours were lysed in RIPA buffer and analyzed by SDS-PAGE. (C) Quantification of transfected cells containing GFP-SOD1 G93C aggregates in shRNA control or OPTNdepleted HeLa cells transiently co-transfected with empty vector (as control), wild-type myc-TBK1 and myc-TBK1 kinase mutant K38A (TBK1 KM). All data are shown as mean values \pm s.e.m.; ***P*<0.01, ****P*<0.001; ns, not significant.

In this work, we report on the colocalization of OPTN, SOD1 and htt protein aggregates as previously described for other proteins involved in autophagy such as ubiquitin, p62 and LC3 (Fig. 1) (Filimonenko et al., 2010). We demonstrate that the interaction of OPTN with aggregation-prone proteins is indirect and occurred already with soluble monomeric or oligomeric forms of aggregating proteins (Fig. 1). The identity of the protein that functions as an adaptor between OPTN and aggregating protein remains unknown. A potential candidate is ubiquitin. Indeed, OPTN is an autophagy cargo receptor with the ability to bind LC3 through an LC3-interacting motif (LIR) and ubiquitin through ubiquitin-binding domain (UBAN). The importance of the UBAN domain in autophagic clearance of cytosolic Salmonella enterica has been shown (Wild et al., 2011). Here we show that OPTN interacts with p62 bodies through its UBAN domain recognizing polyubiquitin chains. However, OPTN interaction with SOD1 and htt aggregates is ubiquitinindependent. The mutation in the UBAN domain (E478G) that completely abolishes OPTN binding to polyubiquitin chains (Wild et al., 2011) had no impact on OPTN interaction with SOD1 aggregates (Fig. 2). It is possible that ubiquitin is not the signal/motif of protein aggregates recognized by OPTN to be recruited to the autophagy machinery, as ubiquitination might not be a universal marker of protein aggregates. Indeed, ubiquitinnegative inclusions were found in ALS patients (Allen et al., 2002; Osawa et al., 2011). It was also shown that p62 interaction with SOD1 protein aggregates (Gal et al., 2009) as well as with a non-ubiquitinated substrate STAT5A_ Δ E18, an aggregationprone isoform of STAT5A (Watanabe and Tanaka, 2011), could be independent of ubiquitin signals. Moreover, as recently shown co-chaperone BAG3 mediated aggresome targeting and selective autophagy of SOD1 aggregates do not depend on substrate ubiquitination (Gamerdinger et al., 2011). OPTN interacts with SOD1 and htt aggregates via its C-terminal coiled-coil domain (454-520 amino acids). However, future studies will have to determine which specific residues of OPTN

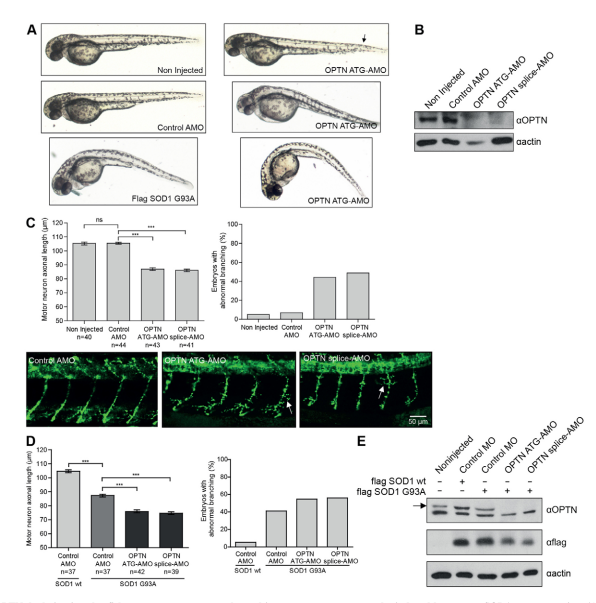


Fig. 7. OPTN depletion in zebrafish causes motor axonopathy and increases motor axonopathy induced by mutant SOD1 overexpression. (A) Zebrafish injected with OPTN-specific translation blocking (OPTN ATG-AMO) morpholino showed a phenotype (curved tail indicated by the arrow) similar to that observed in zebrafish overexpressing SOD1 G93A at 48 hours after fertilization in comparison with non-injected zebrafish or zebrafish injected with control morpholino (control AMO). A comparable phenotype was also observed in zebrafish injected with splice blocking (OPTN Splice-AMO) morpholino (data not shown). (B) Immunoblot analysis of protein extracted from zebrafish using antibodies against OPTN and actin showed efficiency of OPTN silencing. (C) Quantification of motor axon length and branching in zebrafish injected with control AMO. Abnormal branching is expressed as percentage of embryos showing premature branching in two or more axons. Representative images of motor neuron axons at 48 hours after fertilization of motor axon length and branching in zebrafish with overexpressed FLAG-SOD1 G93A and control AMO showed a decrease in motor neurons length and increase in premature branching compared with zebrafish overexpressing wild-type FLAG-SOD1 G93A and OPTN AMO showed an additional decrease in motor neuron length and additional increase in premature branching. (E) Immunoblot analysis of protein extracts from zebrafish with RIPA buffer using antibodies against FLAG, OPTN and actin shows efficiency of OPTN silencing and FLAG-SOD1 expression. All data are shown as mean values \pm s.e.m.; **P<0.01; ns, not significant.

are involved in its interaction with protein inclusions and which protein is recognized by OPTN on protein inclusions.

Furthermore, OPTN depletion in HeLa cells resulted in a significant increase in aggregation of SOD1 G93C or htt ex1 Q103, suggesting a role of OPTN in the degradation of aggregation-prone proteins (Fig. 3). OPTN was modulating SOD1 G93C or htt ex1 Q103 degradation by the autophagy-lysosome system. This effect of OPTN was regulated by

phosphorylation of Ser177 by TBK1, necessary for OPTN binding to LC3, following recruitment of the autophagy machinery during the degradation of *Salmonella enterica* (Wild et al., 2011). Even though non-phosphorylated OPTN was detected in protein inclusions, phosphorylation of OPTN by TBK1 is a rate-limiting step in autophagic degradation of protein aggregates. Therefore, we hypothesize that OPTN is first recruited to the protein aggregates following recognition and

phosphorylation by TBK1, allowing the subsequent recruitment of LC3 and the autophagy machinery.

Interestingly, we also showed that even in the absence of OPTN, overexpression of TBK1 could still decrease the number of cells containing SOD1 G93C aggregates (Fig. 6C). This observation suggests that in addition to OPTN, TBK1 potentially could phosphorylate other proteins involved in the degradation of protein aggregates. Potential candidates might be other previously identified autophagy receptors such as p62, Alfy and NBR1. As TBK1 depletion caused decreased incorporation of LC3 into autophagosomal membranes and a decreased autophagic activity, potential candidates could also be components of autophagy initiation ULK1-dependent complex or members of the PI3K complex involved in vesicle nucleation (Behrends et al., 2010).

Our cellular observations were confirmed in vivo as we colocalized TBK1 and phosphorylated OPTN with SOD1 G93A protein aggregates in the spinal cord of a murine model of ALS (Gurney et al., 1994). Most of the SOD1 protein inclusions were imunoreactive for phosphorylated OPTN and TBK1 as well as for ubiquitin even at the non-symptomatic stage when the spinal cord contained almost no inclusions and the SOD1 G93A mice had not developed any phenotype. This observation concurs with our in vitro results showing OPTN interaction with soluble monomeric or oligomeric mutant SOD1 G93C. There was a slight discrepancy between ubiquitin and phospho S177 OPTN staining, as some ubiquitinated aggregates were not co-stained with phospho S177 OPTN or TBK1. Recently, it was shown that OPTN was absent or rare in ALS patients (Hortobágyi et al., 2011), which might be in agreement with our observation that some SOD1 aggregates are not recognized by phospho S177 OPTN or TBK1 in the mutant SOD1 G93A mouse. However, we also demonstrated that ubiquitin is not the signal recognized by OPTN on the protein aggregates and therefore not surprisingly some ubiquitinated SOD1 aggregates were not co-stained with phospho S177 OPTN or TBK1 in the mutant SOD1 G93A mouse. This might be an in vivo confirmation that the presence of ubiquitin is not attracting OPTN to the protein aggregates. On the other hand, it might also indicate that the phospho S177 OPTN/TBK1-negative aggregates are lacking the protein recognized by OPTN on aggregation-prone proteins. Further studies are needed to identify the adaptor protein between OPTN and protein inclusions as its presence might facilitate the degradation of aggregation-prone proteins. Although controversial results have been obtained on the presence of OPTN in protein aggregates in different neurodegenerative diseases (Deng et al., 2011; Ito et al., 2011; Schwab et al., 2012; Hortobágyi et al., 2011) the localization of phospho S177 OPTN or TBK1 in inclusions in human cases of neurodegenerative diseases still remains to be determined.

We also investigated the effect of OPTN depletion in zebrafish and showed that silencing of OPTN in wild-type zebrafish induced a phenotype similar to the one displayed by a zebrafish model of ALS overexpressing SOD1 G93A. The fact that OPTN deletion caused an ALS motor axonopathy phenotype in wild-type zebrafish is further supported by the description of OPTN deletion mutations in ALS patients (Maruyama et al., 2010; Del Bo et al., 2011; Sugihara et al., 2011; Iida et al., 2012; Tumer et al., 2012; van Blitterswijk et al., 2012). Furthermore, OPTN depletion in zebrafish expressing FLAG-SOD1 G93A provoked an even more severe motor axonopathy phenotype. This observation of an additive effect of OPTN deletion and SOD1 G93A overexpression might suggest that the pathological mechanisms involved in OPTN and SOD1 G93A are complementary. However, as the severity of the motor axonopathy phenotype in ALS zebrafish model overexpressing SOD1 G93A is dose dependent (Lemmens et al., 2007), OPTN depletion might cause a lack of autophagic clearance and a subsequent accumulation of SOD1 G93A aggregates. We have showed that OPTN is an important player in autophagy clearance of misfolded and aggregation-prone proteins and taking this into consideration, neurodegenerative pathology caused by mutation or complete deletion of OPTN in zebrafish or in human ALS cases can be explained as a dysfunctional autophagy clearance of misfolded and aggregation-prone proteins. However, in ALS patients without OPTN mutation OPTN is recruited to the protein aggregates as an attempt to degrade the aggregation-prone proteins but an increase stimulation of this potentially protective pathway might be needed to avoid the accumulation of proteins in the nervous system.

In conclusion, we provided evidence that OPTN regulates protein aggregation through autophagy pathway by recognizing protein aggregates via the C-terminal coiled-coil domain and by recruiting LC3 to protein aggregates via its LIR motif. Furthermore, TBK1 regulates protein aggregation by phosphorylating OPTN and also various other proteins involved in the regulation of autophagy. These findings support a potential protective role of OPTN and TBK1 in neurodegenerative disorders associated with intracellular protein aggregation.

Materials and Methods

Plasmids, reagents and cell lines

pcDNA3.1-HA-OPTN (wild-type, F178A and E478G), pMAL-c2x-OPTN, pGEX4T1-OPTN, pGEX4T1-tetraUb and pcDNA3.1-myc-TBK1 (wild-type, K38A) plasmids were described previously (Ikeda et al., 2007; Wild et al., 2011). Homodimer GFP-SOD1 G93C was subcloned in pcDNA3.1 vector from vector pEGFP-N1-SOD1 G93C/G93C previously described (Witan et al., 2008). pcDNA3-GFP-htt ex1 Q103 was a kind gift from Dimitri Krainc (Institute for Neurodegenerative Disease, Harvard Medical School). Plasmids encoding FLAG-SOD1 (wild-type and G93A) were kindly provided by Ryosuke Takahashi (Department of Neurology, Kyoto University Hospital and Graduate School of Medicine, Japan). HA-OPTN 1-315 and HA-OPTN 454-520 were subcloned into BamHI-XhoI sites of the pcDNA3.1 vector. HA-OPTN Δ454-520 was generated by in vitro mutagenesis (Stratagene). Plasmids encoding wild-type His-SOD1 and G93C mutant or GST-SOD1 and G93C mutant were subcloned into BamH1 site of the pRSET vector or pGEX4T1, respectively. All constructs were verified by sequencing before use. GeneJuice (Merck) and Lipofectamine RNAiMAX were used for mammalian cells transfection. GFP-Trap was purchased from ChromoTek, puromycin, MG132 and rapamycin from Sigma-Aldrich, bafilomycin A1 from LC Laboratories, Protein G Agarose from Roche, Glutathione-Sepharose 4B from GE Healthcare, Amylose resin from New England Biolabs, Ni NTA Agarose from Qiagen.

Cell lines and transfection of siRNA and plasmids

HeLa cells were obtained from ATCC. Stable HeLa cells expressing CFP-htt ex1 O103 were kindly provided by Anne Simonsen (Institute of Basic Medical Sciences, University of Oslo, Norway) (Filimonenko et al., 2010). HeLa cell lines stably expressing a control non-targeting shRNA (ctrl shRNA) or shRNAs directed against OPTN (shOPTN #3 and shOPTN #4) were previously described (Wild et al., 2011). All cells were grown in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 100 U/ml penicillin/ streptomycin at 37°C, 5% CO2. For siRNA experiments, HeLa cells were transfected with siRNA using Lipofectamine RNAiMAX in reverse transfection according to the manufacturer's protocol. After 24 hours cells were transfected with plasmids using GeneJuice and all analyses were performed 48 hours later. Sequences of siRNA used in this study: Non-Targeting siCONTROL #2 from Thermo Fisher, siOPTN #1 5'-GCACGGCAUUGUCUAAAUA-3', siOPTN #2 5'-GAAGUUUACUGUUCUGAUU-3', siTBK1 5'-GACAGAAGUUGUGAUC-ACA-3'. A final concentration of 10 (ctrl siRNA, siTBK1, siOPTN #2) or 20 nM (siOPTN #1) of siRNAs was used for silencing.

Antibodies

The following antibodies were used in the study: rabbit polyclonal anti-OPTN (Abcam, ab23666), mouse monoclonal anti-GFP (Roche), mouse monoclonal anti-ubiquitin

(FK2, Enzo Life Science), mouse monoclonal anti-FLAG M2 (Sigma-Aldrich), mouse monoclonal anti-actin (Sigma-Aldrich), mouse monoclonal anti-vinculin (Sigma-Aldrich), mouse monoclonal anti-myc (Santa Cruz), rabbit polyclonal anti-LC3 (MBL, PM036), rabbit polyclonal anti-LC3B (a kind gift from Zvulun Elazar, the Weizmann Institute of Science, Israel), mouse monoclonal anti-synaptic vesicle 2 (DSHB), rabbit polyclonal anti-TBK1 (Cell Signaling), rabbit polyclonal anti-p62 (Enzo Life Sciences), mouse monoclonal anti-HA (Covance), mouse monoclonal anti-MBP (Sigma), mouse monoclonal anti-GST (Santa Cruz Biotechnology), mouse monoclonal anti-His (Novagen). Rabbit polyclonal anti-SOD1 (Enzo) was conjugated to DyLight 650 using the DyLight antibody labeling kit (Thermo Scientific). The phospho S177 antibody was generated by immunoGLobe as previously described (Wild et al., 2011). Secondary horseradish peroxidase (HRP)-conjugated goat anti-mouse and goat antirabbit IgGs antibodies were purchased from Santa Cruz Biotechnology. Cy3, Cy5 and Alexa-Fluor-488-conjugated secondary antibodies were purchased from Jackson Immuno Research.

Immunoprecipitation, binding assay and dot blot assay

HeLa cells were transfected with the indicated plasmids using GeneJuice according to the manufacturer's instructions. After 24 (Fig. 1D) or 48 hours (Fig. 1B; Fig. 2C), cells were lysed in lysis buffer (50 mM Tris, 150 mM NaCl, 1 mM EDTA, 5 mM MgCl₂, 0.5% NP40, pH 7.4) supplemented with Complete Protease Inhibitor Cocktail (Roche). Lysates were incubated with GST fusion protein bound beads for 4 hours in binding assay or were incubated either with 3 µl of anti-GFP-coupled beads for 2 hours or 5 µl anti-OPTN antibody for 4 hours followed by a 1-hour incubation with Protein G agarose in immunoprecipitation assay. The washed beads were subjected to SDS-PAGE gels for immunoblot analysis. GST, MBP or His fusion proteins were expressed in E. coli BL21 cells in LB medium and purified using glutathione Sepharose, amylose resin or Ni-NTA agarose, respectively. MBP-tagged proteins were eluted from the amylose resin by 10 mM maltose, GST-tagged proteins with 20 mM reduced glutathione and His-tagged proteins with 200 mM imidazole. Supernatant containing eluted MBP or His-tagged proteins were used for subsequent GST pull-down assays with GST-proteins bound to beads. GST pull-down assays for recombinant protein-protein interactions were performed in 1× incubation buffer (20 mM Tris-HCl, pH 8.4; 150 mM NaCl; 2.5 mM CaCl₂; 10% glycerol, 1% Triton X-100, 1 mM DTT) for 4 hours at 4°C. Beads were washed three times with incubation buffer and subjected to SDS-PAGE gels for immunoblot analysis.

Dot blot assay was performed as previously described (Yerbury et al., 2013). Purified His-SOD1 G93C mutant (20 μ M) was incubated in 10 mM potassium phosphate buffer, pH 7.4 containing 5 mM EDTA and 10 mM DTT at 37°C for 20 hours for inducing *in vitro* aggregation. Aggregated His-SOD1 G93C (20 μ M) was incubated in absence or presence of GST-OPTN (20 μ M) for 1 hour at 37°C. All samples were centrifuged for 30 minutes at 4°C, soluble fraction were separated and pellet containing insoluble fraction was washed three times with potassium phosphate buffer. Soluble and insoluble fractions were spotted onto a nitrocellulose membrane and membrane was blotted with either His or GST antibody.

Immunofluorescence

HeLa cells were grown on glass coverslips and transfected with indicated plasmids. At 48 hours post-transfection cells were fixed with 2% paraformaldehyde (PFA) for 20 minutes. Subsequently, cells were permeabilized with 0.2% Triton X-100 solution in PBS at room temperature for 10 minutes. All cells were blocked in PBS containing 5% BSA at 4°C overnight. Primary and secondary antibodies were diluted in the blocking solution and incubated for 1 hour at room temperature. Washes were performed in 0.1% Tween 20 in PBS. The coverslips were mounted with aqueous mounting medium (Mowiol) containing DAPI (Molecular Probes). Images were acquired by the LSM 510 META laser-scanning microscope (Zeiss) and processed using ImageJ software.

Transgenic mice and tissue preparation

Transgenic C57/B6 mice expressing human mutant SOD1 G93A (Gurney et al., 1994) were kept in the local animal facility under a 12-hour light cycle and food and drinking water ad libitum. Mice were housed, anesthetized and sacrificed according to the European and German guidelines for the care and use of laboratory animals. The animals were sacrificed at 5 (no phenotype), 7 (early-stage of the disease) and 10 months (end-stage of the disease, i.e. when the mice were not able to right themselves after 10 seconds when put on their side) (Gurney et al., 1994). Animals were perfused transcardially with 4% PFA. The lumbar spinal cord was removed and post-fixed overnight. The tissue was then incubated in 20% sucrose in phosphate buffer overnight for cryoprotection. 15 µm spinal cord sections were cut on a cryostat and immunostaining was performed on floating sections. Briefly, spinal cord sections were permeabilized with 0.5% Triton X-100 in PBS and then washed with 0.1% Triton X-100 in PBS (PBST). After a 2-hour blocking with PBST containing 3% BSA, the sections were incubated with the primary antibodies directed against ubiquitin and TBK1 or phospho S177 OPTN at 4°C for 48 hours. Sections were then washed and incubated for 2 hours with the secondary antibodies. After washing, the sections were incubated overnight at 4°C with the Cy5conjugated anti-SOD1 antibody. Finally, sections were washed, mounted and observed under the LSM 510 META laser-scanning microscope (Zeiss).

Cell fractionation and filter trap assay

Cells were lysed in NP40 lysis buffer (50 mM Tris, 150 mM NaCl, 1 mM EDTA, 5 mM MgCl₂, 0.5% NP40, pH 7.4 supplemented with Complete Protease Inhibitor Cocktail). After centrifugation (10 minutes, 5000 g) the supernatant was collected and constituted the soluble fraction. The pellet was resuspended in NP40 buffer containing Bensonaze (Merck) and 2% SDS which represented the insoluble fraction. The soluble and insoluble fractions were analyzed by SDS-PAGE. The insoluble fraction was applied on dot-blot apparatus. Total cell lysate was obtained by lysing cells in RIPA buffer (50 mM Tris, 150 mM NaCl, 0.5% sodium deoxycholate, 1% SDS, 1% Triton-X 100, pH 7.4) supplemented with Complete Protease Inhibitor Cocktail at room temperature for 15 minutes.

Quantification of aggregate formation and statistics

Cells were analyzed on a Leica CTR7000 HS epifluorescence microscope. A minimum of 500 transfected cells were counted for each condition. The experiments were performed in triplicate and repeated twice. An example of cells containing SOD1 G93A aggregates is shown in Fig. 3A. Statistical significance was determined by Student's *t*-test for comparison between two samples. To compare three or more ungrouped or grouped samples were used respectively one-way ANOVA or two-way ANOVA followed by Bonferroni multiple comparison test as post-hoc analysis. All statistical analysis was performed using Graph PadPrism (Prism Software).

Zebrafish (Danio rerio)

The zebrafish were maintained, mated, and raised according to established procedures (Westerfield, 2000). The Tuebingen wild-type strain was used for all experiments. Morpholinos used in this study were purchased from Gene Tools (Philomath): Standard control morpholino 5'-CCTCTTACCTCAGTTACAATTTATA)-3', translation blocking morpholino targeted to start codon (OPTN ATG-AMO) 5'-CGATGATCCAGATGCCATGCTTTCT-3' and splice blocking morpholino exon2intron2 (OPTN splice-AMO) 5'-AAATTTCTCTCACCTCAGCTCCACT-3'. In vitro transcription of wild-type FLAG-SOD1 and FLAG-SOD1 G93A was performed with the MessageMachine T7 Kit (Ambion). Morpholinos were diluted in ultrapure water with 0.05% phenol red and injected into 2-8 cell stage zebrafish embryo as previously described (Laird and Robberecht, 2011). Dose-dependent curves of AMO and mRNA toxicity were performed and 1 nl of 0.5 mM AMOs (ctrl AMO, OPTN splice-AMO) or 0.4 mM OPTN ATG-AMO and 200 pg wild-type FLAG-SOD1 and FLAG SOD1 G93A mRNA were injected to minimize morpholino or cDNA-induced developmental delay and toxicity. After injections, embryos were stored in E3 medium (5 mM NaCl, 0.17 mM KCl, 0.33 mM CaCl₂, 0.33 mM MgSO₄) at 28°C. At 48 hours after fertilization, zebrafish embryos were fixed with 4% PFA in PBS and immunostained using SV2 antibody in order to visualize the motor neurons. Brightfield images were acquired using a Leica MZ165 dissecting microscope and fluorescent images were acquired using a LSM 510 META laser-scanning microscope (Zeiss). Motor neuron length was measured in ImageJ and the average measure of the first ten motor axons after the yolk sac was calculated in each embryo. Embryos were marked as embryos with abnormal branching when two or more axons per embryo showed premature branching. Zebrafish lysates were prepared in RIPA lysis buffer and analyzed by immunoblotting.

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