SSH1 impedes SQSTM1/p62 flux and MAPT/Tau clearance independent of CFL (cofilin) activation

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**ABSTRACT**

Accumulation of toxic protein assemblies and damaged mitochondria are key features of neurodegenerative diseases, which arise in large part from clearance defects in the Macroautophagy/autophagy-lysosome system. The autophagy cargo receptor SQSTM1/p62 plays a major role in the clearance of ubiquitinated cargo through Ser403 phosphorylation by multiple kinases. However, no phosphatase is known to physiologically dephosphorylate SQSTM1 on this activating residue. RNAi-mediated knockdown and overexpression experiments using genetically encoded fluorescent reporters and defined mutant constructs in cell lines, primary neurons, and brains show that SSH1, the canonical CFL (cofilin) phosphatase, mediates the dephosphorylation of phospho-Ser403-SQSTM1, thereby impairing SQSTM1 flux and phospho-MAPT/tau clearance. The inhibitory action of SSH1 on SQSTM1 is fully dependent on SQSTM1 Ser403 phosphorylation status and is separable from SSH1-mediated CFL activation. These findings reveal a unique action of SSH1 on SQSTM1 independent of CFL and implicate an inhibitory role of SSH1 in SQSTM1-mediated clearance of autophagic cargo, including phospho-MAPT/tau.

**Abbreviations:** AAV: adeno-associated virus; Aβ42\textsubscript{O}: amyloid β1-42 oligomers; AD: Alzheimer disease; CA3: cornu Ammonis 3; CSNK2/CK2: casein kinase 2; FCCP: 2-[2-[4-(trifluoromethoxy)phenyl]hydrazinyliden]-propanedinitrile; FTLD: frontotemporal lobar degeneration; GFP: green fluorescent protein; MAP1LC3/LC3: microtubule associated protein 1 light chain 3; SQSTM1/p62: sequestosome-1; PLA: proximity ligation assay; RFP: red fluorescent protein; RIPA: radioimmunoprecipitation assay; shRNA: short hairpin RNA; siRNA: small interfering RNA; Ser403: Serine403; SSH1: slingshot protein phosphatase 1; TBK1: TANK-binding kinase 1; ULK: unc-51 like kinase 1
Introduction

Accumulation of toxic protein assemblies and dysfunctional mitochondria are associated with synaptic and neuronal loss in multiple neurodegenerative disorders. Such accumulations are thought to arise in large part from clearance deficits in the autophagy-lysosome system [1–7]. Selective autophagy is a major type of autophagy characterized by the recognition and degradation of specific cargo, such as damaged organelles, misfolded proteins, or invading pathogens [8–14]. Mitophagy is a form of selective autophagy that specifically removes damaged mitochondria from cells, which is vital to mitochondrial quality control and cellular health [15–17].

SQSTM1/p62, hereafter referred to as SQSTM1, is a major selective autophagy cargo receptor that recognizes polyubiquitin conjugates on cellular organelles (i.e. mitochondria) and misfolded proteins to target them for autophagic clearance through its interaction with MAP1LC3/LC3, hereafter referred to as LC3 [18]. The activity of SQSTM1 is positively regulated by phosphorylation on Ser403 within the UBA domain, which serves to activate SQSTM1 and increase its affinity for polyubiquitin chains, thereby enhancing its ability to recognize misfolded proteins and dysfunctional mitochondria ubiquitin-tagged for clearance [18–20]. While kinases TBK1 (TANK-binding kinase 1), ULK (unc-51 like kinase 1), and CNK2/CK2 (casein kinase 2) are known to phosphorylate SQSTM1 on Ser403 [18,19,21–23], no phosphatase is known to physiologically dephosphorylate SQSTM1 on this activating residue. It has been reported that Ser403 phosphorylated SQSTM1 is significantly reduced in brains of Alzheimer disease (AD) patients [2], suggesting a failure of SQSTM1 to clear substrates as part of the disease mechanism. Moreover, ARRB2/β-arrestin2 oligomers were reported to suppress SQSTM1-mediated autophagy and promote tauopathy in frontotemporal lobar degeneration (FTLD-MAPT/tau) [24], and multiple ALS-FTLD-linked mutations in Tbk1 or SQSTM1 were also shown to reduce SQSTM1 phosphorylation on Ser403 and nearby Ser407 [25], implicating reduced SQSTM1 activity in the pathogenesis of AD, FTLD, and ALS.

Multiple studies have implicated the involvement of the F-actin-severing protein CFL (cofilin) in amyloid β (Aβ)-induced dendritic spine shrinkage [26,27], accumulation of CFL-actin aggregates/rods in AD brains [28,29], and increased CFL activity in brains of AD patients [27,30] and AD mouse models [31]. Specifically, Aβ42 oligomers (Aβ42O) rapidly induce the activation/dephosphorylation of CFL [30–33], resulting in depletion of F-actin-associated synaptic proteins and translocation of mitochondria to cytoplasm [31]. Interestingly, experimental reduction of SSH1 (slingshot protein phosphatase 1), a major CFL phosphatase [34], mitigates CFL activation and prevents Aβ42O-induced mitochondrial CFL translocation and toxicity [31]. In this study, we report the identification of SSH1 as the first phosphatase that dephosphorylates SQSTM1 on p-Ser403, resulting in significant physiological and pathological consequences. This function of SSH1 on SQSTM1, which is modular and separable from its activity on CFL, impairs SQSTM1-mediated selective autophagy, thereby impeding the clearance of damaged mitochondria and phospho-MAPT/tau.

Results

SSH1 impairs LC3 recruitment to mitochondria and impedes mitophagy

We previously showed that SSH1 reduction prevents Aβ42 oligomer (Aβ42O)-induced translocation of CFL to mitochondria, mitochondrial dysfunction, and toxicity [31,35,36]. As damaged mitochondria are targeted for clearance by mitophagy, we looked for evidence of mitophagy in hippocampus derived HT22 neuroblastoma cells treated with Aβ42O. LC3 is a central protein in mitophagy where it functions in autophagosome biogenesis and hence is used as an autophagosome marker [37]. Aβ42O was prepared as previously reported [31,38]. Aβ42O treatment (1 μM) weakly increased the number of endogenous LC3 puncta and their association with mitochondria (TOMM20) at 8 h (Figure 1A–C), indicative of mild mitophagy induction. However, both LC3 puncta and their association with mitochondria gradually subsided below control levels after 24–48 h (Figure 1A–C). In contrast, RNAi-mediated knockdown of SSH1 increased the number of endogenous LC3 puncta and their association with mitochondria at steady state, both of which were further increased at 8 h and remained highly elevated for up to 48 h post Aβ42O treatment (Figure 1A–C). Knockdown of SSH1 by siRNA was confirmed by immunocytochemical staining for SSH1 (Figure 1A) and western blotting (Figure 1D). To confirm this finding in a different way, we transfected HT22 cells with vector control or SSH1-Flag and depolarized mitochondria with the mitochondrial uncoupler FCCP for up to 4 h. As expected, FCCP treatment increased the number of endogenous LC3 puncta (Figure 1E,F), indicative of autophagosome formation. This was proportionately reflected in increased LC3 puncta colocalized with mitochondria (Figure 1E,F). SSH1 overexpression reduced the number of endogenous LC3 puncta at steady state and dramatically suppressed FCCP-induced increase in LC3 puncta (Figure 1E,F), which was also reflected in proportionately decreased endogenous LC3 puncta associated with mitochondria (Figure 1E,F). Similar results were obtained when co-transfected with RFP-LC3 to directly visualize RFP-LC3 puncta (Fig. S1AB). As the reduction of LC3 recruitment to mitochondria is expected to impair mitochondrial clearance, we assessed mitophagy using the mito-QC mitophagy flux reporter, in which the pH-sensitive mCherry-GFP protein is fused to FIS1 mito-targeting sequence [39]. Under steady-state conditions, the mitochondrial network fluoresces both red and green; however, upon mitophagy, mitochondria are delivered to lysosomes where mCherry fluorescence remains stable, but GFP fluorescence is quenched by the acidic environment. Hence acidified mCherry-only puncta represent mitochondria that are associated with lysosomes. Indeed, we confirmed that SSH1 overexpression significantly reduces mCherry-only mitochondria (Fig. S1C,D), whereas SSH1 siRNA significantly increases mCherry-only mitochondria (Fig. S1E,F).

SSH1 suppresses LC3-II biogenesis but does not alter LC3 flux

LC3-II is the active lipid modified form of LC3 involved in autophagosome membrane expansion and fusion [40,41]. As SSH1 overexpression significantly reduced the number of LC3
puncta, we tested if SSH1 alters LC3-II levels. SSH1 overexpression significantly reduced LC3-II at steady state (Fig. S2A,B). FCCP treatment for 4 h increased LC3-II levels as expected, and SSH1 overexpression also significantly suppressed FCCP-induced LC3-II generation (Fig. S2A,C). Conversely, SSH1 knockdown by siRNA increased LC3-II both at steady state and after FCCP treatment (Fig. S2D-F). The changes in LC3-II by SSH1 may be due to altered
biogenesis or clearance by the lysosome. To determine if the reduction in LC3-II by SSH1 is due to accelerated lysosomal clearance, we treated HT22 cells with FCCP (4 h) or rapamycin (16 h) with or without the lysosome inhibitor bafilomycin A1 (final 4 h). FCCP markedly increased LC3-II after 4 h but did not further increase with bafilomycin co-treatment, suggesting that FCCP-induced LC3-II generation is far greater than its turnover during this time period (Figure 2A,C). SSH1 transfection significantly reduced FCCP-induced LC3-II both in the presence and absence of bafilomycin A1 (Figure 2A,B). Rapamycin treatment over 16 h did not show a substantial increase in LC3-II; however, co-treatment of rapamycin with bafilomycin during the final 4 h treatment dramatically increased LC3-II in control transfected cells (Figure 2C,D), indicating rapamycin-induced LC3-II flux and clearance by the lysosome. This bafilomycin-induced accumulation of LC3-II was significantly blunted in SSH1 transfected cells (Figure 2C,D). SSH1 overexpression also significantly reduced LC3-II levels upon bafilomycin treatment alone (Fig. S2G,J). Hence, these results suggest that SSH1 suppresses LC3-II biogenesis at a step prior to fusion of LC3+ autophagosomes with lysosomes.

To assess the effects of SSH1 on LC3 in a dynamic way, we used the mCherry-GFP-LC3 reporter construct, which measures both total LC3 puncta and the rate of conversion of total LC3+ puncta (autophagosomes) to mCherry-only LC3+ puncta (autolysosomes & lysosomes). This reporter takes advantage of GFP’s sensitivity to low pH (quenching), mCherry’s insensitivity to low pH, and LC3, which allows for the tracking of LC3 to lysosomes [42–44]. Therefore, perfectly colocalized red and green puncta (yellow) are indicative of non-lysosomal LC3. However, upon fusion with lysosomes (autolysosomes), red puncta persist while green puncta disappear (red-only). HT22 cells were co-transfected with mChERRY-GFP-LC3 with vector control or SSH1-Flag and treated with or without FCCP for up to 4 h. As expected, FCCP increased the total number of LC3 puncta (Figure 2E,F) and red-only puncta (Figure 2E,G) in a time-dependent manner in control vector transfected cells. SSH1 overexpression significantly reduced the total number of LC3 puncta (Figure 2E,F) and red-only LC3 puncta (Figure 2E,G). However, the percentage of red-only puncta was not significantly altered by SSH1 overexpression (Figure 2H), indicating that SSH1 overexpression does not directly inhibit the rate of LC3+ autophagosome conversion to LC3+ autolysosomes (i.e. LC3 flux). Bafilomycin A1 treatment similarly increased the total number of LC3 puncta in control and SSH1 transfected cells (Fig. S2K,L), confirming that SSH1 overexpression does not alter lysosomal LC3 clearance. Converse experiments with Ssh1 siRNA showed that the loss of SSH1 similarly increases both total LC3 puncta and red-only LC3 puncta from 0 to 2 h of FCCP treatment (Figure 2I–K), thereby enhancing LC3 biogenesis but not significantly altering lysosomal LC3 flux during this period. After 4 h FCCP, however, we observed a significant increase in the percentage of red-only LC3 puncta (Figure 2L), indicating that the loss of SSH1 can promote LC3 flux after prolonged FCCP exposure. Transfection of mCherry-GFP displayed diffuse nuclear and cytoplasmic fluorescence without observable puncta (Fig. S2M), unlike those seen with mCherry-GFP-LC3.

**SSH1-mediated inhibition of SQSTM1 flux underlies LC3 suppression**

The observation that SSH1 significantly inhibits LC3-II generation and LC3 puncta formation while having little to no effects on LC3 flux led us to hypothesize that such SSH1-mediated effects originate upstream of LC3. The autophagosome protein LC3 binds to autophagy cargo receptors, such as SQSTM1 and OPTN (optineurin). These cargo receptors recognize polyubiquitinated cargo via the ubiquitin-binding domain and recruit LC3 through the LIR domain [18]. Interestingly, a recent study showed that the cargo-bound SQSTM1 also acts as an autophagy inducer, enhancing the synthesis of LC3 and its conversion to LC3-II to promote autophagosome formation [45]. Hence, we initially co-transfected HT22 cells with GFP-Sqstm1 or GFP-OPTN with vector control or SSH1-Flag. Neither total GFP-OPTN puncta number nor its recruitment to TOMM20+ mitochondria was altered by SSH1 transfection at steady state or after FCCP treatment (Fig. S3A–C). However, SSH1 overexpression significantly reduced the total number of GFP-SQSTM1 puncta and decreased GFP-SQSTM1 recruited to TOMM20+ mitochondria at steady state and after FCCP (Fig. S3D–F). To investigate SSH1-induced changes in SQSTM1 further, we assessed SQSTM1 flux using the mCherry-GFP-SQSTM1 reporter [44,46,47], which allows tracking of SQSTM1 puncta flux according to the same principle as mCherry-GFP-LC3. In vector control transfected HT22 cells, FCCP treatment generally increased total SQSTM1 puncta and red-only acidified SQSTM1 puncta (Figure 3A–C). However, SSH1 significantly reduced total SQSTM1 puncta and even more robustly reduced red-only SQSTM1 puncta even with FCCP treatment (Figure 3A–C). Hence, SSH1 significantly reduced the percentage of red-only SQSTM1 puncta compared to vector control (Figure 3A,D), indicating that SSH1 impedes SQSTM1 flux. Ssh1 siRNA transfection, in contrast, significantly increased the number of SQSTM1 puncta and those that were recruited to TOMM20+ mitochondria in the absence and presence of FCCP (Fig. S3G–I), indicating that endogenous SSH1 normally inhibits SQSTM1 function. To determine if SQSTM1 is required for the effects of SSH1 on LC3, we co-transfected control or Sqstm1 siRNA together with control or Ssh1 siRNA (Fig. S3J). As expected, Sqstm1 siRNA greatly reduced the number of endogenous LC3 puncta (Figure 3E,F) and their recruitment to mitochondria (Figure 3E,G,H), while Ssh1 siRNA dramatically increased total LC3 puncta (Figure 3F,E) and their recruitment to mitochondria (Figure 3E,G,H). However, Ssh1 siRNA failed to increase LC3 puncta or recruitment to mitochondria when SQSTM1 was knocked down (Figure 3E–H), indicating that the inhibitory effects of SSH1 on LC3 require endogenous SQSTM1.

**SSH1 decreases SQSTM1 Ser403 phosphorylation and inhibits SQSTM1-ubiquitin binding**

The activity of SQSTM1 is positively regulated by its phosphorylation on Ser403 within the C-terminal ubiquitin-associated (UBA) domain by TBK1, ULK, or CSNK2/CK2 [18,19,21–23]. This phosphorylation is known to activate SQSTM1 and
SSH1 suppresses LC3-II biogenesis but does not alter LC3 flux. (A) Representative blots from HT22 cells transfected with vector control or Flag-SSH1, treated ± 5 μM FCCP ± 200 nM bafilomycin A1 for 4 h, and subjected to western blotting for LC3, SSH1, and ACTB/actin. (B) Quantitative graph of figure A expressed as means ± SEM, two-way ANOVA, group F(1, 24) = 68.35, df = 1, P < 0.0001, followed by Sidak’s posthoc test, n = 4/condition, *p < 0.05, #p < 0.0001. (C) Representative blots from HT22 cells transfected with vector control or Flag-SSH1, treated ± 200 nM rapamycin for 16 h ± 200 nM bafilomycin A1 for 4 h, and subjected to western blotting for LC3, SSH1, and ACTB/actin. (D) Quantitative graph of figure C expressed as means ± SEM, two-way ANOVA, group F(1, 24) = 48.13, df = 1, P < 0.0001, followed by Sidak’s posthoc test, n = 4/condition, *p < 0.05, #p < 0.0001. (E) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with mCherry-GFP-LC3 and vector control (p3xFlag) or Flag-SSH1, treated with 5 μM FCCP for 4 h, 2 h, 0.5 h, or 0 h (DMSO), and subjected to staining for Flag-M2 (light blue) and direct fluorescence imaging (mCherry: red, GFP: green). (F–H) Quantitative graphs of figure E expressed as means ± SEM, 2-way ANOVA: (F)
increase SQSTM1 affinity for polyubiquitin chains, thereby enhancing its ability to recognize misfolded proteins or dysfunctional mitochondria tagged for clearance [20]. Importantly, the ratio of soluble Ser403-phosphorylated SQSTM1 (p-Ser403-SQSTM1) to total SQSTM1 is significantly reduced in brains of AD patients [2]. As SSH1 is a phosphatase, we hypothesized that the inhibitory action of SSH1 on SQSTM1 might be associated with SQSTM1 dephosphorylation on Ser403. Overexpression of SSH1-Flag in HT22 cells resulted in significantly reduced endogenous p-Ser403-SQSTM1 without altering total SQSTM1 (Figure 4A,B). Conversely, Ssh1 siRNA significantly increased endogenous p-Ser403-SQSTM1 without altering total SQSTM1 (Figure 4C-D).

To determine if SSH1-mediated reduction in p-Ser403-SQSTM1 alters SQSTM1 binding to ubiquitin, we assessed SQSTM1-ubiquitin complexes in 3 ways. First, we co-transfected HT22 cells with HA-ubiquitin, GFP-Sqstm1, and/or SSH1-Flag, and performed co-immunoprecipitation (co-IP) experiments to assess SQSTM1-ubiquitin complexes. Co-IP studies confirmed that SSH1 transfected cells contain significantly reduced GFP-SQSTM1 in HA-ubiquitin immune complexes (Figure 4Ei,Eii), despite equivalent levels of HA-ubiquitin, GFP-SQSTM1, and SSH1-Flag expression (Figure 4Ei,Eii). Second, we detected the colocalization of ubiquitin with SQSTM1 puncta by immunocytochemistry (ICC) from HT22 cells co-transfected with GFP-Sqstm1 with/without SSH1-Flag. We found significantly reduced numbers of GFP-SQSTM1 puncta (Figure 4F,G), number of GFP-SQSTM1 puncta colocalized with ubiquitin (Figure 4F,H), and percentage of GFP-SQSTM1 puncta colocalized with ubiquitin (Figure 4F,I) in SSH1-Flag transfected cells in the presence or absence of FCCP treatment. In contrast, Ssh1 shRNA transfection increased GFP-SQSTM1 puncta per cell (Figure 4J,K), GFP-SQSTM1 puncta colocalized with ubiquitin (Figure 4J,L), and percentage of GFP-SQSTM1 puncta colocalized with ubiquitin (Figure 4M) in FCCP treated cells. Third, we detected SQSTM1-ubiquitin complexes by DuoLink in situ proximity ligation assays (PLA) (Millipore Sigma), a powerful tool that allows in situ detection of protein-protein interactions with high specificity and sensitivity. In PLA, two antibodies (raised against different species) for corresponding proteins are applied, after which a pair of oligonucleotide-labeled secondary antibodies bind to primary antibodies. If in close proximity to each other (~40 angstroms) [48], connector oligos join the PLA probes and are ligated to form circular DNA that are amplified by DNA polymerase. This allows up to 1000-fold amplification, and complementary detection oligos coupled to a fluorochrome are hybridized to detect the PLA signal. Using PLA and two antibodies directed against SQSTM1 and ubiquitin, we detected endogenous SQSTM1-ubiquitin complexes, which were significantly reduced by ~70% in SSH1 transfected cells (Figure 4N,P; red PLA puncta) and increased by ~65% in Ssh1 shRNA transfected cells (Figure 4O,Q). Exclusion of 1 primary or 1 secondary antibody yielded no detectable PLA signal, indicating specificity of the assay (Figure S4). Therefore, these data collectively show that SSH1 reduces SQSTM1 phosphorylation on Ser403, which inhibits SQSTM1 binding to ubiquitinated substrates and downstream LC3 activation.

**SSH1-mediated SQSTM1 inhibition requires p-Ser403 modification of SQSTM1**

The C-terminal region of SQSTM1 contains the LC3-interacting LIR domain and the ubiquitin-interacting UBA domain (Figure S5A). To determine if these regions of SQSTM1 are required for the inhibitory effects of SSH1, we co-transfected the GFP-Sqstm1ΔC (lacking both LIR and UBA domains) with vector control or SSH1-Flag. SSH1 had no significant inhibitory effect on the number of GFP-SQSTM1ΔC puncta with or without FCCP treatment (Figure S5B,C), indicating that the C-terminal region is required for the SSH1-induced inhibition of SQSTM1. Next, we used the GFP-SQSTM1-LIR mutant, in which the LC3-binding site in the LIR domain of SQSTM1 has been mutated to prevent LC3 binding [49] (Figure S5A). In contrast to the GFP-SQSTM1ΔC mutant, SSH1 overexpression significantly reduced the number of GFP-SQSTM1-LIR puncta and number of GFP-SQSTM1-LIR puncta colocalized with TOMM20+ mitochondria (Figure S5D-F), indicating that SQSTM1-LC3 binding per se is not required for the inhibitory effect of SSH1 on SQSTM1. The N-terminal PBl domain is required for initial SQSTM1 self-interaction to allow the formation of SQSTM1 helical filaments arranged in a head to tail configuration [50]. Because SSH1 reduced the number of SQSTM1 puncta, we assessed whether SSH1 inhibits SQSTM1 self-interaction at the molecular level. However, we found no evidence that SSH1 affects SQSTM1 self-interaction by co-IP (Figure S5G) or PLA (Figure S5H) experiments, suggesting that SSH1 does not directly alter the PBl self-interaction domain.

To determine if the inhibitory actions of SSH1 on SQSTM1 depend on the UBA domain Ser403 phosphorylation status, we generated 2 artificial Sqstm1 constructs: GFP-Sqstm1S403A and GFP-Sqstm1S403E, the former that cannot be phosphorylated and the latter that mimics phosphorylation on Ser403 (Figure S5A). The SQSTM1S403E mutant has been shown to increase SQSTM1 binding affinity for ubiquitin, resulting in efficient targeting of polyubiquitinated substrates in sequestosomes for autophagosome entry [20]. Upon expression of these constructs in HT22 cells, we observed a significant ~65% increase in the number of GFP-SQSTM1S403E puncta compared to equivalent transfection of GFP-Sqstm1S403A at steady state (Figure 5A,B). Manders split colocalization with TOMM20+ mitochondria demonstrated an even more significant ~3-fold increase in GFP-SQSTM1S403E recruitment to mitochondria compared to GFP-SQSTM1S403A

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*group F(1, 91) = 51.64, df = 1, P < 0.0001; (G) group F(1, 91) = 44.65, df = 1, P < 0.0001; (H) group F(1, 91) = 2.190, df = 1, P = 0.1424, followed by Sidak’s posthoc test, n ≥ 10/condition from 3 different experiments, *p < 0.05, **p < 0.005, **p < 0.0001. (I) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with mCherry-GFP-LC3 and control shRNA or Ssh1-shRNA, treated with 5 μM FCCP for 4 h, 2 h, 0.5 h, or 0 h (DMSO), and subjected to staining for SSH1 (blue), DAPI (light blue) and direct fluorescence imaging (mCherry: red, GFP: green). (J–L) Quantitative graphs of figure I expressed as means ± SEM, 2-way ANOVA: (I) group F(1, 88) = 35.09, df = 1, P < 0.0001; (K) group F(1, 88) = 35.12, df = 1, P < 0.0001; (L) group F(1, 88) = 2.464, df = 1, P = 0.1201, followed by Sidak’s posthoc test, n ≥ 10/condition from 3 different experiments, *p < 0.05, **p < 0.005.*
SSH1-mediated inhibition of SQSTM1 underlies LC3 suppression. (A) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with mCherry-GFP-Sqstm1 and control (p3xFlag) or Flag-SSH1, treated with 5 μM FCCP for 4 h, 2 h, 0.5 h, or 0 h (DMSO), and subjected to staining for Flag-M2 (dark blue) or direct imaging (mCherry: red, GFP: green). (B-D) Quantitative graphs of figure A expressed as means ± SEM, 2-way ANOVA: (B) group F(1, 86) = 61.16, df = 1, P < 0.0001; (C) group F(1, 86) = 71.52, df = 1, P < 0.0001; (D) group F(1, 86) = 104.6, df = 1, P < 0.0001, followed by Sidak’s posthoc test, n ≥ 10/condition from 3 different experiments, *p < 0.05, **p < 0.005, ***p < 0.0005, #p < 0.0001.

(E) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells transfected with control siRNA, Ssh1 siRNA, Sqstm1 siRNA or both Ssh1 and Sqstm1 siRNA, treated with 5 μM FCCP for 4 h, 2 h, 0.5 h, or 1 h (DMSO), incubated with MitoTracker-Green for 45 min before fixation, and subjected to staining for LC3 (red), SQSTM1 (light blue), and SSH1 (dark blue) or direct imaging (LC3: red). (F-H) Quantitative graphs of figure E expressed as means ± SEM, 2-way ANOVA: (F) group F(3, 251) = 23.63, df = 3, P < 0.0001; (G) group F(3, 251) = 207.8, df = 3, P < 0.0001; (H) group F(3, 251) = 115.6, df = 3, P < 0.0001, followed by Sidak’s posthoc test, n ≥ 15/condition from 3 different experiments, *p < 0.05, **p < 0.005, #p < 0.0001.
SSH1 decreases SQSTM1 Ser403 phosphorylation and inhibits SQSTM1-ubiquitin binding. (A) Representative blots of p-S403-SQSTM1 from HT22 cells transfected with vector control (p3xFlag) or Flag-SSH1. (B) Quantitative graph of figure A expressed as means ± SEM, 2-tailed t-test, *p < 0.05. (C) Representative blots of p-S403-SQSTM1 from HT22 cells transfected with control siRNA or Ssh1-siRNA. (D) Quantitative graph of figure C expressed as means ± SEM, 2-tailed t-test, *P = 0.0115, n = 4/condition. (E) Representative blots showing co-immunoprecipitation of HA-ubiquitin and GFP-SQSTM1 in HT22 cells co-transfected with HA-ubiquitin, GFP-Sqsm1 and control (p3xFlag) or Flag-SSH1. Figure Eiii shows quantitative graph of figure Ei (SQSTM1-ubiquitin co-IP) expressed as means ± SEM, 2-way ANOVA: (G) group F(1, 124) = 116.2, df = 1, P < 0.0001; (H) group F(1, 124) = 741.3, df = 1, P < 0.0001; (I) group F(1, 124) = 484.6, df = 1, P < 0.0001, followed by Sidak’s posthoc test, n ≥ 20/condition from 3 different experiments, *p < 0.05, #p < 0.0001. (F) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with GFP-Sqsm1 and vector control (p3xFlag) or Flag-SSH1, treated with 5 μM FCCP for 4 h, 0.5 h, or 0 h (DMSO), and subjected to staining for Flag-M2 (light blue), and ubiquitin (red) and direct fluorescence imaging (green: GFP). (G-I) Quantitative graphs of figure F expressed as means ± SEM, 2-way ANOVA: (G) group F(1, 124) = 116.2, df = 1, P < 0.0001; (H) group F(1, 124) = 741.3, df = 1, P < 0.0001; (I) group F(1, 124) = 484.6, df = 1, P < 0.0001, followed by Sidak’s posthoc test, n ≥ 20/condition from 3 different experiments, *p < 0.05, #p < 0.0001. (J) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells
(Figure 5A,B), indicating that Ser403 “phosphorylated” SQSTM1 is indeed more active in enhancing puncta number and recruitment to mitochondria. FCCP treatment (4 h) further significantly increased the number of total GFP-SQSTM1\textsuperscript{5403E} puncta and association with mitochondria compared to GFP-SQSTM1\textsuperscript{5403A} (Figure 5A,B). GFP-SQSTM1\textsuperscript{5403A} was completely unresponsive to FCCP in both measures (Figure 5B), indicating that mitochondrial depolarization recruits only Ser403-phosphorylated SQSTM1. Overexpression of SSH1-Flag had no effect on either GFP-SQSTM1\textsuperscript{5403E} or GFP-SQSTM1\textsuperscript{5403A} on the number of SQSTM1 puncta or their recruitment to mitochondria in the absence or presence of FCCP (Figure 5A,C,D). Likewise, shRNA-mediated knockdown of SSH1 had no significant effect on either GFP-SQSTM1\textsuperscript{5403E} or GFP-SQSTM1\textsuperscript{5403A} on the number of puncta or recruitment to mitochondria in the absence or presence of FCCP (Fig. S5J–O), collectively demonstrating that the inhibitory action of SSH1 on SQSTM1 requires SQSTM1 modification on Ser403.

Ser403 phosphorylation of SQSTM1 has been shown to inhibit the dimerization of the UBA domain independent of the PR1 domain [50]. Hence, we set out to measure the sizes of mutant GFP- SQSTM1\textsuperscript{5403A/E} puncta. Due to the wide range of puncta sizes, we first assessed puncta size by 4 increasing size categories. The SQSTM1\textsuperscript{5403A} mutant significantly increased the distribution of puncta to the 2 largest size categories (>5 μm\textsuperscript{2} & 1–5 μm\textsuperscript{2}) compared to the SQSTM1\textsuperscript{5403E} mutant (Figure 5E), and the average size of SQSTM1\textsuperscript{5403A} puncta in these categories were significantly larger than SQSTM1\textsuperscript{5403E} puncta (Figure 5F). We also assessed wild type GFP-SQSTM1 puncta sizes with or without SSH1-Flag transfection. Like the SQSTM1\textsuperscript{5403A} mutation, SSH1 significantly increased the distribution of GFP-SQSTM1 puncta to the 2 largest size categories compared to vector control (Figure 5G,H), and SSH1 transfected cells contained significantly larger GFP-SQSTM1 puncta compared to vector control in these size categories (Figure 5I). As expected, SSH1 also significantly reduced the total number of GFP-SQSTM1 puncta and those colocalized with TOMM20\textsuperscript{+} mitochondria (Fig. S5P,Q). These data therefore demonstrate that SSH1-mediated reduction in p-Ser403 phosphorylation recapitulates the SQSTM1\textsuperscript{5403A} mutation in all measures and suggest that loss of p-Ser403 phosphorylation decreases SQSTM1 puncta number by coalescing smaller puncta into larger puncta.

**SSH1\textsuperscript{ΔN} dephosphorylates p-Ser403-SQSTM1 but not p-Ser3-CFL and inhibits SQSTM1 activity**

SSH1 contains an N-terminal phospho-CFL binding site (p-CFL) within the N-terminal region, followed by the catalytic domain (CAT) and a sizable C-terminal region of unknown function (Figure 6A) [51]. It has been shown that SSH1\textsuperscript{ΔC} (or N461), containing the N-terminal 461 residues but lacking C-terminal residues 462–1049, is sufficient to dephosphorylate CFL on Ser3 (p-S3-CFL), whereas the C395S mutation in the catalytic site destroys SSH1 phosphatase activity [51]. To determine which domains of SSH1 are required for pSer403 SQSTM1 dephosphorylation, we obtained the SSH1\textsuperscript{ΔN} construct [51] and generated the SSH1\textsuperscript{ΔN} and SSH1\textsuperscript{ΔN,C395S} constructs lacking the N-terminal 307 amino acids (p-CFL binding region) but containing the catalytic domain and the C-terminal region (Figure 6A). As expected, overexpression of SSH1\textsuperscript{ΔN} robustly reduced p-Ser3-CFL (Figure 6B); however, SSH1\textsuperscript{ΔC} failed to alter pS403-SQSTM1 (Figure 6B,C). In contrast, neither SSH1\textsuperscript{ΔN} nor SSH1\textsuperscript{ΔN,C395S} altered p-Ser3-CFL (Figure 6B), and SSH1\textsuperscript{ΔN} but not SSH1\textsuperscript{ΔN,C395S} strongly reduced p-Ser403-SQSTM1 (Figure 6B,C). To determine if SSH1 can directly dephosphorylate p-Ser403-SQSTM1, we expressed and purified SSH1\textsuperscript{ΔN-His, SSH1\textsuperscript{ΔN-His, and SSH1\textsuperscript{ΔN-His} produced no effect on the activity of SSH1\textsuperscript{ΔN-His} detected by co-immunoprecipitation of GFP-SQSTM1, whereas little to no SSH1\textsuperscript{ΔN-Flag} was present when no GFP-Sqstm1 was transfected or pulled down with IgG beads alone (Fig. S6Bii,C).

To test whether SSH1\textsuperscript{ΔN} and SSH1\textsuperscript{ΔC} differentially alters SQSTM1 activity, we co-transfected SSH1\textsuperscript{ΔC-Flag} or SSH1\textsuperscript{ΔN-Flag} with GFP-Sqstm1 in HT22 cells. SSH1\textsuperscript{ΔN} significantly reduced both the number of GFP-SQSTM1 puncta and GFP-SQSTM1 puncta recruited to TOMM20\textsuperscript{+} mitochondria at steady state and after treatment with FCCP (Figure 6E–G), essentially identical to that seen with full-length SSH1. In contrast, the catalytically dead SSH1\textsuperscript{ΔN,C395S} mutant failed to alter GFP-SQSTM1 puncta or their association with mitochondria (Fig. S6D–F), indicating that the catalytic phosphatase activity is required for these effects. Moreover, SSH1\textsuperscript{ΔC} had no significant effect on either GFP-SQSTM1 puncta number or its recruitment to mitochondria (Figure 6H–I), indicating that the sole ability of SSH1 to dephosphorylate CFL is insufficient for inhibition of SQSTM1. As expected, SSH1\textsuperscript{ΔN} significantly reduced endogenous SQSTM1-ubiquitin complexes in PLA experiments (Figure 6K,L). Hence, these data collectively demonstrate the modular activity of SSH1 on SQSTM1 via dephosphorylation, which requires catalytic activity on SQSTM1 and is separable from that on CFL.
SSH1-mediated SQSTM1 inhibition requires p-Ser403 modification of SQSTM1. (A) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with GFP-Sqstm1^S403E or GFP-Sqstm1^S403A and mito-dsRed ± vector control or Flag-SSH1, treated with 5 μM FCCP for 4 h or 0 h (DMSO), and subjected to staining for Flag-M2 (dark blue) and DAPI (light blue) or direct imaging (mito-dsRed; red, GFP-SQSTM1: green). (B-D) Quantitative graphs of figure A expressed as means ± SEM, 2-way ANOVA: (B) upper graph, group F(1, 62) = 70.97, df = 1, P < 0.0001; lower graph, group F(1, 62) = 63.82, df = 1, P = 0.0036; (C) upper graph, group F(1, 60) = 0.002671, df = 1, P = 0.9590; lower graph F(1, 60) = 1.19, df = 1, P = 0.2855; (D) upper graph F(1, 60) = 0.3904, df = 1, P = 0.5350, followed by Sidak’s posthoc test, n ≥ 15/condition from 3 different experiments, **p < 0.005, ***p < 0.0005, #p < 0.0001. (E) Quantitative graphs of figure A showing the distribution of SQSTM1 puncta number across size categories, χ^2 test for trend, n ≥ 15/condition from 3 different experiments, χ^2 = 9.570, df = 1, P = 0.0019 (FCCP 0 h), χ^2 = 10.13, df = 1, P = 0.0015 (FCCP 4 h). (F) Quantitative graphs of figure A expressed as means ± SEM in different size categories, 2-way ANOVA, group F(1, 53) = 24.82, df = 1, P < 0.0001 (>1μm^2), group F(1, 53) = 4.206, df = 1, P = 0.0452 (<1μm^2) followed by Sidak’s posthoc test, n ≥ 12/condition from 3 different experiments, *p < 0.05, **p < 0.005. (G) Representative images at 4 h FCCP condition with white insets.
Endogenous SSH1 and SSH1ΔN impair SQSTM1 flux and ubiquitin binding in primary neurons

To confirm the effects of SSH1 on SQSTM1 in primary neurons, we generated and purified high titer (>10^{12} vg/mL) control and SSH1ΔN rAAV9, control and SSH1-shRNA rAAV9, as well as mCherry-GFP-Sqstm1 rAAV9. On DIV4, we transduced wild type neurons with mCherry-GFP-Sqstm1 rAAV9 and control or SSH1ΔN rAAV9. In a different set of experiments, we also transduced neurons with mCherry-GFP-Sqstm1 rAAV9 with control or SSH1-shRNA rAAV9. On DIV20, we quantified SQSTM1 puncta, acified red-only SQSTM1 puncta, and percentage of acified red-only SQSTM1 puncta with or without FCCP treatment for 4 h. Expression of SSH1ΔN significantly decreased total SQSTM1 puncta (Figure 7A,B), red-only puncta (Figure 7A,C), and percentage of red-only puncta (Figure 7A,D) in both the absence and presence of FCCP. Conversely, SSH1 shRNA significantly increased total SQSTM1 puncta (Figure 7E,F), acified red-only SQSTM1 puncta (Figure 7E,G), and percentage of red-only SQSTM1 puncta (Figure 7E,H) together with loss of SSH1 expression (Figure 7E) in both the absence and presence of FCCP, indicating that SSH1 reduction promotes SQSTM1 flux to lysosomes in neurons. Assessment of endogenous SQSTM1-ubiquitin complexes by in situ PLA confirmed that SSH1ΔN significantly reduces the SQSTM1-ubiquitin complex (Figure 7I,J), whereas SSH1 shRNA significantly enhances the SQSTM1-ubiquitin complex (Figure 7K,L). Western blotting from primary neuronal lysates confirmed the expression of SSH1ΔN with the expected reduction in p-Ser403-SQSTM1 (Fig. 5A), as well as efficient knockdown of endogenous SSH1 by SSH1-shRNA with the expected increase in p-Ser403-SQSTM1 (Fig. 5B). An example of high titer rAAV9 transduction efficiency with SSH1ΔN is shown in primary neurons (Fig. S7C).

SSH1ΔN exacerbates and SSH1 reduction partially rescues SQSTM1 flux in MAPT/tau^{P301S} brains

Previous studies have shown that SQSTM1 targets misfolded MAPT/tau for autophagic clearance. Specifically, Sqstm1^{−/−} mice display severe neurodegeneration as well as hyperphosphorylated tau and neurofibrillary tangles [52], and SQSTM1 overexpression strongly reduces pathogenic MAPT/tau in transfected cells and in vivo [53]. Conversely, accumulation of pathogenic MAPT/tau adversely deregulates autophagy, as evidenced by abnormal accumulation of autophagosomes and LC3 [7,54–56], suggestive of a maladaptive feed-forward cycle of tauopathy and autophagy deregulation. However, SQSTM1 flux in the setting of tauopathy in vivo has not been investigated. Thus, we used the MAPT/tau^{P301S} mice (PS19), in which tauopathy starts at 3 months and progressively becomes more filamentous at 6 months of age [57]. To assess whether SQSTM1 flux is altered in MAPT/tau^{P301S} brains and whether the non-CFL component of SSH1 (SSH1ΔN) alters SQSTM1 flux in vivo, we injected high titer (>10^{12} vg/mL) mCherry-GFP-Sqstm1 rAAV9 with control or SSH1ΔN-Flag rAAV9 into the hippocampus of 3-month-old MAPT/tau^{P301S} mice. We also injected control wild type littermate hippocampus with mCherry-GFP-Sqstm1 rAAV9 and vector control rAAV9. Two months post-injection, we processed brains for histological detection and quantification of mCherry, GFP, SSH1, and phospho-MAPT/tau (p-S199/p-S202) in the CA3 region of the hippocampus where mCherry-GFP-SQSTM1 was homogeneously expressed. Total SQSTM1 puncta number per cell was significantly higher in MAPT/tau^{P301S} CA3 compared to wild type CA3 by ~40% (Figure 8A,B). The percentage of acified red-only SQSTM1 puncta was close to 80% in wild type CA3 (Figure 8A,Bii), indicating healthy SQSTM1 flux; however, the percentage of acified red-only SQSTM1 puncta was significantly reduced to ~50% in CA3 of MAPT/tau^{P301S} brains (Figure 8A,Bii), indicative of SQSTM1 flux impairment in MAPT/tau^{P301S} brains. Similar SQSTM1 flux impairment was observed in DIV20 MAPT/tau^{P301S} primary neurons compared to littermate WT neurons (Fig. S8A,B), indicating an early phenotype intrinsic to MAPT/tau^{P301S} expression. Expression of SSH1ΔN in MAPT/tau^{P301S} brains significantly reduced total SQSTM1 puncta number per cell (Figure 8A,Bii) as well as percentage of acified red-only SQSTM1 puncta such that less than 30% of SQSTM1 puncta were red-only (Figure 8A,Bii), indicating that SSH1ΔN worsens SQSTM1 flux in MAPT/tau^{P301S} brains. This exacerbated impeded SQSTM1 flux by SSH1ΔN was accompanied by a significant 30% increase in p-S199/p-S202-MAPT/tau immunoreactivity in CA3 (Figure 8C,D), indicating that SSH1ΔN impairs SQSTM1-mediated phospho-MAPT/tau clearance. Indeed, western blotting from dissected hippocampus confirmed the expression of SSH1ΔN-Flag together with markedly increased total MAPT/tau and phospho-MAPT/tau but equivalent levels of mCherry-GFP-SQSTM1 (GFP antibody) and ACTB/actin (Figure 8E). As seen in MAPT/tau^{P301S} brains, overexpression of full-length SSH1 or SSH1ΔN but not SSH1AC significantly increased MAPT/tau levels in Hela-V5-MAPT/tau cells [58] stably expressing human wild type MAPT/tau (4R0N) (Fig. S8E-H), confirming that the SQSTM1 but not CFL component of SSH1 increases MAPT/tau levels.

We next assessed whether knockdown of endogenous SSH1 can rescue the SQSTM1 flux impendence in MAPT/tau^{P301S} brains. We produced a new batch of high titer magnified. HT22 cells co-transfected with GFP-Sqstm1-1WT, mito-dsRed, and vector control (p3AFlag) or Flag-SSH1, treated with 5 μM FCCP for 4 h or 0 h (DMSO), and subjected to staining for Flag-M2 (dark blue) and DAPI (light blue) or direct imaging (GFP-SQSTM1: green, mito-dsRed: red). (H) Quantitative graphs of figure G showing the distribution of SQSTM1 puncta number across size categories. χ^2 test for trend, χ^2 = n ≥ 15/condition from 3 different experiments, χ^2 = 33.43, df = 1, P < 0.0001 (FCCP 0 h), χ^2 = 9.467, df = 1, P = 0.0021 (FCCP 4 h). (I) Quantitative graphs of figure H expressed as mean ± SEM in different size categories, 2-way ANOVA, group F(1, 66) = 19.89, df = 1, P < 0.0001 (>1μm²); group F(1, 66) = 0.7743, df = 1, P = 0.3821 (<1μm²), followed by Sidak’s posthoc test, n ≥ 15/condition from 3 different experiments, *p < 0.05, **p < 0.005.
**Figure 6.** Direct dephosphorylation of p-Ser403-SQSTM1 by SSH1 is separable from CFL dephosphorylation. (A) Schematic of full length SSH1 and SSH1 mutants. (B) Representative blots of total SQSTM1, p-Ser403-SQSTM1 and p-Ser3-CFL from HT22 cells transfected with vector control (p3xFlag), Flag-SSH1ΔN, Flag-SSH1ΔN(393S) or CFP-SSH1ΔC. (C) Quantitative graph of figure B expressed as means ± SEM, 1-way ANOVA, F(3, 12) = 54.20, df = 3, P < 0.0001, followed by Tukey’s posthoc test, n = 4/condition, #p < 0.0001 (control vs. SSH1ΔN). (D) Representative blots from recombinant SQSTM1 purified from Sf9 insect cells incubated with recombinant SSH1 variants for the indicated times and immunobotted for p-S403-SQSTM1, showing dephosphorylation of SQSTM1 by SSH1ΔN but not SSH1ΔC. (E) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with GFP-Sqstml and vector control (p3xFlag) or Flag-SSH1ΔN, treated with 5 μM FCCP for 4 h, 2 h, 0.5 h, or 0 h (DMSO), and subjected to staining for Flag-M2 (dark blue) and TOMM20 (red) or direct imaging for GFP-SQSTM1 (green). (F and G) Quantitative graphs of figure E expressed as means ± SEM, 2-way ANOVA: (F) group F(1, 95) = 24.58, df = 1, P < 0.0001; (G) group F(1, 95) = 39.51, df = 1, P < 0.0001, followed by Sidak’s posthoc test, n ≥ 15/condition from 3 different experiments, *P < 0.05**P < 0.0005. (H) Representative images at 4 h FCCP condition with white insets magnified. HT22 cells co-transfected with GFP-Sqstml and vector control (p3xFlag), Flag-SSH1, or Flag-SSH1ΔC, treated with 5 μM FCCP for 4 h, 2 h, 0.5 h, or 0 h.
(>10^{12} \text{ GC/mL}) mCherry-GFP-Sqstm1 rAAV9 and injected mCherry-GFP-Sqstm1 rAAV9 together with control or Ssh1-shRNA rAAV9 into the hippocampus of MAPT/tau-P301S mice at 3 months of age and processed brains for assessment at 5 months of age. The percentage of red-only SQSTM1 puncta were again close to 50% in control rAAV9 injected MAPT/tau-P301S hippocampal CA3 (Figure 8F,Gii). However, Ssh1-shRNA rAAV9 significantly increased the percentage of red-only puncta to ~70% and markedly reduced endogenous Ssh1 staining in CA3 (Figure 8F,Gii), indicating that Ssh1 reduction partially rescues the SQSTM1 flux impairment in MAPT/tau-P301S brains. This enhancement of SQSTM1 flux by Ssh1 shRNA was accompanied by a significant 30% decrease in p-S199/p-S202-MAPT/tau immunoreactivity in CA3 (Figure 8H,I), indicating that Ssh1 reduction enhances SQSTM1-mediated phospho-MAPT/tau clearance. Western blotting from dissected hippocampus confirmed that Ssh1-shRNA lowers endogenous Ssh1 by ~70% together with marked reduction in total MAPT/tau and phospho-MAPT/tau but equivalent levels of mCherry-GFP-SQSTM1 (GFP antibody) and ACTB/actin (Figure 8J). Therefore, these results taken together demonstrate that while MAPT/tau-P301S expression impedes SQSTM1 flux in vivo, Ssh1ΔN expression significantly exacerbates SQSTM1 flux and increases phospho-MAPT/tau levels, whereas knockdown of endogenous Ssh1 partially rescues SQSTM1 flux and significantly lowers phospho-MAPT/tau levels in MAPT/tau-P301S brains.

**Discussion**

Accumulation of misfolded proteins and damaged mitochondria constitute pathological signatures of neurodegenerative diseases. These features, in large part, are attributed to impairments in selective autophagy and mitophagy, vital cellular processes that remove polyubiquitinated substrates tagged for clearance [59–61]. SQSTM1 is a major selective autophagy cargo receptor that plays a key role in targeting misfolded MAPT/tau and damaged mitochondria for degradation. This is achieved by the activating phosphorylation of SQSTM1 on Ser403 by ULK1, TBK1, or CSNK2/C2K, which increases SQSTM1 affinity for polyubiquitinated substrates [18,19,21] and activates the recruitment of LC3 [18,37]. This phosphorylation is transient and subject to rapid dephosphorylation [19]. However, to date, the identity of phosphatases that physiologically mediate SQSTM1 dephosphorylation on p-Ser403 has remained mysterious. In this study, we identified Ssh1 as the first phosphatase that deactivates SQSTM1 by dephosphorylation of p-Ser403 (Figure 9, schematic model). This event resulted in fewer but larger SQSTM1 puncta and reduced binding to ubiquitinated substrates, thereby significantly slowing SQSTM1 flux. As cargo-bound SQSTM1 acts as an autophagy inducer by enhancing the synthesis of LC3 and its conversion to LC3-II [45], Ssh1-mediated dephosphorylation of SQSTM1 also resulted in impaired downstream LC3 puncta formation, reduced LC3-II generation, and slowed mitophagy without directly altering lysosomal LC3 flux. The inhibitory actions of Ssh1 on SQSTM1 were dependent on the UBA domain Ser403 SQSTM1 phosphorylation site, and Ssh1ΔN but not Ssh1ΔC was sufficient to directly dephosphorylate p-Ser403-SQSTM1 and inhibit SQSTM1 activity, thereby separating the canonical function of Ssh1 on CFL from SQSTM1. This inhibitory action of Ssh1 on SQSTM1 was validated in primary neurons and *in vivo*. Specifically, SQSTM1 flux was impaired in MAPT/tau-P301S primary neurons and brains, which was exacerbated by expression of Ssh1ΔN in brain resulting in increased phospho-MAPT/tau accumulation. Conversely, shRNA-mediated knockdown of endogenous Ssh1 partially rescued SQSTM1 flux in MAPT/tau-P301S brains resulting in reduced phospho-MAPT/tau levels. These results taken together identify a novel function of Ssh1 as an upstream inhibitor of SQSTM1, separate from its action on CFL, and implicate the Ssh1-SQSTM1 pathway in impaired clearance of misfolded MAPT/tau.

**Inhibition of SQSTM1 by Ssh1-mediated dephosphorylation of p-Ser403**

The ability of SQSTM1 to function as a selective autophagy cargo receptor depends on three essential domains. N-terminal PB1 domain mediates the self-interaction of SQSTM1 allowing the formation of SQSTM1 helical filaments arranged in a head to tail configuration [50]. These filaments function as scaffolds for the binding to multiple LC3 through the LIR domain, thereby allowing the nucleation of the autophagosome membrane together with SQSTM1 [18]. The UBA domain of SQSTM1 mediates the binding to ubiquitinated cargo [20], and such binding is regulated by several processes. First, homodimerization of the UBA domain precludes polyubiquitin binding and contributes to SQSTM1 puncta abundance [62,63], as deletion of the UBA domain impairs both ubiquitin binding and formation of SQSTM1 bodies even in the presence of the PB1 domain [20,50]. Second, Ser403 phosphorylation within the UBA domain destabilizes UBA homodimerization and liberates the UBA domain from dimeric repression [18], thereby facilitating the binding to polyubiquitinated cargo. A third regulatory process via direct ubiquitination of the UBA domain on K420 has also recently been shown to modulate binding to ubiquitinated cargo [64,65]. Our observation that Ssh1 dephosphorylates p-Ser403 of SQSTM1, reduces SQSTM1 puncta number while increasing puncta size, and decreases ubiquitin binding is consistent with the expected increase in UBA domain

(DMSO), and subjected to staining for Flag-M2 (light blue) and TOMM20 (red) or direct imaging for GFP-SQSTM1 (green). (I) Quantitative graph of figure H expressed as means ± SEM, 2-way ANOVA: (I) group F(2, 187) = 30.81, df = 2, P < 0.0001; (J) group F(2, 187) = 49.31, df = 2, P < 0.0001, followed with Sidak’s posthoc test, n ≥ 20/condition from 3 different experiments, *p < 0.05, **p < 0.005, ***p < 0.0001. (K) Representative images from HT22 cells transfected with vector control (p3XFlag) or Flag-SHSH1ΔN, subjected to *in situ* PLA for SQSTM1-ubiquitin complexes (red), and staining for Flag-M2 (green) and DAPI (light blue). (L) Quantitative graph of figure K expressed as means ± SEM, 2-tailed t-test, P < 0.0001, n ≥ 20/condition from 3 different experiments, ***p < 0.0001.
homodimerization by p-Ser403 dephosphorylation. Our co-IP and PLA experiments to detect SQSTM1 self-interaction at the molecular level, however, did not find a measurable change in overall SQSTM1 self-interaction, which is mechanistically distinct to the recently reported role of ARRB2/β-arrestin2 oligomers in inhibiting SQSTM1 self-interaction [24]. This is likely due to the sufficiency of the PB1 domain to maintain SQSTM1 self-interaction at the molecular level, whereas puncta size is governed by PB1 oligomerization and UBA dimerization, the latter which is inhibited by Ser403 phosphorylation [18]. In addition, a recent study showed that polyubiquitin binding to the UBA domain promotes fragmentation of SQSTM1 filaments [50]. Hence, SSH1-induced inhibition of SQSTM1 ubiquitin binding may also account for the observed reduction in SQSTM1 puncta number but increased puncta size.

Consistent with either or both scenarios, we validated our findings using the SQSTM1S403E phosphomimetic mutant, which produced an identical phenotype to that seen with SSH1 knockdown resulting in increased SQSTM1 puncta number and reduced the puncta size. Conversely, SQSTM1S403A “inactive” mutant produced an identical phenotype to that seen with SSH1 overexpression, resulting in reduced SQSTM1 puncta number and increased puncta size. Neither SSH1 knockdown nor overexpression could alter these molecular signatures of SQSTM1S403E or SQSTM1S403A mutants, demonstrating the requirement of Ser403 modification in SSH1-induced inhibition of SQSTM1. These results therefore indicate that SSH1-mediated dephosphorylation of p-Ser403-SQSTM1, like the SQSTM1S403A mutant, decreases the number of active SQSTM1 puncta by coalescing smaller puncta into larger

Figure 7. Endogenous SSH1 and SSH1ΔN impair SQSTM1 flux and SQSTM1-ubiquitin binding in primary neurons. (A) Representative images at 4 h FCCP condition with white insets magnified. WT primary hippocampal neurons co-transduced with rAAV9-mCherry-GFP-Sqstm1 and rAAV-Flag or Flag-SSH1ΔN, treated with 2 μM FCCP for 4 h or 0 h (DMSO) and subjected to staining for Flag-M2 (dark blue) and DAPI (light blue) or direct imaging (mCherry: red, GFP: green). (B-D) Quantitative graph of figure A expressed as means ± SEM, 2-way ANOVA: (B) group F(1, 56) = 12.71, df = 1, P = 0.0008; (C) group F(1, 56) = 35.44, df = 1, P < 0.0001; (D) group F(1, 56) = 19.58, df = 1, P < 0.0001, followed by Sidak's posthoc test, n ≥ 15/condition from 3 different experiments, *p < 0.05, ***p < 0.0005 (control vs. SSH1ΔN). (E) Representative images at 4 h FCCP condition with white insets magnified. WT primary hippocampal neurons co-transduced with rAAV9-mCherry-GFP-Sqstm1 and rAAV9-control shRNA or SSH1-shRNA, treated with 2 μM FCCP for 4 h or 0 h (DMSO) and subjected to staining for SSH1 (dark blue) and DAPI (light blue) or direct imaging (mCherry: red, GFP: green). (F-H) Quantitative graphs of figure E expressed as means ± SEM, 2-way ANOVA: (F) group F(1, 44) = 26.20, df = 1, P < 0.0001; (G) group F(1, 44) = 32.77, df = 1, P < 0.0001; (H) group F(1, 44) = 5.638, df = 1, P = 0.0220, followed by Sidak’s posthoc test, n ≥ 15/condition from 3 different experiments, *p < 0.05, **p < 0.005, ***p < 0.0001 (control-shRNA vs. SSH1-shRNA). (I) Representative images with white insets magnified. WT primary hippocampal neurons transduced with rAAV9-Flag-control or Flag-SSH1ΔN and subjected to in situ PLA for SQSTM1-ubiquitin complexes (red) and staining for M2-Flag (green) and DAPI (light blue). (J) Quantitative graph of figure I expressed as means ± SEM, 2-tailed t-test, *P < 0.0001, n ≥ 15/condition from 3 different experiments. (K) Representative images from WT primary hippocampal neurons transduced with rAAV9-GFP or Ssh1-shRNA-GFP and subjected to in situ PLA for SQSTM1-ubiquitin complexes (red) and staining for DAPI (light blue) or direct imaging for GFP (green). (L) Quantitative graph of figure K expressed as means ± SEM, 2-tailed t-test, **P = 0.0007, n = 10/condition from 3 different experiments.
Figure 8. SSH1ΔN exacerbates and SSH1 reduction partially rescues SQSTM1 flux in MAPT/tauP301S brains. (A–D) 3-month old WT and MAPT/tauP301S mice co-injected with rAAV9-mCherry-GFP-Sqstm1 and rAAV9-Flag-control or Flag-SSH1ΔN and sacrificed 2 months post-injection at 5 months of age. (A) Representative images showing staining for Flag-M2 (dark blue) and DAPI (light blue) or direct fluorescence for mCherry-GFP-SQSTM1 (mCherry: red, GFP: green) in the hippocampus CA3 region. (B) Quantitative graphs of figure A expressed as means ± SEM, 1-way ANOVA: (Bi) F(2, 15) = 4.956, df = 2, P = 0.0223; (Bii) F(2, 15) = 24.86, df = 2, P < 0.0001, followed by Tukey’s posthoc test, n = 6 mice/condition *p < 0.05, **p < 0.005. (C) Representative images showing staining for p-S199/p-S202-MAPT/tau (dark blue) and DAPI (light blue) and direct imaging for mCherry-GFP-SQSTM1 (mCherry: red, GFP: green) in the hippocampus CA3 region of MAPT/tauP301S mice. (D) Quantitative graph of figure C (pS199/pS202-MAPT/tau) expressed as means ± SEM, 2-tailed t-test, P = 0.0117, n = 6 mice/condition *p < 0.05. (E) Representative blots of total MAPT/tau, p-S199/p-S202-MAPT/tau, Flag and GFP from dissected hippocampus of injected MAPT/tauP301S mice. (F–J) 3-month old MAPT/tauP301S mice co-injected with rAAV9-mCherry-GFP-Sqstm1 and rAAV9-control-shRNA or rAAV9-Ssh1-shRNA and sacrificed 2 months post-injection at 5 months of age. (F) Representative images showing staining for SSH1 (dark blue) and DAPI (light blue) or direct fluorescence for mCherry-GFP-SQSTM1 (mCherry: red, GFP: green) in the hippocampus CA3 region.
particles containing dimerized UBA domain. We found no evidence that SSH1 alters the activating phosphorylation (p-Ser317) ofULK (not shown), a key upstream kinase that phosphorylates Ser403-SQSTM1 and a kinase shown to be essential for LC3 lipidation [66]. Moreover, our observation that SSH1ΔN, but not SSH1ΔC, dephosphorylates p-Ser403-SQSTM1 using purified recombinant proteins provides cogent evidence for a direct effect of SSH1 in SQSTM1 inhibition.

**Regulation of LC3 biogenesis and recruitment by SSH1**

The interaction of SQSTM1 with LC3 is essential for SQSTM1-mediated selective autophagy. Under conditions of proteotoxic stress or mitochondrial damage, autophagy or mitophagy is initiated and regulated by the cargo. Recent studies indicate that cargo-bound SQSTM1 stimulates LC3-positive puncta formation colocalized with SQSTM1, increases the synthesis of LC3 and its conversion to a lipitated form, LC3-II [44,45]. Our observation that SSH1-mediated SQSTM1 dephosphorylation reduces LC3-II generation, LC3 puncta, and LC3 recruitment to mitochondria upon depolarization is consistent with this notion. Indeed, we validated that these effects on LC3 are completely dependent on the presence of SQSTM1, as SQSTM1 knockdown abrogated the effects of SSH1 on LC3 puncta and recruitment. The reduction in LC3-II level by SSH1 was not due to faster removal by the lysosome, as SSH1 suppressed LC3-II level both in the presence or absence of rapamycin, FCCP, and/or bafilomycin. Hence, despite the changes in LC3 puncta formation, SSH1 did not alter its flux as seen by the mCherry-GFP-LC3 flux reporter. In contrast, SSH1 significantly slowed FCCP-induced mitophagy flux as seen by the mito-QC flux reporter, which is reflected in reduced LC3 puncta formation and its overall recruitment to mitochondria. As SSH1 did not alter LC3 flux per se, we looked for upstream effectors that may account for changes in LC3. It is notable that OPTN, a SQSTM1-related autophagy cargo receptor of similar function, was not affected by SSH1. This confirms the specificity of SSH1 on SQSTM1, which was validated by SQSTM1 deletion and phosphorylation mutants.

**Divergent activities of SSH1 on SQSTM1 and CFL: Implications for tauopathy**

SSH1 is a phosphatase classically known for its ability to dephosphorylate and activate CFL [34], a major F-actin severing protein implicated in a wide array of biological processes [67]. We found that SSH1 dephosphorylates p-Ser403 of SQSTM1, which was modular and separable from that on p-Ser3 of CFL. This discovery not only identified SSH1 as the first phosphatase with this activity, but also indicated that 2 major and apparently unrelated pathways (CFL & SQSTM1) are regulated by a single phosphatase. The CFL binding site is located within the N-terminal region [51], while the SQSTM1 binding site is in the C-terminal region, with the shared catalytic domain located in between. At present, it is unknown whether both activities of SSH1 are regulated simultaneously or separately. However, previous studies have shown that SSH1 is activated by oxidative stress. Specifically, the binding and sequestration of SSH1 by YWHAZ/14-3-3 ζ and YWHAQ/14-3-3 τ heterodimers inhibits SSH1 activity; however, upon oxidation of YWHA/14-3-3, SSH1 is released from YWHA/14-3-3 inhibitory control, thereby allowing SSH1 to dephosphorylate and activate CFL [68,69]. We and others have previously shown that Aβ42 oligomers activate CFL [31,35,70], a process that requires reactive oxygen species (ROS) generation by NOX [32] and dephosphorylation of CFL by SSH1 [31]. Aβ42 oligomer-induced activation of CFL has been shown to destabilize F-actin at the synapse and translocate to mitochondria to promote mitochondrial dysfunction [31]. As SSH1 reduction mitigates Aβ42 oligomer-induced mitochondrial translocations of CFL, our observation that SSH1 knockdown robustly promotes an early step in mitophagy, LC3 recruitment to mitochondria, is rather surprising and suggests that oxidative stress may simultaneously induce mitochondrial damage and impair mitochondrial clearance through SSH1 activation, a hypothesis that requires validation.

The implications of the dual and divergent functions of SSH1 in MAPT/tau biology and tauopathy are multi-fold. We recently reported that CFL reduction mitigates tauopathy through reduced displacement of MAPT/tau from microtubules, as activated/dephosphorylated CFL promotes MAPT/tau displacement from microtubules; however, activated CFL does not directly regulate MAPT/tau levels [71]. This was also evidenced in this study, in which SSH1ΔC, which selectively dephosphorylates CFL, failed to increase MAPT/tau, whereas SSH1ΔN, which selectively dephosphorylates SQSTM1, significantly increased MAPT/tau as effectively as full-length SSH1. Indeed, we confirmed that SSH1ΔN enhances MAPT/tau and phospho-MAPT/tau levels in brain. By using the mCherry-GFP-SQSTM1 flux reporter, we found that MAPT/tauP301S brains and primary neurons exhibit significantly impaired SQSTM1 flux, in line with observations that p-Ser403-SQSTM1 per total SQSTM1 is dramatically reduced in brains of AD patients [2]. This impairment was exacerbated by SSH1ΔN, resulting in a corresponding increase in phospho-MAPT/tau, while MAPT/tauP301S-induced impairment in SQSTM1 flux was partially rescued by SSH1-shRNA, resulting in a corresponding decrease in phospho-MAPT/tau. Hence, in addition to the SSH1-CFL activation pathway in promoting MAPT/tau displacement from microtubules [71], this study highlights the divergent SSH1-SQSTM1 inhibitory pathway in impairing autophagic clearance of misfolded MAPT/tau. The SQSTM1
pathway was also recently shown to be suppressed by the accumulation of ARRB2/β-arrestin2 oligomers in FTLD-MAPT/tau, leading to normal clearance through the autophagy-lysosome system. Deactivation of SQSTM1 by SSH1-mediated dephosphorylation of Ser403 renders SQSTM1 unable to activate LC3 and bind to ubiquitinated cargo, leading to their accumulation and toxicity.

Materials and methods

Cell lines

Mouse hippocampus-derived HT22 neuroblastoma cells (Professor David Schubert; Salk Institute, La Jolla, CA [72]), human embryonic kidney cell line HEK293 cells (ATCC, CRL-1573), human embryonic kidney cell line HEK293T cells (ATCC, CRL-3216) and stably transfected Hela-V5-MAPT/tau cells (Dr. Chad Dickey, USF Health, Tampa, FL, USA [58]) overexpressing wild-type 4R0N human MAPT/tau were cultured in Dulbecco’s modified Eagle’s medium (DMEM,) supplemented with 10% fetal bovine serum (FBS) and 1% penicillin-streptomycin (P/S).

Mice

WT and MAPT/tauP301S [57] mice were obtained from the Jackson Laboratory and bred in the C57BL/6 background. Mice were housed together until the time they were introduced with rAAV9 at 3 months of age. Water and food were supplied ad libitum with 12 h light/dark cycle under standard vivarium conditions. All mice were sacrificed at 5 months of age.

Primary neurons

Hippocampal and cortical primary neuron from WT and MAPT/tauP301S mice were prepared from P0 pups as
previously described \[73,74\]. In brief, both hippocampus and cortex were dissected separately in ice-cold HBSS and digested with trypsin (Gibco, 25,200,056). Mouse neurons were plated on glass coverslips or plates coated with poly-
d-lysine (Sigma-Aldrich, 27,964–99-4) in neurobasal medium (Invitrogen, 21,103,049), 2% GlutaMAX (Invitrogen, 35,050,061), and 2% B27 supplement (Invitrogen, 17,504,044). All the cells were cultured in a humidified atmosphere (5% CO₂) at 37°C.

**Antibodies**

Rabbit polyclonal Anti-LC3A/B (Cell Signaling Technology, 4108); rabbit polyclonal anti-SSH1/Slingshot-1 L (C-terminal region) (ECM Biosciences, SP1713); rabbit monoclonal anti-SSH1 (E1K3W) (Cell Signaling Technology, 13,578); rabbit monoclonal anti-TOMM20/Tom20 (D8T4N) (Cell Signaling Technology, 42,406); mouse monoclonal anti-FLAG* M2 (Sigma-Aldrich, F3165); mouse monoclonal anti-ACTB/β-Actin (Sigma-Aldrich, A5316); rabbit polyclonal anti-SQSTM1/p62 (Cell Signaling Technology, 5114); rabbit polyclonal anti-phospho-SQSTM1/p62 (Ser403) (D.5.1) (Cell Signaling Technology, 39,786); mouse monoclonal anti-HA-Tag (6E2) (Cell Signaling Technology, 2367); rabbit monoclonal anti-GFP (D.5.1) XP* (Cell Signaling Technology, 2956); mouse monoclonal anti-ubiquitin (P4D1) (Santa Cruz Biotechnology, sc-8017); rabbit monoclonal anti-CFL/Cofilin (D3F9) XP* (Cell Signaling Technology, 5175); rabbit monoclonal anti-phospho-CFL/Cofilin (Ser3) (77G2) (Cell Signaling Technology, 3313); rabbit monoclonal anti-His-Tag (D310) XP* (Cell Signaling Technology, 12,698); mouse monoclonal anti-MAPT/tau (A10) (Santa Cruz Biotechnology, sc-390,476); rabbit polyclonal anti-phospho-MAPT/tau (pSer199/202) (Sigma-Aldrich, T6819); mouse polyclonal anti-Horseradish Peroxidase (Jackson ImmunoResearch Inc., Code: 223–005-024, RRID: AB_2339261); rabbit polyclonal anti-horseradish peroxidase (Jackson ImmunoResearch Inc., Code: 323–005-024, RRID: AB_2315781); goat polyclonal anti-herpesvirus peroxidase (Jackson ImmunoResearch Inc., Code: 123–005-021, RRID: AB_2338952); donkey anti-rabbit IgG (H + L) highly cross-adsorbed secondary antibody, Alexa Fluor Plus 594 (Invitrogen, A32754); donkey anti-mouse IgG (H + L) highly cross-adsorbed secondary antibody, Alexa Fluor Plus 594 (Invitrogen, A32744); goat anti-rabbit IgG (H + L) cross-adsorbed secondary antibody, Alexa Fluor 405 (Invitrogen, A-31,556); goat anti-mouse IgG (H + L) cross-adsorbed secondary antibody, Alexa Fluor 405 (Invitrogen, A-31,553); donkey anti-rabbit IgG (H + L) highly cross-adsorbed secondary antibody, Alexa Fluor Plus 647 (Invitrogen, A32795); donkey anti-mouse IgG (H + L) highly cross-adsorbed secondary antibody, Alexa Fluor Plus 647 (Invitrogen, A32787); donkey anti-rabbit IgG (H + L) highly cross-adsorbed secondary antibody, Alexa Fluor Plus 488 (Invitrogen, A32790); donkey anti-mouse IgG (H + L) highly cross-adsorbed secondary antibody, Alexa Fluor Plus 488 (Invitrogen, A32766).

**Reagents, Chemicals, Peptides, Recombinant Proteins and Oligonucleotides**

Agarose linked antisera anti-mouse IgG kit (American Qualex, G1060); agarose linked antisera anti-rabbit IgG kit (American Qualex, G1360B); MitoTracker* red CMXRs (Cell Signaling Technology, 9082); DAPI (Sigma-Aldrich, D9542); Duolink* In Situ PLA* probe anti-mouse PLUS (Sigma-Aldrich, DUO92001); Duolink* In Situ PLA* probe anti-rabbit MINUS (Sigma-Aldrich, DUO92005); Duolink* In Situ detection reagents red (Sigma-Aldrich, DUO92008); Duolink* In Situ wash buffers, fluorescence (Sigma-Aldrich, DUO82049); Duolink* In Situ mounting medium with DAPI (Sigma-Aldrich, DUO82040); Pierce BCA protein assay kit (Thermo Scientific, 23225); Aligent QuikChange II site-directed mutagenesis kit (Agilent, 200521); FCCP (Sigma-Aldrich, C2920); rapamycin (Sigma-Aldrich, R8781); bafilomycin A₁ (Santa Cruz Biotechnology, sc-201550); Benzonase* nuclease (Sigma-Aldrich, E1014); OptiPrep™ density gradient medium (Sigma-Aldrich, D1556); lipofectamine 2000 (Invitrogen, 11668019); fluorochrome mounting solution (Thermo scientific, TA-030-FM); Opti-MEM I (1x) (Gibco, 31985–070); parafomaldehyde, 96% extra pure. (PFA) (Acros Organics, 41678003); PBS (Gibco, 10010049) DMEM (Dulbecco’s modified eagle medium) (Corning, 10–013-CV); fetal bovine serum (FBS) (Sigma-Aldrich, 12306 C); pen strep (P/S) (Gibco, 15140–1222); BM-cyclin (Roche, 1079905001); SF-900° II SFM (Gibco, 10902096); GE Healthcare Ni sepharose (GE Healthcare, 17526801); Xhol restriction enzyme (NEW ENGLAND Biolabs, R0146S); Agel restriction enzyme (NEW ENGLAND Biolabs, R0552S); Ascl restriction enzyme (NEW ENGLAND Biolabs, R0558S); NheI restriction enzyme (NEW ENGLAND Biolabs, R0131S); NotI restriction enzyme (NEW ENGLAND Biolabs, R0189S); Sall restriction enzyme (NEW ENGLAND Biolabs, R0138S); Spel restriction enzyme (NEW ENGLAND Biolabs, R0133S); Beta-Amyloid 1–42, TFA (GenicBio Limited, A-42-T-1); SSH1 siRNA: GAG CAG CUG UCC CGA UGA C (Dharmacon GE Healthcare, CTM-395590); SignalSilence® SQSTM1/p62 siRNA II (Cell Signaling Technology, 6399); SHH1ΔC-his, SHH1ΔN-his, SQSTM1/p62-his were generated by this paper (see method details).

**Recombinant DNA**

pN3-3xFlag-Control (Addgene, 10771, Guntran Suske Lab [75]); pmRFP-LC3 (Addgene, 21075, Tamotsu Yoshimori Lab [43]); pMXs-puro GFP-Sqstm1/p62 (Addgene, 38277, Noboru Mizushima Lab [49]); pBABE-puro mCherry-EGFP-LC3B (Addgene, 22418, Jayanta Deb Nath Lab [76]); pBABE-EGFP (Addgene, 36999, Debu Chakravarti Lab [77]); ECFP-SHSH1ΔC (N461) (Dr. Mizuno lab [51]); EGFPSHSH1 (Dr. Storz lab [69]); mKeima-Red-Mito-7 (Addgene, 56018); pOPTN-EGFP (Addgene, 27052Beatrice Yue Lab [78]); pMXs-puro GFP-Sqstm1/p62ΔC (Addgene, 38282, Noboru Mizushima Lab [49]); pMXs-puro GFP-Sqstm1/p62 D337, 338, 339A (GFP-Sqstm1-LIR) (Addgene, 38280, Noboru Mizushima Lab [49]); HA-Ubiquitin (Addgene, 18712, Edward Yeh Lab [79]); pDR125 (Addgene, 37150, Dale Ramsden Lab); pN3-
3xFlag-SSH1, mCherry-EGFP-Sqstm1/p62, mCherry-EGFP-Sqstm1S403A, mCherry-EGFP-Sqstm1S403E, pN3-Flag-SSH1ΔC, pN3-Flag-SSH1ΔN and pN3-Flag-SSH1ΔN5393S were generated in this work (see methods below).

DNA transfections and adenoviral transductions

DNA plasmids were transiently transfected in HT22 cells or HEK293T cells using Lipofectamine 2000 and Opti-MEM I. SiRNA was transfected twice every 24 h. After four to six hours post transfection, the medium was replaced with new complete medium. Generally, cells were grown for 48 h after transfection prior to experimental assays. rAAV9 variants were transduced to primary neurons on DIV4 and were assayed for experiments on DIV18.

Cell/tissue lysis and immunoblotting

Cultured cells were lysed with RIPA lysis buffer (50 mM Tris-Cl, pH 7.4, 150 mM NaCl, 2 mM EDTA, and 1% Triton X-100 [Amresco, 0694–1 L], 0.1% SDS) plus protease inhibitor (GeneDEPOT, P3100-010) and phosphatase inhibitors (GeneDEPOT, P3200-005). Brain homogenates were lysed with RIPA buffer plus protease and phosphatase inhibitors with sonication. Proteins were extracted and centrifuged at 17,000 g for 15 min at 4°C, and supernatants were used for western blotting. Protein quantification was performed with a colorimetric detection reagent (BCA protein assay, Pierce, 23225). Equal amounts of total protein were subjected to SDS-PAGE and transferred to nitrocellulose membranes (GE Healthcare, 10600002) for immunoblotting. After probing with the primary antibody overnight at 4°C, the corresponding peroxidase-conjugated secondary antibody was detected by ECL western blot reagents (Pierce, 34578). ECL images were captured with the Fuji LAS-4000 imager (LAS-4000, Pittsburgh, PA, USA) and quantified using the ImageJ software. All protein amounts were normalized to actin prior to comparison between experimental conditions.

Immunofluorescence

For immunocytochemistry (ICC), cells were washed with PBS and fixed at room temperature for 15 min with 4% paraformaldehyde (PFA). After washing with PBS, fixed cells were incubated with blocking solution containing 0.2% Triton X-100, 3% normal goat serum (Vector Laboratories Inc., S-1000) for 1 h, followed by overnight incubation at 4°C with related primary antibodies. After three washes with PBS, cells were incubated for 1 h with Alexa Fluor 488-, Alexa Fluor 594- or Alexa Fluor 647-conjugated secondary IgG antibodies. Slides were then washed three times with PBS and mounted with fluorochrome mounting solution. For immunohistochemistry (IHC), mice were perfused with PBS, and half brains were immediately stored at −80°C for biochemical analyses, and the other half was fixed with 4% paraformaldehyde at 4°C for 24 h followed by cryoprotection in 30% sucrose. Thirty-micron sections were blocked using normal goat serum for 1 h and subjected to primary antibodies at 4°C overnight, followed by secondary antibody (Alexa Fluor 488, Alexa Fluor 594 or Alexa Fluor 647) incubation for 1 h at room temperature prior to mounting. For PLA assays, Duolink® In Situ PLA* reagents were purchased from Sigma-Aldrich. PLA assay was performed following the manufacturer’s instructions. All images from ICC, IHC, and PLA experiments were captured with the Olympus FV10i confocal microscope (Tokyo, Japan).

Plasmids and constructs

The mCherry-EGFP-Sqstm1 construct was generated by replacing the LC3B sequence with the Sqstm1 sequence (pMXs-puro GFP-SQSTM1) from the pBABE-puro mCherry-EGFP-LC3B construct using EcoRI and Sall restriction sites. The mCherry-EGFP-Sqstm1S403A, mCherry-EGFP-Sqstm1S403E constructs were generated using the Agilent QuikChange II Site-Directed Mutagenesis Kit from mCherry-EGFP-Sqstm1. SSH1, SSH1ΔC, SSH1ΔN were subcloned into pN3-3XFlag vector using HindIII and Sall sites by PFU-based PCR amplification of SSH1 from EGFP-SSH1 as a template. Flag-SSH1ΔN5393S was generated using the Agilent QuikChange II Site-Directed Mutagenesis Kit from Flag-SSH1ΔN. Constructs used for recombinant proteins SSH1ΔC-his, SSH1ΔN-his, Sqstm1-his were inserted into the backbone of pFastbac1 with Xbal and XhoI sites (addgene pDR125 removed LIG4 with Xbal and XhoI). All primers were custom designed as following and ordered from IDT:

- mCherry-EGFP-Sqstm1-Flag SSH1ΔN-his: 5’-TTTTGGAATATCAGGCTTACCTCAGGGAAGTACG-3’
- mCherry-EGFP-Sqstm1-Flag SSH1ΔC-his: 5’-TTTTTCTCGAGCTGCTGACGCCACAGCTT-3’
- mCherry-EGFP-Sqstm1S403A-Flag SSH1ΔN-his: 5’-CTTCCCAGATGTCGCTGAGGCTTCTCGCGAT-3’
- mCherry-EGFP-Sqstm1S403E-Flag SSH1ΔN-his: 5’-ATCCGAGAATATCGCAGCTGAGGCTTCTCGCGAT-3’
- mCherry-EGFP-Sqstm1S403A-Flag SSH1ΔC-his: 5’-CTTCCCAGATGTCGCTGAGGCTTCTCGCGAT-3’
- mCherry-EGFP-Sqstm1S403E-Flag SSH1ΔC-his: 5’-ATCCGAGAATATCGCAGCTGAGGCTTCTCGCGAT-3’
Generation of rAAV9 and stereotaxic injections in mice

To generate the pTR12.1-MCSW-Flag-SSH1ΔN construct, Flag-SSH1ΔN, was digested by Agel and Sall restriction enzymes and subcloned into the pTR12.1-MCSW vector using Agel and XhoI (destroyed after ligation) cloning sites. To generate the pTR12.1-MCSW-ECFP-SSH1ΔC construct, the ECFP-SSH1ΔC was cloned into pTR12.1-MCSW vector at Agel and NotI cloning sites. To generate the pTR12.1-MCSW-mCherry-EGFP-Sqstm1 construct, the mCherry-EGFP-Sqstm1, was digested by SpeI and Sall restriction enzymes and cloned into the pTR12.1-MCSW vector using SpeI and XhoI (destroyed after ligation) cloning sites. To generate H1rSc-Ssh1-shRNA and H1rSc-Ssh1-shRNA-hrGFP constructs, the duplex of mouse Ssh1 short hairpin RNA with the target sequence 5’-CCCGTITTAGATCAGACCGAGCTA-3’ was cloned into H1rSC and H1rSC-hrGFP vectors at ASCI and NheI cloning sites. Recombinant AAV9 viruses were generated by co-transfection of serotype vector expressing the gene of interest with pAAV9 and pXX6 in HEK293 cells. Cells were lysed in the presence of 0.5% sodium deoxycholate, 1 mM MgCl₂ and 50 U/ml Benzonase by freeze thawing after 48 h incubation with transfected vectors, and the virus was isolated using a discontinuous iodixanol (OptiPrepTM Density Gradient Medium) gradient and purified with ApolloTM 20 mL High-Performance Centrifugal quantitative concentrators (Orbital Biosciences, 2,015,010) [74,80]. For brain injections, isoflurane anesthetized mice (3-month old, equally balanced for gender per condition) were bilaterally injected with a 26-gauge needle attached to a 10-μl syringe (Hamilton, 80,330) at the following coordinates: anteroposterior 2.7 mm, lateral 2.7 mm, and vertical 3.0 mm. A total volume of 2 μl purified rAAV9 (1.3 × 10¹² vg/ml) was injected over a 2-min period using the convection enhanced delivery method [74,80]. Mice were sacrificed 8 weeks post injection.

Recombinant proteins

pFast-SSH1ΔC-his, pFast-SSH1ΔN-his, pFast-Sqstm1-his constructs were transformed into DH10Bac competent cells. After blue-white screening, DH10Bac strains were chosen to express and amplify recombinant Bacmids. S9 insect cells transfected with Bacmid (midiprep from DH10Bac competent cells) were cultured for 3 days with S900 II SFM medium, then P1 generation virus in medium was collected and added to new S9 cells. After 2 days culture, S9 cells were harvested and lysed with lysis buffer (Tris 20 mM, pH7.4, NaCl 150 mM, Triton X-100 1%, 10 mM imidazole, with protease inhibitors). After centrifugation at 12,000 g for 15 minutes, supernatant was collected and shaken for 1 h at 4°C with Ni Sepharose (GE Healthcare, 17–5268-01). Bound proteins on sepharose were washed 3 times with ice-cold lysis buffer, and recombinant proteins were eluted with ice-cold elution buffer (Tris 20 mM, pH 8.5, NaCl 150 mM, 200 mM imidazole), after which proteins were dialyzed in dialysis buffer (Tris 20 mM, pH 8.0, NaCl 150 mM, DTT 1 mM) at 4°C overnight.

Quantification and statistical analysis

Immunofluorescence images were quantified by using the Image J software (National Institutes of Health, Bethesda, MD). All comparison images were acquired with identical laser intensity, exposure time, and filter. Adjustments to the brightness/contrast and threshold were applied equally to all comparison images. Regions of interest (ROI) for cell lines and primary neurons were cell body (excluded cell nuclei). ROI for brain tissue were chosen randomly throughout the CA3 region of the hippocampus. All puncta numbers were quantified manually with the Image J cell counter plugin. All percentages of co-localization were measured with Image J COLOC Threshold Manders’ Correlation with in-program threshold. Then number of puncta that colocalized with mitochondria were quantified manually using the Colocalized Pixel Map Image (generated by Image J COLOC Threshold) with the Image J cell counter plug-in. Puncta sizes were measured as particle area with Image J Analyze Particles. Investigators were blinded to genotypes of mice and experimental conditions during image acquisition and quantification. Statistical data were analyzed by the GraphPad Prism 6.0 software using Student’s 2-tailed t-test, χ² test for trend, 1-way ANOVA, or 2-way ANOVA. One-way ANOVA was followed by Tukey’s post hoc test, and 2-way ANOVA was followed by Sidak’s post hoc test. ANOVA post hoc tests were performed when initial ANOVA tests showed significance at p < 0.05. All quantitative graphs were expressed as means ± S.E.M. Differences were deemed significant when P < 0.05.

Ethics approval

All experimental methods and protocols involving mice were approved by Institutional Animal Care and Use Committee (IACUC) at the University of South Florida (USF). IACUC and Institutional Biosafety Committees (IBC) at the USF have also approved that all the methods used in this study were performed in accordance with the relevant guidelines and regulations.

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Data availability
The data that support the findings of this study are available from the corresponding authors on reasonable request.

Disclosure statement
No potential conflict of interest was reported by the authors.

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