Defective recognition of LC3B by mutant SQSTM1/p62 implicates impairment of autophagy as a pathogenic mechanism in ALS-FTLD

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ABSTRACT
Growing evidence implicates impairment of autophagy as a candidate pathogenic mechanism in the spectrum of neurodegenerative disorders which includes amyotrophic lateral sclerosis and frontotemporal lobar degeneration (ALS-FTLD). SQSTM1, which encodes the autophagy receptor SQSTM1/p62, is genetically associated with ALS-FTLD, although to date autophagy-relevant functional defects in disease-associated variants have not been described. A key protein-protein interaction in autophagy is the recognition of a lipid-anchored form of LC3 (LC3-II) within the phagophore membrane by SQSTM1, mediated through its LC3-interacting region (LIR), and notably some ALS-FTLD mutations map to this region. Here we show that although representing a conservative substitution and predicted to be benign, the ALS-associated L341V mutation of SQSTM1 is defective in recognition of LC3B. We place our observations on a firm quantitative footing by showing the L341V-mutant LIR is associated with a ~3-fold reduction in LC3B binding affinity and using protein NMR we rationalize the structural basis for the effect. This functional deficit is realized in motor neuron-like cells, with the L341V mutant EGFP-mCherry-SQSTM1 less readily incorporated into acidic autophagic vesicles than the wild type. Our data supports a model in which the L341V mutation limits the ability of the SQSTM1 protein, have over the past 4 years been identified as a cause of familial ALS-FTLD. Autophagy is a critical pathway for the removal of damaged and aggregation-prone proteins and organelles; for example, in the ALS context mediating the turnover of aggregated forms of the major pathological protein TARDBP (TAR DNA binding protein), and is essential for neuronal survival. ALS-FTLD-associated mutants of the autophagy receptor OPTN are defective in the selective degradation of mitochondria via autophagy with a mechanism that may involve compromised maturation of autophagosomes. Further, depletion of optn in zebrafish is associated with a motor axonopathy, which is phenotypically similar to a model of ALS expressing mutant SOD1. Similarly, numerous missense and truncating mutations affecting the SQSTM1 gene, which encodes the SQSTM1 protein, have over the past 4 years been identified in patients with ALS and FTLD. Curiously SQSTM1 mutations are also found in patients with the skeletal disorder Paget disease of bone (PDB), including some variants that are common to both disease phenotypes and polymorphisms in other

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autophagy genes have recently been associated with PDB in a Spanish cohort.\textsuperscript{14} SQSTM1 is a multidomain scaffold protein involved in various signaling pathways and more recently characterized as a cargo receptor for ubiquitin-mediated autophagy.\textsuperscript{15} At least in osteoclasts derived from mice expressing the equivalent of a PDB and ALS-associated mutant SQSTM1\textsuperscript{P392L}, markers indicative of dysregulation of autophagy are evident.\textsuperscript{16} In zebrafish, knockdown of Sqstm1 is associated with a locomotor phenotype that can be rescued by the autophagy activator rapamycin, as well as wild-type human SQSTM1 but not the P392L mutant, suggesting that changes in SQSTM1-mediated autophagy are linked to the phenotype.\textsuperscript{17}

ALS-FTLD-associated missense mutations of SQSTM1 are distributed in different functional domains throughout the primary sequence of the 440-residue protein\textsuperscript{13} and notably a subset of these mutations map to the LC3-interacting region (LIR) of the protein.\textsuperscript{18} The LIR is a critical motif that allows SQSTM1 to couple to the developing phagophore through the recognition of lipid-anchored LC3 proteins (mammalian family of orthologs of yeast Atg8), within the phagophore membrane. This highly conserved region is located between amino acids 321 to 342 of human SQSTM1\textsuperscript{15,19} with structural analyses indicating that the motif DDDWTHL (corresponding to residues 335 to 341) is directly involved in the interaction with LC3. One particular SQSTM1 mutation, an L341V missense mutation identified in a Chinese patient with late-onset sporadic ALS,\textsuperscript{18} is of interest since it affects a residue within the LIR of SQSTM1. Indeed the crystal structure of an LIR peptide (residues 332 to 342) in complex with LC3B revealed that L341 forms direct contacts with LC3B within the surface binding cleft.\textsuperscript{19} Consistent with this observation an artificial mutation at the same position (L341A) in the 332 to 342 LIR peptide results in significantly reduced SQSTM1-LC3B complex in qualitative protein binding assays,\textsuperscript{19} as is also the case in a slightly longer LIR peptide (332 to 347).\textsuperscript{20} Here we show that the ALS-associated L341V mutation, although previously predicted to represent a benign substitution,\textsuperscript{18} actually exerts a quantifiable effect on LC3B binding in vitro and affects SQSTM1 recruitment into acidic autophagic vesicles in living cells. Our data supports a model in which certain SQSTM1 mutations limit a critical step in autophagy and provides important functional evidence for a role of the autophagic pathway in ALS-FTLD.

Results

\textbf{The L341V mutation affects the recognition of LC3B by SQSTM1}

The L341V missense mutation of SQSTM1, identified in a late-onset sporadic ALS patient, represents a conservative amino acid change, which has previously been predicted (based on SIFT and PolyPhen-2 prediction tools) to represent a benign substitution.\textsuperscript{18} We first used biochemical approaches to examine any impact of the L341V mutation on well-characterized protein-protein interactions relevant to SQSTM1-mediated autophagy. Protein affinity isolation assays in which interaction partners LC3B or ubiquitin covalently immobilized on beads were used to capture recombinant GST-SQSTM1 fusion proteins, followed by detection of bound protein by western blotting. The same assay has previously been used to demonstrate ubiquitin-binding defects associated with PDB-linked UBA domain mutations of SQSTM1.\textsuperscript{21} Consistent with its location within the SQSTM1 LIR, defined by the 338 to 341 WXXL motif, we found that the L341V mutant selectively reduced capture of GST-SQSTM1 by LC3B, relative to the wild type, whereas binding to ubiquitin was unaffected (Fig. 1 and Fig. S1). As a control the previously characterized PDB-linked UBA domain SQSTM1 mutation, G425R (SQSTM1\textsuperscript{G425R}),\textsuperscript{21} also now reported in cases of ALS,\textsuperscript{22} was found to selectively impact only on ubiquitin-binding, with no evidence of this mutation affecting LC3B recognition. Thus, L341V exhibits a domain-specific effect on the interaction of SQSTM1 with LC3B in vitro.

\textbf{ESI-MS confirms weaker binding of the L341V mutant LIR to LC3B}

We next used a native ESI-MS approach, where under carefully optimized nondenaturing conditions noncovalent protein complexes representative of solution binding can be maintained in the gas phase, in order to determine the effects of the L341V mutation on LC3B recognition by the LIR. Synthetic peptides with LIR (L341V) or without the mutation (WT LIR), representing an extended LIR sequence (residues 332 to 351) compared to those previously characterized in structural detail (332 to 342\textsuperscript{19} and 332 to 347\textsuperscript{20}), were generated by solid-phase synthesis. Mass differences associated with the leucine to valine substitution meant that multiply charged species of the 2 peptides could be readily distinguished by ESI-MS (Fig. 2A). We set up a competitive binding mixture including 5 \(\mu\)M of purified LC3B along with equimolar concentrations of the 2 LIR peptides (i.e., a mixture of 1:1:1) and assessed complex formation by ESI-MS. Notably, peaks in the spectra showed evidence for a more significant population of the LC3B-WT LIR complex compared to LIR (L341V) (Fig. 2B, 2+ complex of free LIRs and 9+ complex of LIR-LC3B complexes indexed in lower panel) indicative of a reduction in binding affinity for the mutant peptide. Under these conditions the abundance ratio of LC3B bound to WT LIR compared to LIR (L341V) was 3:1, entirely consistent with a reduced binding affinity of the mutant LIR.

\begin{figure}[h]
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\caption{ALS-FTLD-associated SQSTM1 mutations impact on the recognition of LC3B (SQSTM1\textsuperscript{L341V}) or ubiquitin (SQSTM1\textsuperscript{G425R}) in vitro. Mutations as indicated (or wild type, WT) were introduced into the full-length GST-SQSTM1 sequence and affinity isolation assays (LC3B and ubiquitin on beads) were performed at 37°C. Bacterial lysates containing the GST-SQSTM1 fusions were incubated with glutathione- (G), control- (C), LC3B (LC3), and ubiquitin-Sepharose (Ub) beads and captured proteins were detected by western blotting (anti-SQSTM1 antibodies). A representative blot is shown; see Fig. S1 for quantification of 3 independent experiments.}
\end{figure}
Quantifying the magnitude of the effects of the L341V mutation on LC3B binding

We used isothermal titration calorimetry (ITC) to corroborate the MS data and determine binding affinities (K_d). Separate titrations were performed with starting concentration of the LC3B at 15 μM, with addition of aliquots of the LIR peptides (WT LIR or LIR (L341V) at 350 μM, at 25°C in 25 mM phosphate buffer, 150 mM NaCl, pH 7.5). High quality binding isotherms (Fig. 3) fitted well to a one-site binding model with a K_d value for WT LIR in the low μM range (3.2 ± 1.1 μM). However, the LIR (L341V) peptide was associated with a ~3-fold reduction in LC3B binding affinity (10.9 ± 1.1 μM) and a substantially reduced enthalpy of interaction. The latter would appear to reflect possible steric effects and unfavorable van der Waals contacts from forcing a ‘square peg into a round hole.’ These binding affinities are within the same order of magnitude as previously published affinities from Novak et al.23 which use a similar-length peptide but covering a different set of residues around the LIR (residues 321 to 342).

NMR reveals the structural basis for the effects of the L341V mutation on LC3B recognition

To understand the mechanism for impairment of binding of the LIR (L341V) at the molecular level we used NMR studies at 800 MHz to map differences in the binding interactions of LC3B, using the same WT LIR and LIR (L341V) peptide sequences used for MS and ITC studies. We first carried out an extensive amide backbone NMR assignment for LC3B (> 95% assignment of the nonprolyl residues) using double-isotope labeling with 13C/15N and a combination of 3D heteronuclear NMR spectra using well established methodologies (see Methods; Fig. S2, gray). Subsequently, we were able to use the sensitivity of 1H/15N-HSQC spectra of the assigned amide NHs sequences used for MS and ITC studies. We first carried out an extensive amide backbone NMR assignment for LC3B (> 95% assignment of the nonprolyl residues) using double-isotope labeling with 13C/15N and a combination of 3D heteronuclear NMR spectra using well established methodologies (see Methods; Fig. S2, gray). Subsequently, we were able to use the sensitivity of 1H/15N-HSQC spectra of the assigned amide NHs

Figure 2. ESI-MS indicates weaker binding of the LIR (L341V) to LC3B compared to WT LIR. (A) Native ESI-MS spectrum of an equimolar mixture of WT LIR and LIR (L341V) peptides (5 μM, residues 332 to 351). (B) LIR peptide mixture titrated with 5 μM LC3B. Top, full spectrum indicating free LIR (gray filled circle, mixture of WT LIR and LIR [L341V]), free LC3B and LC3B-LIR complexes of indicated (multiple) charge states. Below, zoomed-in spectra showing m/z values of the free and bound LIR complexes at the indicated charge states.

Figure 3. Isothermal titration calorimetry (ITC) binding isotherms from titrating LC3B with LIR peptides at 25°C. Binding isotherms were fitted to a one-site binding model and K_d values determined. Starting concentration of the LC3B was approximately 30 μM, and of LIR peptide (residues 332 to 351) stocks approximately 350 μM. Black diamond, LIR (L341V); gray circle WT LIR.
at the individual residue level to map the perturbations to LC3B associated with ligand (LIR) binding (Fig. 4, Fig. S2, green and blue). Titration studies with WT LIR revealed significant changes to protein chemical shifts (> 0.5 ppm) for nearly 20 key residues; a subset of 9 of these demonstrated perturbations of > 1.0 ppm (Fig. 5A) indicative of substantial structural re-organization around the surface cleft to accommodate the bound ligand. These residues can be mapped to the surface of LC3B (Fig. 5C) and correspond very clearly with those identified as LIR contact residues in the X-ray structure of the LC3B-LIR (332 to 342) complex.19 Of particular note was the use of a shorter LIR-containing peptide in the X-ray studies (SGGDDWDTHSS versus SGGDDWDTHSSKEVDPSTG used here). Given that NMR CSP effects were observed over a substantial number of LC3B residues both in direct contact, and in some cases on the more remote nonbinding face, we compared NMR data with a truncated LIR peptide of only 11 residues (amino acids 335 to 345). The 2 titrations with the WT LIR-containing motifs highlighted a few small differences (data not shown), however, the largest perturbations were quantitatively very similar, indicating that the C-terminally extended peptide sequence used in our studies, which accommodates the additional VDPSTG motif corresponding to the KEAP1 interaction region (KIR), forms no additional significant interactions across the surface of LC3B.

We repeated the titration experiments with LIR (L341V) in the longer peptide sequence (332 to 351) and observed CSP effects of similar magnitude to WT LIR, showing the same extensive interaction (data not shown). By taking the difference in CSP effects between WT LIR and LIR (L341V) sequences we could readily identify 8 residues that were substantially different between the 2 complexes, with the largest effects involving V33, F52, L53, V54 and R70 (Fig. 5B). When these were specifically highlighted on the LC3B protein (Fig. 5D) they correlated remarkably closely with those constituting the binding cleft in proximity to L341. The perturbations demonstrate that binding of the mutant sequence is associated with some reorganization of the pocket to accommodate the different steric demands of the β-branched valine residue. We modeled the L341V substitution (data not shown), and observed significant steric clashes for all side chain rotamers of the substituted valine side chain, consistent with some loss of binding complementarity, leading to reduced affinity.

**Functional effects of the L341V mutation in a cellular context**

To determine whether the L341V mutation affects autophagy in living cells we made use of an mCherry-EGFP double-tagged SQSTM1 construct which allows acidic autophagic vesicles to be distinguished from neutral SQSTM1-positive inclusion bodies and autophagosomes.15 When ectopically expressed in cells SQSTM1 can incorporate into cytoplasmic bodies with a similar phenotype to those observed when cells are stained for the endogenous protein.24 The inclusion bodies represent non-membrane enclosed structures whereas the autophagosomes are membrane-limited; upon live cell imaging of both structures, red and green fluorescence (combined yellow) can be detected as each represent pH neutral microenvironments. However, following fusion of phagophores with lysosomes the EGFP (green) signal is quenched in the resulting acidic milieu, and only red fluorescence is detected, allowing transit of ectopically expressed SQSTM1 from phagophores to acidic autophagic vesicles to be monitored.

Previous studies demonstrate that deletion of the LIR strongly reduces the ability of SQSTM1 to be recruited into phagophores and subsequently acidic vesicles.15 To assess the impact of the more subtle L341V missense mutation, we introduced this change in to the mCherry-EGFP-SQSTM1 construct and transiently expressed this in the motor neuron-like cell line NSC-34. Notably endogenous levels of SQSTM1 are previously reported to be low (almost undetectable) in these cells.25 However, our more rigorous analysis using different SQSTM1 antibodies (Fig. S3) indicated that although NSC-34 cells do appear to

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**Figure 4.** NMR titrations indicate selective perturbations of LC3B residues upon binding of LIR peptides. (A) 1H,15N-HSQC spectrum of LC3B (0.25 mM dark gray) overlaid with the spectrum of the complex of LC3B with WT LIR (0.5 mM blue, residues 332 to 351) (ratio of 1:2) showing extensive chemical shift perturbations (CSPs) across the spectrum induced by ligand binding at 298 K; (B) Expansion of the region highlighted in (A) showing overlap between the spectrum of LC3B (dark gray), LC3B + WT LIR (blue) and LC3B + LIR (L341V) (green) illustrating a number of key residues within the LIR binding pocket that are perturbed to different extents by the WT LIR and LIR (L341V). Arrows identify the shifts of V33, F52 and V54 in the 2 complexes, while the majority of other residues show only small differences between the spectra of the 2 complexes (overlap of blue and green peaks), demonstrating selective effects on residues in direct contact with the LIRs.
express lower steady-state levels of SQSTM1 than other lines (adventitiously limiting the contribution of endogenous protein to any observed phenotype) these levels are not unusually low. Further, levels of endogenous SQSTM1 and LC3B increased in response to bafilomycin A1 (BafA1) treatment with a similar profile to HeLa cells (Fig. S3) indicating NSC-34 have normal autophagy machinery. We confirmed that steady-state levels of transfected mCherry-EGFP-SQSTM1 (wild type and L341V) were similar by western blot (Fig. S4). In cells 48 h after transfection with wild-type mCherry-EGFP-SQSTM1 we noted obvious yellow and red structures (Fig. 6), with the red acidic vesicles most prominent within the characteristic short neurites. Treatment with the autophagy inhibitor BafA1 visually reduced the number of red acidic structures (Fig. 6), which was quantified by determining the increase in the mean Pearson correlation coefficient (PCC) value of mCherry and EGFP overlap (Fig. 7). In contrast, in cells transfected with mutant mCherry-EGFP-SQSTM1L341V there were visibly fewer red acidic vesicles in the absence of BafA1 (Fig. 6), evidenced by a mean PCC value which was greater than that for the wild type (Fig. 7). Treatment of the L341V mutant-expressing cells with BafA1 further increased the mean PCC value (Fig. 7), indicating that the cells were still functional with respect to autophagy, consistent with reversal of the mutant phenotype (reduction in PCC values) observed following treatment of mutant-expressing cells with the autophagy-enhancer rapamycin (Fig. 7). We also confirmed the L341V mutant phenotype in an additional neuronal cell line, mouse neuroblastoma Neuro-2a cells (Fig. S5); although the difference in mean PCC values between wild type- and L341V mutant-expressing cells was smaller in these cells this was still statistically significant. Overall these data are consistent with mutant mCherry-EGFP-SQSTM1L341V being less readily incorporated into acidic vesicles than the wild type.

**Discussion**

Our findings, using multiple independent approaches, of defective LC3B recognition by an ALS-associated variant of SQSTM1, and a reduced ability of the mutant protein to incorporate into acidic autophagic vesicles in living cells, adds to the accumulating evidence that disturbances in autophagy may underlie some cases of ALS-FTLD. Other studies to date have presented histological and genetic evidence for such a disturbance, but importantly we now add functional evidence by describing on a firm quantitative and structural footing a defect in an autophagy-relevant protein-protein interaction. Specifically we confirm that the ALS-associated L341V mutation of SQSTM1 exerts a quantifiable effect on LC3B binding in vitro. While a 3-fold reduction in binding affinity may appear modest, we suggest that over the lifetime of a neuron it may become biologically significant (see later). Indeed the change in binding affinity is sufficient to give a clear autophagy phenotype in our NSC-34 cell model as well as in Neuro-2a cells, i.e., two different neuronal cell lines, one of which is motor neuron-like. In this cellular context it is also possible that post-translational modification (e.g. phosphorylation of the LIR) may directly
or indirectly exaggerate the differences in binding affinities between wild-type and mutant sequences. However, a further important consideration in this respect is the structural organization of SQSTM1 in forming flexible oligomeric assemblies critical to its functional role in autophagy. Recent cryo-EM studies have established that the N-terminal (100 residue) PB1 domain of SQSTM1 and nearby sequences (also associated with ALS-related mutations) drive the formation of polymeric SQSTM1 filaments. These SQSTM1 filaments are proposed to present multiple LC3 binding sites to aid the growth of the phagophore onto the filaments. Thus, we speculate that avidity effects between multiple LIRs on the surface of the assembled SQSTM1 scaffold and membrane-tethered LC3B could provide a mechanism for amplifying any small detrimental effects on binding affinities from individual LIR mutations and these effects are ultimately manifested as a clear phenotype in the NSC-34 and Neuro-2a cell models. Similarly, in vitro affinity isolation with GST-SQSTM1, dimerization of GST may further promote avidity effects. Notably in the NSC-34 cells rapamycin is able to rescue the effects of the mutation on incorporation of SQSTM1 into phagophores, via currently undefined changes downstream of MTOR-dependent signaling, which must overcome the relatively modest reduction in LC3B-binding activity associated with the mutation; this observation supports the notion that autophagy-enhancing strategies may be beneficial in some cases of ALS.

The particular ALS-associated variant of SQSTM1 that we analyzed, identified in a late-onset sporadic ALS patient, is a
conservative amino acid change (leucine to valine) which has previously been predicted by web-based prediction tools to represent a benign substitution.14 Such a prediction probably arose since L341 of SQSTM1, although highly conserved in mammals, in *Xenopus* is substituted by the valine amino acid found in the ALS patient. However it is not unusual for a deleterious missense mutation in one species to be found as the wild-type residue in orthologs of other species without effecting on the missense mutation in one species to be found as the wild-type in the ALS patient. However it is not unusual for a deleterious mutation to be found as the wild-type in another species, for example, an aspartic acid to glutamic acid (D337E) mutation, also identified in a Chinese patient with sporadic ALS,18 is predicted to be benign. Although the artificial mutation of aspartic acid to alanine (D337A) reveals little effect on LC3B binding in affinity isolation assays using GST-tagged LIR sequences,56 subtle but biologically-relevant effects on LC3B recognition cannot be discounted. It is also notable that many ALS-associated SQSTM1 mutations affect other functional regions outside of the LIR that have significance for autophagy, for example the UBA domain and PB1 domains which are both necessary for successful cargo delivery to the phagophore.3,15 Therefore, while LIR mutations appear to limit the crucial step in selective autophagy of cargo receptor delivery to the phagophore, a model emerges where other SQSTM1 mutations may have an impact on, for example, cargo recognition, and the ability of the PB1 domain to regulate SQSTM1 assembly into the larger helical scaffolds required to nucleate the growing autophagosomal membrane.28 The role of avid interactions in amplifying the effects of individual mutations is intriguing in the light that SQSTM1 not only forms oligomeric scaffolds, but that the UBA domain also forms a dimeric structure that also plays a role in regulating binding to ubiquitinated cargo.30 In a similar context Lee et al. have recently highlighted how other ALS-associated genetic alterations may impact on different steps of autophagy, for example, vesicle nucleation (SOD1), autophagosome trafficking (DCTN [dynactin], DYNClH1 [dynein cytoplasmic 1 heavy chain1], SOD1) and phagophore-lysosome fusion (VCP [valosin containing protein], CHIP28 [charged multivesicular body protein 2B]).31 Dysregulation of autophagy in ALS-FTLD may represent a wider mechanistic aberration that crosses over between other mutational and perhaps even nonmutational causes. The notion of autophagic dysfunction is entirely consistent with the emerging thinking that ‘lysosomal dysfunction’ underlies a range of neurodegenerative disorders including dementia with Lewy bodies and frontotemporal dementia (FTD, the most common FTLD syndrome), supported from recent genome-wide association study (GWAS) analyses.32,33

We speculate that subtle effects on autophagy associated with SQSTM1 (and other) variants predispose neurons to a vulnerability, which over time may become exposed by, for example, an age-related decline in autophagy pathway activity34,35 or ‘second-hit’ in another protein clearance pathway such as the ubiquitin-proteasome system.6,36 Causal mutations in other autophagy-related ALS-linked genes likely have more dominant effects. In this regard, Hardy and coworkers have recently speculated that motor neurons and cortical pyramidal neurons exist close to the edge of a ‘catastrophic cliff.’ Any events which expose neuronal vulnerability, for example mutations or other factors impacting on protein quality control, could be sufficient to push the cell over the cliff i.e., ultimately resulting in cell death. In other words, subtle variants such as the L341V mutation of SQSTM1, over the lifetime of a neuron affect protein quality control, which in combination with other insults ultimately tip the balance from cell survival toward cell death. If the L341V mutant SQSTM1 were to accumulate over time (due to impaired turnover) then it would also have the potential to impact on the clearance of ubiquitinated proteins destined for proteasomal degradation by delaying their delivery to the proteasome.37 Presumably in such models, at the mechanistic level proteostasis defects ultimately drive the accumulation of proteins such as TARDBP in ALS and FTLD. If the model is found to be viable, therapies directed at clearing excess TARDBP would likely involve targeting a combination of these pathways.38

**Materials and methods**

**Plasmids and peptides**

The plasmids for expression of full-length wild-type and G425R mutant SQSTM1 protein (residues 1 to 440) as GST fusion proteins (pGEX-4T-1; GE Healthcare, 28-9545-49) in *E. coli* have been described previously.21,39,40 The L341V mutant construct was produced from the wild-type plasmid by site-directed mutagenesis (QuikChange II kit; Agilent Technologies, 200523) and subsequently verified by DNA sequencing. The L341V mutation was introduced into the wild-type pDEST-mCherry-EGFP-SQSTM1 construct via site-directed mutagenesis and verified by DNA sequencing. Custom peptides (residues 332 to 351 of human SQSTM1, SGGDDWDTHSSKEVDPSTG (WT LIR) and SGGDDWDTHSSKEVDPSTG (LIR [L341V])) were synthesized by PeptideSynthetics with a purity of > 98%. The cDNA of human LC3B was cloned into the pGEX-4T-3 plasmid (GE Healthcare, 28-9545-52) between the EcoRI and SalI cloning sites. The cDNA of human UBC [ubiquitin C] was cloned into pKK233-3 (Promega, 27-4935-01) between EcoRI and HindIII sites. Both the LC3B and ubiquitin were expressed by growing transformed BL21 (DE3) *E. coli* cells in either M9 minimal media or LB as described previously.41

**Protein purification**

Ubiquitin was expressed and purified as described previously.41 The unlabeled,15N-labeled and 13C/15N doubly-labeled LC3B was expressed as described previously for UBA domain constructs of SQSTM1.30,41 Thrombin cleavage was used to remove the GST-tag from glutathione-Sepharose-immobilized LC3B fusions. Purification of the eluted proteins using a salt gradient (0 to 2 M NaCl) on a cationic column (GE Healthcare, 17115201) in 5 mM potassium phosphate, pH 7, allowed fractions of purified LC3B to be concentrated by lyophilization. Fractions were then subsequently desalted on a 5 x 5 ml
HiTrap™ desalting column (GE Healthcare, 17-1408-01) in 25 mM ammonium acetate before further lyophilization.

**Ubiquitin and LC3B affinity isolation assays**

Ubiquitin and LC3B affinity isolation assays of GST-SQSTM1 proteins were performed as described previously. Briefly, the GST fusion proteins were expressed in 10 ml cultures of *E. coli* and cells were lysed by sonication in 1 ml TBS-T buffer (10 mM Tris, 150 mM NaCl, 0.1% v/v Triton-X100 [Thermo Scientific™, 85112], pH 7.5). The lysed cells were centrifuged at 16060 g for 10 min at room temperature and the cleared supernatant fractions were diluted 1:10 in TBS-T buffer. One ml of each diluted lysate was incubated at 37°C with excess glutathione-Sepharose (GE Healthcare, 17-0756-01), ubiquitin-Sepharose (10 mg/ml human ubiquitin immobilized on CNBr-activated Sepharose 4B [Sigma-Aldrich, C9142]), LC3B-Sepharose (1 mg/ml LC3B immobilized on CNBr-activated Sepharose 4B) or control-Sepharose (CNBr-activated Sepharose prepared without ubiquitin or LC3B). The unbound proteins were then removed and the beads washed with 50 mM NaCl, pH 7.5. The mixtures were directly infused into the mass spectrometer as equal (v/v) solution mixtures in 25 mM ammonium acetate (oa-TOF). Three replicates of 6 titration samples were prepared as data processing was performed using MassLynx™ 4.1 software. Spectra were generated from the raw data acquired.

**Isothermal titration calorimetry (ITC)**

Methods for isothermal titration calorimetry (ITC) have been described previously. In brief ITC was carried out using a MicroCal VP-ITC instrument. Ten μl of WT LIR or LIR (L341V) stock solution (350 μM) in 25 mM phosphate buffer, 150 mM NaCl, pH 7.5, was sequentially injected into the ITC cell containing 30 μM LC3B at 298 K in the same buffer solution, and the exothermic heat pulse was measured. Data were corrected for the effects of buffer mixing by including an enthalpy of mixing as an iterated variable. The data were analyzed using the MicroCal Origin software to determine a Kd using a global fit standard nonlinear least-squares regression analysis.

**Protein NMR experiments**

3D NMR backbone assignments of LC3B (1H, 13Ca, 13Cb and 15N for > 95% of non-prolyl residues) were carried out on Bruker AV(III)800 (Bruker UK Limited, Coventry, UK) and Bruker AV(III)600 NMR (Bruker UK Limited, Coventry, UK) spectrometers with a QCI cryoprobe and TXI room temperature probes respectively. Standard Bruker pulse sequences with Watergate solvent suppression at 298 K were used on a 13C, 15N isotopically labeled LC3B sample of 1 mM concentration in NMR buffer solution consisting of 25 mM potassium phosphate buffer, 50 mM NaCl, 5% (v/v) D2O (Sigma-Aldrich®, 151882), 0.02% (w/v) sodium azide, pH 7.0. The assignment was achieved using a combination of 2D and 3D experiments (15N-HSQC, HNCO, HN(CA)CO, HNCA, HN(CO)CA, HN(CACB) and HN(CO)CAB) using established assignment procedures.

NMR titration experiments were executed solely on the Bruker AV(III)800 NMR spectrometer using a QCI cryoprobe and standard pulse sequences with watergate solvent suppression. NMR titration studies of 15N-labeled LC3B (0.25 mM) with unlabeled LIR-peptides (WT LIR or LIR [L341V]) were performed at 298 K and 43 Chemical shift perturbations were calculated as $\Delta\delta = [\Delta\delta_H]^2 + [\Delta\delta_N(5)]^{1/2}$, where $\Delta\delta_H$ and $\Delta\delta_N$ are the observed shift perturbations in the 1H and 15N dimension of the HSQC spectrum.

**Fluorescence microscopy analyses**

For live cell imaging the motor neuron-like cell line NSC-34 (a kind gift from Dr. Neil Cashman, University of Sheffield), a hybrid line produced by fusion of neuroblastoma with mouse motor neuron-enriched primary spinal cord cells, and the mouse neuroblastoma cell line Neuro-2a (a kind gift from...
Dr. Lynn Bedford, University of Nottingham) were used. Cells were cultured in Dulbecco’s modified Eagle’s medium (Sigma-Aldrich, D6429) supplemented with 1% (v/v) penicillin/streptomycin (Sigma-Aldrich, P0781) and 10% (v/v) FBS (Sigma-Aldrich, F9665) and kept in humidified 37°C, 5% CO₂ incubators. Undifferentiated NSC-34 or Neuro-2a cells grown in glass bottom dishes (WillCo) were transfected using Lipofectamine 3000 (Invitrogen™, L3000-08) according to manufacturer’s instructions with pDEST-mCherry-EGFP-SQSTM1 constructs (wild type or the L341V mutant). Thirty-two h after transfection cells were treated with constructs (wild type or the L341V mutant). Thirty-two h after treatment the media was removed from the cells, the cells were washed in PBS (Sigma-Aldrich, D8537) and then placed in Ringer buffer (2 mM CaCl₂, 10 mM glucose, 10 mM HEPES, 5 mM KCl, 2 mM MgCl₂-6H₂O, 155 mM NaCl, 2 mM NaH₂PO₄·H₂O, pH 7.2) and directly examined by wide-field fluorescence microscopy at 37°C for up to one h. Cells undergoing rapamycin treatment were treated as for cells without BafA1 but with 20 nM rapamycin (VIVA Bioscience Ltd., VB2112) added 3 h prior to imaging. Z-stacks were collected (200 nm spacing) using a DeltaVision Elite microscopy system (GE Healthcare Life Sciences, Buckinghamshire, UK) with a x60, 1.42 numerical aperture objective (Olympus, Southend-on-Sea, UK), images were recorded on a Photometric CoolSnap camera (Photometrics, Arizona, USA). The live cell filter wheel was used with the Quad-m channel polyychroic. EGFP fluorescence was excited at 475 nm (bandwidth 28 nm) and the emission was recorded at 525 nm (bandwidth 25 nm). mCherry fluorescence was excited at 575 nm (bandwidth 25 nm) and the emission recorded at 632 nm (bandwidth 60 nm). Images of GFP and mCherry fluorescence were acquired sequentially for each z-stack slice. Single-cell image stacks underwent iterative restoration in either Velocity (Perkin Elmer) or Huygens Professional (Scientific Volume Imaging). Velocity was used to determine a Pearson correlation coefficient (PCC) value for each cell using the Costes method for estimating thresholds. Images of each condition were taken on a minimum of 3 independent occasions and a minimum of 50 cells per condition were analyzed. For every cell analyzed the PCC value was individually determined and the mean PCC values were calculated for all cells in each condition. GraphPad Prism 6 was used to determine if the PCC values between different conditions were statistically different using a one-way ANOVA analysis with a multiplicity adjusted P value for each comparison (significance set at 0.05).

**Western blotting for autophagy markers**

HeLa and NSC-34 cells were seeded at a density of 2 × 10⁵ cells/well in 12-well plates. Twenty-four h later cells were treated with varying concentrations of BafA1 (0, 1, 10 and 20 nM). Sixteen h after treatment the media was removed from the cells, the cells were washed in PBS and then harvested in fresh PBS. Cells were then pelleted at 380 g for 5 min in a table-top centrifuge and the PBS was removed. Cells were lysed in 300 μL of RIPA buffer (150 mM NaCl, 1% IGEPA® CA-630 [Sigma-Aldrich®, 18896], 0.5% sodium deoxycholate [Sigma-Aldrich®, D6750], 0.1% SDS [Fisher Scientific, BP166], 50 mM Tris, pH 8.0, including protease [Sigma-Aldrich, P8340] and phosphatase [Sigma-Aldrich, P5726] inhibitors [at a 1:1000 dilution]). A BCA assay (ThermoFisher Scientific, 23225) determined the total protein concentration of each lysate and then lysates were separated by SDS PAGE on a 5–20% acrylamide gels with 20 μg of total protein loaded per well. Gels were transferred overnight and autophagy marker proteins were revealed by western blotting with the mouse anti-SQSTM1/lck ligand antibody, rabbit anti-SQSTM1 antibody (Enzo Life Sciences, BML-PW9860) and rabbit anti-LC3B antibody (MBL International, PM036). The blots were also stained for ACTB/β-actin (Sigma-Aldrich, A1978) to confirm equal protein loading.

**Abbreviations**

ALS amyotrophic lateral sclerosis  
MAP1LC3B/LC3B microtubule-associated protein 1 light chain 3 β (a mammalian ortholog of yeast Atg8)  
CSP chemical shift perturbation  
EGFP enhanced green fluorescent protein  
ESI-MS electrospray ionization mass spectrometry  
FTLD frontotemporal lobar degeneration  
HSQC heteronuclear single quantum correlation  
ITC isothermal titration calorimetry  
KEAP1 interacting region  
LIR LC3-interacting region  
NMR nuclear magnetic resonance  
PDB Paget disease of bone  
PB1 phox and bem1 domain  
PBS phosphate-buffered saline  
PCC Pearson correlation coefficient  
SOD1 superoxide dismutase 1, soluble  
SQSTM1/p62 sequestosome 1  
TAR DNA binding protein  
UBA ubiquitin-associated domain  
WT wild type

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

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