TBK1-mediated phosphorylation of LC3C and GABARAP-L2 controls autophagosome shedding by ATG4 protease

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Abstract

Autophagy is a highly conserved catabolic process through which defective or otherwise harmful cellular components are targeted for degradation via the lysosomal route. Regulatory pathways, involving post-translational modifications such as phosphorylation, play a critical role in controlling this tightly orchestrated process. Here, we demonstrate that TBK1 regulates autophagy by phosphorylating autophagy modifiers LC3C and GABARAP-L2 on surface-exposed serine residues (LC3C S93 and S96; GABARAP-L2 S87 and S88). This phosphorylation event impedes their binding to the processing enzyme ATG4 by destabilizing the complex. Phosphorylated LC3C/GABARAP-L2 cannot be removed from liposomes by ATG4 and are thus protected from ATG4-mediated premature removal from nascent autophagosomes. This ensures a steady coat of lipidated LC3C/GABARAP-L2 throughout the early steps in autophagosome formation and aids in maintaining a unidirectional flow of the autophagosome to the lysosome. Taken together, we present a new regulatory mechanism of autophagy, which influences the conjugation and de-conjugation of LC3C and GABARAP-L2 to autophagosomes by TBK1-mediated phosphorylation.

Keywords ATG4; ATG8; autophagy; phosphorylation; TBK1

Subject Categories Autophagy & Cell Death; Post-translational Modifications & Proteolysis

Introduction

The recycling of redundant cytosolic components and damaged organelles is termed autophagy. It is a highly conserved process, which increases during starvation conditions or during other cellular stresses [1,2]. Autophagy involves the formation of a phagophore, a double-membrane cup-shaped structure, which expands to enwrap and enclose the designated cellular cargo to form the autophagosome, which then fuses with lysosomes to enable enzymatic degradation of its cargo along with its inner membrane [3]. Upon induction of autophagy, small ubiquitin-like LC3 proteins (autophagy modifiers) are conjugated to phosphatidylethanolamine (PE) anchoring them to the growing phagophore membrane (LC3-II). This conjugation is carried out by the lipidation cascade enzymes (ATG3, ATG5, ATG7, ATG12, and ATG16L1) and allows cargo selection and autophagosome formation [4,5]. In humans, there are six autophagy-modifier proteins, grouped into two subfamilies: (i) LC3A, LC3B, and LC3C, and (ii) GABARAP, GABARAP-L1, and GABARAP-L2/GATE-16 [6]. Both subgroups play different roles during autophagy. The LC3 subgroup mainly mediates the elongation of the phagophore membrane, whereas the GABARAP subfamily acts at later stages during autophagosome sealing and autophagosome–lysosome fusion [7,8]. LC3 proteins undergo two processing steps, (i) an initial proteolytic cleavage of the peptide bond responsible for the conversion of pro-LC3 to active LC3 (LC3-I) and (ii) the subsequent cleavage of the amide bond for de-lipidation of LC3-II from autophagosomes to regenerate a free cytosolic LC3 pool. Both processing steps are catalyzed by ATG4 [9]. Among the four mammalian paralogs of ATG4, ATG4B is the most active protease followed by ATG4A and ATG4C/D, which exhibit minimal protease activity [10,11]. LC3s are bound by the ATG4B enzyme body and through LC3-interacting regions (LIRs) located at the
N- and C-terminal flexible tails of ATG4B [12]. ATG4s are cysteine proteases that cleave peptide bonds of pro-LC3 to expose the C-terminal glycine and allow conjugation with PE. ATG4 also de-conjugates LC3-II from the outer membrane of autophagosomes preceding, or just after autophagosome–lysosome fusion by cleaving the amide bond between PE and the C-terminal glycine residue of LC3s [13]. Phosphorylation of ATG4B at S383/392 increases its protease activity [14], especially during the LC3 de-lipidation phase, whereas ULK1-mediated phosphorylation of ATG4B at S316 (in humans) [15] or at S307 (in yeast) [16] reduces pro-LC3 binding and C-terminal tail cleavage. Likewise, oxidation of ATG4 by H2O2 attenuates its activity and blocks LC3 de-lipidation [17].

The serine-threonine kinase TBK1 has been implicated in the selective degradation of depolarized mitochondria (mitophagy) and intracellular pathogens (xenophagy) [18–20]. Specific autophagic cargo marked with an ubiquitin signal is recognized by autophagy receptor proteins such as optineurin (OPTN), p62, or NDP52 [5,21]. They physically bridge the cargo to the nascent phagophore by binding to ubiquitin via their UBD and to LC3 family proteins via their conserved LIR motifs, respectively. These autophagy receptors also recruit TBK1 to the site of autophagosome formation. Local accumulation of active TBK1 phosphorylates p62 and OPTN, which increases their binding affinity for polyubiquitin chains and the LC3 family proteins, thereby driving autophagy [19,20,22–27].

In this study, we expand the role of TBK1 in the autophagic process by demonstrating its ability to directly phosphorylate LC3C on S93/96 and GABARAP-L2 on S87/88. We study the consequences of LC3 phosphorylation during autophagy and show that this phosphorylation primarily impedes ATG4-mediated processing of LC3 on the liposomes, adding a new layer of regulation.

Results

TBK1 phosphorylates LC3C and GABARAP-L2 in vitro

The serine–threonine kinase TBK1 has previously been shown to phosphorylate autophagy receptors such as OPTN and p62 [19,20,26]. To test whether recombinant TBK1 can also directly phosphorylate autophagy modifiers, we performed an in vitro kinase assay. LC3A, LC3C, GABARAP-L1, and GABARAP-L2 are directly phosphorylated by TBK1 in vitro, and the phosphorylation sites of LC3 family proteins were identified by mass spectrometry (Fig 1A and B). Since LC3A is only weakly phosphorylated and the detected phosphorylation site of GABARAP-L1 is a tyrosine residue (most likely an in vitro assay artifact), we decided to further investigate the phosphorylation sites of LC3C at S93 and S96 and GABARAP-L2 at either S87 and S88, which could not be unambiguously assigned. The TBK1-mediated phosphorylation sites of LC3C (Fig EV1A) and GABARAP-L2 (Fig EV1B) are topologically equivalent and are present in surface-exposed loops (depicted in red). The position of the loop is in between β-strand 3 and α-helix 3 in both LC3C and GABARAP-L2, on the opposite face of the LIR binding pocket. This indicates that LIR-mediated interactions of LC3C might not be affected directly upon phosphorylation. The TBK1-mediated phosphorylation sites of LC3C (Fig 1C) and GABARAP-L2 (Fig 1D), marked with red triangles, are mostly conserved in orthologs from higher eukaryotes and fit to the general TBK1 consensus phosphorylation motif (Fig 1E), which prefers a hydrophobic residue (mostly leucine) after the target phosphoserine site [28]. Moreover, these phosphorylation sites are situated in solvent-accessible regions of LC3C and GABARAP-L2 (Fig 1F).

TBK1 phosphorylates and binds LC3C and GABARAP-L2 in cells

To test whether TBK1 also phosphorylates LC3C in cells, HEK293T cells were SILAC-labeled and either WT TBK1 (heavy label) or TBK1 kinase dead (K38A; light label) was overexpressed along with GFP-LC3C or with GFP-GABARAP-L2. GFP proteins were immuno-precipitated and analyzed by mass spectrometry. Phosphorylation at positions S93 and S96 of LC3C was enhanced in the presence of WT TBK1 (factors 10 and 6, respectively; Fig 2A), as compared to TBK1 K38A. Similarly, the presence of WT TBK1 resulted in enhanced phosphorylation of GABARAP-L2 at S87 and S88 (by factors 13 and 2, respectively; Fig 2A). To confirm this phosphorylation event on an endogenous level, we treated cells with control or TBK1 siRNA and induced TBK1 kinase activity with CCCP treatment (Fig 2B). siRNA-mediated depletion of TBK1 in cells resulted in a twofold reduction of S96 LC3C and S87 GABARAP-L2 phosphorylation. Unfortunately, our efforts to generate phospho-specific antibodies against GABARAP-L2 S87-PO4 and GABARAP-L2 S87/88-PO4 failed (Fig EV1C).

Next, we visualized the phosphorylation event by using Phos-tag™ polyacrylamide gels, where phosphorylated proteins are retained by the Phos-tag reagent and appear at a higher molecular weight. Overexpression of WT TBK1, but not TBK1 kinase dead,
**Figure 1.**

(A) **in vitro kinase assay**

\[ \gamma^{32} P \text{ATP, GST-TBK1} \]

(B) | protein     | site | PEP value | localization probability |
|----------|------|-----------|-------------------------|
| LC3A     | S29  | 1.49E-03  | 1.00                    |
| LC3A     | S61  | 1.47E-02  | 1.00                    |
| LC3C     | T48  | 1.04E-13  | 1.00                    |
| LC3C     | T56  | 1.04E-13  | 1.00                    |
| LC3C     | S93  | 3.6E-273  | 1.00                    |
| LC3C     | S96  | 2.13E-17  | 0.98                    |
| GABARAP-L1| Y25  | 1.05E-02  | 1.00                    |
| GABARAP-L2| S10  | 3.16E-66  | 1.00                    |
| GABARAP-L2| S39  | 8.28E-14  | 1.00                    |
| GABARAP-L2| S87  | 2.22E-09  | 0.66                    |
| GABARAP-L2| S88  | 5.61E-03  | 0.54                    |

(C) Residue conservation

(D) Residue conservation

(E) TBK1 substrate specificity

(F) Residue conservation

**Figure 2.**

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### Figure 2. TBK1 phosphorylates and binds LC3C and GABARAP-L2 in cells.

**A, B** Identification of phosphosites by mass spectrometry following GFP-LC3C or GFP-GABARAP-L2 immunoprecipitation. (A) TBK1 WT was overexpressed in heavy- and light-labeled SILAC HEK293T cells. (B) HEK293T cells were treated with control or TBK1 siRNA and CCCP (3 h, 40 μM). Phosphosite intensities were normalized to total protein intensity. Data are presented as mean ± SD, n = 3 biological replicates.

**C** SDS–PAGE and Western blot of HEK293T cell lysates. Cells were transfected with HA-LC3C, TBK1 WT or K38A, and GFP-Parkin and left untreated or treated with CCCP (3 h, 40 μM) to induce mitophagy. The ratio of phosphorylated to unphosphorylated LC3C was quantified. Data are presented as mean ± SD, n = 3 biological replicates, *P < 0.05, as analyzed by Student’s t-test.

**D** SDS–PAGE and Western blot of HEK293T and HeLa cell lysates and GST-LC3C or GST-GABARAP-L2 immunoprecipitations.

**E** SDS–PAGE and Western blot of HEK293T cell lysates transfected with full-length TBK1, a C-terminal truncation mutant (TBK1 ΔC), a kinase-dead version (TBK1 K38A), and GST-LC3C or GST-GABARAP-L2 immunoprecipitations.

Source data are available online for this figure.
induced an upward shift and retention of phosphorylated LC3C (Fig 2C). The ratio of phosphorylated to unphosphorylated LC3C from three independent experiments was quantified and is higher upon the induction of mitophagy, by the addition of Parkin, an E3 ligase, and CCCP, a mitochondrial depolarization agent (Fig 2C). The induction of mitophagy leads to TBK1 recruitment to autophagosomes with concomitant kinase activation and results in increased LC3C phosphorylation.

Endogenous TBK1 from HeLa or HEK293T cell lysate binds to GST-LC3C and GST-GABARAP-L2 (Fig 2D) indicating direct physical interaction. The binding of TBK1 to LC3C and GABARAP-L2 is independent of its catalytic activity (Fig 2D) and could be mediated through its C-terminal coiled-coil region (Fig 2E), which is known to bind to OPTN [29].

**Phospho-mimetic LC3C impedes ATG4 cleavage and binding**

To understand the consequences of phosphorylated-LC3C, we looked at its phosphosites in more detail. The LC3C phosphorylation sites S93 and S96 are situated on the face opposite to the hydrophobic pocket enabling LIR binding (Figs 1F and EV1A) and are therefore less likely to influence the direct binding of LC3C to autophagy receptors or adaptors. However, they are in close proximity to the C-terminal tail of LC3C which is proteolytically processed. ATG4-mediated processing of the LC3C C-terminal tail allows lipid conjugation and adherence to autophagosomes. To test whether phosphorylation of these residues could impair the proteolytic cleavage of the LC3C C-terminal tail by ATG4, an in vitro cleavage assay was performed. Double-tagged His-LC3C-Strep WT, S93/96A, or phospho-mimetic LC3C S93/96D were incubated with ATG4B for indicated times, and the C-terminal cleavage of LC3C was monitored by detecting the appearance of truncated His-LC3C protein (Fig 3A).

ATG4B cleaves the entire pool of LC3C WT or S93/96A within 10 min, whereas only half of the phospho-mimetic LC3C S93/96D pool is cleaved (Fig 3A). When LC3 proteins are overexpressed in HEK293T cells, they are rapidly processed by endogenous ATG4 proteins. The C-terminal tail of LC3C that is cleaved by ATG4s is considerably larger (21 residues) than that of other LC3 family proteins. Hence, a pro-form of LC3C S93/96D can be visualized by separating the cell lysate on a 15% polyacrylamide gel (Fig 3B).

The inability of ATG4 to process phosphorylated LC3C might be distinct during stress conditions. To test this, we induced mitophagy in HEK293T cells by adding CCCP. Upon induction of mitophagy, LC3C S93/96D could not be completely processed by ATG4s (Fig 3C). Similarly, GABARAP-L2 phospho-mimetic (S88D) could not be cleaved by endogenous ATG4s, impairing subsequent lipidation (Fig 3D). We reasoned that this inability of ATG4 to process LC3C S93/96D and GABARAP-L2 S87/88D could be due to an impediment in direct protein binding, and therefore tested this by co-expressing GFP-LC3 proteins with Flag-ATG4A, Flag-ATG4B, Flag-ATG4C, and Flag-ATG4D in HEK293T cells and subjected them to GFP immunoprecipitation (Fig EV2A). Phospho-mimetic mutants LC3C S93/96D and GABARAP-L2 S87/88D displayed reduced binding to ATG4A, ATG4B, ATG4C, and ATG4D, and the Kd of WT GABARAP-L2 binding to ATG4A is half the Kd of GABARAP-L2 S87/88D (Fig EV2B). In addition, we performed mass spectrometry analysis of GABARAP-L2 WT, S87/88A, and S87/88D binding partners. Differential binding of GABARAP-L2 interaction partners (significantly enriched according to a FDR 0.05 corrected Student’s t-test with a minimal enrichment factor of 2) to S87/88D and S87/88A mutants is depicted in Fig 3E. The binding of endogenous ATG4B is significantly decreased (P < 0.05) in immunoprecipitated samples of GABARAP-L2 S87/88D compared to WT. Other well-known GABARAP-L2 binding partners such as TBK1, p62, NBR1, and ATG7 bind equally well to WT and S87/88D or S87/88A mutants (Fig 3E). To understand this reduced binding, we modeled the full-length LC3C-ATG4B complex based on the core crystal structure [30] (see Materials and Methods). We tested the effect of phosphorylation at both these sites (S93 and S96) by modeling phosphate groups onto serine residues in the LC3C-ATG4B complex and performed molecular dynamics (MD) simulations (up to 1.5 μs). We found that the WT LC3C-ATG4B complex with and without additional LIR interactions between ATG4B and LC3C remained stable. The C-terminal tail of LC3C remained bound and strongly anchored to the active site of ATG4B throughout the simulation. In the complex, the phosphosites S93 and S96 of LC3C (red cartoon in Fig 3F) are in close proximity to the ATG4B interacting surface (gray surface). LC3C S96 forms a hydrogen bond interaction with ATG4B E350, and LC3C S93 is close to a network of hydrogen bonds and salt bridges. In MD simulations, double phosphorylation of S93 and S96 interfered with these interactions and disrupted the binding interface between ATG4B and LC3C (Movies EV1 and EV2). The phosphorylated serine residues detached from the ATG4B surface and partially dislodged the LC3C, resulting in partial retraction of the LC3C C-terminal tail from the ATG4B active site. The negative charge introduced by phosphorylation severely weakens complex stability based on calculated binding energies (Table 1), with electrostatic interactions as the dominant factor. Figure EV2C–E shows residue-wise contributions to the binding energy mapped onto the LC3C structure. According to these calculations, phosphorylated S93 and S96 are strongly destabilizing (Fig EV2C–E; red thick cartoon), whereas unphosphorylated S93 and S96 are favorable (Fig EV2C–E; blue thin cartoon). The MD simulations and binding energy calculations indicate that phosphorylation disrupts the LC3C-ATG4B interface and destabilizes the complex.

ATG4 acts on pro-LC3s upon ribosomal release to activate and process them to LC3-1 under basal conditions, as well as after autophagosome formation to recycle lipidated LC3s. However, under basal conditions TBK1 is not active to phosphorylate LC3s. TBK1 kinase activity is induced at autophagy initiation [31,32] (e.g., during mitochondrial depolarization), which is when phosphorylation potentially disrupts the LC3-ATG4 interface.

**Phosphorylation at S93 and S96 affects LC3C C-terminal tail structure and thereby impedes ATG4-mediated cleavage**

Based on the simulation results for the LC3C-ATG4B complex, we hypothesized that phosphorylation of unbound LC3C could affect its C-terminal tail structure and prevent binding to the ATG4B active site. In MD simulations (see Materials and Methods) of free LC3C, we found that the C-terminal tail of LC3C (126–147) was disordered and highly dynamic (Movie EV3). By contrast, in the phosphorylated LC3C variants (S93-PO4 and S96-PO4), the C-terminal tail structure and prevent binding to the ATG4B active site. In MD simulations (see Materials and Methods) of free LC3C, we found that the C-terminal tail of LC3C (126–147) was disordered and highly dynamic (Movie EV3). By contrast, in the phosphorylated LC3C variants (S93-PO4 and S96-PO4), the C-terminal tail adopted more ordered conformations (Movies EV4 and EV5; Fig 4A–C). The phosphoserines formed intramolecular salt bridges with R134 (Fig 4A and B) that pulled the C-terminal tail of LC3C toward the protein, structuring it locally. In repeated simulations
Figure 3.
Figure 3. Phospho-mimetic LC3C and GABARAP-L2 impede ATG4 cleavage and binding.

A SDS–PAGE and Western blot of in vitro ATG4 cleavage assay. Purified double-tagged His-LC3C-Strep WT and mutants were incubated with ATG4B for indicated time points. LC3C S93/96D mutation slows down C-terminal cleavage of LC3C by ATG4B.

B SDS–PAGE and Western blot of HEK293T cell lysates transfected with LC3C WT or mutants. S93/96D mutation of LC3C impedes cleavage of pro-LC3C by endogenous ATG4s.

C, D SDS–PAGE and Western blot of HEK293T cell lysates transfected with LC3C WT or mutants (C) or GABARAP-L2 WT or mutants (D). Cells were left untreated or treated with 40 μM CCCP for 3 h to induce mitophagy.

E Mass spectrometry interactome analysis of HEK293T cells with overexpressed HA–GABARAP-L2 WT, S88A, and S87/R88D. Data were analyzed with Perseus, and binding partners that specifically bind GABARAP-L2 in comparison with mock IP are depicted (n = 3 biological replicates).

F Full-length LC3C (red cartoon) binds to ATG4B (gray surface) with its C-terminal tail accessible to the active site of ATG4B. Phosphorylation of LC3 at S93 and S96 (sticks) affects binding to ATG4B. Zoom-up showing the side and top view of LC3C–ATG4B interface. S96 of LC3C and E350 of ATG4B form direct hydrogen bonds across the interface. S93 position is central to a network of polar interactions (blue dashed lines; side chains shown as sticks) across the interface.

Source data are available online for this figure.

Table 1. Binding free energy computations for LC3C–ATG4B complexes.

| System/Energy terms (kJ/mol) | LC3C–ATG4B (unphosphorylated) | LC3C–ATG4B (unphosphorylated + LIR) | LC3C–ATG4B (S93/596–PO4 + LIR) |
|-----------------------------|-------------------------------|--------------------------------|--------------------------------|
| van der Waals               | −1,036.2 ± 110.2               | −985.9 ± 91.6                  | −970.1 ± 128.4                  |
| Electrostatic               | −4,652.7 ± 536.4               | −4,495.8 ± 315.7               | −1,610.2 ± 293.4                |
| Polar solvation             | 2,940.0 ± 406.0                | 2,883.0 ± 339.2                | 2,672.4 ± 311.0                 |
| SASA                        | −136.2 ± 145                   | −134.8 ± 10.2                  | −125.2 ± 14.0                   |
| Total binding energy        | −2,885.1 ± 305.2               | −2,733.5 ± 204.0               | −331.2 ± 202.4                  |
| ΔΔG                         | −                              | 151.6 ± 367.1                 | 2,851.9 ± 966.2                 |

The table lists different energetic contributions (mean ± SD) to the binding of LC3C and ATG4B in different phosphorylation states and with modeled LIR–WXXL interactions. The different binding energy contributions were computed using the MM–PBSA approach implemented in g_mmpbsa (see Materials and Methods) from MD simulations of LC3C–ATG4B complexes. The non-bonded energy terms (van der Waals and electrostatic) contribute significantly to the molecular mechanics interaction energy of the complex, whereas changes in the bonded terms (bond length, angle, and dihedral terms) do not contribute significantly to the interaction energy during complex formation.

Table 2. Comparison of ΔG values for LC3C and GABARAP-L2 with different modifications.

| Modification       | ΔG (kcal/mol) |
|--------------------|---------------|
| WT                 | −20.4         |
| S93–D              | −20.5         |
| S93–D, S96–D       | −20.5         |
| S93–D, S96–D, R134A| −20.5         |
| R134A              | −20.5         |
| S93–D, S96–D, R142A| −20.5         |
| R142A              | −20.5         |

The table compares the binding free energy (ΔG) values for LC3C and GABARAP-L2 with different modifications. The ΔG values are calculated using the MM–PBSA approach implemented in g_mmpbsa (see Materials and Methods) from MD simulations of LC3C–ATG4B complexes.

(n = 6 each) of unphosphorylated and phosphorylated variants of LC3C (Fig EV3A–C), we observed a total of six salt-bridge formation events, indicating that the intramolecular salt-bridge formation between the phosphoserines and R134 is robust. We observed the salt-bridge formation on a sub-microsecond time scale (Fig 4D and E). To confirm this finding and the role of R134, we performed ATG4-mediated in vitro cleavage experiments of double-tagged LC3C WT, S93/96D, S93/96D R134A, and S93/96D R142A (a control mutation in the C-terminal tail). The LC3C C-terminal cleavage was monitored by the disappearance of its C-terminal Strep-tag. The mutation of S93/96D delayed the cleavage of the C-terminal tail of LC3C by ATG4B (Fig 4F). The R134A mutation could partially rescue this phenotype of S93/96D, whereas the other C-terminal tail mutation, R142A, could not (Fig 4F). The results of the ATG4-mediated cleavage assay are thus consistent with R134-phosphoserine interactions sequestering the LC3C C-terminal tail and preventing access to ATG4B and subsequent cleavage.

Phospho-mimetic LC3C and GABARAP-L2 cannot localize to autophagosomes

GABARAP-L2 lacks the C-terminal tail, and the ATG4B-mediated processing removes only a single C-terminal residue (F117), which exposes G116 for lipidation. Therefore, we hypothesized that phosphorylating S87 and S88 in GABARAP-L2 weakens binding to ATG4B and in turn slows down proteolytic processing. Accordingly, we tested whether phospho-mimetic GABARAP-L2 S88D and LC3C S93/96D can localize to autophagosomes, despite not being processed by ATG4B. U2OS cells were co-transfected with HA–Parkin and GFP–GABARAP-L2 WT, S88D, S88A (Fig 5A and B) or GFP–LC3C WT, S93/96D, S93/96A (Fig 5C and D), and mitophagy was induced by the addition of CCCP for 3 h. Upon induction of mitophagy, GABARAP-L2 WT and S88A localized to autophagosomes. By contrast, the phospho-mimetic GABARAP-L2 S88D remained dispersed throughout the cell and no autophagosome formation was observed (Fig 5A and B). Likewise, phospho-mimetic LC3C S93/96D did not localize to autophagosomes upon induction of mitophagy unlike WT and S93/96A LC3C (Fig 5C and D).

Phospho-mimetic Δ C-terminal LC3C and GABARAP-L2 are not lipidated and do not localize to autophagosomes

Since LC3 family proteins can only be integrated into autophagosomes after C-terminal cleavage by ATG4, we tested whether artifically truncated LC3C or GABARAP-L2 (Δ C-term: LC3C (1–126) and GABARAP-L2 (1–116)) could circumvent ATG4-mediated processing, undergo lipidation, and form autophagosomes. U2OS cells were co-transfected with HA–Parkin and GFP–GABARAP-L2 Δ C-term WT, S88A or S88D (Figs 6A and EV4A), or GFP–LC3C Δ C-term WT, S93/96A or S93/96D (Figs 6B and EV4B), and mitophagy was induced by the addition of CCCP for 3 h. Upon induction of mitophagy, GABARAP-L2 or LC3C Δ C-term WT and alanine mutants localized to autophagosomes, while phospho-mimetic mutants with truncated C-terminus (GABARAP-L2 Δ C-term S88D or LC3C Δ C-term S93/96D) remained dispersed throughout the cell and no autophagosome formation could be observed (Fig 6A and B). Upon induction of
mitophagy, GABARAP-L2 lipidation can be observed by the appearance of a lower band on Western blots (Fig 3D), which can also be observed during mitophagy induction of GABARAP-L2 Δ C-term WT and S88A, but not with phospho-mimetic GABARAP-L2 Δ C-term S88D (Fig EV4C). Hence, the phosphorylation of LC3C or GABARAP-L2 not only impedes their C-terminal cleavage by ATG4, but also their lipidation by the lipidation cascade enzymes ATG12-ATG5-ATG16L1. ATG7 function is similar to ubiquitin-activating (E1) enzymes; it recruits ATG3 (an E2-like enzyme), which then catalyzes the conjugation of the lipid moiety (PE) to the C-terminal exposed glycine of the truncated LC3 Δ C-term. Binding of the ATG12-ATG5-ATG16L1 complex (E3-like enzyme) to ATG3 enhances the lipidation of LC3, since the ATG12-ATG5-ATG16L1 complex ensures that nascently lipidated LC3 is incorporated into the phagophore membrane [4]. In order to test whether phosphorylation of LC3C/GABARAP-L2 Δ C-term impedes their processing by the lipidation cascade enzymes, we also performed an in vitro lipidation assay (Fig 6C). LC3C Δ C-term WT and LC3C Δ C-term S93/96D or GABARAP-L2 Δ C-term WT and GABARAP-L2 Δ C-term S87/88D were incubated with ATP, liposomes, hATG3, hATG7, and hATG12-ATG5-ATG16L1 (reaction mix). WT LC3C and GABARAP-L2 could be successfully lipidated, while S93/96D LC3C and S87/88D GABARAP-L2 could not efficiently be conjugated to membrane in vitro (Fig 6C and D), indicating that phosphorylation of LC3s also affects lipid conjugation.

**TBK1-mediated GABARAP-L2 phosphorylation impedes its premature cleavage from autophagosomes by ATG4**

TBK1 is recruited to the site of autophagosome formation by autophagy receptor proteins, where TBK1 phosphorylates OPTN and p62 to promote autophagy flux [19,20,22–27,31,32]. Hence, it is most likely that LC3C and GABARAP-L2 are phosphorylated by TBK1 during autophagosome formation and not during the initial processing step of pro-LC3 cleavage post-ribosomal release. In order to test whether the TBK1-mediated phosphorylation of
Figure 5. Phospho-mimetic LC3C and GABARAP-L2 cannot localize to autophagosomes.

A–D U2OS cells were transfected with GFP-GABARAP-L2 (A) or GFP-LC3C (C) WT or mutants and HA-Parkin. Mitophagy was induced by the addition of 40 μM CCCP for 3 h. WT, S87/88A GABARAP-L2 and WT, S93/96A LC3C localize to autophagosomes, whereas S87/88D GABARAP-L2 and S93/96D LC3C remain dispersed in the cytosol. Scale bar represents 10 μm. (B, D) GFP-expressing cells were counted and segregated into classes with greater and less than 10 autophagosomes per cell. Data are presented as mean ± SD, n = 3 biological replicates, **P < 0.01, ***P < 0.001, ns = not significant, as analyzed by Student's t-test.
LC3C and GABARAP-L2 has an impact on ATG4-mediated de-lipidation of LC3s from the mature autophagosome, an in vitro de-lipidation assay of LC3C-II and GABARAP-L2-II proteins was performed. The fractions of PE-conjugated LC3C Δ C-term WT, phospho-mimetic S93/96D and PE-conjugated GABARAP-L2 Δ C-term WT and phospho-mimetic S87/88D were enriched (see Materials and Methods) and used as substrates for the de-lipidating enzymes ATG4A or ATG4B (Fig 7A, D and E). WT and S87/88A GABARAP-L2-II could be de-lipidated from liposomes by ATG4B and ATG4A (at a slower rate) (Fig 7A and D) in a dose-dependent manner (Fig EV5A). On the contrary, we found that GABARAP-L2 S87/88D is not a target of ATG4A or ATG4B (Fig 7A, D and E). In addition, we tested whether TBK1 is able to phosphorylate GABARAP-L2 and LC3C that are already incorporated into liposomes (Fig 7B). 10 μM LC3C/GABARAP-L2-I or LC3C/GABARAP-L2-II, or 5 μM of each LC3C/GABARAP-L2-I and LC3C/GABARAP-L2-II were incubated with TBK1, and phosphorylation was monitored by autoradiography. TBK1 phosphorylates PE-lipidated forms (-II) equally well as unlipidated (-I) LC3s (Fig 7B). In order to verify that the phospho-mimetic S87/88D GABARAP-L2-II mutant behaves similar to TBK1-phosphorylated GABARAP-L2-II, we have also tested the ability of ATG4A (Fig EV5B) and ATG4B (Figs 7C and D, and EV5B) to cleave TBK1-phosphorylated GABARAP-L2 from PE and observed that the cleavage rate is decreased in comparison with unphosphorylated protein.
Figure 7. TBK1-mediated GABARAP-L2 phosphorylation impedes its premature cleavage from autophagosomes by ATG4.

A. GABARAP-L2 WT-, S87/88D-, or S87/88A-conjugated liposomes were treated or not with 2 μM ATG4A or ATG4B for 1 h at 37°C. Samples were then subjected to SDS–PAGE together with unconjugated GABARAP-L2-I.

B. Coomassie stain and autoradiography of SDS–PAGE after an in vitro kinase assay with GST-TBK1, PE-conjugated (II), and unconjugated (I) GABARAP-L2 and LC3C.

C. PE-conjugated GABARAP-L2 WT was incubated in phosphorylation assay buffer in the presence or absence of GST-TBK1 for 4 h and treated or not with 2 μM ATG4B for 30 min. Samples were then subjected to SDS–PAGE together with unconjugated GABARAP-L2-I.

D. The extent of de-lipidation in (A) and (C) was quantified and plotted as percentage of total protein (conjugated and unconjugated). W = WT, D = S87/88D, A = S87/88A, pW = TBK1-phosphorylated WT GABARAP-L2. Data are presented as mean ± SD, n = 3 biological replicates, *P < 0.05, **P < 0.01, ***P < 0.001, ns = not significant, as analyzed by Student’s t-test.

E. LC3C WT-, LC3C S93/96D-, GABARAP-L2 WT-, or GABARAP-L2 S87/88D-conjugated liposomes were treated or not with 2 μM ATG4A, ATG4B, or RavZ for 1 h at 37°C. Samples were then subjected to SDS–PAGE.

Source data are available online for this figure.
Finally, we also tested whether the phosphorylation of GABARAP-L2 has an impact on its de-lipidation from the phagophore by other proteases such as RavZ (Figs 7E and EV5A). RavZ is a bacterial effector protein from the intracellular pathogen Legionella pneumophila that interferes with autophagy by directly and irreversibly uncoupling GABARAP-L2 attached to PE on autophagosome membranes [33,34]. We found that small amounts of RavZ could remove GABARAP-L2 WT and S87/88D mutant from liposomes (Figs 7E and EV5A), indicating its effectiveness in circumventing Legionella growth restriction via xenophagy (when TBK1 is also activated). Likewise, RavZ is also able to cleave LC3C WT and S93/96D mutant from liposomes in vitro (Fig 7E), whereas ATG4B is able to de-lipidate and cleave small amounts of LC3C Δ C-term WT from liposomes, but has no activity toward LC3C Δ C-term S93/96D (Fig 7E).

**TBK1-mediated LC3 phosphorylation ensures steady autophagy flux**

To assess the physiological relevance of these phosphorylation sites, we tested whether phosphorylated LC3C or GABARAP-L2 adhered to autophagosomes are still functional to perform downstream reactions. LC3 family proteins interact with autophagosome receptors such as p62, which link the growing autophagosome to cargo [35,36]. Both LC3 C-terminal WT and phospho-mimetic S93/96D can bind to p62 (Fig 8A). Similarly, p62 and OPTN can be recruited to autophagosomes by WT as well as S87/88D GABARAP-L2 (Fig 8B) [20,37]. Once all of the cargo has been engulfed by the autophagosome, degradation can take place through the fusion with lysosomes [38]. GABARAP family proteins mediate autophagosomal-lysosomal fusion by binding to the autophagy adaptor protein PLEKHM1 [39,40]. Phosphorylation of GABARAP-L2 by TBK1 does not interfere with its ability to bind to PLEKHM1 (Fig 8B). In order to assess the physiological significance of this phosphorylation event, we have generated tandem monomeric mCherry-GFP-LC3C WT (Fig 8C) and alanine mutant cells to measure autophagic flux by FACS (Fig 8D) and high-content microscopy (Fig EV5C). Phagophores or autophagosomes that have not fused with lysosomes exhibit GFP and mCherry signal. In contrast, amphisomes or autolysosomes do not exhibit GFP fluorescence, since the GFP signal is sensitive to the acidic conditions of the lysosome lumen and only the mCherry signal is stabilized. Cells treated with bafilomycin A1 exhibit “yellow” (mCherry and GFP colocalized) puncta due to the inhibition of lysosomal acidification of the autophagosomes. LC3C S93/96A mutant cells, which cannot be phosphorylated by TBK1 upon mitophagy, exhibit fewer GFP puncta (Figs 8C and D, and EV5C). This indicates that WT LC3C (which is phosphorylated during mitophagy) stays longer on the autophagosome and is protected from ATG4 premature de-lipidation, whereas the alanine mutant LC3C is recycled quickly and fuses with lysosomes.

Hence, TBK1-mediated phosphorylation of GABARAP-L2 and LC3C does not interfere with cargo binding but protects them from premature removal from autophagosomes by ATG4 and ensures efficient cargo engulfment with subsequent degradation.

**Discussion**

The autophagy pathway is tightly regulated to ensure proper recycling and disposal of cellular material during nutrient shortage. Here, we present a new regulatory mechanism of autophagy, which influences the conjugation and de-conjugation of LC3C and GABARAP-L2 to autophagosomes. The kinase TBK1 fulfills several roles during selective autophagy. Upon autophagy induction, TBK1 is recruited to the site of autophagosome formation and gets activated by trans-autophosphorylation after accumulation [28,41]. We show that, at this stage, TBK1 can phosphorylate LC3C and GABARAP-L2 at specific serine residues to protect them from ATG4-mediated premature removal from autophagosomes.

ATG4 mediates regular processing of pro-LC3s post-ribosomal release and de-lipidation of incorrectly lipidated LC3-II on other endomembranes and favors incorporation of LC3s into autophagosomes by ATG12-ATG5-ATG16L1. Spatial and temporal regulation of recruitment and dissociation of LC3 family proteins to and from autophagosomes is achieved through regulation of ATG4 activity [13–17]. ATG4 constitutively de-conjugates LC3 family proteins from all endomembranes except from autophagosomes, to maintain a pool of unlipidated LC3 [42]. This suggests that LC3-II conjugated to autophagosomes is protected from premature de-lipidation by a timely regulatory mechanism. This regulation is achieved through the phosphorylation and dephosphorylation of ATG4 itself [15,16] and the phosphorylation of LC3s by TBK1.

The kinase activity of TBK1 is tightly regulated, and during xenophagy and mitophagy, TBK1 phosphorylates autophagy receptor proteins [19,20,22,23,25–27]. The active recruitment of TBK1 to the sites of autophagosome formation by NDP52 [19,23,31,32] hints that TBK1-mediated phosphorylation occurs on nascent phagophores, resulting in phosphorylated forms of membrane-embedded LC3s. Phosphorylation prevents premature removal of lipidated LC3C/GABARAP-L2 from growing autophagosomes by ATG4. Molecular modeling and atomistic simulations of the ATG4B-LC3

**Figure 8.** TBK1-mediated LC3 phosphorylation ensures steady autophagy flux.

A, B SDS–PAGE and Western blot of HEK293T cell lysates and GFP IPs. (A) Cells were transfected with GFP, GFP-LC3C Δ C-terminal WT or S93/96D, and lysates used for GFP IPs WT and S93/96D LC3C bind endogenous p62. (B) Cells were transfected with GFP, GFP-GABARAP-L2 Δ C-terminal WT or S87/88D, and lysates used for GFP IPs WT and S87/88D GABARAP-L2 bind endogenous p62, optineurin (OPTN), and PLEKHM1.

C U2OS cells were seeded onto coverslips and transfected with mCherry-GFP-LC3C WT or S93/96A or mCherry-GFP-GABARAP-L2 WT or S87/88A. Mitophagy was induced by the addition of 40 μM CCCP for 3 h, and lysosomal degradation of GFP was blocked by the addition of 200 nM bafilomycin A1 for 3 h. Scale bar represents 10 μm.

D HeLa cells expressing doxycycline-inducible Parkin and stable mCherry-GFP-LC3C WT or S93/96A were treated with 1 μg/ml doxycycline and 10 μM CCCP for 16 h. mCherry and GFP fluorescence was measured by FACS, and the fold change of median fluorescence intensity of GFP over mCherry was calculated for each sample. Data are presented as mean ± SD. n = 6 biological replicates. **P < 0.01, ns = not significant, as analyzed by Student’s t-test.

Source data are available online for this figure.
Figure 8.
complex revealed that LC3C phosphorylation impedes binding to ATG4. The weakened binding slows down de-lipidation, which ensures that a steady coat of lipidated LC3C/GABARAP-L2 is maintained throughout the early steps in autophagosome formation (Fig 9). The phosphorylation of LC3C/GABARAP-L2 does not impede their binding to autophagy receptors such as p62 or PLEKHM1 [39], which promotes unhindered downstream steps, i.e., cargo engulfment and subsequent autophagosome–lysosome fusion. Thus, phosphorylation of LC3s aids in efficient cargo engulfment and maintaining an unperturbed and unidirectional flow of the autophagosome to the lysosome.

At later stages of autophagosome formation, this process could be slowed down or reversed either by TBK1 dissociation from autophagosomes or by diminished catalytic activity. Alternatively, action of phosphatases could reverse this effect and allow de-lipidation by ATG4 prior to autophagosomal–lysosomal fusion, thereby efficiently recycling LC3 proteins.

**Materials and Methods**

**Expression constructs**

Expression constructs of indicated proteins were cloned into indicated vectors using PCR or the gateway system. Site-directed mutagenesis was performed by PCR to introduce desired amino acid substitutions. All expression constructs were sequenced by Seqlab.

**Protein expression and purification**

GST or His-tagged fusion proteins were expressed in *Escherichia coli* strain BL21 (DE3). Bacteria were cultured in LB medium supplemented with 100 µg/ml ampicillin at 37°C in a shaking incubator (150 rpm) until OD600 ~ 0.5–0.6. Protein expression was induced by the addition of 0.5 mM IPTG, and cells were incubated at 16°C for 16 h. Bacteria were harvested by centrifugation (1,800 g) and lysed by sonication in GST lysis buffer (20 mM Tris–HCl, pH 7.5, 10 mM EDTA, pH 8.0, 5 mM EGTA, 150 mM NaCl, 0.1% β-mercaptoethanol, 1 mM PMSF) or His lysis buffer (25 mM Tris–HCl, pH 7.5, 200 mM NaCl, 0.1% β-mercaptoethanol, 1 mM PMSF, 1 mg/ml lysozyme). For the purification of ATG4, the use of PMSF was omitted. Lysates were cleared by centrifugation (16,800 g), 0.05% of Triton X-100 was added, and the lysates were incubated with glutathione-Sepharose 4B beads (GE Life Sciences) or Ni-NTA agarose beads (Thermo Fisher) on a rotating platform at 4°C for 1 h. The beads were washed five times either in GST wash buffer (20 mM Tris–HCl, pH 7.5, 10 mM EDTA, pH 8.0, 150 mM NaCl, 0.5% Triton X-100, 0.1% β-mercaptoethanol, 1 mM PMSF) or in His wash buffer (25 mM Tris–HCl, pH 7.5, 200 mM NaCl, 0.05% Triton X-100, 10 mM imidazole). The immobilized proteins

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**Figure 9. Model of TBK1-mediated LC3C and GABARAP-L2 phosphorylation.**

TBK1 recruitment to autophagosomes promotes phosphorylation of membrane-embedded LC3C and GABARAP-L2. Phosphorylation prevents premature removal of LC3C and GABARAP-L2 from autophagosomes, ensuring an unperturbed and unidirectional flow of the autophagosome to the lysosome.
were reconstituted in GST storage buffer (20 mM Tris–HCl, pH 7.5, 0.1% NaN₃, 0.1% β-mercaptoethanol) or eluted with His elution buffer (25 mM Tris–HCl, pH 7.5, 200 mM NaCl, 300 mM Imidazole) and dialyzed in 25 mM Tris–HCl, pH 7.5, 200 mM NaCl at 4°C overnight.

Recombinant GST-TBK1 and GST-ATG4A were obtained from the MRC PPU DSTT in Dundee, UK.

Purification of proteins used for in vitro lipiddation/de-lipiddation: Full-length hATG7, hATG3, and hATG5-12-ATG16L1 complex was expressed and purified from HEK suspension cells (HEK-F, Invitrogen) as previously described [43]. To purify mATG3, LC3C, LC3C S93/96D, GABARAP-L2 WT, GABARAP-L2 S87/88D, and RavZ, pGEX-6P-1 plasmid containing the corresponding cDNA was transcribed into BL21-Gold (DE3) grown for three additional hours before they were collected by centrifugation. The supernatant was incubated at 4°C overnight.

The supernatant was incubated at 4°C with Glutathione Beads (Sigma) for 4 h. Beads were collected and washed twice with NT350 buffer before HRV 3C protease was added and allowed to cut at 4°C overnight. The next morning, protein fractions were collected and stored at −80°C in 20% glycerol. (mATG3 and RavZ plasmids were a gift from Thomas Melia (Yale University)).

Cell culture

HEK293T, HeLa, and U2OS cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco) supplemented with 10% fetal bovine serum, 2 mM L-glutamine, and 1% penicillin/streptomycin and maintained at 37°C in a humidified atmosphere with 5% CO₂.

Recombinant GST-TBK1 and GST-ATG4A were obtained from the MRC PPU DSTT in Dundee, UK.

Immunoprecipitation and protein binding assays

Cleared cell extracts were mixed with glutathione-Sepharose beads (GE Healthcare) conjugated to LC3 family proteins or GST, FLAG-, or HA-agarose beads (Sigma-Aldrich) or GFP-Trap_A beads (ChromoTek) for 2 h at 4°C on a rotating platform. The beads were washed four times in lysis buffer. Immunoprecipitated and input samples were reduced in SDS sample buffer (50 mM Tris–HCl, pH 6.8, 10% glycerol, 2% SDS, 0.02% bromophenol blue, 5% β-mercaptoethanol) and heated at 95°C for 5 min [44,45].

Western blot

For immunoblotting, proteins were resolved by SDS-PAGE and transferred to PVDF membranes. Blocking and primary antibody incubations were carried out in 5% BSA in TBS-T (150 mM NaCl, 20 mM Tris, pH 8.0, 0.1% Tween-20), and secondary antibody incubations were carried out in 5% low-fat milk in TBS-T and washings in TBS-T. Blots were developed using Western Blot Luminol Reagent (sc-2048; Santa Cruz).

Antibodies

The following antibodies were used in this study: anti-HA-tag (11867423001; Roche), anti-FlagM2-tag (F3165; Sigma-Aldrich), anti-GFP-tag (ab23666; Abcam). Secondary HRP-conjugated antibodies goat anti-rabbit (sc-2030; Santa Cruz), goat anti-rat (sc-2048; Santa Cruz), and goat anti-mouse (sc-2031; Santa Cruz). IgGs were used for immunoblotting.

Kinase assays

LC3-I or LC3-II family proteins were incubated in 20 μl phosphorylation buffer (50 mM Tris–HCl, pH 7.5, 10 mM MgCl₂, 0.1 mM EGTA, 20 mM β-glycerophosphate, 1 mM DTT, 0.1 mM Na₃VO₄, 0.1 mM ATP, or γ³²P ATP (500 cpm/pmol)) with 1 μg of recombinant GST-TBK1 for 30 min at 30°C. The kinase assay was stopped by adding SDS sample buffer containing 1% β-mercaptoethanol and heating at 95°C for 5 min. The samples were resolved by SDS-PAGE, and the gels were stained with InstantBlue (Expe-deon) and dried. The radioactivity was analyzed by autoradiography [46].

Phosphopeptide identification

Cells were maintained in custom-made SILAC DMEM (heavy: R10, K8/light R0, K0) for 14 days, treated accordingly, and lysed...
(as stated above). Incorporation of labeled amino acids to more than 95% was verified by mass spectrometry. Lysates of SILAC-labeled cells expressing GFP-tagged LC3C or GFP-tagged GABARAP-L2, TBK1 WT (heavy labeled), and TBK1 K38A (light labeled) were combined at equal amounts and incubated with GTF-Trap beads for 1 h, followed by washes under denaturing conditions (8 M Urea). Bound proteins were incubated with GFP-Trap beads for 1 h, followed by washes and loaded onto 4–20% gradient SDS–PAGE gels [19]. Alternatively, in vitro phosphorylated LC3 family proteins (as stated above) were used to determine TBK1-dependent phosphorylation sites. Proteins were stained using InstantBlue and digested in-gel with trypsin. Label-free phosphopeptide identification was performed with cells maintained in normal media and treated with control or TBK1 siRNA. HEK293T cells were treated with 20 nM TBK1 siRNA (sequence: 5’-GACAGAGUUGUGAATACCTT-3’) for 24 h. Prior to cell lysis in total cell lysis buffer, cells were treated with 40 μM CCCP for 3 h and 100 nM Calyculin A ( (#9902; Cell Signaling Technology) for 15 min. Lysates of cells expressing GTF-tagged LC3C or GFP-tagged GABARAP-L2 were incubated with GTF-Trap beads for 1 h, followed by washes under denaturing conditions (8 M Urea). Bound proteins were reduced and alkylated with 5 mM TCEP and 20 mM chloroacetamide prior to digestion with trypsin. SILAC, in vitro phosphorylated or label-free peptides were desalted on reversed-phase C18 StageTips and analyzed on an Orbitrap Elite™ or Q Exactive HF mass spectrometer (Thermo Fisher). The raw data were analyzed using MaxQuant 1.6.5.0. The database used to identify the peptides was the sequence of either GABARAP-L2 or LC3C downloaded from UniProt and a database with common laboratory contaminants. The FDR was set to 1% on protein, PSM, and site decoy level. Phosphorylation sites were validated using manual spectra interpretation.

**Interactome analysis by mass spectrometry**

HEK293T cells were transfected with control plasmid, HA-GABARAP-L2 WT, HA-GABARAP-L2 S87/88A, or HA-GABARAP-L2 S87/88D plasmids, lysed in IP Lysis Buffer, and immunoprecipitated as stated above. Bound proteins were eluted in NuPAGE LDS Sample Buffer (Life Technologies) supplemented with 1 mM DTT, boiled at 70°C for 10 min, alkylated, and loaded onto 4–20% gradient SDS–PAGE gels. Proteins were stained using InstantBlue and digested in-gel with trypsin. Peptides were desalted on reversed-phase C18 StageTips and analyzed on a Q Exactive HF mass spectrometer (Thermo Fisher). The raw data were analyzed using MaxQuant 1.6.5.0 with standard settings and activated LFQ quantification. The database used to identify the peptides was the human reference protein database (UniProt downloaded December 2017), and the FDR was set to 1% on protein, PSM, and site decoy level. Statistical analysis was done with Perseus 1.6.5. Proteins were defined as interactors, if they passed a 5% FDR-corrected one-sided two-sample Student’s t-test with a minimal enrichment factor of two compared to a HA-only IP. For interactors, differential binding of S to D/A mutants compared to WT was detected by two-sided two-sample t-tests.

**Phos-tag™ SDS–PAGE**

Phos-tag™ acrylamide (Wako) gels were used as indicated by the supplier. Gels were prepared with 10% acrylamide, 50 μM Phos-tag™, and 100 μM MnCl₂. Cells were lysed in SDS sample buffer supplemented with 100 μM MnCl₂. Western blot lanes were quantified using ImageJ.

**Bio-layer interferometry**

GST-ATG4A was obtained from the MRC PPU DSTT in Dundee, UK, and His-GABARAP-L2 WT and S87/88D proteins were purified as described above. GST-ATG4A was diluted to 1 μg/ml in bio-layer interferometry (BLI) assay buffer (300 mM NaCl, 0.004% Tween-20, 0.05% BSA in PBS) and immobilized onto anti-GST biosensors (Pall ForteBio). His-GABARAP-L2 WT and S87/88D proteins were diluted in BLI assay buffer to 20, 10, 5, and 0.625 μM concentrations in a black flat-bottom 96-well plate. The kinetic binding assay was performed on Octet RED96 instrument (Pall ForteBio). Reference wells were subtracted from sample wells, and a 1:1 global fitting model was used to determine the steady-state curve with K_d values.

**ATG4 cleavage assay of unlipidated LC3C**

His-LC3C-Strep double-tag proteins were purified as described above and incubated in buffer (50 mM Tris–HCl, pH 7.5, 1 mM EDTA, 150 mM NaCl, 1.5 mM DTT) at 37°C for indicated time points. The assay was stopped by adding SDS sample buffer containing 1% β-mercaptoethanol and heating at 95°C for 5 min. The samples were resolved by SDS–PAGE and imaged by Western blot.

**Modeling and MD simulations with analysis**

Starting from the human LC3C (8–125) structure (PDB id: 3WAM; [47]), we added N- and C-terminal overhangs using MODELLER [48] to construct full-length LC3C (1–147). The LC3C-ATG4B complex was modeled using the core complex structure of LC3B(1–124)-ATG4B(1–357) [PDB id: 2Z0E; [30] as template using MODELLER [48]. The C-terminal tails of both LC3C and ATG4B were modeled in extended conformations without steric clashes across their interface. Additional unresolved loops in ATG4B were modeled using the loop modeling protocol of MODELLER [48]. ATG4B contains an N- and a C-terminal LIR, both of which can, in principle, interact with the WXXL-binding sites on either non-substrate or substrate LC3. Therefore, we modeled an alternative WT complex structure including the interaction between the C-terminal LIR of ATG4B and the WXXL-binding site on substrate LC3C. Phosphoserines at positions S93 and S96 were modeled using CHARMM-GUI [49]. All structures were solvated in TIP3P water and 150 mM NaCl. After energy minimization, MD simulations of different phosphorylation states were performed using GROMACS v5.1 [50], with position-restrained NVT equilibration and NPT equilibration runs for 1,000 ps each. Production runs at 310 K and 1 atm were simulated for different times (see Table 2). We used the CHARMM36m force field [51], the Nosé–Hoover thermostat [52], the Parrinello–Rahman barostat [53], and a time step of 2 fs. For each of the LC3C systems (Table 2), six replicates were simulated with different initial velocities. We also used the molecular mechanics Poisson–Boltzmann surface area (MM-PBSA) to compute the binding
energies of the phosphorylated and unphosphorylated LC3C-ATG4B complexes as implemented in g_mmpbsa [54]. These binding energies contain molecular mechanical (MM), polar, and non-polar solvation energies. MM energies depend on bonded and non-bonded terms including electrostatic (E_{ele}) and van der Waals (E_{vdw}) contributions. The polar solvation energies were computed at an ionic strength of 150 mM, a solvent dielectric constant of 80, and a protein dielectric constant of 2 by solving the linearized Poisson–Boltzmann equation with a fine grid width of 0.5 Å and a coarse grid width of 1.5 times the long axis of the complex, as implemented in Assisted Poisson–Boltzmann Solver (APBS) [55]. The non-polar solvation contributions were estimated with the SASA model using a probe radius of 1.4 Å, a surface tension of $\gamma = 0.0226$ kJ/mol/Å$^2$, and an offset of 3.84 kJ/mol [56]. Binding free energies were estimated as the difference energies between bound and free states,

$$
\Delta G_{\text{binding}} = G_{\text{LC3C-ATG4B}} - G_{\text{LC3C}} - G_{\text{ATG4B}},
$$

where the free energy contributions of the protein complex and free proteins are decomposed into a sum of molecular mechanics, solvent, and configurational entropy contributions,

$$
G = \Delta G_{\text{MM}} + \Delta G_{\text{solv}} - T \Delta S
$$

$$
G = \Delta E_{\text{bnd}} + \Delta E_{\text{vdw}} + \Delta E_{\text{ele}} + \Delta G_{\text{polar}} + \Delta G_{\text{non-polar}} - T \Delta S
$$

The binding energies were evaluated at intervals of 10 ns from the 1,000-ns MD trajectories and averaged (see Table 1). Double differences between unphosphorylated and phosphorylated complexes minimize systematic errors caused by possible energy-function inaccuracy. For the dynamic LC3C-ATG4B protein complexes studied here, these calculated free energy differences point to trends, but should not be interpreted in terms of dissociation constants.

### Liposome and proteoliposome preparation

All lipids were purchased and dissolved in chloroform from Avanti Polar Lipids (Alabaster, AL). Liposomes were prepared by combining 55 mol% 1,2-dioleoyl-sn-glycero-3-phosphoethanola- mine (DOPE), 35 mol% 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC), and 10 mol% bovine liver phosphoinositol (PI). The lipids were dried under nitrogen gas, and the lipid film was further dried under vacuum for 1 h. The lipids were reconstituted in NT350 buffer (350 mM NaCl, 20 mM Tris–HCl pH 7.4) and subjected to seven cycles of flash-freezing in liquid nitrogen and thawing in a 37°C bath. Liposomes were further sonicated immediately prior to the lipidation reaction.

### Lipidation reaction of LC3C and GABARAP-L2

For a full lipidation reaction, LC3C, LC3C S93/96D, GABARAP-L2 WT, or GABARAP-L2 S87/88D (10 μM) were mixed with hATG7 (0.5 μM), hATG3 (1 μM), ATG12-ATG5-ATG16L1 (0.25 μM), sonicated liposomes (3 mM), and 1 mM DTT. Lipidation was initiated by adding 1 mM ATP, and reactions were incubated at 30°C for 90 min. Samples were mixed with LDS loading buffer and immediately boiled to stop further lipidation. The reactions were run on a 4–20% SDS-PAGE gel and visualized by Coomassie blue stain and analyzed with Image Lab 6.0 (Bio-Rad).

### De-lipidation reaction of LC3C and GABARAP-L2

For a full de-lipidation reaction, LC3C, LC3C S93/96D, GABARAP-L2 WT, or GABARAP-L2 S87/88D (10 μM) were mixed with hATG7 (0.3 μM), mAtg3 (containing an extended N-terminal amphipathic helix that permits lipidation in the absence of ATG12-ATG5-ATG16L1) (1 μM), sonicated liposomes (3 mM), and 1 mM DTT. Lipidation was initiated by adding 1 mM ATP, and reactions were incubated at 30°C for 90 min. After the reaction was complete, the lipidation reaction was run on a Nycode buz density gradient. The bottom layer of the gradient consisted of 150 μl of 80% Nycodez and 150 μl of the lipidation reaction. The second layer consisted of 250 μl of 30% Nycodez, while the top layer was 50 μl of NT350 buffer. Gradients were centrifuged at 82,000 g at 4°C for 4 h in a
Beckman SW55 rotor. Liposomes with the conjugated LC3C or GABARAP-L2 protein were collected from the top of the tube before use in subsequent de-lipidation experiments. To measure the activity of proteases, 10 μM of proteoliposomes (concentration estimated by Coomassie blue stain) were mixed with NT350 buffer and kept on ice until activity assays were initiated by the addition of 2 μM (or indicated amounts) of either ATG4A, ATG4B, or RavZ. Reactions were incubated at 37°C for 1 h.

De-lipidation reactions of phosphorylated GABARAP-L2 were performed by adding GST-TBK1 and phosphorylation buffer to the samples for 4 h prior to the addition of 2 μM ATG4A or ATG4B for 30 min. Samples were mixed with SDS loading buffer and immediately boiled to stop proteolysis. The reactions were run on a 4–20% SDS–PAGE gel and visualized by Coomassie blue stain and analyzed with Image Lab 6.0 (Bio-Rad).

**Yokogawa CQ1 microscopy imaging**

HeLa cells expressing doxycycline-inducible Parkin and stable mCherry-GFP-LC3C WT or S93/96A HeLa cells were seeded onto black, clear flat-bottom 24-well plates and treated with 1 μg/ml doxycycline and 10 μM CCCP for 16 h. Cells were incubated with Hoechst 33342 (R37605; Thermo Fisher) and washed in PBS before fixation (2.6*10⁵ cells/well) and treated with 1 μM mCherry-GFP-LC3C WT or S93/96A were seeded onto 6-well plates. After Yokogawa with 4% paraformaldehyde for 15 min at room temperature. Plates were mixed with SDS loading buffer and immediately boiled to stop proteolysis. The reactions were run on a 4–20% SDS–PAGE gel and visualized by Coomassie blue stain and analyzed with Image Lab 6.0 (Bio-Rad).

**Flow cytometry (FACS)**

HeLa cells expressing doxycycline-inducible Parkin and stable mCherry-GFP-LC3C WT or S93/96A were seeded onto 6-well plates (2.6*10⁵ cells/well) and treated with 1 μg/ml doxycycline and 10 μM CCCP for 16 h. mCherry and GFP fluorescence was measured on a FACS Canto II machine and analyzed using BD FACSDiva 8.0.1 software. The fold change of median fluorescence intensity of GFP/mCherry was calculated for each sample.

**Statistical analysis**

Data are presented as the mean with error bars indicating the SD (standard deviation). Statistical significance of differences between experimental groups was assessed with Student’s t-test. Statistical analysis for in vitro lipidation assays was also performed by one-way ANOVA followed by Bonferroni’s multiple comparison test. Differences in means were considered significant if $P < 0.05$. Differences with $P < 0.05$ are annotated as $^*$, $P < 0.01$ are annotated as $^{**}$, $P < 0.001$ are annotated as $^{***}$, and $P < 0.0001$ are annotated as $^{****}$. All Western blots shown are representative of biological replicates.

**Data availability**

All data are available in the main manuscript text or the supplementary materials and will be made available upon request. The mass spectrometry proteomic data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the following dataset identifier: in vitro phosphorylation sites: PXD015065, www.ebi.ac.uk/pride/archive/simpleSearch?q=PXD015065; GABARAP-L2 SILAC phosphorylation sites: PXD015075, www.ebi.ac.uk/pride/archive/simpleSearch?q=PXD015075; LC3C SILAC phosphorylation sites: PXD015076, www.ebi.ac.uk/pride/archive/simpleSearch?q=PXD015076; endogenous phosphorylation sites (si/TBK1 or control): PXD015077, www.ebi.ac.uk/pride/archive/simpleSearch?q=PXD015077; and GABARAP-L2 interactome data: PXD015155, www.ebi.ac.uk/pride/archive/simpleSearch?q=PXD015155.

**Expanded View** for this article is available online.

**Acknowledgements**

We thank Andreas Ernst and Suchithra Guntrum from Fraunhofer Institute, Stefan Stein and Annette Trzmiel from the FACS facility of the Georg-Speyer-Haus Frankfurt for technical help with FACS measurements, and Masato Akutsu and David McEwan for very valuable comments. This work was supported by grants from the DFG (SFB 1177 on selective autophagy), The Cluster of Excellence “Macromolecular Complexes” of the Goethe University Frankfurt (EXC 115). L.H. is supported by a European Molecular Biology Organization (EMBO) long-term postdoctoral fellowship (ALTF 1200-2014, LTFCOFUND2013, GA-2013-609409). R.M.B. and G.H. acknowledge support by the Max Planck Society and computational resources at MPMDC, Garching, A.H.L. and A.S. were supported by the Research Council of Norway (project number 221835) and through its Centers of Excellence funding scheme (project number 262652), as well as the Norwegian Cancer Society (project number 171318).

**Author contributions**

LH and ID conceived the study. LH designed and performed most of the experiments to generate Fig. 8C. FB analyzed mass spectrometry data. LH and ID wrote the manuscript with contribution from all authors. All authors approved the final version of the manuscript.

**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**

1. Xie W, Zhou J (2018) Aberrant regulation of autophagy in mammalian diseases. Biol Lett 14: 20170540
2. Dikic I (2017) Proteasomal and autophagic degradation systems. Annu Rev Biochem 86: 193–224
3. Yang Z, Klionsky DJ (2010) Eaten alive: a history of macroautophagy. Nat Cell Biol 12: 814–822
4. Nakatogawa H (2013) Two ubiquitin-like conjugation systems that mediate membrane formation during autophagy. Essays Biochem 55: 39–50
5. Stolz A, Ernst A, Dikic I (2014) Cargo recognition and trafficking in selective autophagy. Nat Cell Biol 16: 495–501
6. Cemma M, Grinstein S, Brumell JH (2016) Autophagy proteins are not universally required for phagosome maturation. *Autophagy* 12: 1440—1446

7. Weidberg H, Shvets E, Shpilka T, Shimron F, Shinder V, Elazar Z (2010) LC3 and GATE-16/GABARAP subfamilies are both essential yet act differently in autophagosome biogenesis. *EMBO J.* 29: 1792—1802

8. Nguyen TN, Padman BS, Usher J, Oorschot V, Ramm G, Lazarou M (2016) Atg8 family LC3/GABARAP proteins are crucial for autophagosome-lysosome fusion but not autophagosome formation during PINK1/Parkin mitophagy and starvation. J Cell Biol 215: 857—874

9. Zhang L, Li J, Ouyang L, Liu B, Cheng Y (2016) Unraveling the roles of Atg4 proteins from autophagy modulation to targeted cancer therapy. *Cancer Lett* 373: 19—26

10. Li M, Hou Y, Wang J, Chen X, Shao ZM, Yin XM (2011) Kinetics comparisons of mammalian Atg4 homologues indicate selective preferences toward diverse Atg8 substrates. *J Biol Chem* 286: 7327—7338

11. Lystad AH, Simonsen A (2019) Mechanisms and pathophysiologica roles of the ATG8 conjugation machinery. Cells 8: 697

12. Maruyama T, Noda NN (2017) Autophagy-regulating protease Atg4: structure, function, regulation and inhibition. *J Antibiot (Tokyo)* 71: 72—78

13. Yu ZQ, Ni T, Hong B, Wang HY, Jiang FJ, Zou S, Chen Y, Zheng XL, Klon-sky DJ, Liang Y et al. (2012) Dual roles of Atg8–PE deconjugation by Atg4 in autophagy. *Autophagy* 8: 883—892

14. Yang Z, Wilkie-Grantham RP, Yanagi T, Shu CW, Matsu-zawa S, Reed JC (2015) ATG4C (Autopagin-1) phosphorylation modulates autophagy. J Biol Chem 290: 26549—26561

15. Pengo N, Agrotis A, Prak K, Jones J, Ketteler R, Cemma M, Grinstein S, Brumell JH, Herhaus L, Dikic I (2015) The structure of Atg8 family LC3/GABARAP reveals a mechanism for deconjugation of LC3. *J Cell Sci* 128: 309—314

16. Wild P, Farhan H, McEwan DG, Wagner S, Rogov VV, Brady NR, Richter J, Waidmann I, Choudhary C et al. (2016) Phosphorylation of the autophagy receptor optineurin restricts Salmonella growth. Science 333: 228—233

17. Richter B, Sliter DA, Herhaus L, Stolz A, Wang C, Beli P, Zaffagnini G, Wild P, Martens S, Wagner SA et al. (2016) Phosphorylation of OPTN by TBK1 enhances its binding to Ub chains and promotes selective autophagy of damaged mitochondria. *Proc Natl Acad Sci USA* 113: 4039—4044

18. Wild P, Farhan H, McEwan DG, Wagner S, Rogov VV, Brady NR, Richter B, Korac J, Waidmann I, Choudhary C et al. (2011) Phosphorylation of the autophagy receptor optineurin restricts Salmonella growth. Science 333: 228—233

19. Herhaus L, Dikic I (2015) Expanding the ubiquitin code through post-translational modification. *EMBO Rep* 16: 1071—1083

20. Heo JM, Orduereau A, Paulio JA, Rinehart J, Harper JW (2015) The PINK1–PARKIN mitochondrial ubiquitylation pathway drives a program of OPTN/NDPS2 recruitment and TBK1 activation to promote mitophagy. *Mol Cell* 60: 7—20

21. Lazarou M, Sliter DA, Kane LA, Sarraf SA, Wang C, Burman JL, Sideris DP, Fogel AI, Youle RJ (2015) PINK1 phosphorylation of ubiquitin recruits autophagy receptors to mitochondria and upstream autophagy machinery for mitophagy. Nature 524: 309—314

22. Matsumoto G, Wada K, Okuno M, Kurosawa M, Nukina N (2011) Serine 403 phosphorylation of p62/SQSTM1 regulates selective autophagic clearance of ubiquitinated proteins. *Mol Cell* 44: 279—289

23. Orduereau A, Heo J, Duda DM, Olzewski JL, Yanishevski D, Rinehart J, Schulman B, Harper JW (2015) Defining roles of PARKIN and ubiquitin phosphorylation by PINK1 in mitochondrial quality control using a ubiquitin replacement strategy. *Proc Natl Acad Sci USA* 112: 6637—6642

24. Pilli M, Arko-Mensah J, Poppuak M, Roberts E, Master S, Mandell MA, Dupont N, Ormatowski W, Jiang S, Bradfute SB et al. (2012) TBK1 promotes autophagy-mediated antimicrobial defense by controlling autophagosome maturation. *Immunity* 37: 223—234

25. Thurston TL, Ryzhakov G, Bloor S, von Muñihinen N, Randow F (2009) The TBK1 adaptor and autophagy receptor NDP52 restricts the proliferation of ubiquitin-coated bacteria. *Nat Immunol* 10: 1215—1221

26. Ma X, Helgason E, Phung QT, Quan CL, Iyer RS, Lee MW, Bowman KK, Starovasnik MA, Dubeur EC (2012) Molecular basis of Tank-binding kinase 1 activation by transautophosphorylation. *Proc Natl Acad Sci USA* 109: 9378—9383

27. Freischmidt A, Wieland T, Richter B, Ruf W, Schaeffer V, Muller K, Marro-quin N, Nordin F, Hubers A, Weydert P et al. (2015) Haploinsufficiency of TBK1 causes familial ALS and fronto-temporal dementia. *Nat Neurosci* 18: 631—636

28. Satoo K, Noda NN, Kurmata H, Fujioka Y, Mizushima N, Ohsumi Y, Inagaki F (2009) The structure of Atg4B–LC3 complex reveals the mechanism of LC3 processing and delipidation during autophagy. *EMBO J.* 28: 1341—1350

29. Vargas JNS, Wang C, Bunker E, Hao L, Maric D, Schiavo G, Randow F, Youle RJ (2019) Spatiotemporal control of ULK1 activation by NDP52 and TBK1 during selective autophagy. *Mol Cell* 74: 347—362 e346

30. Ravenhill BJ, Boyle KB, von Muñihinen N, Ellison CJ, Masson GR, Otten EG, Foeglein A, Williams R, Randow F (2019) The cargo receptor NDP52 initiates selective autophagy by recruiting the ULK complex to cytosol-invasive bacteria. *Mol Cell* 74: 320—329 e326

31. Choy A, Dancourt J, Mugo B, O'Connor TJ, Isberg RR, Melia TJ, Roy CR (2012) The Legionella effector RavZ inhibits host autophagy through irreversible Atg8 deconjugation. *Science* 338: 1072—1076

32. Kwon DH, Kim S, Jung YO, Roh K-H, Kim L, Kim B-W, Hong SB, Lee YJ, Song H, Lee WC et al. (2017) The 1.2 complex between RavZ and LC3 reveals a mechanism for deconjugation of LC3 on the phagophore membrane. *Autophagy* 13: 70—81

33. Pankiv S, Clausen TH, Lamark T, Brech A, Bruun JA, Outzen H, Overvatn A, Bjørkøy G, Johansen T (2007) p62/SQSTM1 binds directly to Atg8/LC3 to facilitate degradation of ubiquitinated protein aggregates by autophagy. *J Biol Chem* 282: 24131—24145

34. Zheng YT, Shahnazari S, Brech A, Lamark T, Johansen T, Brumell JH (2009) The adaptor protein p62/SQSTM1 targets invading bacteria to the autophagy pathway. *J Immunol* 183: 5909—5916

35. Wong YC, Holzbaeur EL (2014) Optineurin is an autophagy receptor for damaged mitochondria in parkin-mediated mitophagy that is disrupted by an ALS-linked mutation. *Proc Natl Acad Sci USA* 111: E4439—E4448

36. Nakamura S, Yoshimori T (2017) New insights into autophagosome-lysosome fusion. *J Cell Sci* 130: 1209—1216

37. McEwan DG, Popovic D, Cubas A, Terawaki S, Suzuki H, Stadel D, Coxon FP, de Stegmann DM, Bhogaraju S, Maddi K et al. (2015) PLEKH1M regulates autophagosome-lysosome fusion through HOPS complex and LC3/GABARAP proteins. *Mol Cell* 57: 39—54
40. Wang H, Sun HQ, Zhu X, Zhang L, Albanesi J, Levine B, Yin H (2015) GABARAPs regulate PI4P-dependent autophagosome-lysosome fusion. Proc Natl Acad Sci USA 112: 7015 – 7020
41. Shu C, Sankaran B, Chaton CT, Herr AB, Mishra A, Peng J, Li P (2013) Structural insights into the functions of TBK1 in innate antimicrobial immunity. Structure 21: 1137 – 1148
42. Nakatogawa H, Ishii J, Asai E, Ohsumi Y (2012) Atg4 recycles inappropriately lipidated Atg8 to promote autophagosome biogenesis. Autophagy 8: 177 – 186
43. Lystad AH, Carlsson SR, de la Ballina LR, Kauffman KJ, Nag S, Yoshimori T, Melia TJ, Simonsen A (2013) Atg8 Lipidation in Autophagy-related Processes. Nat Cell Biol 21: 372 – 383
44. Herhaus L, Al-Salihi M, Macartney T, Weidlich S, Sapkota GP (2013) OTUB1 enhances TGFbeta signalling by inhibiting the ubiquitylation and degradation of active SMAD2/3. Nat Commun 4: 2519
45. Herhaus L, Al-Salihi MA, Dingwell KS, Cummins TD, Wasmus L, Vogt J, Ewan R, Bruce D, Macartney T, Weidlich S et al (2014) USP15 targets ALK3/BMPRIa for deubiquitylation to enhance bone morphogenetic protein signalling. Open Biol 4: 140065
46. Herhaus L, Perez-Oliva AB, Cozza G, Gourlay R, Weidlich S, Campbell DG, Pinna LA, Sapkota GP (2015) Casein kinase 2 (CK2) phosphorylates the deubiquitylase OTUB1 at Ser16 to trigger its nuclear localization. Sci Signal 8: ra35
47. Suzuki H, Tabata K, Morita E, Kawasaki M, Kato R, Dobson RC, Yoshimori T, Wakatsuki S (2014) Structural basis of the autophagy-related LC3/Atg13 LIR complex: recognition and interaction mechanism. Structure 22: 47 – 58
48. Sali A, Blundell TL (1999) Comparative protein modelling by satisfaction of spatial restraints. J Mol Biol 234: 779 – 815
49. Jo S, Kim T, Iyer VG, Im W (2008) CHARMM-GUI: a web-based graphical user interface for CHARMM. J Comput Chem 29: 1859 – 1865
50. Pronk S, Pall S, Schulz R, Larsson P, Bjelkmar P, Apostolov R, Shirts MR, Smith JC, Kasson PM, van der Spoel D et al (2013) GROMACS 4.5: a high-throughput and highly parallel open source molecular simulation toolkit. Bioinformatics 29: 845 – 854
51. Huang J, Rauscher S, Nawrocki G, Ran T, Feig M, de Groot BL, Grubmuller H, Mackerell AD Jr (2017) CHARMM36m: an improved force field for folded and intrinsically disordered proteins. Nat Methods 14: 71 – 73
52. N65e S (1984) A unified formulation of the constant temperature molecular dynamics methods. J Chem Phys 81: 511 – 519
53. Parrinello M, Rahman A (1981) Polymorphic transitions in single crystals: a new molecular dynamics method. J Appl Phys 52: 7182 – 7190
54. Kumar R, Kumar R, Open Source Drug Discovery Consortium, Lynn A (2014) g_mmpbsa--a GROMACS tool for high-throughput MM-PBSA calculations. J Chem Inf Model 54: 1951 – 1962
55. Koncercy R, Baker NA, McCammon JA (2012) iAPBS: a programming interface to Adaptive Poisson-Boltzmann Solver (APBS). Comput Sci Discov 5: 015005
56. Lee MR, Duan Y, Kollman PA (2000) Use of MM-PB/SA in estimating the free energies of proteins: application to native, intermediates, and unfolded villin headpiece. Proteins 39: 309 – 316
57. Ashkenazy H, Abadi S, Martz E, Chay O, Mayrose I, Pupko T, Ben-Tal N (2016) ConSurf 2016: an improved methodology to estimate and visualize evolutionary conservation in macromolecules. Nucleic Acids Res 44: W880 – W885

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