HSF1 stress response pathway regulates autophagy receptor SQSTM1/p62-associated proteostasis

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ABSTRACT
Proteostasis is important for protecting cells from harmful proteins and is mainly controlled by the HSF1 (heat shock transcription factor 1) stress response pathway. This pathway facilitates protein refolding by molecular chaperones; however, it is unclear whether it functions in autophagy or inclusion formation. The autophagy receptor SQSTM1/p62 is involved in selective autophagic clearance and inclusion formation by harmful proteins, and its phosphorylation at S349, S403, and S407 is required for binding to substrates. Here, we demonstrate that casein kinase 1 phosphorylates the SQSTM1 S349 residue when harmful proteins accumulate. Investigation of upstream factors showed that both SQSTM1 S349 and SQSTM1 S403 residues were phosphorylated in an HSF1 dependent manner. Inhibition of SQSTM1 phosphorylation suppressed inclusion formation by ubiquitinated proteins and prevented colocalization of SQSTM1 with aggregation-prone proteins. Moreover, HSF1 inhibition impaired aggregate-induced autophagosome formation and elimination of protein aggregates. Our findings indicate that HSF1 triggers SQSTM1-mediated proteostasis.

Introduction
The accumulation of harmful proteins, such as misfolded proteins and protein aggregations, leads to cellular dysfunction and cell death. In order to avoid such accumulations, protein quality control is performed via the protein homeostasis (proteostasis) network, a multicompartmental system that coordinates protein synthesis, folding, disaggregation, and degradation. HSF1 (heat shock transcription factor 1) is the master regulator of the proteostasis network and is present normally in a negatively regulated state as an inert monomer. Upon exposure to a variety of stresses, HSF1 is derepressed; it then trimerizes and translocates to the nucleus. Subsequently, activated HSF1 binds to the heat shock element (HSE), a stress-responsive promoter, resulting in the promotion of expression of heat shock proteins (HSPs) such as the molecular chaperones HSPA1/HSP70 and HSP90AA/HSP90. The degradation of harmful proteins is principally mediated by the ubiquitin-proteasome system (UPS) and the autophagy-lysosome pathway. Although macroautophagy (the degradation of cellular contents, hereafter referred to as autophagy) has formerly been considered to be a nonselective degradation mechanism, a number of studies have shown that some substrates, such as protein aggregates and impared organelles, are selectively eliminated by this process. A range of autophagic receptor proteins are involved in this selective process, e.g., SQSTM1/p62, BNIP3, BNIP3L, FUNDC1, NBR1, CALCOCO2/NDP52, and OPTN. SQSTM1, which has an LC3-interacting receptor domain, is involved in selective autophagic clearance of protein aggregates, KEAP1, and cytosolic pathogens. Recent studies have revealed that SQSTM1 must be phosphorylated at S349, S403, or S407 in order to bind to its substrates. S349 phosphorylation is mediated by Mtorc1 (mechanistic target of rapamycin [serine/threonine kinase] complex 1) and is required for interaction with KEAP1. Phosphorylation at S403 by TBK1 (TANK binding kinase 1) and CSNK2/Ck2 (casein kinase 2) is necessary for SQSTM1 involvement in the elimination of protein aggregates and mycobacteria. In addition to its function as an autophagy receptor, SQSTM1 plays an important role in inclusion formation by ubiquitinated proteins. The accumulation of ubiquitinated protein inclusions by impairment of autophagy or proteasomes is suppressed by the loss of SQSTM1. The formation of protein inclusions and aggresomes is assumed to be a cytoprotective mechanism to sequester harmful proteins. Thus, SQSTM1 is one of the key molecules for proteostasis.

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unknown how these processes are initiated and regulated. Recently, it was reported that several stress sensor proteins, such as XBP1, HSF1, and STAT3, are involved in the regulation of autophagy-related protein expression. For example, the endoplasmic reticulum stress sensor XBP1 triggers an autophagic response via autophagic vesicle formation and expression of BECN1/Beclin1 and LC3B.\textsuperscript{16} Moreover, HSF1 cooperatively binds with PSMD10/Gankyrin (26S non-ATPase regulatory proteasome subunit 10) on the ATG7 promoter in cells at an advanced stage of starvation and upregulates ATG7 expression.\textsuperscript{17} These findings lead us to speculate that stress response pathways are involved in autophagy regulation; thus, it is important to elucidate the process of autophagy regulation in detail.

In this study, we investigated the mechanism of SQSTM1 phosphorylation and the role of phosphorylated SQSTM1 in inclusion formation by ubiquitinated proteins. We found that SQSTM1 phosphorylation was mediated by the HSF1 stress response pathway, and resulted in inclusion formation and autophagic clearance of harmful proteins (aggrephagy). These results suggest that the HSF1 stress response pathway is involved in SQSTM1-associated proteostasis functions.

**Results**

**Colocalization of phosphorylated SQSTM1 with protein aggregates**

SQSTM1 is phosphorylated at one site in the KEAP1-interaction region domain (S349) and 2 sites in the ubiquitin associated domain (S403 and S407) (Fig. 1A); these phosphorylation events regulate selective autophagic clearance of ubiquitinated proteins, KEAP1, and mycobacteria.\textsuperscript{9,12} We investigated whether phosphorylation of SQSTM1 was induced by the generation of different types of protein aggregate such as ubiquitinated proteins, nonubiquitinated aggregation-prone protein, and a disease-associated protein. Proteasome inhibition by MG132 led to the accumulation of ubiquitinated proteins and induction of SQSTM1 phosphorylation at both S349 and S403 (Fig. 1B). When phosphorylated SQSTM1 was treated with \( \lambda \) protein phosphatase, these bands were not detected by anti-phosphorylated SQSTM1 antibodies (Fig. S1A). Immunocytochemical analysis showed that S349-phosphorylated SQSTM1 and S403-phosphorylated SQSTM1 colocalized with polyubiquitinated protein inclusions (Fig. 1C). To examine SQSTM1 phosphorylation further, we used expression of the nonubiquitinated aggregation-prone protein EGFP-STAT5A(E18\( \Delta \)), a C-terminal frameshift mutant of STAT5A.\textsuperscript{18} EGFP-STAT5A(E18\( \Delta \)) (E18\( \Delta \)) was transiently expressed in HeLa cells and phosphorylation of SQSTM1 at S349 and S403 was measured by an immunoblotting analysis. Phosphorylation at these residues was induced by generation of EGFP-STAT5A(E18\( \Delta \)) aggregates, as also occurs for ubiquitinated inclusions (Fig. S2A), and EGFP-STAT5A(E18\( \Delta \)) aggregates colocalized with both S349- and S403-phosphorylated SQSTM1 (Fig. S2B).

We previously showed that SQSTM1 was colocalized with SNCA\( \alpha \)-synuclein aggregates, a common neuropathological hallmark of \( \alpha \)-synucleinopathy, using a cell culture model of \( \alpha \)-synucleinopathies such as Parkinson disease and dementia with Lewy bodies.\textsuperscript{19} Here, we examined SQSTM1 phosphorylation states in this model. When SNCA fibrils were introduced into HEK293 cells that exogenously expressed SNCA, Lewy body-like phosphorylated SNCA aggregates were formed intracellularly (Fig. 2A). The SNCA aggregates colocalized with SQSTM1 and S439-phosphorylated SQSTM1 (Fig. 2A). However, S403-phosphorylated SQSTM1 was not detected at SNCA aggregates (Fig. 2A). Similarly, immunoblotting analysis demonstrated that SQSTM1 phosphorylation at S349 was induced not only by the appearance of SNCA aggregates but also by SNCA overexpression, although S403-phosphorylation was not detected in either case (Fig. 2B). Since SNCA overexpression induces an increase in its oligomer form,\textsuperscript{20} S349-phosphorylation probably resulted from the accumulation of its oligomer.

We further investigated the colocalization of phosphorylated SQSTM1 with SNCA aggregates using an \( \alpha \)-synucleinopathy mouse model produced by the microinjection of SNCA fibrils into the striatum.\textsuperscript{21} At 6 mo after SNCA fibril injection, phosphorylated SNCA-positive inclusions were observed in cell bodies and neurites of the striatum (Fig. 2C). Interestingly, SQSTM1 mainly colocalized with the inclusions only at cell bodies (Fig. 2C). Likewise, S349 (S351 in mice)-phosphorylated SQSTM1 was predominantly detected in cell bodies rather than neurites (Fig. 2C). Phosphorylation of SQSTM1 at S403 (S405 in mice) site did not occur in the in vivo model, similarly to our results using the cell culture model (Fig. 2C). These results indicate that SQSTM1 phosphorylation status changes depending on the nature of the protein aggregates.

**S349 site of SQSTM1 is phosphorylated by CSNK1 (casein kinase 1)**

Although it has been reported that S349 and S403 of SQSTM1 are phosphorylated by MTORC1 and CSNK2-TBK1, respectively,\textsuperscript{9,11,12} it is also possible that other kinases can mediate phosphorylation of these sites. We explored this possibility using various protein kinase inhibitors. This analysis identified 2 novel pathways for SQSTM1 S349-phosphorylation: one directly mediated by CSNK1/CK1 (casein kinase 1) and the other indirectly by CSNK2. CKI-7, a CSNK1 and SGK (serum glucocorticoid regulated kinase) inhibitor and TBCA, a CSNK2 inhibitor, inhibited MG132-induced S349-phosphorylation. Both inhibitors had a greater effect than rapamycin, an MTORC1 inhibitor (Fig. 3A). Moreover, the combination of rapamycin and CKI-7 or rapamycin and TBCA additively inhibited S349-phosphorylation compared to rapamycin alone (Fig. 3A). Phosphorylation was completely blocked by the combination of all 3 inhibitors (Fig. 3B). These results suggested that the SQSTM1 (S349)-phosphorylation pathway mediated by CSNK1 and CSNK2 was distinct from that via MTORC1. Similarly, SQSTM1 (S403)-phosphorylation was also examined, but the compounds did not have an inhibitory effect (Fig. 3A).

Next, we performed in vitro phosphorylation assays of SQSTM1. Wild-type SQSTM1 (WT) and S349A and S403A mutants were transiently expressed in SQSTM1 KO HEK293 cells and the proteins were purified by immunoprecipitation. Immunoprecipitation products were incubated with CSNK1 or CSNK2 in the presence of ATP for 1h at 37\(^{\circ}\)C, followed by an immunoblotting analysis. The S349 residue of WT and S403A
mutant SQSTM1 was phosphorylated by CSNK1 but not in the S349A mutant (Fig. 3C). λ protein phosphatase treatment reduced its phosphorylation (Fig. S1B). For CSNK2, S349-phosphorylated bands were detected at a low level in WT and S403A mutant SQSTM1 (Fig. 3C). We did not detect S403-phosphorylation when SQSTM1 WT was incubated with either CSNK1 or CSNK2 (Fig. 3C). Our results indicate that SQSTM1 S349 was directly phosphorylated by CSNK1, and that CSNK2 was indirectly involved in this phosphorylation.

We investigated the effect of blocking S349-phosphorylation on SQSTM1 localization with protein aggregates. After a combined treatment with rapamycin, CKI-7 and TBCA, ubiquitinated protein inclusions were no longer formed in MG132-treated HeLa cells; additionally, SQSTM1- and
S349-phosphorylated SQSTM1-positive inclusions also disappeared (Fig. 3D). An immunoblotting analysis confirmed that ubiquitinated proteins and SQSTM1 in the Triton X-100 insoluble fraction were markedly reduced in kinase inhibitor treated cells (Fig. S3). As shown in Figure 3A, ubiquitinated proteins were normally increased in these cells, proving that the ubiquitination process was not prevented by treatment with the protein kinase inhibitors. We obtained a similar result for the aggregation-prone protein. The inhibition of S349-phosphorylation by kinase inhibitors

Figure 2. Generation of SNCA aggregates induces the phosphorylation of SQSTM1 on S349 but not on S403. (A) SNCA fibrils (+Fibril) or PBS-mock (Control) were introduced using a transfection reagent into HEK293 cells exogenously expressing human SNCA. After 24 h, the cells were subjected to immunocytochemical analysis using anti-phosphorylated SNCA (p-SNCA), anti-SQSTM1 (SQSTM1), and anti-phosphorylated SQSTM1 (p-SQSTM1 [S349] and p-SQSTM1 [S403]) antibodies. Cell nuclei were counterstained blue with DAPI. Scale bar: 10 μm. Lewy body-like SNCA aggregates are indicated by arrowheads. (B) Lysates of HEK293 (lane 1), HEK293 exogenously expressing human SNCA (lane 2), and HEK293 introduced with SNCA fibrils (lane 3) were analyzed by immunoblotting with anti-SNCA, anti-SQSTM1, anti-phosphorylated SQSTM1, and anti-actin antibodies. Arrowheads indicate molecular species of SNCA including SDS-soluble forms (open arrowhead), SDS-insoluble dimer (closed black arrowhead), and SDS-insoluble aggregates (closed gray arrowhead). Band intensities were measured, and phosphorylated-SQSTM1 values were normalized to total SQSTM1. The data are reported as means ± SD (n = 4). Statistical analyses were performed using one-way ANOVA, followed by the Tukey post-hoc test. *P < 0.05, **P < 0.01. (C) SNCA fibrils were microinjected into the striatum of mice. After 6 mo, the striatum was immunohistochemically examined using anti-phosphorylated SNCA (p-SNCA), anti-SQSTM1 (SQSTM1), and anti-phosphorylated SQSTM1 (p-SQSTM1 [S351] and p-SQSTM1 [S405]) antibodies. The insets show colocalization images of p-SNCA and SQSTM1. Scales are indicated on the images. Middle panels are high-magnification images of upper panels.
treatment reduced SQSTM1-colocalization to EGFP-STAT5A(E18Δ) aggregates (Fig. S2C) even though this protein aggregates in a SQSTM1-independent manner.18 Next, we examined whether S403 phosphorylation was involved in the formation of inclusions by ubiquitinated proteins. HeLa cells were treated with MG132 alone or with a
combination of MG132 and the TBK1 inhibitor BX795. In cells treated with MG132 alone, S403-phosphorylated SQSTM1 was detected by immunoblotting analysis (Fig. 4A). The combined MG132/BX795 treatment resulted in a 90% reduction in S403 phosphorylation but had no effect on S349 phosphorylation (Fig. 4A). Ubiquitin- and SQSTM1-positive inclusions were detected after MG132 treatment; this was also seen in cells treated with BX975 to inhibit S403 phosphorylation (Fig. 4B). We confirmed that these inclusions colocalized with S349-phosphorylated SQSTM1 (Fig. 4B). Next, we investigated the inclusion formation of ubiquitinated proteins in SQSTM1-mutant-expressing cells. SQSTM1 WT or -mutant (S349A or S403A) was stably expressed in SQSTM1 KO HeLa cells (Fig. S4A). Triton X-100-soluble and -insoluble fractions from control and MG132-treated cells were assessed by immunoblotting with anti- SQSTM1 and anti-poly-ubiquitin antibodies. Triton-insoluble poly-ubiquitinated proteins were significantly decreased in SQSTM1 KO (Vector) and S349A mutant-expressing cells, compared with SQSTM1 WT or -S403 mutant-expressing cells (Fig. S4B and C). Together, our results indicate

Figure 4. Colocalization of SQSTM1 with ubiquitinated inclusions is not affected by inhibition of SQSTM1 (S403)-phosphorylation. (A) HeLa cells were treated with a BX795 TBK1 inhibitor (1 μM) and MG132 (10 μM) for 12 h. Cell lysates were analyzed by immunoblot analysis. Band intensities were measured, and phosphorylated-SQSTM1 values were normalized to total SQSTM1. The data are reported as means ± SD (n = 4). Statistical analyses were performed using one-way ANOVA, followed by the Tukey post-hoc test. *P < 0.01. (B) Colocalization of SQSTM1 with ubiquitinated inclusions were immunocytochemically assessed in cells treated with MG132 alone (left) or MG132 and BX795 (right). Cell nuclei counterstained blue with DAPI. Scale bar: 10 μm.
that S349 phosphorylation is required for inclusion formation by ubiquitinated proteins.

**Regulation of SQSTM1 phosphorylation by the HSF1 stress response pathway**

When misfolded and unfolded proteins accumulate intracellularly following heat or oxidative stresses, HSF1 is activated and upregulates the expression of proteostasis-related proteins such as HSPs. In a similar manner, the accumulation of ubiquitinated proteins after inhibition of proteasomes enhanced HSF1 activation. Here, HSF1 activity was measured by an affinity isolation assay using a biotinylated oligonucleotide that included the HSE sequence that is a target sequence for activated HSF1. HSF1 activity was induced within 1 h after MG132 treatment and achieved a maximum level after 9 h (Fig. S5). We hypothesized that SQSTM1 phosphorylation was modulated through the HSF1 stress response pathway. To test this hypothesis, HSF1 was inhibited by KRIBB11, an HSF1-specific inhibitor. Immunoblotting analysis showed that HSF1 inhibition strongly suppressed both S349- and S403-phosphorylation although ubiquitinated proteins were increased by the MG132 treatment (Fig. 5A). This inhibitor did not influence SQSTM1 or MTOR expression (Fig. 5A). Immunocytochemical analysis showed that inclusion formation by ubiquitinated proteins did not occur in HeLa cells treated with MG132 and KRIBB11 (Fig. 5B), similarly to the effect of kinase inhibitory treatment. Immunoblotting analysis also demonstrated that KRIBB11-treatment markedly reduced Triton X-100-insoluble ubiquitinated proteins and SQSTM1 (Fig. S3). The same result was obtained for the aggregation-prone protein EGFP-STAT5A (E18Δ). Although S349- and S403-phosphorylated SQSTM1 were significantly increased in HeLa cells expressing EGFP-STAT5A(E18Δ), HSF1 inhibition by KRIBB11 reduced their levels (Fig. 5C). Immunocytochemical analysis also showed that SQSTM1 was not localized to EGFP-STAT5A(E18Δ) aggregates in KRIBB11-treated cells (Fig. 5D).

Next, we used HSF1 knockout (HSF1 KO) cells to investigate whether the HSF1 stress response pathway modulates the SQSTM1 phosphorylation pathway. We designed a single guide RNA vector that targeted exon 2 of the HSF1 gene and generated HSF1 KO HeLa cells using the CRISPR/Cas9 system. HSF1 KO HeLa cells were cloned, and gene knockout was validated by an immunoblotting analysis (Fig. 6A). In comparison with WT cells, expression of SQSTM1 was unaltered in HSF1 KO HeLa cells although HSPA1 expression was slightly decreased (Fig. 6A). When HSF1 KO cells were treated with MG132, phosphorylation of S349 and S403 was reduced to 40% and 3%, respectively, compared with WT cells (Fig. 6B). However, exogenous expression of HSF1 in the HSF1 KO cells restored the levels of phosphorylation (Fig. 6B). Thus, our observations were not due to off-target effects of the CRISPR/Cas9 system.

Next, we screened HSF1 KO cells immunocytochemically for inclusion formation by ubiquitinated proteins and for SQSTM1 localization and found no evidence for ubiquitin-positive inclusions (Fig. 6C); this finding is consistent with the outcome of the inhibitor experiment. Curiously, SQSTM1 was localized to the perinuclear region when cells were treated with MG132 (Fig. 6C and Fig. S6). Perinuclear SQSTM1 was not detected by either of the anti-phospho SQSTM1 antibodies (Fig. 6C), indicating that perinuclear SQSTM1 was unphosphorylated. We then determined whether SQSTM1-perinuclear localization in MG132-treated HSF1 KO cells could be prevented by KRIBB11 treatment (Fig. S6). We found that KRIBB11-treatment suppressed perinuclear localization of SQSTM1 in MG132-treated HSF1 KO cells (Fig. S6). KRIBB11 is known to inhibit HSF1-dependent recruitment of P-TEFb (positive transcription elongation factor b complex) to the HSPA1 promoter, but our results here indicate it must have other properties. It has been reported that SQSTM1 shuttles continuously between nuclear and cytosolic compartments at a high rate, therefore, SQSTM1 nuclear transport might be impaired in HSF1-deficient cells resulting in the perinuclear accumulation of SQSTM1.

**HSF1 stress response pathway regulates autophagic clearance of protein aggregates**

It was previously reported that SQSTM1 phosphorylation regulates selective autophagic clearance of ubiquitinated proteins, bacteria, and KEAP1. We postulated that HSF1 might regulate this process via SQSTM1 phosphorylation. To test this proposal, we performed a metabolic analysis of EGFP-STAT5A (E18Δ) aggregates using the tetracycline (Tet)-On system. It was previously shown that EGFP-STAT5A(E18Δ) undergoes SQSTM1-independent aggresome formation and SQSTM1-dependent autophagic clearance. Here, we generated EGFP-STAT5A(E18Δ) aggregates in HeLa TREX-EGFP-STAT5A (E18Δ) cells and placed the cells in tetracycline-free medium for 0, 12, or 24 h. We found that EGFP-STAT5A(E18Δ) aggregates were mostly eliminated within 24 h in control cells (Fig. 7A); treatment with kinase inhibitors (rapamycin, CKI-7, and TBCA) or KRIBB11 delayed this elimination (Fig. 7A). EGFP intensity was quantified by high-content image analysis using an IN Cell Analyzer 2200 to accurately measure the level of the remaining EGFP-STAT5A(E18Δ) aggregates. Higher EGFP-intensities were present in kinase inhibitor- or KRIBB11-treated cells after 24 h (Fig. 7B). The average EGFP intensities in control, kinase inhibitor, and KRIBB11-treated cells after 24 h were 437 ± 424 (arbitrary units), 822 ± 1027, and 1814 ± 1138, respectively (Fig. 7A).

Our data raise the question of whether delayed elimination of protein aggregates is due to inhibition of proteasome activity or autophagy. We therefore examined the effects of impairment of HSF1-mediated stress responses on the proteasome and autophagy. Kinase inhibition and HSF1-deficiency did not affect proteasome activity (Fig. S7B and C). Meanwhile, MG132-induced autophagosome formation was not detected in KRIBB11-treated cells or HSF1 KO cells using immunostaining with an anti-LC3 antibody (Fig. 7D). Moreover, LC3 conversion from LC3-I to LC3-II was monitored to assess autophagy flux. However, we could not measure it because combined treatment with MG132-bafilomycin A1 or epoxomicin-bafilomycin A1 to HeLa cells had an undesirable side effect (data not shown). We then performed an autophagy flux assay using an mCherry-EGFP-LC3 reporter system. In HeLa-WT cells, MG132 treatment increased mCherry-positive and
EGFP-negative LC3-puncta, whereas autophagy induction was inhibited by KRIBB11 treatment (Fig. S8). Similarly, MG132-induced autophagy flux was not detected in HSF1 KO cells (Fig. S8), indicating that the processes of autophagy are impaired in these cells. However, there was a possibility that this inhibition was due to reduced activity in the basic autophagic machinery since it has previously been reported that HSF1 regulates autophagy by directly binding to the ATG7 promoter to upregulate its expression.\textsuperscript{17,27} We measured ATG7 and ATG5 mRNA expression in these cells and found that there was no significant change in the expression of either gene (Fig. 7C). This finding was supported by the outcome of inducing autophagy in starving cells of amino acids: LC3 puncta appeared even in KRIBB11-treated or HSF1 KO cells (Fig. 7D and Fig. S8), indicating that the components of the basic autophagic machinery were not affected by an impairment of HSF1. Therefore, we concluded that HSF1 was specifically required for MG132-induced autophagy. However, triple kinase

![Figure 5](image-url)

**Figure 5.** Phosphorylation of S349 and S403 on SQSTM1 is suppressed by a KRIBB11 HSF1 inhibitor. (A) HeLa cells were treated with 10 μM MG132 and 50 μM KRIBB11 for 12 h. Cell lysates were subjected to immunoblot analysis. Band intensities were measured, and phosphorylated SQSTM1 values were normalized to total SQSTM1. The data are reported as means ± SD (n = 4). Statistical analyses were performed using one-way ANOVA, followed by the Tukey post-hoc test. *P < 0.01. (B) Colocalization of SQSTM1 with ubiquitinated inclusions were immunocytochemically assessed in cells treated with MG132 alone (left) or MG132 and KRIBB11 (right). Cell nuclei were counterstained blue with DAPI. Scale bar: 10 μm. (C) HeLa cells were transiently transfected with mock control or pEGFP-STAT5A(E18Δ) (STAT5A[E18Δ]), and then treated with or without 50 μM KRIBB11. After 12 h, cell lysates were examined by immunoblot analysis. Band intensities of S349- and S403-phosphorylated SQSTM1 were measured, and the data are reported as means ± SD (n = 4). Statistical analyses were performed using one-way ANOVA, followed by the Tukey post-hoc test. *P < 0.01, **P < 0.05. (D) Colocalization of SQSTM1 with EGFP-STAT5A(E18Δ) were immunocytochemically assessed in cells treated with DMSO control (left) or KRIBB11 (right). Cell nuclei were counterstained blue with DAPI. Scale bar: 10 μm.
inhibitor treatment constitutively induced autophagosome formation after MTORC1 inhibition by rapamycin (Fig. 7D). Triple kinase inhibitor treatment probably accelerated bulk-degradation by autophagy, and thereby mitigated against the accumulation of EGFP-STAT5A(E18Δ) aggregates compared with KRRB11-treated cells (Fig. 7B and S7). Overall, HSF1 is a key player in triggering aggrephagy and the impairment of HSF1 causes a reduction in the clearance of protein aggregates.

Figure 6. Suppression of SQSTM1 S349-phosphorylation in HSF1 KO cells. (A) HSF1 was targeted using the CRISPR/Cas9 system. Cell lysates of HSF1-WT and individual HSF1-targeted clones (#6 and #12) were subjected to immunoblot analysis with anti-HSF1, anti-SQSTM1, anti-HSPA1, and anti-actin. (B) HeLa WT, HeLa HSF1 KO, and HSF1 KO exogenously expressing HSF1 (HeLa HSF1 KO + pEF1-HSF1) were treated with 10 μM MG132 for 12 h. Cell lysates were examined by immunoblot analysis. Band intensities were measured, and phosphorylated-SQSTM1 values were normalized to total SQSTM1. The data are reported as means ± SD (n = 4). Statistical analyses were performed using one-way ANOVA, followed by the Tukey post-hoc test. *P < 0.01, **P < 0.05. (C) Colocalization of SQSTM1 with ubiquitinated inclusions were immunocytochemically assessed in cells treated with MG132. Cell nuclei were counterstained blue with DAPI. Scale bar: 10 μm.
Discussion

The phosphorylation of SQSTM1 at S349 and S403 is required for stable binding to substrates such as KEAP1 and ubiquitinated proteins. Proteasome inhibition was found to be induced by phosphorylation of S349 or S403 in agreement with previous reports. In the present study, we found that CSNK1 could directly phosphorylate SQSTM1 at S349; this novel observation adds to the previous reports that MTORC1, CSNK2, and TBK1 can phosphorylate SQSTM1. There are no canonical consensus sequences for CSNK1 substrates, such as phospho-S/T-X-S/T or D/E-X-X-S/T, around the S349 phosphorylation site. However, CSNK1 phosphorylates noncanonical sequences (S-L-S and K/R-X-K/R-X-S/T); thus, S349 is a noncanonical targeting site for CSNK1. In addition to CSNK1, the specific CSNK2
inhibitor TBCA also suppressed phosphorylation; this effect was achieved indirectly according to the results of the in vitro phosphorylation assay. Importantly, phosphorylation of S349 of SQSTM1 was completely blocked by triple kinase inhibitor treatment (rapamycin, CKI-7, and TBCA), suggesting that these phosphorylation pathways are independent. Phosphorylation of S403 of SQSTM1 was blocked by treatment with the TBK1 inhibitor but not TBCA in HeLa cells, in agreement with a previous report by Pilli et al.9 However, it has previously been reported that CSNK2 phosphorylates S403 of SQSTM1 in MG132-treated Neuro-2A and MEF cells.11 Although we cannot deny the possibility that the S403-phosphorylation process is different among cell lines, it is essential to resolve this discrepancy. In the present study, we examined the phosphorylation status of S349 and S403 simultaneously and found that inhibition of phosphorylation of one site did not interfere with phosphorylation at the other.

We demonstrated here that HSF1 regulates SQSTM1 phosphorylation at both sites upstream of kinase activities. It remains unclear how HSF1 activates MTORC1, CSNK1, CSNK2, and TBK1. A genome-wide analysis shows that genes encoding the kinases are not directly driven by HSF1.22 Also, we found that MTOR expression was not influenced by HSF1 deficiency. Thus, a further downstream factor(s) must be involved in the activation of the kinases. The HSPA1-cochaperone BAG3 gathers misfolded proteins into SQSTM1-containing aggregates, and its expression is mediated by HSF1.31,32 This evidence leads us to speculate that the HSPA1-BAG3 complex may be involved in recruitment of kinases downstream of HSF1. Alternatively, the molecular chaperone HSP90AA may participate in this pathway. Previously, it has been reported that HSP90AA protects CSNK2 from aggregation and dramatically activates CSNK2.33 In addition, TBK1 interacts with HSP90AA, resulting in its protection from proteasome-mediated degradation.34 We speculate that the induction of HSP90AA expression causes an increase in CSNK2 activities and TBK1 levels and subsequent activation of SQSTM1 phosphorylation. Furthermore, pharmacologic inhibition of HSP90AA leads to a decrease in the MTOR-RPTOR/RAPTOR interaction,35 suggesting that HSP90AA might also regulate MTORC1 activity under stress conditions.

Our study highlighted a novel role for the HSF1 stress response pathway in SQSTM1 function; previously, this pathway was shown to be involved in transcriptional control of proteostasis components such as molecular chaperones.36 SQSTM1 acts as a signaling hub of various transduction pathways and is involved in autophagic degradation and inclusion formation of harmful proteins.7,14,37 Loss of SQSTM1 accelerates cell death that is induced by the overexpression of polyglutamine-expanded huntingtin,38,39 suggesting a key role for SQSTM1 in proteostasis. In the present study, inhibition of SQSTM1 phosphorylation at S349 by kinase inhibitor treatment and loss of HSF1 function suppressed inclusion formation by ubiquitinated proteins and elimination of the aggregation-prone protein; these results indicate the necessity for S349 phosphorylation in SQSTM1 functions and a novel network of HSF1 stress response pathways in proteostasis. We propose multiple functions for HSF1 in proteostasis (Fig. 8). Proteotoxic stresses lead to the activation of HSF1 and the subsequent upregulation of HSP expression, resulting in protein

**Figure 8.** Schematic model of the proteostasis network via the HSF1-SQSTM1 axis. HSF1 is activated by the accumulation of harmful proteins, resulting in the induction of cytoprotective genes including HSPs. Moreover, phosphorylation of SQSTM1 by MTORC1, CSNK1, and TBK1 is also induced via activation of HSF1. Phosphorylated SQSTM1 accelerates inclusion formation and autophagic clearance of harmful proteins.
refolding and protection from proteotoxicity. Additionally, HSF1 accelerates inclusion formation and elimination of harmful proteins via SQSTM1 activation that is mediated by several kinases such as MTORC1 and CSNK1. Protein inclusion formation, including the aggresome, is generally believed to be a cytoprotective process by which harmful proteins are compacted and directed toward autophagic clearance. Therefore, the HSF1 stress response pathway is required for proteostasis systems. Although phosphorylation of SQSTM1 at S349 increases binding affinity for KEAP1, it remains unclear whether this phosphorylation is also required for the direct or indirect interaction with ubiquitinated proteins. It was recently reported that N-terminally arginylated HSPA5/BiP (R-HSPA5), an endoplasmic reticulum chaperone, is a delivery determinant in SQSTM1-mediated autophagic clearance of harmful proteins. R-HSPA5 is associated with misfolded proteins in the cytoplasm upon various stresses and binds to the ZZ domain of SQSTM1. Subsequently, this binding induces SQSTM1-oligomerization and increases the interaction between SQSTM1 and LC3. This observation leads to speculate that phosphorylation of S349 of SQSTM1 may be involved in the interaction of SQSTM1 with the R-HSPA5 cargo complex. In the future, further structural and functional analyses will be necessary to elucidate the roles of this phosphorylation.

HSF1 loss does not directly affect the activity of proteasomes and the expression of the autophagy machinery. However, its loss does cause the impairment of the induction of autophagosome formation in proteasome-inhibited cells while starvation-induced autophagy is normal. These observations agree with the results obtained in SQSTM1 deficient cells, indicating that SQSTM1 phosphorylation is specifically required for aggrephagy induction. Itakura et al. report that SQSTM1 oligomerization is essential for its localization to the autophagosome formation site where SQSTM1 associates with upstream autophagy factors such as ULK1 and VMP1. Moreover, the interaction of SQSTM1 with LC3 is promoted by SQSTM1 oligomerization. Thus, the initiation of aggrephagy might be triggered by SQSTM1 oligomerization and localization to the autophagosome formation site after phosphorylation and substrate capturing of SQSTM1. Other receptor proteins such as OPTN (optineurin) and CALCOCO2/NDP52 have recently been shown to have an important role in the recruitment of the upstream autophagy machinery including ULK1, ZFYVE1/DFCP1, and WIPI1 for selective mitochondrial autophagy. It has been reported that ULK1 phosphorylates SQSTM1 at S407. Based on these data, we suggest that SQSTM1 oligomerization might be required for the assembly of upstream components for autophagosome nucleation upon aggrephagy induction.

Our analyses showed that the generation of various protein aggregates caused SQSTM1 phosphorylation. Accumulation of ubiquitinated proteins and the aggregation-prone protein induced phosphorylation of SQSTM1 at both S349 and S403, whereas SNCA aggregates and oligomers enhanced its phosphorylation only at S349. This result is in agreement with a pathological analysis of α-synucleinopathies including dementia with Lewy bodies and multiple system atrophy, while they are not detected by an anti-phospho-SQSTM1 (S403) antibody. Although it is unclear whether this result can be attributed to phosphorylation or dephosphorylation, the phosphorylation status of S403 may be susceptible to the properties of protein aggregates. This puzzling aspect of SQSTM1 phosphorylation might be implicated in the pathogenic mechanism of α-synucleinopathies, and thus further studies will be necessary to determine whether this is the case.

Materials and methods

Animals

C57BL/6J mice were maintained on a 12-h light/dark cycle at 25°C with ad libitum food and water. All experimental designs and procedures were approved by the Committee for Animal Research, Kyoto Prefectural University of Medicine (M23-241), Kyoto Japan, and followed the guidelines of the Ministry of Education, Culture, Sports, Science, and Technology of Japan.

Reagents

The following reagents were purchased: MG132 (Peptide Institute, 3175-v), rapamycin (Santa Cruz Biotechnology, sc-3504), and KRB11 (Calbiochem, 385570), casein kinase inhibitor CKI-7 (Seikagaku Kogyo; out of production), TBCA (Cayman Chemical, 14831), and BX795 (ChemScene, CS-0259).

Cells

HeLa and HEK293 cells were grown in DMEM medium (Nacalai Tesque, 08458-16) supplemented with 10% fetal bovine serum. Transfection of plasmids DNA was performed using Lipofectamine 2000 (Invitrogen, 11668027) or a CUY21Pro-Vitro electroporator (NEPA GENE, Ichikawa, Japan). Cells were cultured in Hank’s balanced salt solution (Nissui, 05905) for observation of amino acid starvation-induced autophagy.

Antibodies

The complete list of antibodies and sources used is given in Table 1.

Immunoblotting analysis

Cells were washed with PBS (137 mM NaCl, 8.1 mM Na2HPO4, 2.68 mM KCl, 1.47 mM KH2PO4) and extracted with ice-cold PBS containing 0.5% Triton X-100 (Wako Pure Chemical Industries, 168-11805) and a protease inhibitor cocktail (Nacalai Tesque, 25955-11). Protein concentrations in samples were measured using the Lowry protein assay, followed by addition of Laemmli SDS sample buffer. For immunoblotting analysis, samples were separated on 10% or 15% SDS-polyacrylamide gels using a Tris–glycine buffer system (25 mM Tris, 192 mM glycine, 0.1% SDS). After electrophoresis, the separated proteins were transferred to Immobilon-P membranes (Merck Millipore, IPVH00010). The membranes were blocked with 10% skimmed milk, 5% bovine serum albumin (Nacalai

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confocal laser microscope. Confocal laser scanned images were obtained with an LSM 510 META microscope (Carl Zeiss, Oberkochen, Germany). Alexa-Fluor-488 and EGFP were excited using a 488 nm argon laser, and emission was recorded through a 500- to 530-nm band-pass filter. Alexa Fluor 594 was excited with a 543-nm helium-neon laser, and emission was recorded through an LP 560 nm filter. DAPI was excited at 800 nm, with a Mai Tai 2-photon laser (Spectra-Physics, Santa Clara, CA, USA), and fluorescence was recorded through a 390- to 465-nm band-pass filter. Images were obtained through a Plan-Apochromat 1006/1.4 Oil DIC lens (Carl Zeiss). We adjusted the detector gain and laser power settings to avoid oversaturation of SQSTM1-aggregate images. We used lower than usual settings for laser power/gain because the immunofluorescence signal from SQSTM1 aggregates was very bright.

**Immunohistochemistry**

Under deep anesthesia, mice were intracardially perfused with 0.1 M PBS (0.1 M phosphate buffer, pH 7.4, 0.9% NaCl) followed by 4% paraformaldehyde in 0.1 M PB (0.1 M phosphate buffer, pH 7.4). Brains were dissected into blocks including the cerebral cortex and hippocampus, and post-fixed with the same fixative for 12 h at 4°C. Coronal sections (30 μm) were obtained using a vibratome (DSK, Kyoto, Japan). Sections were permeabilized with 0.1% Triton X-100 in PBS for 30 min, and blocked with 5% normal goat serum in PBS for 1 h. Sections were then treated with primary and secondary antibodies following the immunocytochemical procedures described above. After treatment, the sections were washed with PBS and then with 20 mM Tris-HCl buffer, and mounted in FluorSave. Images were obtained as Z stacks (10–20 z-sections, 1 μm apart, 1024 × 1024 pixels) using an LSM 510 META microscope.

### 1. protein phosphatase treatment

HeLa cells were treated with 10 μM MG132 for 12 h and harvested in PBS using a cell scraper. Cells were homogenized by sonication, and the insoluble fraction was recovered (20,000 × g, 15 min, 4°C) and resuspended in 1 × PMP buffer including 1 mM MnCl2 (New England Biolabs, P0753S). Samples were incubated with 400 units of 1. protein phosphatase (New England Biolabs, P0753S) at 37°C for 1 h. SQSTM1 immunoprecipitates were treated with CSNK1 and 1. protein phosphatase at 37°C for 1 h in CSNK1 reaction buffer including 1 mM ATP and 1 mM MnCl2.

### Cell culture and mouse models of α-synucleinopathy

SNCA fibrils were prepared as described previously.19,47,48 pcDNA-SNCA plasmid20 was transiently transfected into HEK293 cells plated onto 24-well plates, and 4 μg SNCA fibrils were introduced on the following day using Lipofectamine 2000 reagent. Cells were incubated in DMEM medium supplemented with 10% fetal bovine serum for 24 h, and used for immunoblotting analysis and immunocytochemistry.

The α-synucleinopathy mouse model was generated as described previously.21 SNCA fibrils (5 μg/μl) were sonicated

### Drug treatment and immunocytochemistry

HeLa cells were treated with 10 μM MG132 for 12 h for proteasome inhibition. The kinase inhibitors rapamycin (1 μM), CKI-7 (100 μM) and TBCA (25 μM) were added at the same time as MG132 for 12 h. For inhibition of HSF1, HeLa cells were treated with 10 μM KRB11 just after pEGFP-STAT5A (E18Δ) transfection or MG132 treatment.46

Immunocytochemical analysis was performed as described previously.18 Cells were fixed with 4% paraformaldehyde (Merck Millipore, 104005) after drug treatment. The fixed cells were permeabilized in PBS containing 0.1% Triton X-100 or 0.01% digitonin (Nacalai Tesque, 123-33) and placed in blocking solution for 30 min. Samples were incubated with the primary antibodies for 12 h at 25°C. After a PBS wash, samples were further incubated with Alexa Fluor 488-labeled goat anti-mouse IgG (Invitrogen, A11029) or Alexa Fluor 594-labeled goat anti-rabbit IgG (Invitrogen, A11037) for 4 h at 25°C. After washing, they were stained with 4’,6-diamidino-2-phenylindole (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523). The cells on cover glasses were mounted on glass slides in an aqueous mounting medium (FluorSave (Dojindo, D523).

### Table 1. Antibodies.

| Antibodies                        | Source (CAT No.)       | Applications            |
|----------------------------------|------------------------|-------------------------|
| Anti-SQSTM1 pAb (rabbit)         | Sigma-Aldrich (P0067)  | IB, IC, IH, IP           |
| Anti-phospho-SQSTM1 (Ser351) pAb (rabbit) | MBL (PM074)           | IB, IC, IH               |
| Anti-phospho-SQSTM1 (Ser351) mAb (mouse) | MBL (M217-3)       | IC                      |
| Anti-phospho-SQSTM1 (S403) pAb (rabbit) | GeneTex (GTX128171) | IB, IC, IH               |
| Anti-poly-ubiquitin mAb (FK2) (mouse) | Nippon Biotest Labs (MFK-004) | IB, IC                  |
| Anti-phospho-SNCA (S129) mAb (mouse) | Wako Pure Chemical Industries (015-25191) | IC, IH                  |
| Anti-phospho-SNCA (S129) pAb (rabbit) | Abcam (ab59264)      | IC, IH                   |
| Anti-SNCA mAb (mouse)            | BD Biosciences (610787) | IB                      |
| Anti-LC3 pAb (rabbit)            | MBL (PM036)           | IH                      |
| Anti-HSF1 mAb (rabbit)           | Cell signaling Technology (12972) | IB                     |
| Anti-MTOR mAb (rabbit)           | Cell signaling Technology (2983) | IB                     |
| Anti-HSPA1 pAb (rabbit)          | StressMarq Biosciences (SPC-103D) | IB                     |
| Anti-GFP pAb (rabbit)            | MBL (598)             | IB                      |
| Anti-Actin mAb (mouse)           | Thermo Fisher (MS-1295-P0) | IB                     |

Note. Immunoblot (IB), Immunocytochemistry (IC), Immunohistochemistry (IH), Immunoprecipitation (IP).
and microinjected (1 μl/site) into the striatum using a 30-gauge Hamilton syringe (Hamilton, Reno, NV, USA) at a rate of 0.25 μl/min. Likewise, PBS was microinjected into the striatum of control mice. The coordinates for the striatum were: AP, +0.2 mm from the bregma; ML, ± 2.0 mm from the midline; and DV, −3 mm below the skull surface. Mice were used for immunohistochemistry 6 mo after injection.

**In vitro phosphorylation assay**

For the SQSTM1 phosphorylation analysis after CSNK1 and CSNK2 treatment, SQSTM1 WT, SQSTM1S349A, or SQSTM1S403A was expressed in HEK293 SQSTM1 KO cells. The proteins were purified by immunoprecipitation with an anti-SQSTM1 antibody and protein G Mag Sepharose (GE Healthcare, 28951379). A reaction buffer containing 1 mM ATP was added to immunoprecipitation products, and then the mixtures were incubated with or without 1000 units of CSNK1 (New England Biolabs, P6030S) or 500 units of CSNK2 (New England Biolabs, P6010S) for 1 h at 37°C. The kinase reaction was stopped by adding 3 × Laemmli SDS sample buffer. Phosphorylation at S349 or S403 was assessed by immuno blotting analysis using anti-phospho-SQSTM1 (Ser351) and anti-phospho-SQSTM1 (Ser403) antibodies.

**Plasmid construction**

For construction of the HSF1 expression vector (pEF1-HSF1), human HSF1 cDNA was cloned into pEF1-mycHisB (Thermo Fisher, A21170) according to the manufacturer's instructions. A reaction buffer containing 1 mM MgCl₂, 50 mM HEPES, pH 7.4, 100 mM NaCl, 5 mM EDTA, 0.5% Triton X-100, and protease inhibitor cocktail. Cell extracts were incubated with 1 μg of biotinylated HSE double strand oligonucleotide (5’-biotin-AACGATACCTCGAGAATGGCTGCT-3’ and 5’-AGCCATTCCTCGAGAATGGCTGCT-3’) for 30 min at room temperature. The oligonucleotides were precipitated with Streptavidin Mammal MagnaSphere Paramagnetic Particles (Promega, Z5481) for 1 h at 4°C. HSF1 pulled down with HSE-oligonucleotides was quantified by an immunoblotting analysis using an anti-HSF1 antibody.

**Gene targeting using CRISPR/Cas9 system**

We designed target sequences for SQSTM1 and HSF1, and cloned double-strand oligonucleotides for SQSTM1 (5’-AAAGATTGGACCGCTCAGCTATTGCCTCTACTGGGTTT-3’/5’-GTAGACGGGTCTCATTGTGCT-3’) and HSF1 (5’-CGCCACCATGGAGCATCTGCAGGCACCATGGGACCGCTCAGCTATTGCCTCTACTGGGTTT-3’/5’-GTAGACGGGTCTCATTGTGCT-3’) into a GeneArt CRISPR Nuclease vector (Thermo Fisher, A21170) according to the manufacturer's instructions. Inserts were confirmed by DNA sequencing analysis. The constructs were transfected into HEla and HEK293 cells using an electroporator. Targeted clones were obtained by limiting dilution and confirmed by immunoblotting analysis. Target genes of individual clones were amplified by PCR, followed by cloning and sequencing. The SQSTM1 KO_HEK293 clone has 16-base pair (bp) (1039 to 1054 bp) and 5-bp (1046 to 1050 bp) deletions within exon 7 of the SQSTM1 gene. The SQSTM1 KO_HeLa clone has a 1-bp insertion (T) at the 36-bp (1014 to 1049 bp) deletion site within exon 7 of the SQSTM1 gene. The HSF1 KO_HeLa clone has a 10-bp insertion (CCCTGGGAAATA) at the 14-bp (131 to 144 bp) deletion site and a 13-bp insertion (TGGAAATTTTGCA) at the 9-bp (143 to 151 bp) deletion site within exon 2 of the HSF1 gene. Although the copy numbers at the SQSTM1 and HSF1 gene loci in our HeLa cells are unknown, the intact DNA sequence of these genes was never detected.

**HSF1 affinity isolation assay**

The HSF1 affinity isolation assay was performed as described previously.²² HeLa cells were treated with 10 μM MG132 for 1 to 12 h at 37°C or heat shocked for 1 h at 42°C. Cells were lysed in ice-cold lysis buffer (25 mM HEPES, pH 7.4, 100 mM NaCl, 5 mM EDTA, 0.5% Triton X-100, and protease inhibitor cocktail). Cell extracts were incubated with 1 μg of biotinylated HSE double strand oligonucleotide (5’-biotin-AACGATACCTCGAGAATGGCTGCT-3’ and 5’-AGCCATTCCTCGAGAATGGCTGCT-3’) for 30 min at room temperature. The oligonucleotides were precipitated with Streptavidin Mammal MagnaSphere Paramagnetic Particles (Promega, Z5481) for 1 h at 4°C. HSF1 pulled down with HSE-oligonucleotides was quantified by an immunoblotting analysis using an anti-HSF1 antibody.

**High-content analysis using IN Cell Analyzer**

We used the Tet-On system in EGFP-STAT5A(E18Δ) cells for analysis of metabolism. The Tet-On construct, pcDNA5-EGFP-STAT5A(E18Δ), was made by inserting a pEGFP-STAT5A(E18Δ) fragment (BglII-SspI) into the BamHI-EcoRV site of pcDNA5 FRT-TO (Invitrogen, V65200). HeLa cells were transfected with pcDNA5-EGFP-STAT5A(E18Δ) and pcDNA6/TR (Invitrogen, V102520), and selected with 40 μg/ml G418 (Calbiochem, 400052) for 2 wk. Single clones (HeLa TReX-EGFP-STAT5A(E18Δ)) were obtained by limiting dilution and confirmed by immunoblotting analysis.

HeLa TReX-EGFP-STAT5A(E18Δ) cells (4 × 10⁴) were plated into 96-well plates and incubated in a CO₂ incubator for 18 h (10 wells/group). For formation of EGFP-STAT5A(E18Δ) aggregates, cells were treated with tetracycline and 2.5 μM MG132 for 12 h. Concurrently, cells were treated with 5 μM KRB11 or kinase inhibitors (1 μM rapamycin, 100 μM CK1-7, and 25 μM TBCA). Cells were washed 3 times with fresh medium, and then continuously incubated in fresh medium containing 5 μM KRB11 or kinase inhibitors (1 μM rapamycin, 100 μM CK1-7, and 25 μM TBCA). After 0, 12, or 24 h, cells were fixed with 4% paraformaldehyde for 15 min and then stained with DAPI. Plates were set in the IN Cell Analyzer 2200 (GE Healthcare, Chicago, IL, USA), and cell images were acquired using a ×10 objective lens (3 fields/well). Excitation and emission wavelengths for DAPI were 390 nm/435 nm, and for EGFP were 475 nm/525 nm. The EGFP intensity in each cell was determined by IN Cell Investigator 1.6 software. For cell segmentation, we chose the collar method and defined a
3 μm collar radius as the cell area. The EGFP intensity per cell was measured and plotted in a graph (n ≥ 5000).

**Autophagy flux assay**

pBABE-puro mCherry-EGFP-LC3B was a gift from Dr. Jayanta Debnath (Addgene, 22418). A DNA region encoding mCherry-EGFP-LC3B was cloned into the pEF1-mycHisB vector (pEF1-mCherry-EGFP-LC3B). HeLa WT or HSF1 KO cells were stably transfected with pEF1-mCherry-EGFP-LC3B, followed by G418 selection. Clones were isolated and used for the autophagy flux assay. Cell images were chosen randomly, and the number of mCherry-only LC3 puncta and mCherry-positive EGFP-positive puncta in individual cells (30 cells each) was measured manually.

**Statistical analyses**

Statistical analyses were conducted using Kaleida Graph 4.1 and GraphPad Prism 5J. Differences between treatments were analyzed for significance using the Student t test or one-way ANOVA with the Tukey post hoc test. Significance was set as p ≤ 0.05. Values in graphs are presented as means ± SD. All experiments were repeated 4 times.

**Abbreviations**

| Abbreviation | Description |
|--------------|-------------|
| CSNK2/CK2 | casein kinase 2 |
| CSNK1/CK1 | casein kinase 1 |
| GPD/G3PDH | glycerol-3-phosphate dehydrogenase |
| HSF1 | heat shock transcription factor 1 |
| HSE | heat shock element |
| HSPs | heat shock proteins |
| KEAP1 | Kelch-like ECH associated protein 1 |
| KIR | KEAP1-interaction region |
| LIR | LC3-interacting region |
| MTERC1 | mechanistic target of rapamycin (serine/threonine kinase) complex 1 |
| PB1 | (Phox and Bem1) domain |
| proteostasis | protein homeostasis |
| SGK | serum- and glucocorticoid-inducible kinase |
| TANK | tank-binding kinase 1 |
| UBA | ubiquitin-associating |
| UPS | ubiquitin-proteasome system |
| ZZ | zinc finger |

**Disclosure of potential conflicts of interest**

No potential conflicts of interest were disclosed.

**Acknowledgments**

We thank Dr. Jayanta Debnath (University of California at San Francisco) for the pBABE-puro mCherry-EGFP-LC3B plasmid.

**Funding**

This work was supported by Grants-in-Aid for Scientific Research from the Japan Society for the Promotion of Science (YW: 15K09320 and 24591272, and MT: 25290014).

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