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Persulfidation of ATG18a regulates autophagy under ER stress

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Abstract

Hydrogen sulfide (H₂S) is an endogenously generated gaseous signalling molecule, which recently has been implicated in autophagy regulation in both plants and mammals through persulfidation of specific targets. Persulfidation has been suggested as the molecular mechanism through which sulfide regulates autophagy in plant cells. ATG18a is a core autophagy component that is required for bulk autophagy and is required for reticulophagy during ER stress. In this research, we have revealed the role of sulfide in ER stress as a negative regulator of autophagy in plants. We demonstrate that sulfide regulates ATG18a phospholipid-binding activity by reversible persulfidation at Cys103, and this modification activates ATG18a binding capacity to specific phospholipids in a reversible manner. Our findings strongly suggest that persulfidation of ATG18a at C103 regulates autophagy under ER stress and the impairment of persulfidation affects both the number and size of autophagosomes.

Significance Statement

Autophagy is an extremely important conserved mechanism involved in multiple biological processes. Latest studies have reported its regulation through different posttranslational modifications. H₂S acts as an inhibitor of autophagy and recently several ATG proteins have been reported to be modified by persulfidation. ATG18a is a core autophagy protein that binds to phosphoinositides, involved in autophagosome biogenesis during phagophore expansion, and ER stress-induced autophagy is dependent on the function of ATG18a. Our data suggest that sulfide negatively regulates autophagy during ER stress through the persulfidation of ATG18a by regulating its PtdIns(3)P-binding affinity.

Main Text

Introduction

Autophagy (from the Greek meaning 'self-eating'), also known as macroautophagy, is a major catabolic process in eukaryotic cells to degrade dysfunctional or unnecessary cellular components, either non selectively or selectively (1). It has conserved functions in development, cellular homeostasis, and stress responses from yeast to plants and mammals. In plants, autophagy is critically important in many aspects of plant life, including seedling establishment, development, stress resistance, metabolism and reproduction. The autophagy mechanism involves the enclosure of a portion of the cytoplasm into a double membrane vesicle, named an autophagosome. The outer membrane of the autophagosome finally fuses with the vacuole (in yeast and plants) to release the inner autophagic body for hydrolytic degradation of the sequestered cargo. About 40 ATG (autophagy-related) genes have been identified in Arabidopsis, which are required for autophagosome formation (2). Autophagy was initially characterized as a bulk degradation pathway induced by nutrient deprivation with a role in nutrient recycling to enable cell survival, but it also contributes to intracellular homeostasis by selectively degrading aggregated proteins, damaged mitochondria, ribosomes, toxic macromolecules, excess peroxisomes, and pathogens to prevent toxicity (3-5). This selective autophagy is mediated by the binding of adaptor proteins, which link a cargo targeted for degradation to the autophagosome machinery (6). These selective autophagy receptors share the feature of interacting with the autophagosome-localized protein ATG8 through an ATG8-interacting motif (AIM), leading to their recruitment into forming autophagosomes.

An increasing number of targets for selective autophagy under different stress conditions have emerged in recent years, but the underlying mechanisms of regulation of their degradation are still so far unknown. The activation of bulk and selective autophagy must be tightly controlled by the cellular conditions. In that sense, ATG4 is the only ATG that has been shown to be redox regulated in animal, yeast, algae and plant systems (7-10). Nevertheless, in the last decade, a growing number of targets involved in autophagy have been shown to be regulated by different posttranslational modifications (PTMs); for example, ATG4b and ATG1 are regulated by S-nitrosylation and phosphorylation (11, 12). Therefore, the ability of the ATG proteins to interact with a number of autophagic regulators is modulated by different PTMs such as phosphorylation, glycosylation, ubiquitination and nitrosylation (13).

Protein persulfidation is another player in the redox regulation of certain proteins. It is the mechanism for sulfide-mediated signaling, and is an oxidative posttranslational modification of cysteine residues caused by hydrogen sulfide (H₂S) in which thiolate (-SH) is transformed to a persulfide group (-SSH). Persulfidation of proteins can affect their function, localization inside the cells, stability, and resistance

to oxidative stress. H₂S is an endogenously generated gaseous signaling molecule, which has been recently implicated in autophagy regulation both in plants and mammals (14-16).

Analysis of the *Arabidopsis des1* mutant, impaired in the cytosolic production of H₂S from cysteine, led to the conclusion that H₂S acts as an inhibitor of autophagy induced by nutrient deprivation (14). Interestingly, its action is independent of ROS and nitrogen starvation and the mechanism of autophagy inhibition by H₂S has been proposed to be through persulfidation of specific targets (17). Recently, regulation of the proteolytic activity of ATG4 by persulfidation has been demonstrated in plants (10). Autophagy induced upon nitrogen starvation or osmotic stress was negatively regulated by sulfide, and the mechanism has been explained through persulfidation of C170 of this ATG4 protease, which inhibits proteolytic activity. Collectively these results suggest that persulfidation may be the molecular mechanism through which sulfide regulates autophagy in plant cells. The susceptibility to persulfidation of the additional autophagy (ATG)-related proteins ATG18a, ATG3, ATG5, and ATG7 was revealed using a high throughput proteomic approach (18), although the role of this modification in these other ATG proteins has not yet been revealed. ATG18a is a core autophagy protein that binds to phosphoinositides (19, 20). It has a 7-bladed β -propeller structure, formed by WD40 repeats that bind phosphatidylinositol 3-phosphate (PtdIns(3)P) or phosphatidylinositol (3,5)-bisphosphate (PtdIns(3,5)P₂). These two phosphoinositide binding sites are located in blades 5 and 6 surrounding and sandwiching the conserved L/FRRG motif. ATG18a forms a complex with ATG2 which is involved in autophagosome biogenesis during phagophore expansion (20), involved in the formation of pre-autophagosomal structures and lipid recruitment. ATG18a is essential for autophagy under several abiotic stresses and (RNAi)-*ATG18a* transgenic plants showed an autophagy-defective phenotype during nutrient stress and senescence (21, 22). *atg18* mutants show defects in autophagosome formation and display an early senescence phenotype (20). In addition, the ER is a target of autophagy during ER stress in plants, and this ER stress-induced autophagy is dependent on the function of ATG18a (23). Thus, ATG18a is likely to be required for autophagosome formation in *Arabidopsis* for bulk autophagy, and also for reticulophagy during ER stress (23).

In this study, we aimed to clarify the role of sulfide in the regulation of autophagy under ER stress, through persulfidation of ATG18a. We found that persulfidation affects ATG18a lipid binding activity, which in turns regulates the number and size of autophagosomes produced upon ER stress.

Results

Persulfidation of ATG18a occurs at C103 and is sulfide-concentration-dependent.

We showed previously in a proteome-scale analysis that *Arabidopsis* ATG18a is a target for persulfidation (18). To elucidate the target site for persulfidation in ATG18a we carried out liquid chromatography (LC)-tandem mass spectrometry (MS/MS) analysis on the recombinant purified ATG18a protein. Protein was purified from an *E.coli* extract and trypsin digested under non-reducing conditions to avoid the reduction of persulfide residues. As disulfide bridges between digested peptides cannot be avoided, 2 missed cleavages were allowed in the search. The digested peptides were analyzed using LC-MS/MS for a 32-Da mass increase plus carbamidomethylation (SS-CAM) in the fragmentation spectrum. ATG18a was identified with a sequence coverage of 79% (Figure 1), and the only peptide showing SS-CAM modification was ILNCDPFR, showing C103 with a persulfide modification. All of the other seven Cys residues were identified in the analysis, but none of them were modified by persulfidation.

To determine whether the modification was sulfide-dose dependent, an in-gel detection of persulfidated ATG18a protein was performed. Purified recombinant GST-ATG18a was incubated with increasing concentrations (100 nM-200 μ M) of the sulfide donors NaHS and Na₂S₄, and with two reducing agents TCEP and DTT. A newly described dimedone switch detection method was then used (24), where Cl-NBF reacts with all Cys residues and amino groups giving a characteristic fluorescence (λ_{ex} =488 nm) (25); and then persulfide adducts were selectively labelled with Daz-2/Cy5-alkyne. Results showed that higher concentrations of either NaHS or Na₂S₄ produced a more intense Cy5-fluorescent band as a result of a higher level of persulfidation in ATG18a (Figure 2). Band quantification demonstrated that protein treated with 1 μ M of either sulfide donor showed a significant increase in protein persulfidation in comparison with untreated protein. Reducing agents efficiently reduced the intensity of the Cy5-fluorescent band, indicating a decrease in ATG18a persulfidation levels when

treated with DTT or TCEP. This shows that persulfidation of ATG18a protein is a reversible modification. Free GST protein was used as negative control to test whether persulfidation of ATG18a was affected by GST persulfidation (Suppl. Fig. 1A), and although a band was observed indicating persulfidation, it was not Na₂S₄ dose dependent. Furthermore, no persulfidated peptides from GST were identified by mass-spectrometry (data not shown). Additionally, no persulfidated band was observed in purified recombinant ATG18a_C103S (Suppl. Fig. 1B).

ER-stress-induced autophagy is negatively regulated by sulfide

ATG18a has been shown to be critical for ER stress-induced autophagy in Arabidopsis (23). The effect of sulfide on autophagy regulation was therefore tested under ER stress. Arabidopsis transgenic seedlings expressing GFP-ATG8e were treated with TM to induce ER-stress and with sulfide to test its role. Numerous autophagosomes were visualized after TM treatment in comparison with the control, where the autophagosomes were very few (Figure 3A). The number of autophagosomes in the root per frame was quantified manually, and roots treated with TM showed significantly more autophagosomes compared with control roots (Figure 3B). Treatment with sulfide had no significant effect on the number of autophagosomes in the absence of ER stress. By contrast, seedlings treated with TM and sulfide showed a drastic decrease in number of autophagosomes compared with TM alone. ER-stress can be alleviated by antioxidant treatments (26) and sulfide is able to increase the antioxidant capacity of the cell by inducing the activity of several antioxidant proteins (27-29). To test if the effect observed was due to the oxidative stress alleviation, treatments with ascorbic acid and glutathione were performed. Compared with control conditions, no effect was observed when treatments were applied under non-stress conditions. In addition, when ER-stress was induced with TM, no significant decrease in autophagosomes was detected upon ascorbic acid or glutathione treatments, indicating that the effect of sulfide is not due to its antioxidant properties.

Another approach was performed in order to corroborate these observations. WT seedlings were treated with TM for 6 hours and for a prolonged period of 12 and 24 h, and treated with sulfide for 1 hour. The effect of sulfide on autophagy was analyzed by quantification of the autophagy marker ATG8 protein by immunoblot using anti-ATG8 antibody. Results showed that seedlings incubated with TM induced autophagy with an induction peak at 12 h of treatment; meanwhile those treated together with sulfide showed a decrease in autophagy induction (Figure 3C). However, treatment with TM for 24h led to a lower induction of autophagy than shorter incubation times, probably due to toxic effect.

These data suggest that autophagy induced under ER-stress is negatively regulated by sulfide, to a much greater extent than by antioxidant treatments.

It was shown previously that the ER is degraded by autophagy during ER stress in Arabidopsis (23). Therefore, we studied the effect of sulfide on ER degradation using the fluorescent ER marker protein, GFP-HDEL, under ER stress conditions (30). Confocal microscopy showed the typical ER networks in the cytoplasm in control treatments. Similarly, in the presence of 1 μM concanamycin A (concaA) to prevent vacuolar degradation, the vacuole lacked GFP fluorescence in control samples (Figure 4). Similar results were observed in plants treated with sulfide. When plants were treated with TM, a clear increase in GFP-HDEL ER-labeling was observed in the vacuole after concaA incubation, demonstrating that ER was being degraded in the vacuole. However, plants treated with both TM and sulfide lacked the increase in fluorescence in the vacuoles, showing ER localization similar to the control treatments. To confirm that TM was causing ER stress and that the sulfide does not prevent or alleviate the ER stress, but on the contrary regulates autophagy-specific ER degradation, we assessed a bZIP60 splicing in plants treated with TM and sulfide for 6, 12 and 24 h. bZIP60 is an ER stress-responsive transcription factor that is spliced only upon ER stress (31), and the splicing event can be detected by PCR. The results showed that sulfide did not alleviate ER stress, suggesting that the effect observed was specific for autophagy; although ER stress was persistent, autophagy was blocked (Suppl. Fig. 2). Collectively, these results suggest that ER degradation by autophagy is regulated by sulfide.

Sulfide enhances ATG18a binding to membranes

Atg18 protein from *Saccharomyces cerevisiae* and WIPI3 and WIPI4 from mammals are homologues of ATG18a and they were reported to bind to PtdIns(3)P and PtdIns(3,5)P₂ (19, 32-34). The role of sulfide in the phosphoinositide-binding activity of ATG18a protein was assayed using PIP strips. PIP strips contain 100 pmol of 15 different lipids (Figure 5A) spotted onto a cellulose blotting membrane and protein-lipid binding specificity can be analysed easily.

To determine the role of persulfidation in the binding activity of ATG18a, recombinant GST-ATG18a protein was preincubated with increasing concentrations of Na₂S₄ and TCEP and then subjected to PIP strip binding. Results showed that untreated protein had specificity for PtdIns(3)P, and a weaker binding to PtdIns (3,5)P₂ was observed (Figure 5B-C). Sulfide enhanced the binding activity to both phosphoinositides, although more specifically to PtdIns(3)P. Binding was sulfide-dose dependent and treatment with 100 μM Na₂S₄ caused a substantial increase in binding. Treatment with TCEP significantly decreased binding to both phospholipids, demonstrating the reversibility of the sulfide induction of the ATG18a binding activity. Binding of the mutant recombinant protein GST-ATG18a_C103S was also assessed. Both untreated and sulfide treated protein showed a very significantly reduced binding to these phosphoinositides in comparison with the wild-type protein (Figure 5D-E). Free GST protein was assayed in PIP strips to corroborate that the tag did not affect binding in the recombinant proteins, and results showed that GST had no PtdIns(3)P -binding ability (Suppl. Fig. 3). These results suggested that the binding of ATG18a to PtdIns(3)P is significantly and reversibly increased by persulfidation. In addition, a weaker affinity toward PtdIns(3,5)P₂ was observed, although the effect of sulfide on this binding is inconclusive.

To confirm this effect *in vivo*, *Δatg18a* YFP-ATG18a_WT and *Δatg18a* YFP-ATG18a_C103S seedlings were treated with TM and sulfide and a whole protein extract was subjected to cellular fractionation, to separate cytosolic proteins and the membrane-bound fraction. Treatment with TM increased the level of YFP-ATG18a_WT in comparison with untreated seedlings, and furthermore, the proportion of YFP-ATG18a_WT protein in the membrane-bound fraction was significantly higher in seedlings treated with TM and sulfide than in those treated with only TM or (Figure 5F-G). This suggests that sulfide treatment increases the proportion of YFP-ATG18a_WT in the membrane fraction. This effect was not observed with the mutant version of the protein, YFP-ATG18a_C103S, where the proportion of bound protein to membrane was even lower than the control or the TM treated sample. These *in vitro* and *in vivo* experiments suggest that sulfide enhances ATG18a binding to membranes by persulfidation of the C103 residue.

Sulfide regulates autophagosome biogenesis by persulfidation of ATG18a

To further examine the subcellular localization of ATG18a and ATG18a_C103S in the plant cell, and to test their colocalization with ATG8e and therefore with autophagosomes and their precursors, we generated transgenic plants coexpressing YFP-ATG18a_WT or YFP-ATG18a_C103S and Cerulean-ATG8e in a *Δatg18a* background. These seedlings were treated with TM and sulfide to study the effect of sulfide during ER stress. By confocal microscopy, ATG18a and ATG8e colocalized, but colocalization time was significantly shorter in those seedlings with the mutant version of ATG18a (Figure 6A). YFP-ATG18a_WT seedlings showed a YFP/CFP colocalization mean time of 66 seconds while for YFP-ATG18a_C103S it was 24 seconds. Surprisingly, treatments with TM or sulfide did not affect the YFP/Cer colocalization time in comparison with their respective control (Figure 6B). The shortening in the colocalization time of ATG8e and the mutated version of ATG18a could be indicative of the involvement of C103 of ATG18a in autophagosome biogenesis. To assess whether the mutation in C103 alters the number or size of autophagosomes, we measured these parameters for ATG8-positive puncta within the cells of YFP-ATG18a_WT and YFP-ATG18a_C103S seedlings coexpressing Cer-ATG8e. Average size was measured for $n > 200$ puncta over three independent experiments for each condition using ImageJ (35). Results showed that the mutated version of ATG18a decreased the size of autophagosomes compared with the WT version (Figure 6C-D), suggesting that the impairment of ATG18a for persulfidation could affect the average size of autophagosomes. Furthermore, as expected, YFP-ATG18a_WT seedlings showed an increase in Cer-ATG8e puncta number when treated with TM, which was reverted after sulfide treatment (Figure 6E). However, untreated YFP-ATG18a_C103S seedlings showed an increase in autophagosome number in comparison with YFP-ATG18a_WT, which was even higher upon treatment with TM. However, sulfide treatment had no significant effect on the number of puncta in YFP-ATG18a_C103S seedlings under ER stress, with no decrease seen. To confirm the effect of YFP-ATG18a_C103S on the extent

of autophagy in comparison with YFP-ATG18a_WT, ATG8 levels were measured by immunoblot assay. Results showed that YFP-ATG18a_C103S seedlings had increased levels of ATG8 compared with WT in control conditions (Suppl. Fig. 4). In addition, TM treatment increased ATG8 levels in both genotypes but sulfide treatment was only able to decrease ATG8 levels in YFP-ATG18a_WT seedlings. These results suggest that persulfidation of ATG18a at C103 regulates autophagy under ER stress and the impairment of persulfidation affects both the number and size of autophagosomes.

Persulfidation of ATG18a affects the cavity containing C103 and the surrounding electrostatic interaction network

To examine the impact of the persulfidation posttranslational modification of C103 in the interaction between AtATG18a and phosphoinositide molecules we have performed 3D homology modelling and structural alignment using the *Homo sapiens* HsWIPI3 – ATG2A:WIR peptide complex (36). AtATG18a shares up to 43.95 % sequence identity with HsWIPI3 (E-value 2.3×10^{-32}), with conserved residues covering the full sequence. In particular, two highly conserved but not identical phosphoinositide-binding sites (site I and site II), and the potential lipid binding site III (34) are shown (Figure 7A). The 3D structural alignment revealed the lack of a β -sheet and the presence of a long flexible extension in AtATG18a compared with HsWIPI3 (Figure 7A). This long flexible loop is negatively charged and likely affects the charge distribution around binding-site I compared with HsWIPI3 (Suppl. Fig. 5). This finding might negatively influence the binding of PtdIns (3,5)P₂ to AtATG18a site I, explaining the weaker binding observed (Figure 5). It is worth mentioning that site I and site II differ in lipid affinity. Site I tends to recognize preferentially PtdIns (3,5)P₂ whereas site II interacts with PtdIns(3)P (34).

The residue C103 is located inside a hydrophobic cavity formed by residues ⁸³FNQD⁸⁶ and F90 which are highly conserved in the WIPI members of PROPPIN protein family (36). In fact, these residues of AtATG18a are at the exact positions of those of human WIPI3 (Figure 7A inset). Studies in the homologues WIPI4 and WIPI3 have demonstrated that the equivalent residues N15/N19, and D17/D21, respectively, to N84 and D86 of AtATG18a are essential for ATG2A binding (36, 37). Interestingly, the residue N84 and the neighbouring residues R128 and S218, equivalents to R62 and S151 of human WIPI3, respectively, and exposed at the exterior of the protein, have been reported to participate in an additional potential lipid binding site III (34). Positive charges around this region (Figure 7B) would govern the electrostatic interaction with other negatively charged partners such as phosphoinositide molecules. Moreover, the comparison of HsWIPI3 with the AtATG18a model reveals an increase of positive charge distribution on the surface of site III in the Arabidopsis protein (Suppl. Fig. 4).

Consistent with this, persulfidation of C103 could potentially affect the molecular recognition in its neighbouring protein region and consequently the function of AtATG18a. The cysteine persulfidation would not result in a charge perturbation, since the additional –SH group is a neutral contribution. However, the addition of a –SH group to C103 could cause a steric perturbation in the cavity (Figure 7C), destabilizing the original hydrophobic interactions and promoting new ones. The additional sulphur atom would contribute with a covalent radius of 1.02 Å, while that of hydrogen is 0.37 Å. It is worth mentioning that the aromatic-thiol π hydrogen bonding interaction is very sensitive to the orientation of the two lone electron pairs on the sulphur atom relative to the π electron cloud of the phenyl ring [48]. In addition, the replacement C103S could also destabilize the hydrophobic cavity site but in a different manner. Although the energetic cost of desolvating serine (-OH) is higher than for a cysteine (-SH) residue, the steric effect due its replacement would be much smaller than that caused by the additional –SH group (Figure 7B).

Altogether, the C103 modification by persulfidation likely could cause a conformational change and intramolecular rearrangement of the cavity favouring the interaction with additional phosphoinositide molecules at site III and/or increasing the binding affinity of PtdIns(3)P for its target at site II through an allosteric cooperative effect.

Discussion

Autophagy regulation in plants has become an important, although challenging, topic in recent years, due to its involvement in plant health and cellular homeostasis, helping plants to tolerate different

stresses such as oxidative stress, pathogen infection, nutrient deprivation, and drought (21, 22, 38, 39). It is now well known that ER stress triggers autophagy both in mammals and plants (23, 40), as a protective mechanism against misfolded proteins accumulating in the ER. Thus, a selective autophagy pathway named ER-phagy is activated to recycle accumulated misfolded proteins in the ER in order to restore ER homeostasis (41). In plants, it has been shown that inositol-requiring enzyme 1 (IRE1) is necessary for ER-phagy, and the delivery of ER to the vacuole for degradation under ER stress depends on the ATG18a protein (23). In mammals, recent studies showed that H₂S may influence ER stress, playing an important role in many heart, neurological, and respiratory diseases (42). In plants, H₂S sulfide was shown to negatively regulate bulk autophagy under nutrient deprivation, independently of redox conditions (14, 17). This mechanism was explained by persulfidation of ATG core proteins (10, 18). In this work, we have demonstrated that sulfide also regulates autophagy induced by ER stress, and this regulation was independent of its indirect antioxidant effect. Furthermore, we showed that sulfide played a signaling role in ER degradation by autophagy.

Although little has been reported about the biochemical function of plant ATG18a protein, in yeast and mammals a function for the Atg2-Atg18 complex in autophagy has been reported, and the binding of Atg18 to PtdIns(3)P is crucial in autophagy induction (43). Post-translational modifications of ATG18 are known to control its activity. Atg18 in the yeast *Pichia pastoris* was shown to be phosphorylated, and phosphorylation enhanced its binding affinity to PtdIns(3)P (44). However, this modification in *P. pastoris* does not affect autophagy activity, while phosphorylation of Arabidopsis ATG18a inhibits its function in autophagy (45). Therefore, posttranslational regulation of ATG18a must be finely controlled, and probably triggered depending on the stress to which the plant is subjected. Furthermore, ATG18a is also required for bulk autophagy and persulfidation could possibly regulate its activity in multiple stresses in addition to ER stress.

In this study, we explored the functional importance of sulfide in regulating ATG18a binding to PtdIns(3)P. Mass spectrometry analysis of recombinant ATG18a showed that only one Cys residue, C103, was persulfidated under our experimental conditions. Protein persulfidation at more than one Cys residue is rarely found in the literature, and the other Cys residues of ATG18a were not detected as modified peptides, indicating that C103 was the only target for persulfidation. Furthermore, this PTM was shown to be H₂S dose dependent at very low concentrations of NaHS and reversible by reducing agents, suggesting this modification may have a biological role in plants. We further showed the potential physiological role of the persulfidation of ATG18a by using WT and C103 mutant recombinant proteins in an *in vitro* assay; one important finding was that persulfidation of ATG18a increased its affinity for PtdIns(3)P, while impairment of persulfidation abolished almost completely this binding to phospholipids. *In vivo*, sulfide enhanced the fraction of ATG18a bound to membranes, while the mutant ATG18a_C103S was not affected by sulfide treatment and in general, the ATG18a_C103S fraction bound to membranes was less than that of the WT ATG18a protein.

We observed that the mutant protein ATG18a_C103S, impaired in persulfidation, was unable to bind properly to PtdIns(3)P, and therefore the colocalization time with an autophagosome marker was shorter than that of the wildtype protein. Furthermore, autophagosomes in plants expressing only the ATG18a_C103S mutated version had a smaller average size than when the WT counterpart was expressed. However, it seems that plants respond to this situation by increasing the number of autophagosomes. ATG18a forms a complex with ATG2 that is involved in the expansion of the membrane of the autophagosome for phagophore elongation (46), therefore, it makes sense than disturbing the binding affinity of ATG18a for phosphoinositides could affect autophagosome size. In addition, the colocalization time of ATG18a_C103S with autophagosomes is shorter and its binding affinity to PtdIns(3)P is weaker than the WT ATG18a, resulting in an outcome of smaller autophagosomes, suggesting that C103 is important in regulating elongation of phagophore membranes. Previous studies in yeast demonstrated that insufficient levels of Atg7 affect both autophagosome size and number (47), and similarly, mutations disturbing Atg3 function also affect autophagosome size (48). The fact that malfunction of ATG18a has an effect on both autophagosome size (expansion) and number (nucleation) of autophagosomes may indicate that ATG18a could have additional roles in autophagosome formation.

In yeast, it was demonstrated that Atg18 is important for the localization of the Atg2-Atg18 complex to the pre-autophagosomal structure (PAS), through its PtdIns(3)P-binding ability (49, 50). In fact, some studies showed that autophagosome formation could efficiently progress in *atg2Δatg18Δ* cells expressing engineered variants of Atg2 that are capable of localizing to the pre-autophagosomal structure (PAS). Engineered yeast expressing Atg2-HG-FYVE, a fusion of Atg2 with the

phosphatidylinositol 3-phosphate-binding FYVE domain, or Atg2–HG–Atg8, expressing Atg2 fused with the core autophagy protein Atg8, allowed a partial recovery of autophagosome formation independently of Atg18 (51). Nevertheless, the partial recovery suggested that Atg18 possesses other functions beyond targeting Atg2 to the PAS. The general structure of this complex is similar to the mammalian homolog ATG2B-WIPI4, suggesting that the overall shape and function is evolutionarily conserved among species (52, 53). Our results obtained from the 3D structural alignment suggested that persulfidation of C103 could modulate the interaction with PtdIns(3)P molecules by inducing a conformational change of the surrounding site that could affect recognition of potential partners such as PtdIns(3)P through the electrostatic interaction network. Therefore, persulfidation of ATG18a could affect autophagy progression by regulating its PtdIns(3)P-binding affinity.

It is noteworthy that previous studies described how sulfide regulates bulk autophagy triggered by nutrient starvation or ABA (10, 17) and in this study, we demonstrate that ER-induced autophagy is also negatively regulated by sulfide. We provide a new level of regulation of autophagy in plant systems by sulfide, through the persulfidation of ATG18a. Together, our data suggest a model in which the dynamics of ATG18a association with forming autophagosomes is critical for autophagy during ER stress. The reversible persulfidation of ATG18a increases its binding to membranes via association with PtdIns(3)P, but potentially delays its release, inhibiting autophagosome maturation. Preventing persulfidation by mutation of C103 decreases its membrane affinity and localization time to phagophores. As ATG18a is thought to be involved in lipid transfer during autophagosome expansion, this decreased phagophore localization leads to the production of smaller autophagosomes. Autophagosome numbers increase, potentially to compensate for this. In this way, persulfidation of ATG18a may modulate the extent of autophagosome production during stress to ensure an appropriate physiological response.

Materials and methods

A detailed description of all materials and methodology is included in SI Appendix, Materials and Methods. This includes the expression and purification of recombinant Glutathione S-transferase (GST)-tagged protein, mass spectrometry, persulfide detection in cell lysates, plant genotypes, protein modelling, protein-lipid binding assay, membrane fractionation, autophagy detection, microscopy, bZIP60 splicing assay and expression and purification of free Glutathione S-transferase (GST) protein.

Data Availability.

All study data are included in the main text and SI Appendix.

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Figures and Tables

A

MATVSSSSWPNPNPNDSTSASDSDSTFPSHRDRVDEPDSLDSFSSMSLNSDEPN
 QTSNQSPSPPTPNLPVMPPPSVLHLSFNQDHACFAVGTDRGFR ILNCDPFER E I F
 RRDFDRGGGVAVVEMLFR **C**NI LALVGGGPDQYPPNKVMIWDDHQRC **C**IGEL SFR
 SDVRSVRLRR **D**R I I VVLEQK I FVYNFSDLKLMHQIETIANPKGL **C**AVSQGVGSMV
 LV **C**PGLQKGQVR I EHYASKRTK FVMAHDSRIA **C**FALTQDGHLLATASSKGTLVRI
 FNTVDGTLRQEVRRGADRAE I YSLAFSSNAQWLAVSSDKGTVHVFGLKVN S G S Q V
 K D S S R I A P D A T P S S P S S L S L F K G V L P R Y F S S E W S V A Q F R L V E G T Q Y I A A F G H Q K
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B

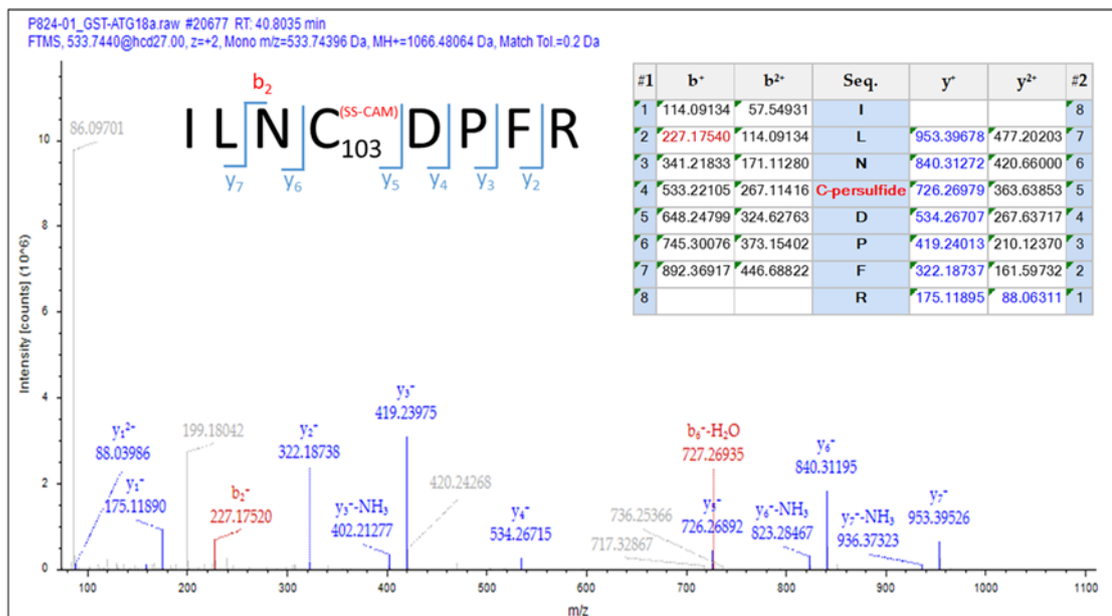


Figure 1. Analysis of AtATG18a using mass spectrometry. (A) The protein was identified with a sequence coverage of 79% highlighted in blue; Cysteines are highlighted in yellow and the peptide containing persulfidated Cys-103 is shown in red underlined. (B) LC-MS/MS analysis of the tryptic peptide containing Cys-103 of ATG18a. The table inside the spectrum contains the predicted ion types for the modified peptide, and the ions detected in the spectrum (Biemann, 1988) are highlighted in red and blue.

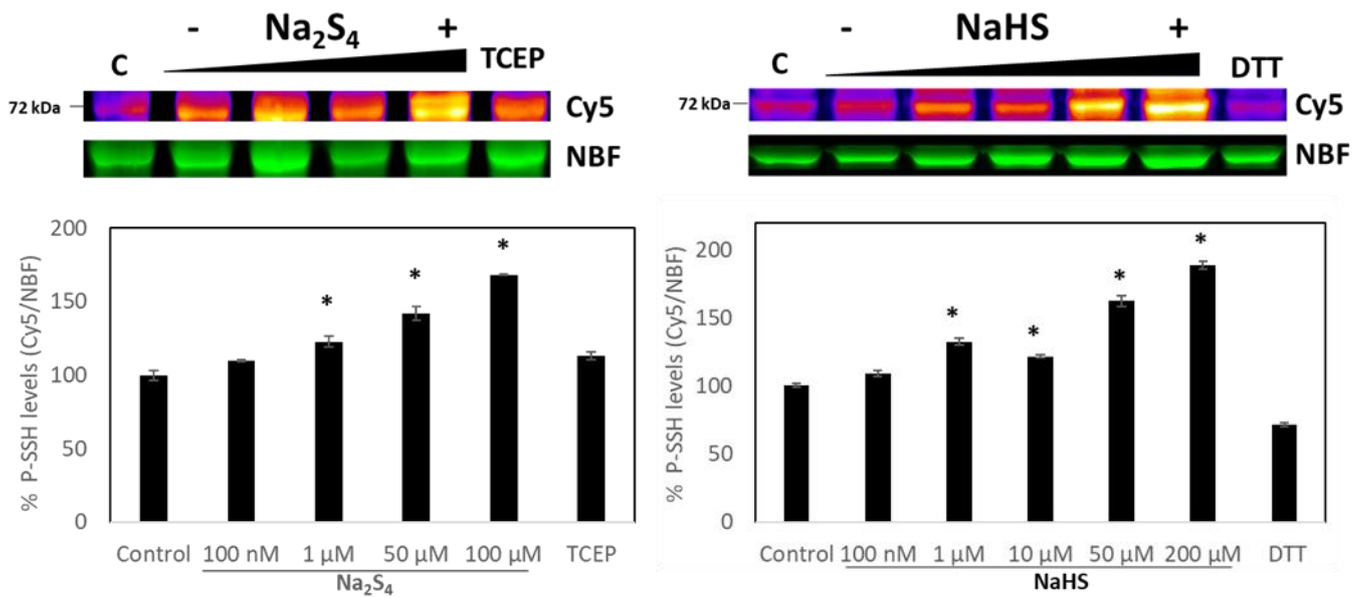


Figure 2. In-gel detection of recombinant AtATG18a protein persulfide labeling. Persulfides labelled with DAz-2/Cy5-alkyne using CuAAC are represented in fire pseudo-coloring to visually enhance the signal. Green fluorescence corresponds to the total protein loaded (NBF-protein adducts). Increasing concentrations of the sulfide donors Na₂S₄ (A) and NaHS (B) were used and two reducing agents, 5mM TCEP or 25mM DTT. Control samples were untreated (C). Ratio of Cy5/NBF signals is used for the quantification. Data is shown as the mean ± SD of 3 individual experiments. *p < 0.01 versus control.

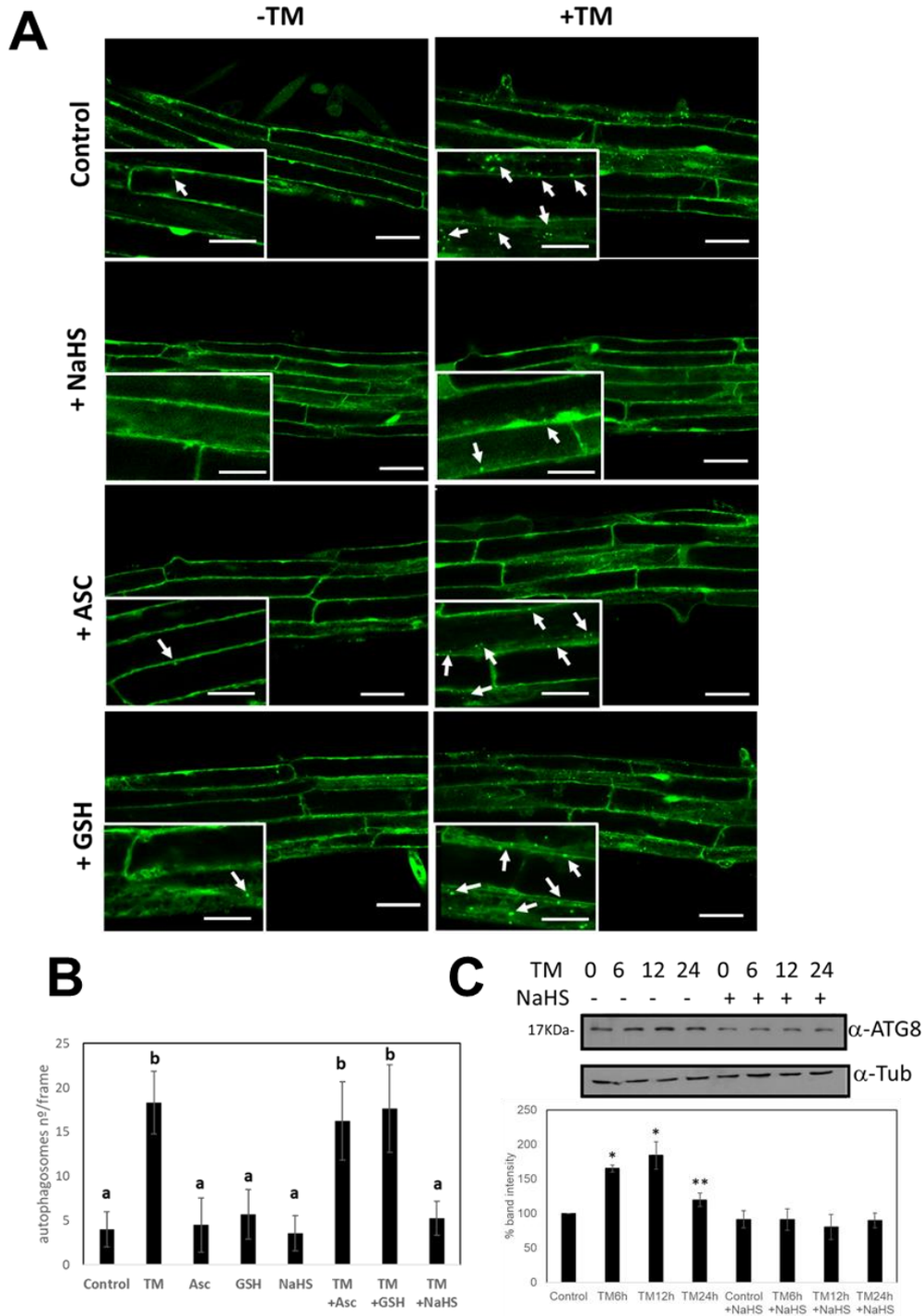


Figure 3. ER-stress induced autophagy is regulated by sulfide in *Arabidopsis* roots. Seven-day-old GFP-ATG8e transgenic plants were transferred to MS liquid medium supplemented with TM to induce ER stress, sulfide (NaHS), ascorbic acid (ASC) and/or glutathione (GSH), or DMSO (Control) (A) GFP-ATG8e-labeled autophagosomes in root epidermal cells were visualized by confocal microscopy. Arrows indicate GFP-labeled autophagosomes or autophagic bodies. Insets show enlargement. Scale bars, 40 μ m for main figure and 20 μ m for insets. (B) Quantification of autophagosomes per frame for each treatment. Values are the mean \pm SD. ($n > 15$) Different letters indicate statistically significant differences (ANOVA, Fisher's least significant difference (LSD) test, $p < 0.05$). (C) Immunoblot analysis using anti-ATG8 antibodies in protein extracts from WT *Arabidopsis* seedlings treated with TM for 6, 12 and 24 hours and NaHS. Graph shows the relative band intensity. Bars represent means \pm SD ($n = 3$). * $p < 0.01$, ** $p < 0.05$.

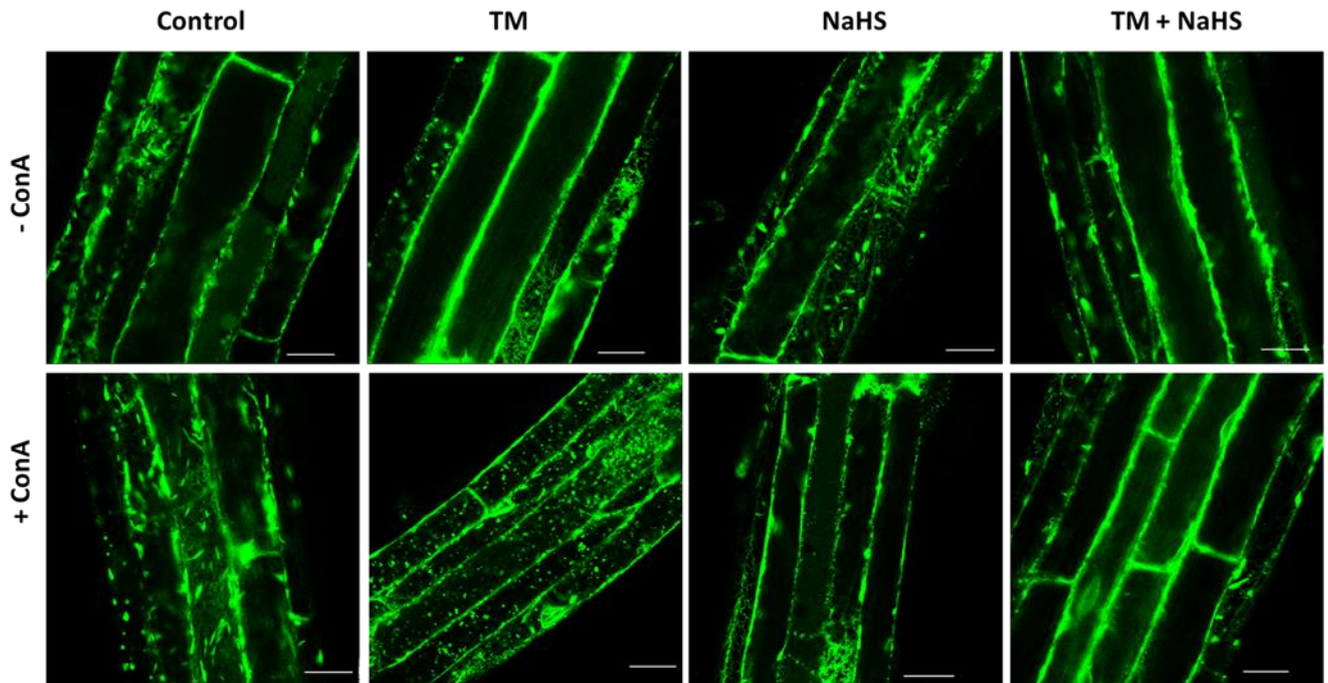


Figure 4. ER degradation by autophagy is regulated by sulfide in *Arabidopsis* roots. Seven-day-old GFP-HDEL transgenic plants were transferred to MS liquid medium supplemented with tunicamycin (TM) for 6 h to induce ER stress, and sulfide (NaHS) for 1 h. DMSO was used as solvent (Control). ER structure in root epidermal cells was visualized by confocal microscopy. Scale bars, 40 μ m.

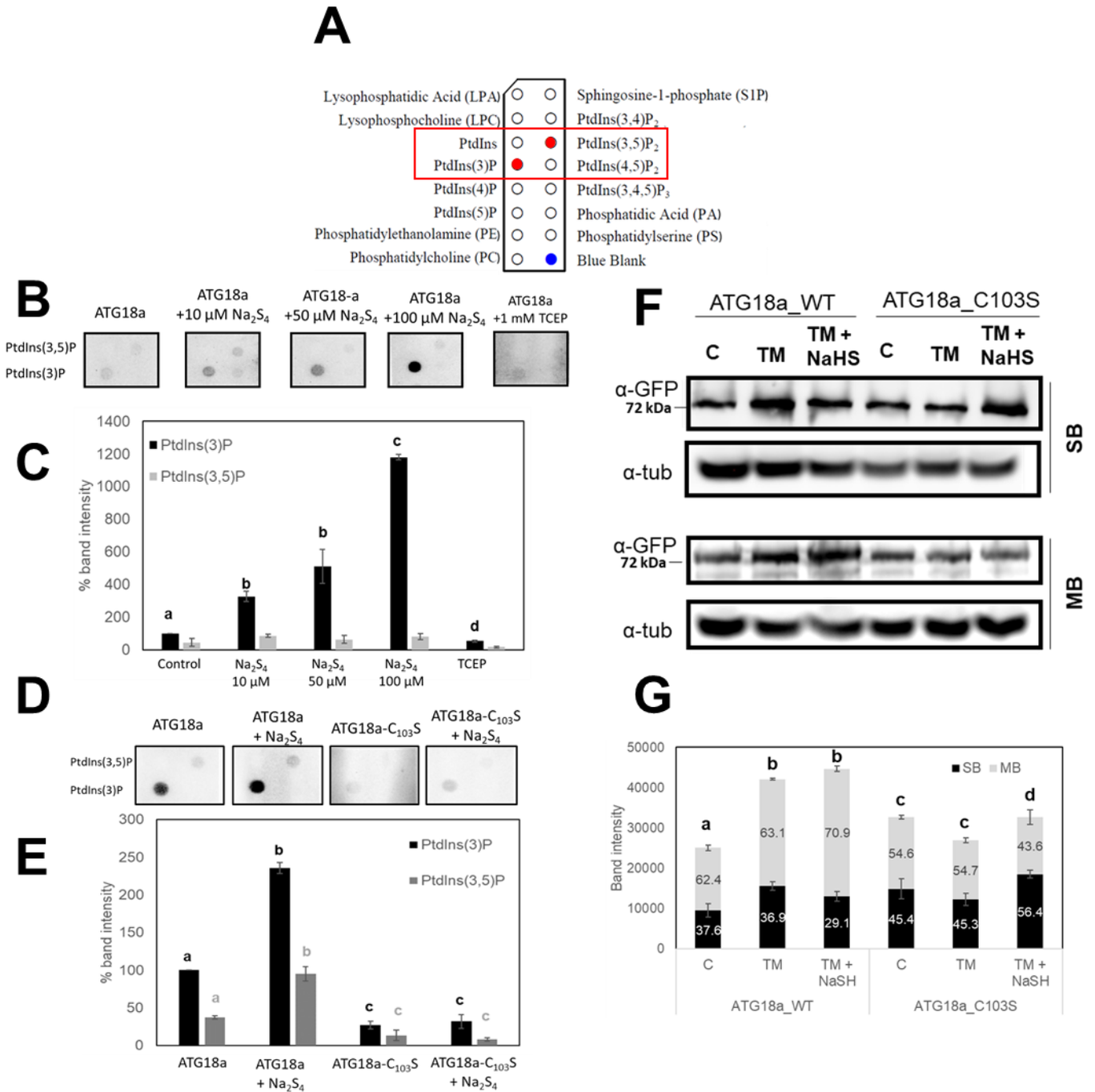


Figure 5. Sulfide enhances ATG18a binding to membranes (A) Schematic representation of the indicated membrane lipids on a PIP strip. Red dots represents the specific lipids PI3P and PI3,5P2 for panels B and D. (B) Binding of recombinant GST-ATG18a proteins to PIP strips. 0.5 μ g of protein was incubated with increasing concentration of NaHS and TCEP, followed by incubation with PIP strips. Bound protein was detected using anti-GST primary antibody and anti-rabbit-HRP secondary antibody. (C) Spot intensity relative quantification of panel B. (D) Comparison of binding affinity to PIP strips of 1 μ g of GST-ATG18a and GST-ATG18a_C103S treated or not with 10 μ M Na₂S₄. (E) Spot intensity relative quantification of panel D. (F) Immunoblot of soluble and membrane-bound fractions of the fusion proteins YFP-ATG18a and YFP-ATG18a_C103S, obtained from transgenic lines treated with tunicamycin (TM) for 6 hours and sulfide (NaHS) for 1 hour. (G) Relative band intensity quantification. Bars represent means \pm SD (n = 3). Different letters indicate statistically significant differences (ANOVA, Fisher's least significant difference (LSD) test, p < 0.05).

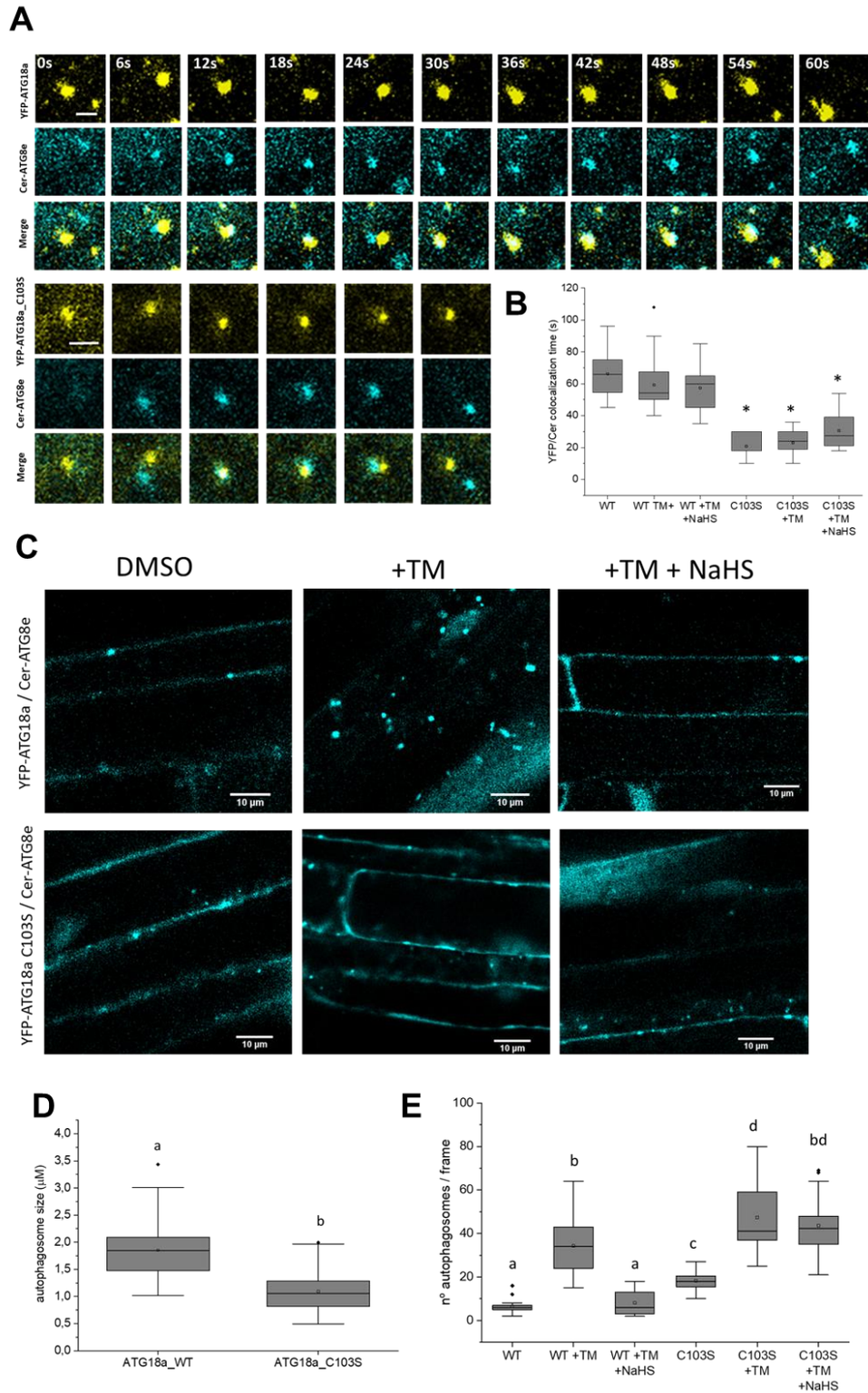


Figure 6. Time-lapse confocal fluorescence microscopy of autophagosomes in roots from 1-week-old seedlings treated with tunicamycin (TM) and sulfide (NaHS). (A) Confocal images for colocalization of YFP-ATG18a and YFP-ATG18a_C103S with Cer-ATG8e. Scale bars, 2 μ m. (B) Analysis of YFP/Cerulean colocalization time. Duration of colocalization is represented in seconds. ($n > 15$). (C) Confocal images for Cer-ATG8e in the double transgenic plants YFP-ATG18a/Cer-ATG8e and YFP-ATG18a_C103S/Cer-ATG8e after treatment with TM and NaHS. (D) Autophagosome size analysis measured with ImageJ, $n = 150$ (E) Autophagosome number quantification counted manually with ImageJ, $n = 15$. Values are the mean \pm SD. Different letters indicate statistically significant differences (ANOVA, Fisher's least significant difference (LSD) test, $p < 0.05$).

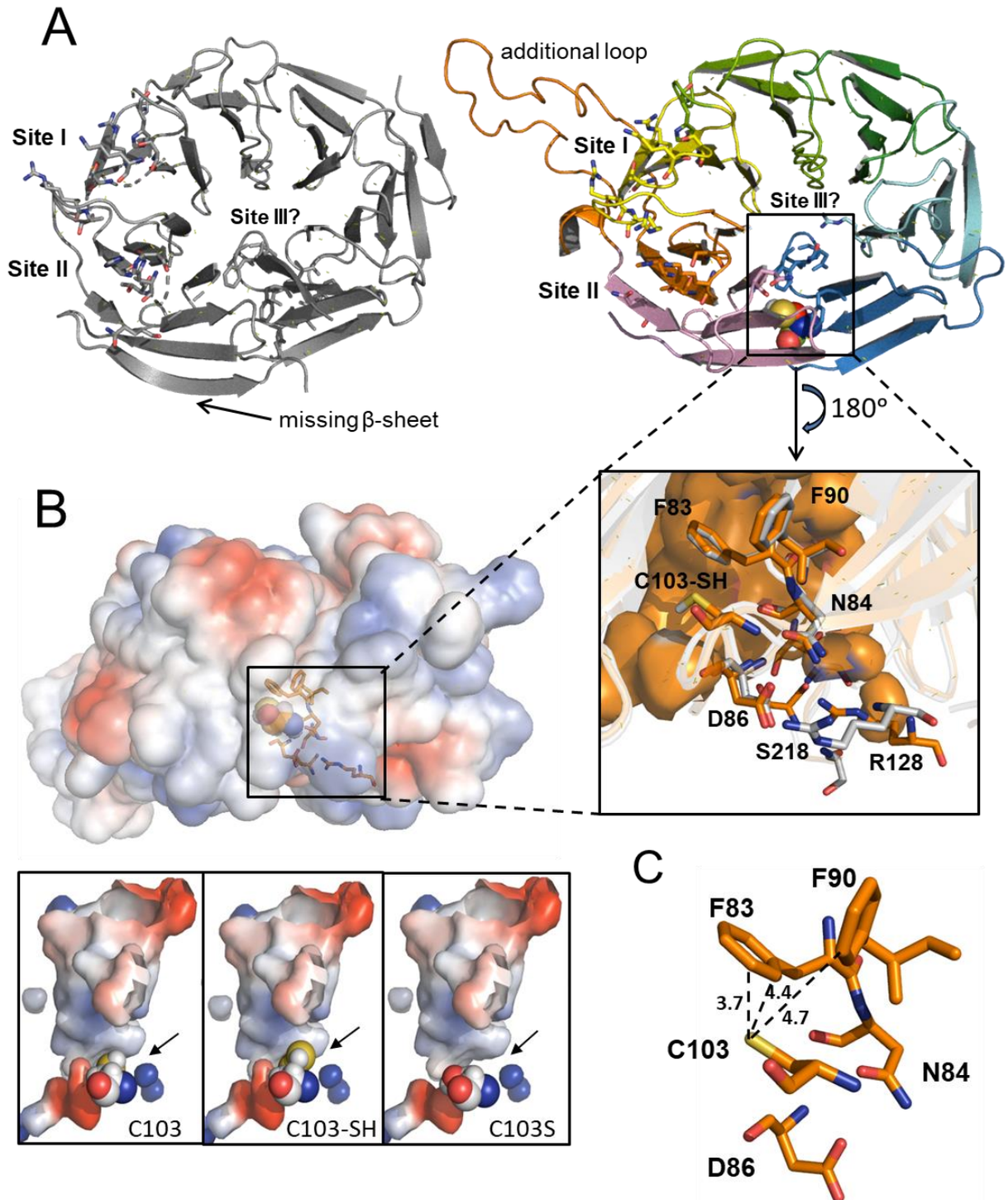


Figure 7.- Predicted structure of AtATG18. (A) Crystal structure of human WIPI3 (PDB ID: 6KRL, left panel) and predicted structural model of Arabidopsis ATG18a (right panel). The proposed lipid binding-sites and the cavity site of C103 are highlighted. The inset shows the residues of AtATG18a surrounding the active C103 site (F83, N84, D86 and F90) and participating in binding-site III (R128 and S218). The equivalent residues in HsWIPI3 are in grey. (B) Representation of surface electrostatic potential distribution in AtATG18a structural model and zoomed into the putative conformation of the active site, showing with spheres the position of C103, the persulfidation C103-SH, and the mutation C103S. Positively and negatively charged regions are depicted in blue and red, respectively. (C) Zoom into the putative conformation of the active site showing distance (\AA) between the catalytic residue C103 and F83 and F90 in AtATG18a.